The Shell Bitumen Handbook

Sixth edition
The Shell Bitumen Handbook

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Principal authors
Dr Robert N. Hunter, Andy Self and Professor John Read
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Dr Robert N. Hunter BSc MSc PhD CEng FICE FIAT MCIArb FCInstCES

Robert has always had an intense interest in asphalt technology and flexible pavements, spending many years on a range of sites supervising highway construction contracts. He undertook two postgraduate research degrees in which he examined various aspects of the behaviour of asphalts. He wrote a computer program that calculates the temperatures in an unchipped asphalt after laying, and wrote the only computer program that calculates the temperatures in a chipped asphalt after laying. In addition, he produced the algorithm on which the only computer program that designs UK compliant pavements and foundations is based.

For much of his career Robert worked for local authorities, and was general manager of a trunk road unit in east and central Scotland. He now runs his own consultancy specialising in providing an expert witness service related to asphalts, pavements and defects in highways. He also advises on contract related issues, and produces contract documents invariably based on NEC3, which he adores. He is also a registered adjudicator with the Institution of Civil Engineers and the Chartered Institute of Arbitrators, and has been appointed as adjudicator in many disputes.


Andy Self BSc(Hons) MBA

Andy has worked in the road construction industry for over 30 years. He began his career working as a laboratory technician, and spent almost 20 years working in various technical roles for asphalt and quarrying companies in the UK. During this time he gained extensive experience in asphalt production, laboratory testing of construction materials, asphalt mixture design and product quality...
management. In 2002, he gained a BSc Honours degree with the Open University.

In January 2003, Andy moved to Shell as asphalt development manager for the bitumen business in UK and Ireland, working closely with customers, suppliers and colleagues on product development. More recently, he worked as bitumen technical manager Europe; managing teams in France, Germany, Norway and the UK, and representing Shell on various industry association committees. Andy completed his MBA during his recent role as skillpool manager in the Shell Specialities (bitumen and sulfur) global technical team, and he now works as global learning advisor for the Shell Specialities business.

Professor John Read PhD BEng(Hons) CEng MIM MIAT MIHT MAAPT
Professor John Read has worked in the bitumen and sulfur industry for nearly 30 years, initially for a consultant testing house, then for an asphalt supplier and more recently for Shell in various roles. He is currently the general manager technology for Shell Specialities, where his principal accountabilities are leading a world class team of technical specialists to deliver research and development and technical services to customers in more than 25 countries around the world.

During his career he has delivered more than 100 public presentations at conferences, industry associations, universities and industry training days, and has published more than 80 papers, articles and technical brochures, including co-authoring the 5th edition of the internationally renowned The Shell Bitumen Handbook.

John has recently finished two terms as the vice-president of Eurobitume, and serves as Shell’s primary director on the Asphalt Institute.

John has two children and lives and works from his home in Derbyshire, UK.
Foreword

It is more than 10 years since I authored the 5th edition of *The Shell Bitumen Handbook* and it gives me great pleasure to write this foreword for the 6th edition. The whole book has been completely reviewed by international experts from Shell, complemented by experts from across the industry, resulting in a much more global approach than the 5th edition, with contributions from Argentina, Australia, China, France, Germany, India, Indonesia, the Netherlands, Malaysia, Thailand, the United Arab Emirates, the USA and the UK. Each chapter has been either edited or completely rewritten. There are several new sections and chapters on recycling, warm asphalt mixtures, foam mixtures, bitumen additives and certification schemes for bitumen and asphalt. The result is a lasting testament to the continued contribution that Shell makes to asphalt technology, as well as confirmation to the wealth of knowledge, expertise and experience that all the authors possess.

Many people have contributed to this edition of the handbook, and I sincerely thank them all, but it would be only fitting for me to give special mention to two individuals in particular: Andy Self and Dr Robert Hunter. As well as being an asphalt expert in his own right, Andy Self has been nursemaid to all the authors, a tyrannical timekeeper and a truly good friend. Without his organisational skills this 6th edition would never have been completed. Dr Robert Hunter agreed to be the technical editor for the second time, and his eye for detail has ensured the quality publication that this book has become. I also need to mention that it was Robert who encouraged me to write the book again, and it is his tenacity that has ensured we have all delivered on time – without him you would not be reading this book. I give my heartfelt thanks to both Andy and Robert.

In closing, I would like to mention all members of the Shell bitumen technology team, who work tirelessly in support of pursuing the advancement of asphalt technology, as well as their families, who are always so understanding and supportive of the amount of time we give – thank you all. You continue to ensure we remain a technology leader, and individually you inspire me each and every day. Lastly, I would like to thank you, the reader, for acquiring this book and taking the time to read it. I hope that you find it a useful resource and are able to benefit from the huge amount of knowledge and experience that has gone into this encyclopaedia of bitumen and asphalt technology.

Professor John Read, General Manager Technology (Bitumen & Sulfur)
May 2014
Acknowledgements

Robert Hunter is delighted to acknowledge the immense support he has enjoyed during the production of this book. First, he would like to thank: the wonderful Debra Francis, librarian at the Institution of Civil Engineers; Professor Ian Walsh, for his immense technical assistance and Bob Allen who, as well as writing a superb chapter dealing with aggregates in asphalts, freely gave enormous assistance on many occasions. Other people he is delighted to thank are Jack Edgar, Lewis Hunter, Russell Hunter, Peter Dick, Eddie Lord, Jeff Farrington, Gordon Steel, Dr Behrooz Saghafi, Andrew Scorer of Miles Macadam Ltd, Rory O’Connor of Tarstone Surfacing Limited, David Merritt of The Transtec Group, Inc, Duncan Tharme of the Curtiss-Wright Corporation, Stephen Collins representing the Road Emulsion Association (REA), Thomas Jennings of The Phoenix Engineering Co Ltd, Peter Wallace of PTS International Ltd and, last but certainly not least, his very good friend Tony Sewell of PTS International Ltd.

Finally, Robert would like to express his gratitude to: Shell, for asking him to work on this book; the authors, many of whom wrote on very complex subjects in a language that is not their native tongue; Andy Self, who is one of the most efficient gentlemen I have ever met and an absolute joy to work with; and Professor John Read, as ever a great friend and a source of substantial support.

Andy Self would like to thank his fellow principal authors and the project team – Professor John Read, Dr Robert Hunter, Diny Rovers, Jayne Davies, Jenny Marsden, Rachel Gerlis and Rebecca Taylor. A very special thanks to Dr Robert Hunter for his tireless and uncompromising commitment to the new edition.

Andy and John would like to acknowledge formally their colleagues at Shell: Dr Richard Taylor and Dr Jia Lu for excellent and timely contributions to several chapters. Others thanked for their support and contributions are: Carlos Maurer, Sonia Hauguel and the team at the Strasbourg Solution Centre, Indra Maizir, Yan Hui, Gary Fitts, Steve Sturridge and John Barlow.

Richard Taylor wishes to thank Dr Mark Bouldin of Oil Re-Refining Specialty Products at Safety-Kleen.
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Frank Beer wishes to thank Jacques Colange, Dr Martin Vondenhof, Jayne Davies, Dr Andreas Opel, Catherine Noireaux and Punith V Shivaprasad, all of Shell.

The authors of Chapter 18 would like to recognise formally the original contribution of Dr Mike Nunn.
An emulsion manufacturing unit located in Bangkok, Thailand

A polymer modified bitumen manufacturing plant in Buenos Aires, Argentina
This book is about refined bitumen, one of the world’s most widely used construction materials. The *Shell Bitumen Handbook* is one of the most well known and highly respected publications on the subject of bitumen and asphalt technology. Thousands of copies have been sold since it was first published. Many people within the construction industry have a copy somewhere in their office or library. Most importantly, it is widely referenced in academic papers.

This new sixth edition covers the same subject matter as previous editions, but a slightly different approach has been taken as to how it has been compiled. For the first time, each chapter has been written by a different subject matter expert. The authors come from many different countries. The intention is to bring a wider, more global focus to the subjects and reflect the fact that growth economies in countries such as China and India are adapting and developing new practices and specifications and driving research in this area.

Shell has a distinguished track record in bitumen research and development, and continues to invest significantly in fundamental research (Table 1.1). Investment in regional technical service laboratories and the recruitment and development of local expertise in many markets has produced a truly global team of subject matter experts. In this edition, this expertise has been used to create something that looks and feels different to previous publications, and is unique in its scope and depth. The aim is to capture all the main new developments in the field since the last edition.

The book is not a review of current academic research and theory, although some chapters refer to recent studies. It is intended to reflect today’s state of the art in terms of practice, knowledge and trends. The intention has been to make it relevant for a wide range of readers and to cover the subject in as much detail as possible.
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<td>Crumb rubber modified bitumen (Shell Mexphalte RM\textsuperscript{\textregistered})</td>
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Readers can use the book in different ways. If you are new to the subject, it makes sense to start at the beginning and read a few chapters before delving into the more complex technical content. Others may like to read specific chapters, or simply use the book as a reference to dip into when required.

The book is structured as follows: Chapters 2 to 9 focus on bitumen technology, covering manufacturing, handling and the main engineering properties and specifications used around the world. The following six chapters (10 to 15) focus on asphalt and its use in flexible pavements – this includes: the main ingredients, design methods, manufacturing and construction. Chapters 16 to 20 cover the testing and performance characteristics of asphalts. The final four chapters cover some other uses of bitumen and asphalt, including two new chapters (for this edition) on alternative approaches to the design and manufacture of asphalts and product certification schemes.

1.1 Origins

Bitumen is defined in the *Oxford English Dictionary* as ‘a tarlike mixture of hydrocarbons derived from petroleum naturally or by distillation, and used for road surfacing and roofing’ (Oxford University Press, 1996).

It is widely believed that the term bitumen originated in the ancient and sacred language of Hindus in India, Sanskrit; in which *jatu* means ‘pitch’ and *jatu-krit* means ‘pitch creating’. These terms referred to the pitch produced by some resinous trees. The Latin equivalent is claimed by some to be originally *gwitu-men* (‘pertaining to pitch’) and by others to be *pixtu-men* (‘bubbling pitch’), which was subsequently shortened to bitumen before passing via French into English.

There are several references to bitumen in the Bible, although the terminology used can be confusing. In Genesis, Noah’s ark is ‘pitched within and without with pitch’, and Moses’ juvenile adventure is in ‘an ark of bulrushes, daubed with slime and with pitch’. Even more perplexing are the descriptions of the building of the Tower of Babel. The Authorised Version of the Bible says ‘they had brick for stone, and slime had they for mortar’; the New International Version states that ‘they used bricks instead of stone and tar instead of mortar’; Moffat’s 1935 translation says ‘they had bricks for stone and asphalt for mortar’; but the New English Bible states that ‘they used bricks for stone and bitumen for mortar’.

The ancient uses of natural bitumens or pitch continued in those inhabited parts of the world where deposits were readily available. However, there seems to have been little development of usage elsewhere. In many countries, none of the present major uses of bitumen were introduced until the end of the nineteenth century. However, there would appear to have been some wider knowledge of large sources of natural bitumen such as lake asphalt in Trinidad. In the middle of the nineteenth century, attempts were made to utilise rock asphalt from European deposits for road surfacing and, from this, there was a slow
The Shell Bitumen Handbook

development of the use of natural products for this purpose, followed by the advent of coal tar and, later, of refined bitumen manufactured from crude oil.

1.2 Definition of bitumen

The term ‘bitumen’ is used in this book to describe refined bitumen, a hydrocarbon product produced by removing the lighter fractions (such as liquid petroleum gas, petrol and diesel) from crude oil during the refining process. In North America, bitumen is commonly known as asphalt binder or asphalt. For the purpose of this book, the term ‘bitumen’ is used.

A comprehensive definition of refined bitumen is used in the industry document The Bitumen Industry – A Global Perspective (Eurobitume and the Asphalt Institute, 2011) and is reproduced here verbatim:

Bitumen is an engineering material and is produced to meet a variety of specifications based upon physical properties. Bitumen is the residual product from the distillation of crude oil in petroleum refining. The basic product is sometimes referred to as ‘straight run’ bitumen and is characterised by CAS# 8052-42-4 or 64741-56-6 which also includes residues obtained by further separation in a deasphalting process. Bitumen can be further processed by blowing air through it at elevated temperatures to alter its physical properties for commercial applications. The general characteristics of oxidized bitumen are described by CAS# 64742-93-4. The vast majority of petroleum bitumens produced conform to the characteristics of these two materials as described in their corresponding CAS definitions.

Bitumen is produced to grade specification either directly by refining or by blending.

Bitumen should not be confused with coal derived products such as coal tar or coal tar pitches. These are manufactured by the high temperature pyrolysis (>800°C) of bituminous coals and differ from bitumen substantially in comparison and physical characteristics. The differences between bitumen and coal-tar products are well defined in the literature.

Similarly, bitumen should not be confused with petroleum pitches (CAS# 68187-58-6), which are often aromatic residues, produced by thermal cracking, coking or oxidation from selected petroleum fractions. The composition of petroleum pitches differs significantly from bitumen.

Bitumen also should not be confused with natural or lake asphalt such as Trinidad Lake Asphalt, Gilsonite, rock asphalt and Selenice. These products are unrefined and not produced by refining of crude oil. They often contain a high proportion of mineral matter (up to 37% by weight) and light components, leading to a higher loss of mass when heated.
Bitumen is manufactured during the distillation of crude oil. It is generally agreed that crude oil originates from the remains of marine organisms and vegetable matter deposited with mud and fragments of rock on the ocean bed. Over millions of years, organic material and mud accumulated into layers some hundreds of metres thick, the substantial weight of the upper layers compressing the lower layers into sedimentary rock. Conversion of the organisms and vegetable matter into the hydrocarbons of crude oil is thought to be the result of the application of heat from within the Earth’s crust and pressure applied by the upper layers of sediments, possibly aided by the effects of bacterial action and radioactive bombardment. As further layers were deposited on the sedimentary rock where the oil had formed, the additional pressure squeezed the oil sideways and upwards through porous rock. Where the porous rock extended to the Earth’s surface, oil seeped through to the surface. Fortunately, the majority of the oil and gas was trapped in porous rock, which was overlaid by impermeable rock, thus forming gas and oil reservoirs. The oil remains here until its presence is detected by seismic surveys and recovered by drilling through the impermeable rock.

1.3 The uses of bitumen

The vast majority of bitumen is used by the construction industry, as a constituent of products used in paving and roofing. Excellent waterproofing characteristics and thermoplastic behaviour make it ideal for a wide range of applications. At elevated temperatures (typically between 100 and 200°C) it acts like a viscous liquid, and can be mixed with other components and manipulated and formed as required. Once cooled, it is an inert solid that is durable and hydrophobic (repels water).

Various terms are used to describe conventional bitumen such as straight run, paving grade and penetration grade (or ‘pen grade’). When people use these terms they normally mean grades of bitumen that can be produced at a conventional refinery in a relatively simple way. The refinery processes used to manufacture bitumen and the effect these processes have on the final characteristics of the product are described in detail in Chapter 2.

The vast majority of bitumen used in asphalt for road construction is conventional bitumen; that is why it is often known as paving grade. The term ‘pen grade’ is short for penetration grade, and reflects the fact that this type of product is often classified (in Europe and parts of Asia) using the penetration test. The term straight run refers to the fact that this type of bitumen is often produced direct from the vacuum distillation process, without any further modification.

Current estimates put the world use of bitumen at approximately 102 million tonnes per year (Eurobitume and the Asphalt Institute, 2011), and about
85% of all the bitumen produced is used in asphalt for the construction of roads and other paved areas. Typically, asphalt will contain approximately 5% by mass of bitumen, with the remaining 95% consisting of a mixture of mineral aggregates and much finer materials such as limestone filler (see Chapters 10, 11 and 12 for more information on asphalt mixtures).

A note on terminology is worth making at this point – ‘asphalt’ is a generic term used to describe a range of road surfacing products containing primarily bitumen and mineral aggregates. A few alternative terms exist including hot mix asphalt (HMA) and asphalt concrete (AC). In this book the term ‘asphalt’ will be used throughout. Asphalt is often referred to incorrectly in the media and in common parlance as tarmac (short for tarmacadam). Tarmacadam is a road surfacing product using coal tar as a binder and has not been used in road construction for over 30 years.

A further 10% of global bitumen production is used in roofing applications, and the remaining 5% is used mainly for sealing and insulating purposes in a variety of building materials, such as pipe coatings, carpet backing, joint sealants and paint. Chapter 24 contains further information on these types of uses.

The extremely wide range of uses for bitumen is demonstrated by the number of registered uses in Europe under the requirements of the Registration, Evaluation, Authorisation and Restriction of Chemicals (REACH) regulations, which require all chemical substances and associated uses to be registered. Table 1.2 provides an overview [Eurobitume, 2013].

Bitumen is available in a variety of grades. Specifications are used across the world to define these grades to meet the needs of the applications, climate, loading conditions and end use. They are usually based on a series of standard test methods that define the properties of each grade such as hardness, viscosity, solubility and durability. Further information about testing and specification for bitumens can be found in Chapters 5, 6 and 7 of this book.

Bitumens are also used to manufacture mixtures or preparations. In these products, bitumen is often the principal component, but they can contain significant proportions of other materials to meet end use requirements. These mixtures are chemically classified as bitumen preparations.

The most commonly used are as follows [Eurobitume and the Asphalt Institute, 2011]:

- **Cut-back bitumen and fluxed bitumen**. Cut-back and fluxed bitumen products are preparations in which the viscosity of the bitumen has been reduced by the addition of a solvent, normally derived from petroleum. Typically the solvents used are white spirit, kerosene and
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<td>Mulches</td>
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<td>Mulching paper</td>
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<td>Joint filler compounds</td>
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<td>Membrane linings, waterproofing</td>
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<td>Papers</td>
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<td>Insulating felts</td>
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<td>Junction box compound</td>
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<td>Buffing compounds</td>
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<td>Well drilling fluid</td>
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<td>Armoured bituminised fabrics</td>
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<td>Acid-proof enamels, mastics, varnishes</td>
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<td>Baking and heat-resistant enamels</td>
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Modified bitumen. Polymer modified bitumens (PMBs) are designed to change the performance properties of straight run bitumen. Properties such as elasticity, adhesive or cohesive strength can be modified by the use of one or more chemical agents. These agents may be polymers, waxes, crumb rubber, sulfur and acids, among other materials. Modified bitumens are now widely used in a variety of applications. More detailed information about PMB is available in Chapter 8.
Bitumen emulsions. Bitumen emulsions are products in which droplets of bitumen or bitumen preparation are dispersed in an aqueous medium. An emulsifier is used to stabilise the mixture. Bitumen emulsions permit the handling, transport and application of bitumen at lower temperatures, and are used mainly in road surfacing applications. Further information on bitumen emulsions and how they are used can be found in Chapters 9 and 21.

1.4 Health, safety and environment

1.4.1 Safety

In service (once applied as an ingredient in a construction product, waterproofing solution, etc.) bitumen is an inert material and represents no hazard to human health. During its storage, transportation and application it is heated to high temperatures, so hazards do exist for those working with the product, although these can be managed by taking appropriate measures to reduce the risk of exposure. Burns remain an immediate hazard, and suitable procedures and safety measures must be used when handling hot bitumen. Hardware such as tanks, pipes and delivery tankers must be specially designed and approved for use. A wealth of industry guidance is available on how to handle hot bitumen. Industry associations such as Eurobitume and the Asphalt Institute are particularly active in this area, and provide user guides that build on many years of improving practice (Eurobitume, 2013; Asphalt Institute, 2013).

Bitumen is not classified as a hazardous substance. However, information on known health, safety and environmental hazards and appropriate measures to reduce any identified risks are captured on safety data sheets (commonly known as SDSs or MSDSs – material safety data sheets). Different country regulators and authorities require certain information to be available, so the format of these can vary, but material safety data sheets are provided to the customer by each bitumen supplier.

1.4.2 Health

In 2013, the International Agency for Research on Cancer (IARC) published a monograph entitled Bitumens and Bitumen Emissions, and Some N- and S-Heterocyclic Polycyclic Aromatic Hydrocarbons (IARC, 2013). The report represents a scientific evaluation of the hazards associated with occupational exposure to bitumen based on available evidence. More details about this report are available in Chapter 3, but the final evaluation is reproduced below:

- Occupational exposures to oxidized bitumens and their emissions during roofing are probably carcinogenic to humans (Group 2A).
- Occupational exposures to hard bitumens and their emissions during mastic-asphalt work are possibly carcinogenic to humans (Group 2B).
Occupational exposures to straight run bitumens and their emissions during road paving are possibly carcinogenic to humans (Group 2B).

The report provides details of the studies reviewed during the official scientific evaluation carried out by the monograph team. It details the evidence used, and explains the reasoning behind the final evaluations for occupational exposure to ‘bitumen and bitumen emissions during roofing, mastic, and paving applications’.

1.4.3 Environment
Many studies are now available concerning the environmental impact and the carbon footprints of oil products, including bitumen. The most widely referenced life cycle inventory report for bitumen was produced by Eurobitume (Eurobitume, 2012). This report was compiled by experts in accordance with the relevant international standards. It was peer reviewed by an independent expert in the field and provides detail on emissions and resource use of producing bitumen at a typical European refinery.

Many other studies have been commissioned and are publicly available concerning the life cycle and carbon footprint of asphalt (EAPA, 2013).

Further information on the handling, safety and environmental aspects of bitumen can be found in Chapter 3.

1.5 Other binders
1.5.1 Lake asphalt
This is the most extensively used and best known form of ‘natural’ asphalt binder. It is found in well-defined surface deposits, the most important of which is located in Trinidad. It is generally believed that this deposit was discovered in 1595 by Sir Walter Raleigh.

There are several small deposits of asphalt on the island of Trinidad, but it is the lake in the southern part of the island that constitutes one of the largest deposits in the world. The lake occupies an area of approximately 35 ha, and is estimated to be some 90 m deep, containing well in excess of 10 million tonnes of material. The surface of the lake is such that it can support the weight of the crawler tractors and dumper trucks that transport excavated material from the surface of the lake to railway trucks that run along the edge of the lake.

Excavated material is refined by heating the material to 160°C, vaporising the water. The molten material is passed through fine screens to remove the coarse, foreign and vegetable matter. This residue is usually termed Trinidad épuré or refined TLA (‘Trinidad lake asphalt’), and, typically, has the following composition (in percentage by weight): binder 54%, mineral matter 36%,
organic matter 10%. This épuré is too hard to be used in asphalts, but is sometimes used as an additive in mastic asphalt production.

1.5.2 Rock asphalt
Natural or rock asphalt (i.e. bitumen-impregnated rock) has been used since the early seventeenth century. Its main uses were for waterproofing, caulkung ships and the protection of wood against rot and vermin. Today, rock asphalt enjoys only minimal use. Applications include waterproofing and mastic asphalts where ageing characteristics are particularly important.

Rock asphalt is extracted from mines or quarries, depending on the type of deposit. It occurs when bitumen, formed by the same concentration processes as occur during the refining of oil, becomes trapped in impervious rock formations.

Many deposits have been successfully mined, processed and used in Europe (Italy, France and Switzerland) and in the USA (Utah, Kentucky).

1.5.3 Buton asphalt
Buton asphalt (asbuton) is a source of natural rock asphalt located in Indonesia. It has been used since 1926 (Kramer, 1989), and is located in South Sulawesi province. The deposit of asbuton is considered large, estimated at more than half a billion tonnes, and has been made available in varying forms.

Asbuton originates from crude oil in the depths of Earth layers in Buton Island that migrated upwards along deep-seated faults and deposited heavier residues in the upper beds after lighter fractions evaporated (Wallace, 1989). Asbuton consists of a mixture of natural asphalt and minerals, where the residues have fused and entered into the pores of the minerals. As a consequence, natural asphalt from asbuton cannot effectively be used in asphalt mixtures (Affandi, 2012).

1.5.4 Gilsonite
The state of Utah in the Midwest of the USA holds a sizeable deposit of natural asphalt. Discovered in vertical deposits in the 1860s, it was first exploited by Samuel H. Gilson in 1880 as a waterproofing agent for timber. The material is very hard, and is sometimes blended with petroleum bitumen. It is sometimes used in bridge and roof waterproofing materials as a means of altering the stiffness of mastic asphalt.

1.5.5 Tar
Tar is a generic word for the liquid obtained when natural organic materials such as coal or wood are carbonised or destructively distilled in the absence
of air. It is customary to prefix the word ‘tar’ with the name of the material from which it is derived. Thus, the products of this initial carbonisation process are referred to as crude coal tar, crude wood tar and so on. Two types of crude coal tar are produced as a by-product of the carbonisation of coal–coke oven tar and low temperature tar.

In the mid-1960s, over 2 million tonnes of crude coal tar was produced per annum, of which around half was manufactured as a by-product of the operation of carbonisation ovens that were used to produce town gas. However, the introduction of North Sea gas in the late 1960s resulted in a rapid reduction of tar production from this source, and by 1975 it had disappeared completely.

1.6 Terminology
Some key terminology introduced in this chapter will be used consistently throughout the book.

**Bitumen**: Hydrocarbon product produced from the refining of crude oil.

**Asphalt**: Road surfacing material consisting of bitumen, mineral aggregates and fillers and may contain other additives, and includes what are described in some areas as hot mix asphalt (HMA).

**Bitumen preparation**: A mixture of bitumen and another ingredient (oil, additive, etc.).

**Bitumen emulsion**: A preparation in which droplets of bitumen or other preparation are dispersed in an aqueous medium.

**Polymer modified bitumen (PMB)**: Widely used preparation in which the primary components are bitumen and polymer(s).

**Straight run bitumen**: Bitumen produced primarily by distillation processes.

**Hard bitumen**: A subset of straight run bitumens that have low penetration values commonly specified using a softening point range.

**Oxidised bitumen**: Bitumen produced by passing air through hot bitumen under controlled temperature and pressure conditions, thus producing a product with specific characteristics.

**References**


Chapter 2

Manufacture and storage of bitumens

Derek Petrauskas
Technical Lead,
Bitumen Manufacturing,
Shell Global Solutions
International. The Netherlands

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2.1 The manufacture of bitumen

Crude oil is the term for ‘unprocessed’ oil, the viscous liquid that comes out of the ground. It is also known as petroleum. Crude oil is a fossil fuel, meaning that it was made naturally from decaying plants and animals living in ancient seas millions of years ago – most places you can find crude oil were once sea beds. Crude oils vary in colour, from clear to jet black, and in viscosity, from water to almost solid.

Crude oils are such a useful starting point for so many different substances because they contain hydrocarbons. Hydrocarbons are molecules that contain hydrogen and carbon, and come in various lengths and structures, from straight chains to branching chains to rings.

Crude oil is a complex mixture of many hydrocarbons, all differing in molecular weight and, consequently, in boiling range. Before it can be used, crude oil has to be separated into component streams, some of which are further chemically or physically changed. These are often then reblended to final products. Bitumen is one of the components produced from this process.

2.2 Fractional distillation of crude oil

The first process in the refining of crude oil is fractional distillation, which physically separates the crude oil into streams varying in boiling point. This is carried out in tall steel towers known as fractionating or distillation columns. The inside of the column is divided at intervals by horizontal steel trays punctured with holes to allow vapour to rise up the column. Over these holes are small domes called bubble caps that deflect the vapour downwards so that it bubbles through liquid condensed on the tray. This improves the efficiency of separation and also has the advantage of reducing the height of the column.
On entering the distillation plant, the crude oil is heated in a furnace to temperatures between 350 and 380°C before being passed into the lower part of the column operating at a pressure slightly above atmospheric. The material entering the column is a mixture of liquid and vapour: the liquid comprises the higher boiling point fractions of the crude oil, and the vapour consists of the lower boiling point fractions. The vapours rise up the column through the holes in the trays, losing heat as they rise. When each fraction reaches the tray where the temperature is just below its own boiling point, it condenses and changes back into a liquid. As the fractions condense on the trays with the required qualities, they are continuously drawn off by means of pipes.

The lightest fractions of the crude oil remain as vapour and are taken from the top of the distillation column; heavier fractions are taken off the column as side-streams; the heaviest fractions remain as liquids, which, therefore, leave at the base of the column. The lightest fractions produced by the crude distillation process include propane and butane, both of which are gases under atmospheric conditions. Moving down the column, naphtha – a slightly heavier material – is produced. Naphtha is used as a feedstock for gasoline production and the chemical industry. Further down the column, kerosene is produced. Kerosene is used primarily for aviation fuel and, to a lesser extent, domestic fuel. Heavier again is gas oil, which is used as a fuel for diesel engines and central heating. The heaviest fraction taken from the crude oil distillation process is known as the long residue, a complex mixture of high molecular weight hydrocarbons, which requires further processing before it can be used as a feedstock for the manufacture of bitumen. Fractional distillation that is carried out under atmospheric pressure is known as atmospheric distillation (Figure 2.1).

The long residue is further distilled at reduced pressure in a vacuum distillation column (Figure 2.2). This is carried out under a vacuum of 10–100 mmHg at a temperature of between 350 and 425°C, to produce gas oil and distillates fractions, plus a bottom residue known as short residue. If this second distillation were carried out by simply increasing the temperature, cracking or thermal decomposition of the long residue would occur, hence the need to lower the pressure. The short residue is the feedstock used in the manufacture of over 20 different grades of bitumen. The viscosity and yield of short residue is a function of both the origin of the crude oil and the temperature and pressure in the vacuum column during processing, and these vary significantly from crude oil to crude oil. The blend of crude oil processed as well as the conditions in the column are adjusted to produce a short residue with a penetration in the decimillimetre range of 35–300 dmm. Figure 2.3 shows the relationship of the distillation process with other fundamental refining processes such as reforming and cracking for the manufacture of saleable
Figure 2.1  Atmospheric distillation (MPA, middle pump-around; TPA, top pump-around)
Figure 2.2 Vacuum distillation (CW, cooling water; MPA, middle pump-around; TPA, top pump-around)
Figure 2.3 Schematic representation of the crude oil distillation process. [Ex, from; nnf, normally no flow; SR, short residue; TC, temperature control; OM, oxygen meter]
products, for example aviation, gasoline and diesel fuels, lube oils, heavy fuel oil and chemical feedstocks.

2.3 Air blowing of short residues

Often by the correct selection of the crude oil feed and operating conditions, bitumen complying with a particular specification can be manufactured directly from the distillation process. If required, the physical properties of the short residue can be further modified by ‘air blowing’. This is an oxidation process that involves passing air through the short residue, either on a batch or a continuous basis, with the short residue raised to a temperature between 240 and 320°C (Figure 2.4). The main effect of blowing is that it converts some of the relatively low molecular weight ‘maltenes’ into relatively higher molecular weight ‘asphaltenes’. The result is a reduction in the penetration of the bitumen with a comparatively greater increase in the softening point, which has the effect of improving the lower temperature susceptibility of the air-blown bitumen.

2.3.1 The continuous blowing process

After preheating, the short residue is introduced into the blowing column just below the normal liquid level. Air is blown through the bitumen by means of an air distributor located at the bottom of the column. The air acts not only as a reactant but also serves to agitate and mix the bitumen, thereby increasing both the surface area and the rate of reaction. The bitumen absorbs oxygen as air ascends through the material. Steam and water are sprayed into the vapour space above the bitumen level, the former to suppress foaming and dilute the oxygen content of waste gases and the latter to cool the vapours in order to prevent after-burning and the resulting formation of coke.

The blown product passes through heat exchangers to achieve the desired ‘rundown’ temperatures (the rundown temperature is the temperature at which bitumen can be taken from the tank) and to provide an economical means of preheating the original short residue feed, before pumping the product to storage. The final penetration and softening point of the blown bitumen are affected by many factors that include the viscosity of the feedstock, the temperature in the blowing column, the residence time in the blowing column, the origin of the crude oil used to manufacture the feedstock and the ratio of air to bitumen fed into the process.

Figure 2.5 shows blowing ‘curves’ for a bitumen feedstock and how these processes can be applied to enable the bitumen to meet the softening point and penetration specification ranges indicated by the boxes. In the blowing processes, the softening point increases and the penetration reduces. However, in the distillation process, the temperature susceptibility for
Figure 2.4 Simplified diagram of a bitumen blowing unit (Ex, from; n nf, normally no flow; SR, short residue; TC, temperature control; OM, oxygen meter)
penetration index) of the material is largely unchanged. Thus, the distillation is a relatively straight line whereas the curves for the blown bitumen flatten substantially as the softening point of the bitumen increases. This demonstrates that the temperature susceptibility of the material is substantially reduced (i.e. the penetration index is increased).

2.3.1.1 Air-rectified or semi-blown bitumens
Many crude oils produce bitumen that requires a limited amount of, ideally, no air blowing in order to generate penetration grade bitumen suitable for road construction. This process is termed semi-blowing or air-rectification. Used judiciously, semi-blowing can be used to reduce the temperature susceptibility of the bitumen (i.e. to increase its penetration index).
2.3.1.2 Fully blown bitumens

Fully blown or oxidised bitumens are often produced by more extensive blowing of a blend of a short residue with a relatively low viscosity flux. The position of the blowing curve in Figure 2.5 is primarily dependent on the viscosity and chemical nature of the feed. The softer, or lower viscosity, the feed, the higher the curve. The amount of blowing that is required depends on the temperature in the column and the air-to-feed ratio. Thus, by selection of a suitable bitumen feedstock, control of the viscosity of the feed and the conditions in the column, a range of blown grades of bitumen can be manufactured.

2.3.1.3 Chemistry of the blowing process

The bitumen blowing process can be described as a conversion process in which oxidation, dehydrogenation and polymerisation take place. Normally, it is effected with the oxygen present in air, and it is this type of process that is considered here.

By subjecting the materials entering and leaving the reaction vessel to a combination of analytical methods, it is possible to draw up an oxygen balance. It has been found that all the oxygen taken up by the bitumen can be accounted for by the formation of hydroxyl, carbonyl, acid and ester groups; no ether oxygen has been detected. The main side products are carbon dioxide, water and some light hydrocarbons. Besides carbon–oxygen bonds, carbon–carbon bonds are also formed.

Of the functional groups mentioned, the esters are particularly important because they serve as a link up of two different molecules and thus contribute to the formation of material of higher molecular weight. This mechanism results (together with the direct formation of carbon–carbon bonds) in an increase in the asphaltene content. In other words, the formation of esters and the direct formation of carbon–carbon bonds are desired reactions; all others are less desirable or even undesirable reactions.

As a typical trend, the amount of oxygen used for condensation reactions decreases from 60 to 20%, whereas the amount of oxygen used for side reactions increases from 40 to 80% with an increase in temperature from 150 to 350°C. In other words, increasing temperature decreases the contribution of the desirable reactions at the cost of the undesirable chemical reactions.

As the formation of asphaltenes – the main aim of the blowing process – is a function of the number of bonds obtained, the number of bonds per mole of reacting oxygen is of paramount importance. The amount of oxygen required for one ester bond is much higher than that required for a carbon–carbon bond, and the amount of ester bonds formed sharply decreases with temperature, while the amount of carbon–carbon bonds increases with
temperature. Consequently, at a certain temperature, an optimum in bond formation may be expected. It has been found that this optimum is reached at a blowing temperature of 250°C.

However, at this preferred temperature the transfer of oxygen from the gaseous to the liquid phase is often not very satisfactory because, due to the relatively high viscosity of the liquid bitumen, the distribution of air bubbles may be poor. An increase in the blowing temperature results in a reduced viscosity and in an increased gas–liquid interface, thus promoting the oxygen uptake (efficiency) by the bitumen. However, this is at the cost of a shift towards a regime where the total number of bonds per mole of reacted oxygen is lower. On balance, the total number of bonds formed per mole of oxygen intake is more favourable at higher temperatures. A shift in the optimum occurs, which explains the trend towards higher blowing temperatures (275–285°C) frequently encountered in practice.

2.4 Delivery, storage and handling temperatures of bitumen

As most bitumen grades are solid at ambient temperature, in order to enable them to be moved through the distribution system as a liquid, they must be heated to temperatures in the range of 140°C to over 200°C, depending on the grade; higher temperatures up to 230°C are applied with an inerted atmosphere for highly oxidised grades. When handled properly, bitumen can be reheated or maintained at elevated temperatures for a considerable time without adversely affecting its properties. However, mistreatment of bitumen by overheating or by permitting the material to be exposed to conditions that promote oxidation can adversely affect the properties of the bitumen and may influence the long term performance of mixtures that contain bitumen. The degree of hardening (or, under certain circumstances, softening) that is produced as a result of mishandling is a function of a number of parameters such as temperature, the presence of air, the surface-to-volume ratio of the bitumen, the method of heating and the duration of exposure to these conditions.

2.4.1 Safe delivery of bitumen

Over 50% of bitumen related accidents and incidents that result in lost working days occur during the delivery of bitumen. In Europe, Eurobitume has produced a detailed code of practice (Eurobitume and the Asphalt Institute, 2011) designed to assist in reducing the frequency of incidents and accidents by raising awareness of their causes. Personnel involved in the delivery and receipt of bitumen are strongly advised to refer to this document.

2.4.2 Bitumen tanks

All bitumens should be stored in tanks specifically designed for the purpose (Eurobitume and the Asphalt Institute, 2011). In order to minimise the
possible hardening of the bitumen during storage, certain aspects of the design of the tank should be considered. In order to minimise the risk of overheating the bitumen, the tank should be fitted with accurate temperature sensors and gauges. These should be positioned in the region of the heaters and preferably be removable to facilitate regular cleaning and maintenance. Oxidation and the loss of volatile fractions from bitumen are both related to the exposed surface-to-volume ratio of the storage tank, which, for a cylindrical vessel, equals the reciprocal height of the filled part of the tank and is given by

$$\frac{\text{surface area}}{\text{volume}} = \frac{\pi r^2}{\pi r^2 h} = \frac{1}{h}$$

where \( h \) is the height of the bitumen and \( r \) is the radius of the tank.

Thus, the dimensions of the bulk storage tank should be such that the surface-to-volume ratio is minimised. Accordingly, vertical storage tanks with a large height-to-radius ratio are preferable to horizontal tanks.

It is common practice for bitumen in bulk storage tanks at mixing plants to be recirculated around a ring main in order to heat the pipework that carries the bitumen to the processing point. Return lines in a recirculation system should re-enter the storage tank below the bitumen surface to prevent hot bitumen cascading through the air. Often, the bitumen is returned to the bulk storage tank through a pipe fitted into the upper part of the tank, flush with the side or roof or protruding just into the air space at the top of the tank. If the bitumen enters the tank above the bitumen, all the factors that promote oxidation are present

- high temperature
- access to oxygen
- high exposed surface-to-volume ratio.

Fortunately, the residence time of bitumen in the tank is usually sufficiently low for any hardening to be insignificant. However, if material is stored for a prolonged period, recirculation should be used only intermittently, and the bitumen should be tested before use to ensure its continued suitability for the proposed application. A recommended layout for a bitumen storage tank is shown in Figure 2.6.

Bitumen storage tanks should be fitted with automatic level indicators together with low and high level alarms to avoid having to dip manually for a reading. Such an approach avoids exposing the hot heater tubes to a potentially combustible or explosive atmosphere should the bitumen level fall below that of the heater tubes. Automatic level control also ensures that the tank is not overfilled. Regardless of whether a high level alarm is fitted, a
maximum safe filled level for the tank should be predetermined, taking into account the effects of thermal expansion of the bitumen in the tank.

Before ordering additional bitumen, it is essential to check that the ullage in the tank is capable of taking the delivery without exceeding the maximum safe working level.

Every tank should be clearly labelled with the grade of bitumen it contains. When the grade of bitumen in a tank is changed, it is important to ensure that the tank is empty and relabelled before the new grade is delivered.

2.4.3 Bitumen storage and pumping temperatures

Bitumen should always be stored and handled at the lowest temperature possible, consistent with efficient use. As a guide, working temperatures for specific operations are given in the Energy Institute’s bitumen safety code (Energy Institute, 2005). These temperatures have been calculated on the basis of viscosity measurements, and are supported by operational experience. For normal operations (i.e. the blending and transferring of liquid bitumen), temperatures of 10–50°C above the minimum pumping
Manufacture and storage of bitumens

temperature are recommended, but the maximum safe handling temperature of 230°C must never be exceeded.

The period during which bitumen resides in a storage tank at elevated temperatures and is recirculated should be minimised, to prevent hardening of the bitumen. If bitumen must be stored for an extended period, say for a period exceeding 1 week without the addition of fresh material, the temperature should be reduced to approximately 20–25°C above the softening point of the bitumen and, if possible, recirculation stopped.

When bitumen is being reheated in bulk storage, care must be taken to heat the bitumen intermittently over an extended period to prevent localised overheating of the product around the heating pipes or coils. This is particularly important where direct flame tube heating is used because surface temperatures in excess of 300°C may be reached. In such installations, the amount of heat that is applied should be limited, sufficient only to raise the temperature of the product to just above its softening point. This will allow the material to soften, after which further heat can be applied to raise the temperature of the product to the required working value. This technique is beneficial because when the bitumen is a fluid, albeit a viscous fluid, convection currents dissipate the heat throughout the material, and localised overheating is thus less of a problem. Circulation of the tank contents should begin as soon as the product is sufficiently fluid, thereby further reducing the likelihood of local overheating. With hot oil, steam or electric heaters that are designed properly, reheating from cold should not cause these problems.

References
Chapter 3

Handling, health, safety and environmental aspects of bitumens

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Bitumen has a long history of being used safely in a wide range of applications. Although it is primarily used in road construction, it is also employed in roofing felts, reservoir linings and as an internal lining for potable water pipes as well as having a myriad of other uses (many of which are given in Table 1.2). Bitumen presents a low order of potential hazard, and provided that good handling practices are observed, the risks of the hazard ever being released are very small. These are described in detail in part of the Model Code of Safe Practice (Energy Institute, 2005), and are also described in the material safety data sheets provided by each supplier. A substantial amount of health, safety and environmental data on bitumen and its derivatives are detailed in CONCAWE Product Dossier No. 92/104 (CONCAWE, 1992). In addition, with the implementation of REACH (Registration, Evaluation, Authorisation and Restriction of Chemicals – the base legislation to which there have been numerous amendments and corrigenda; EC, 2007) a large amount of work around the manufacturing, safe handling and safe delivery of bitumen, focusing on hazard and risk has been documented by Eurobitume (Eurobitume, 2011a) – Eurobitume is the European Bitumen Association, and comprises a number of bitumen companies and bitumen associations in Europe. Notwithstanding the safe use indicated by these various publications, bitumens are generally applied at elevated temperatures, and this brings with it a number of hazards that are considered below.

3.1 Handling and safety

3.1.1 Personal protective equipment as related to occupational exposure limits

The principal hazard from handling hot bitumen is thermal burns resulting from contact with the product. Thus, it is essential to wear clothing that
provides adequate protection (PPE, personal protective equipment), including:

- helmet and neck apron to provide head protection
- visor to protect the face (goggles only protect eyes)
- heat-resistant gloves (with cuffs worn inside coverall sleeves)
- safety boots
- coveralls (with coverall legs worn over boots).

Table 3.1 provides an overview of the occupational exposure limits (OELs) for various countries. Further discussion on these, as evaluated in the International Agency for Research on Cancer (IARC) monograph *Bitumens and Bitumen Emissions, and Some N- and S-Heterocyclic Polycyclic Aromatic Hydrocarbons* (IARC, 2013), is given in section 3.2.1.

Exposure to bitumen fumes can result in irritation to the eyes, nose and respiratory tract, headaches and nausea. The symptoms are usually mild and temporary. Removal of the affected personnel from the source results in rapid recovery. Even though the irritation is usually mild, exposure to bitumen fumes should be minimised and, where there is any doubt, tests (e.g. Drægar tube analysis or personal exposure monitoring) should be undertaken to determine the concentration of bitumen fumes or hydrogen sulfide in the working atmosphere.

Persons affected by the inhalation of bitumen fumes should be removed to fresh air as soon as possible. If the symptoms are severe or if the symptoms persist, medical help should be sought without delay.

### 3.1.2 Personal hygiene

Garments soiled with bitumen should either be replaced or dry cleaned in order to avoid permeation of bitumen into the underclothing. Soiled rags or tools should not be placed in the pockets of overalls, as contamination of the lining of the pocket will result.

Personnel handling bitumen and asphalts should be provided with and use barrier creams to protect exposed skin, particularly hands and fingers. Skin should be thoroughly washed after any contamination, and always before going to the toilet, eating or drinking.

The application of barrier creams prior to handling bitumen assists in subsequent cleaning, should accidental contact occur. However, barrier creams are no substitute for gloves or other impermeable clothing. Consequently, they should not be used as the sole form of protection. Solvents such as petrol, diesel oil, white spirit or similar should not be used for removing bitumen from the skin, as they may spread the contamination as well as being skin sensitisers or carcinogenic or both. An approved skin cleanser together with warm water should be used.
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</table>
### Fire prevention and fire-fighting

The adoption of safe handling procedures will substantially reduce the risk of fire. However, if a fire occurs, it is essential that personnel are properly trained and well equipped to extinguish the fire, thereby ensuring that the risk of injury to personnel and damage to plant is minimised. Detailed advice on fire prevention and fire-fighting is given in the Model Code of Safe Practice (Energy Institute, 2005) and in supplier material safety data sheets.

Small bitumen fires can be extinguished using dry chemical powder, foam, vaporising liquid or inert gas extinguishers, fog nozzle spray hoses and

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**Table 3.1 Continued**

<table>
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<tr>
<th>Country/Advisory body</th>
<th>OEL long term (8 h/day): mg/m³</th>
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<sup>a</sup> Sourced from ACGIH limit  
<sup>b</sup> Quebec uses an 8 hour limit of 5 mg/m³  
<sup>c</sup> Updated with and follows ACGIH limit from 2013  
<sup>d</sup> Ministry of Labour advises use of the ACGIH/NIOSH thresholds for OEL  
<sup>e</sup> Former threshold of 10 mg/kg removed by the employer’s mutual insurance association (Berufsgenossenschaft)  
<sup>f</sup> Units are ppm  
<sup>g</sup> More than one limit listed for UAE; most stringent one recorded in this table  
<sup>h</sup> Advisory organisations based in USA. Limits stated are not regulated unless formally adopted by a country into their OEL legislation
steam lances. Direct water jets must not be used because frothing may occur, which tends to spread the hot bitumen and, therefore, the fire.

Injection of steam or a ‘fog’ of water into the vapour space can extinguish internal tank fires where the roof of the tank is largely intact. However, only trained operatives should use this method, as the water vaporises instantly on contact with the hot bitumen. This initiates foaming, which may result in the tank overflowing, creating an additional hazard. Alternatively, foam extinguishers may be used. The foam ensures that the water is well dispersed, thereby reducing the risk of froth-over. The disadvantage of using this type of extinguisher is that the foam breaks down rapidly when applied to hot bitumen.

Portable extinguishers containing either aqueous film-forming foam or dry chemical powder are suitable for dealing with small bitumen fires, at least initially. In bitumen handling areas, these should be placed at strategic, permanent and conspicuous locations. The type and location of equipment to be used if initial attempts fail should be discussed with the local fire brigade before installation.

3.1.4 Sampling

Sampling of hot bitumen is particularly hazardous because of the risk of heat burns from spills and splashes of the material. It is therefore essential that appropriate protective clothing is worn. The area should be well lit, and safe access to and egress from the sample point should be provided. Gantry access should be provided where samples are required from the tanks of vehicles, as climbing on top of vehicles should be avoided.

3.1.4.1 Dip sampling

In this process, a sample of bitumen is obtained by dipping a weighted can or ‘thief’ on the end of a rope or rod through the access lid into bitumen stored in a bulk tank. The sample is then transferred to a suitable permanent container. The method is simple, but is only appropriate for small samples. Dip sampling from cut-back tanks should be avoided because of the presence of flammable atmospheres in tank vapour spaces. Additionally, where possible, dip sampling should be avoided altogether, as it is also possible to be exposed to hydrogen sulfide when the access lid is opened.

3.1.4.2 Sample valves

Properly designed sample valves are very useful for sampling from pipelines or from tanks. Their design should ensure that they are kept hot by the product in the pipeline or tank in order to avoid blockage when in the closed position, or have a means of flushing them clean.

Sample valves should preferably be the screw-driven plunger type. When closed, the plunger of this type of valve extends into the fresh product.
Thus, when the valve is opened, a representative sample of product is obtained without ‘fore-runnings’. With ball and plug type valves, fore-runnings have to be collected and disposed of before a representative sample can be obtained. Designs for bitumen sample valves are described in detail in the European Norm EN 58:2012 (BSI, 2012).

3.2 Health aspects of bitumen

3.2.1 Hazards associated with bitumen

Some of the key hazards are given below. A more comprehensive review can be found in the joint Eurobitume/Asphalt Institute publication The Bitumen Industry – A Global Perspective (Eurobitume, 2011b).

3.2.1.1 Elevated temperatures

The main hazard associated with bitumen is that the product is held at elevated temperatures during transportation, storage and processing. Thus, it is critical that appropriate PPE is worn and any skin contact with hot bitumen is avoided. Some guidance is given below, but detailed advice is available in a number of publications (CONCAWE, 1992; Energy Institute, 2005; Eurobitume, 2011a) and the relevant supplier material safety data sheets.

3.2.1.2 Vapour emissions

Bitumens are complex mixtures of hydrocarbons that do not have well defined boiling points because their components boil over a wide temperature range. Visible emissions or fumes normally start to develop at approximately 150°C. The amount of fumes generated doubles for each 10–12°C increase in temperature. Fumes are mainly composed of hydrocarbons (Brandt and De Groot, 1996) and small quantities of hydrogen sulfide. Bitumen fumes also contain small quantities of polycyclic aromatic compounds (PACs) and more specifically polycyclic aromatic hydrocarbons (PAHs). These chemicals consist of a number of benzene rings that are grouped together. Some of these with three to seven (usually four to six) fused rings are known to cause or are suspected of causing cancer in humans. However, the concentrations of these carcinogens in bitumen are extremely low (CONCAWE, 1992). The most common PAHs included in this category by the US Environmental Protection Agency (2008) by name and CAS number, are

- benzo[\(a\)]anthracene, 56-55-3
- benzo[\(a\)]phenanthrene (chrysene), 218-01-9
- benzo[\(a\)]pyrene, 50-32-8
- benzo[\(b\)]fluoranthene, 205-99-2
- benzo[\(j\)]fluoranthene, 205-82-3
- benzo[\(k\)]fluoranthene, 207-08-9
A public statement on PAHs by the US Agency for Toxic Substances and Disease Registry states that

One of the most common ways PAHs can enter the body is through breathing contaminated air. PAHs get into your lungs when you breathe them. If you live near a hazardous waste site where PAHs are disposed, you are likely to breathe PAHs. If you eat or drink food and water contaminated with PAHs, you could be exposed. Exposure to PAHs can also occur if your skin contacts PAH-contaminated soil or products like heavy oils, coal tar, roofing tar, or creosote. Creosote is an oily liquid found in coal tar and is used to preserve wood. Once in your body, PAHs can spread and target fat tissues. Target organs include the kidneys and liver. However, PAHs will leave your body through urine and faeces in a matter of days.

Therefore, it is exposure to PAHs over an extended period that causes concerns to health, and measures such as reducing temperature and avoiding enclosed spaces should be practised as a matter of good product stewardship.

In 2013, a comprehensive review was carried out in IARC Monograph 103 that concluded that there was an increased probability of carcinogenicity when exposed to fumes from fully oxidised bitumen versus straight run paving and hard bitumens (IARC, 2013). The IARC Monograph found that there was limited evidence in humans for the carcinogenicity of occupational exposures to bitumens and bitumen emissions during roofing and mastic asphalt work and that there was sufficient evidence in experimental animals for carcinogenicity of fume condensates generated from oxidised bitumens, and on this basis they classified bitumen into three groups as follows.
Occupational exposures to oxidised bitumens and their emissions during roofing are probably carcinogenic to humans (Group 2A).

Occupational exposures to hard bitumens and their emissions during mastic-asphalt work are possibly carcinogenic to humans (Group 2B).

Occupational exposures to straight run bitumens and their emissions during road paving are possibly carcinogenic to humans (Group 2B).

Hence, in contrast to the previous monograph published in 1985 (IARC, 1985), oxidised bitumen moved from a classification of possibly carcinogenic (Group 2B) to probably carcinogenic (Group 2A) and this is now reflected in supplier materials safety data sheets. It must be noted that IARC monographs only look at whether the hazard is present, and take no account of the risk of exposure: that is, whether the hazard will ever be released.

3.2.1.3 Hydrogen sulfide

Hydrogen sulfide is of particular concern, as it can accumulate in enclosed spaces, such as the tops of storage tanks, and exposure to this gas at concentrations of as little as 500 ppm can be fatal, so it is essential that any space where hydrogen sulfide may be present is tested and approved as being gas free before anyone enters the area (Table 3.2). In addition, with the increase in the use of sulfur as a cross-linking agent in the production of stable polymer modified bitumens, it is ever more important to actively manage hydrogen sulfide, which potentially can be generated throughout the supply chain.

3.2.1.4 Combustion

Very high temperatures are required to make bitumen burn. Certain materials, if hot enough, will ignite when exposed to air, and are sometimes described as being ‘pyrophoric’. In the case of bitumens, this auto-ignition temperature is generally around 400°C. However, despite storage and use of temperatures well below the auto-ignition temperature, fires have very occasionally occurred. Under conditions of low oxygen content, hydrogen sulfide from bitumen can react with rust (iron oxide) on the roof and walls of storage tanks, to form ‘pyrophoric iron oxide’. This material reacts readily with oxygen and can self-ignite if the oxygen content of the tank increases suddenly, which, in turn, can ignite coke deposits on the roof and walls of the tank. Coke deposits are the result of condensate from the bitumen that has been deposited on the roof and walls of a tank degrading over a period of time, forming carbonaceous material. Under conditions of high temperature and in the presence of oxygen or a sudden increase in available oxygen, an exothermic reaction, leading to the risk of fire or explosion, can occur. Accordingly, manholes in bitumen tanks should be kept closed and access to tank roofs should be restricted at all times.
3.2.1.5 Contact with water
It is essential that water does not come into contact with hot bitumen. If this does happen, the water is converted into steam. In the process, its volume increases by a factor of 1673 at atmospheric pressure, resulting in spitting, foaming and, depending on the amount of water present, possible boil-over of the hot bitumen.

3.2.1.6 Potential hazards through skin contact
Other than thermal burns, the hazards associated with skin contact of most bitumens are negligible. Studies (Bofetta, 2001) concluded that there was no direct evidence to associate bitumen with long term skin disorders in humans despite bitumens having been widely used for many years. Nevertheless, it is prudent to avoid intimate and prolonged skin contact with bitumen.

Cut-back bitumens and bitumen emulsions are handled at lower temperature, which increases the chance of skin contact. If personal hygiene is poor, regular skin contact may occur. However, studies carried out by Shell (Brandt et al., 1999; Deygout, 2011; Potter et al., 1999) have demonstrated that bitumens are unlikely to be bio-available (skin penetration and body uptake) and bitumens diluted with solvents are unlikely to present a carcinogenic risk. Nevertheless, bitumen emulsions can cause irritation to the skin and eyes, and can produce allergic responses in some individuals.

3.2.2 First aid for skin burns
The following is taken from the Eurobitume notes for the guidance of first aid and medical personnel (Eurobitume, 2011a). This advice is produced in the form of an A5-sized card, copies of which are available from Eurobitume in a number of different languages. It is intended that the card should accompany a burns victim to the hospital, to provide immediate advice on proper treatment.

3.2.2.1 First aid
Bitumen burns should be cooled for at least 15 min, first with cool water to reduce pain, then with warm water to prevent hypothermia if the burned surface is larger than the size of a hand. Burns to the eyes should be irrigated for at least 5 min.

NO ATTEMPT SHOULD BE MADE TO REMOVE THE BITUMEN AT THE WORK SITE.

3.2.2.2 Medical care
(If in doubt do not hesitate to contact a burns centre.)

Measures to remove the bitumen layer from the skin should be taken as soon as possible under the supervision of a doctor, or at a hospital. However, this
### Table 3.2 Various hydrogen sulfide OELs in force (March 2014)

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<th>Country/Advisory body</th>
<th>OEL long term (8 h/day): ppm</th>
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EU OEL; List of Indicative OEL values 12/2009

DECOS 2010/06OSH

Nye administrative normer for forurensning i arbeidsatmosfaere; version 12/2011

Ministra Pracy i Polityki Społecznej (Poland, 07/2009)

Instituto Português da Qualidade

Nariadenie Vlady Slovenskej Republiky

Límites de Exposición Profesional para Agentes Químicos en España, 2012; Ministerio de Trabajo e Inmigración, INSHT

AFS 2005:17

SuvaPro Grenzwerte am Arbeitsplatz 2009

Health and Safety Executive EH40/2005

US Department of Labor

ACGIH

NIOSH

Scientific Committee on Occupational Exposure Limits

\(^a\) North American value

\(^b\) OSHA PEL (permissible exposure limit) (General Industry): ‘Exposures shall not exceed 20 ppm (ceiling) with the following exception: if no other measurable exposure occurs during the 8-hour work shift, exposures may exceed 20 ppm, but not more than 50 ppm (peak), for a single time period up to 10 minutes.’
treatment should be carried out with caution because careless removal of the bitumen may result in the skin being damaged further, bringing with it the risk of infection and the possibility of complications.

Initially, it is not important to know whether the burn is superficial or deep. The priority should be to remove the bitumen without causing further damage.

3.2.2.3 Removal of bitumen adhering to the burned areas
Different methods can be recommended.

- The bitumen layer should be left in place and covered with thick gauze containing paraffin or a paraffin-based antibiotic cream such as Flammazine (silver sulfadiazine). Such treatment will have the effect of softening the bitumen, enabling it to be gently removed after a few days.
- Alternatively, olive oil (new bottle) should be applied and left to soak the affected areas for a few hours. Thereafter, the bitumen can be removed by rubbing gently with some gauze. Any remaining bitumen can be removed by wrapping the affected areas with gauze soaked in olive oil. The dressing should be changed every 4 h. After 24 h, any remaining bitumen can be removed, and the burn may be disinfected and treated conventionally.

3.2.2.4 Circumferential burns with tourniquet effect
When bitumen completely encircles a limb, or other body part, the cooled and hardened bitumen may cause a tourniquet effect due to oedema (swelling) in the burn. In the event of this occurring, the bitumen must be softened as soon as possible and/or split, to prevent restriction of blood flow.

3.2.2.5 Eye burns
No attempt should be made to remove the bitumen by unqualified personnel. The patient should be referred urgently to an ophthalmologist or hospital with an ophthalmology unit for diagnosis and appropriate treatment.

3.3 Bitumen and the environment

3.3.1 Life cycle assessment of bitumen
Life cycle assessment (LCA) is a tool to investigate the environmental aspects and potential impact of a product, process or activity by identifying and quantifying energy and material flows. LCA covers the entire life cycle, including extraction of the raw material, manufacturing, transport and distribution, product use, service and maintenance, and disposal (recycling, incineration or landfill). It is a complete cradle-to-grave analysis focusing on the environmental input (based on ecological effects) and resource use.
LCA can be divided into two distinct parts – life cycle inventory (LCI) and life cycle impact. Eurobitume has carried out a life cycle inventory of bitumen (Eurobitume, 2012) to generate inventory data on the production of paving grade bitumen for future LCI studies where bitumen is used. In addition there are many other bitumen, asphalt and construction material LCAs, but to be comparable they need to meet the standard of the ISO 14040 series (ISO, 2006).

### 3.3.2 Use of bitumen as a potable water lining

Bitumen and asphalt have been used for many years for applications in contact with water, such as reservoir linings, dams and dykes (Schönian, 1999). A number of studies have been carried out both in the USA and Europe to determine if components are leached out of asphalt and bitumen when in prolonged contact with water.

Shell has carried out laboratory studies of leaching on a range of bitumens and asphalts (Brandt and De Groot, 2001). These concluded that although prolonged contact with water will result in PACs being leached into water, the levels rapidly reach an equilibrium level that is well below the surface water limits that exist in a number of EU countries and more than an order of magnitude below the EU limits given in the Drinking Water Directive and listed in Table 3.3 (Council Directive 98/83/EC) (EC, 1998).

### Table 3.3 EU chemical limits for drinking water

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<thead>
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<th>Parameter</th>
<th>Parametric value</th>
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</tr>
<tr>
<td>Benzene</td>
<td>1.0 µg/l</td>
<td>Nickel</td>
<td>20 µg/l</td>
</tr>
<tr>
<td>Benzo[a]pyrene</td>
<td>0.010 µg/l</td>
<td>Nitrate</td>
<td>50 mg/l</td>
</tr>
<tr>
<td>Boron</td>
<td>1.0 mg/l</td>
<td>Nitrite</td>
<td>0.50 mg/l</td>
</tr>
<tr>
<td>Bromate</td>
<td>10 µg/l</td>
<td>Pesticides</td>
<td>0.10 µg/l</td>
</tr>
<tr>
<td>Cadmium</td>
<td>5.0 µg/l</td>
<td>Pesticides (total)</td>
<td>0.50 µg/l</td>
</tr>
<tr>
<td>Chromium</td>
<td>50 µg/l</td>
<td>PAHs</td>
<td>0.10 µg/l</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Sum of concentrations of specified compounds</td>
<td>0.10 µg/l</td>
</tr>
<tr>
<td>Copper</td>
<td>2.0 mg/l</td>
<td>Selenium</td>
<td>10 µg/l</td>
</tr>
<tr>
<td>Cyanide</td>
<td>5 µg/l</td>
<td>Tetrachloroethene and trichloroethene</td>
<td>10 µg/l</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Sum of concentrations of specified parameters</td>
<td>100 µg/l</td>
</tr>
<tr>
<td>1,2-Dichloroethane</td>
<td>3.0 µg/l</td>
<td>Trihalomethanes (total)</td>
<td>100 µg/l</td>
</tr>
<tr>
<td>Epichlorohydrin</td>
<td>0.10 µg/l</td>
<td>Vinyl chloride</td>
<td>0.50 µg/l</td>
</tr>
</tbody>
</table>
The objective of the Drinking Water Directive is to protect human health from the adverse effects of any contamination of water intended for human consumption by ensuring that it is wholesome and clean. The directive applies to:

- all distribution systems serving more than 50 people or supplying more than 10 m³/day, but also distribution systems serving less than 50 people or supplying less than 10 m³/day if the water is supplied as part of an economic activity
- drinking water from tankers
- drinking water in bottles or containers
- water used in the food processing industry, unless the competent national authorities are satisfied that the quality of the water cannot affect the wholesomeness of the foodstuff in its finished form.

References


Constitution and structure of bitumens

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Bitumen is a mixture of organic compounds composed of linear aliphatic, cycloaliphatic and aromatic derivatives. Linear aliphatic compounds are saturated linear carbon compounds (e.g. n-heptane). Cycloaliphatic compounds are ring structures that are composed of saturated carbon atoms (e.g. cyclohexane), or cyclic structures that have a low number of unsaturated groups (e.g. cyclohexene), and aromatic compounds are those compounds that have at least one aromatic ring (e.g. benzene). The interaction between molecules and the dilution effect of other molecules in the mixture are reflected in the properties of bitumen. The wide range of compounds and the varying nature of these compounds from bitumen to bitumen make their individual isolation and identification a challenging proposition. The constituents of bitumen are usually separated into classes based on parameters such as solubility, polarity or hydrodynamic size. The hydrodynamic size refers to the volume that a molecule or molecular aggregate (e.g. a colloid particle) occupies when in a solvent. This volume includes the adherence of solvent molecules onto the molecule or aggregate. This volume will be dependent on the molecular structure as well as the solvent used.

The most common method to characterise bitumen is by means of its rheology. Rheology is the science that deals with the flow and deformation of matter. The rheological characteristics of a bitumen at a particular temperature are determined by both the constitution (chemical composition) and the structure (physical arrangement) of the molecules in the material. Changes to the constitution, structure or both will result in a change to the rheology.

Thus, to understand changes in bitumen rheology, it is essential to understand how the structure and constitution of a bitumen interact to influence its rheology. Rheology is considered in detail in Chapter 7.
4.1 Bitumen constitution

4.1.1 Elemental analysis of bitumen

Bitumen is largely composed of hydrocarbon molecules, with some heterocyclic species and functional groups containing sulfur, nitrogen and oxygen atoms (Romberg et al., 1959; Traxler, 1936; Traxler and Coombs, 1936). Bitumen also contains trace amounts of metals such as nickel, vanadium, iron, calcium and magnesium, which occur in the form of metallic salts, oxides or in porphyrin structures. Porphyrins are complex organic compounds that occur naturally: for example, haemoglobin, found in blood, and chlorophyll, found in green plants, are examples of porphyrins associated with metal atoms. Porphyrins contain four nitrogen atoms, each of which can bond with a metal atom to result in a metalloporphyrin. Elemental analysis of bitumen manufactured from a variety of crude oils shows that most bitumen contains:

- carbon: 82–88%
- hydrogen: 8–11%
- oxygen: 0–1.5%
- sulfur: 0–6%
- nitrogen: 0–1%

It has been known since the 1930s that bitumens contain metalloporphyrins, from analytical work that identified iron and vanadium and, consequently, established the link between marine plant chlorophyll and petroleum genesis. Ash from fuel oil on examination by ultraviolet emission spectrography has been shown to contain the following metallic elements (Crump, 1981):

- aluminium
- barium
- calcium
- chromium
- copper
- gallium
- iron
- lanthanum
- lead
- magnesium
- manganese
- molybdenum
- nickel
- potassium
- silicon
- silver
- sodium
- strontium
- tantalum
- tin
- uranium
- vanadium
- zinc
- zirconium

These elements occur principally in the heavier or involatile components of the oils, some as inorganic contaminants, possibly in colloidal form, but also as salts (e.g. of carboxylic acids), transition metal complexes and porphyrin-type complexes. An analysis of bitumen from various sources is shown in Table 4.1.
The predominant metals present in most fuel oils are sodium, vanadium, iron, nickel and chromium, with most of the sodium present as sodium chloride. Vanadium and nickel are largely present as porphyrin structures, which also represent large numbers of different molecules depending on ring substituents (various organic groups that have been added to the core structure of a molecule) and structural isomerism (repositioning of organic groups in different positions within a molecule, with the same molecular formula but with a different chemical or physical behaviour). Bitumen and fuel oil are related products, and Goodrich surmises that similar elements will be present in bituminous compounds (Goodrich et al., 1986).

The precise composition varies according to the source of the crude oil from which a bitumen originates and chemical modification induced during the manufacturing process and ageing in service.

The chemical composition of bitumen is extremely complex. Thus, a complete analysis of bitumen (if it was possible) would be extremely laborious and would produce such a large quantity of data that correlation with the rheological properties would be impractical, if not impossible. In addition, the dataset obtained would be valid for a particular bitumen sample only and not for bitumen in general.

Traditionally, bitumen has been divided into two broad chemical groups called asphaltenes and maltenes. The maltenes can be further subdivided into saturates, aromatics and resins. The four groups are not well defined, and there is some overlap between them. However, this classification does enable bitumen rheology to be set against broad chemical composition.

Solvent extraction is attractive as it is a relatively rapid technique (Traxler and Schweyer, 1953), but the separation obtained is generally poorer than that which results from using chromatography, where a solvent effect is combined with selective adsorption. Similarly, simple adsorption methods (Marcusson, 1916) are not as effective as column chromatography, in which the eluting solution is constantly re-exposed to fresh adsorbent and different equilibrium conditions.

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**Table 4.1** Elemental analysis of bitumen from various sources (Crump, 1981)

<table>
<thead>
<tr>
<th>Element</th>
<th>Carbon: % w</th>
<th>Hydrogen: % w</th>
<th>Nitrogen: % w</th>
<th>Sulfur: % w</th>
<th>Oxygen: % w</th>
<th>Nickel: ppm</th>
<th>Vanadium: ppm</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Range</strong></td>
<td>80.2–84.3</td>
<td>9.8–10.8</td>
<td>0.2–1.2</td>
<td>0.9–6.6</td>
<td>0.4–1.0</td>
<td>10–139</td>
<td>7–1590</td>
</tr>
<tr>
<td><strong>Average</strong></td>
<td>82.8</td>
<td>10.2</td>
<td>0.7</td>
<td>3.8</td>
<td>0.7</td>
<td>83</td>
<td>254</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Range</strong></td>
<td>5–147</td>
<td>0.1–3.7</td>
<td>1–335</td>
<td>1–134</td>
<td>6–159</td>
<td>1.42–1.50</td>
</tr>
<tr>
<td><strong>Average</strong></td>
<td>67</td>
<td>1.1</td>
<td>118</td>
<td>26</td>
<td>63</td>
<td>1.47</td>
</tr>
</tbody>
</table>

---

The chemical composition of bitumen is extremely complex. Thus, a complete analysis of bitumen (if it was possible) would be extremely laborious and would produce such a large quantity of data that correlation with the rheological properties would be impractical, if not impossible. In addition, the dataset obtained would be valid for a particular bitumen sample only and not for bitumen in general.
conditions as it progresses down the column. (An eluting solution is one that is used to remove an adsorbed substance by washing.)

In 1987, the US Congress authorised the Strategic Highway Research Program (SHRP) – a 5-year applied research initiative – to develop and evaluate techniques and technologies to combat the deteriorating conditions of US highways and to improve their performance, durability, safety and efficiency (Halladay, 1998). As part of the SHRP study to investigate the chemistry of bitumen, methods were developed to separate the constituents based on the nature of their functional groups and molecular weight.

4.1.2 Bitumen classification based on solubility

Chromatographic techniques are the most widely used methods to define bitumen constitution (Corbett and Swarbick, 1958; Schweyer et al., 1955). The basis of these methods is initially to precipitate asphaltenes using n-heptane followed by chromatographic separation of the remaining constituents/fractions. Figure 4.1 shows a schematic representation of the chromatographic method. Using this technique, bitumen can be separated into the four groups: saturates, aromatics, resins and asphaltenes (sometimes described by the acronym SARA). The main characteristics of these four broad component groups and the metallic constituents are now discussed.

**Figure 4.1** Schematic of the saturates, aromatics, resins and asphaltenes (SARA) chromatographic method
4.1.2.1 Saturates
Saturates consist of straight and branched-chain aliphatic hydrocarbons together with cycloaliphatic compounds. They are non-polar viscous oils that are straw coloured or colourless. The hydrogen-to-carbon molar ratio (H/C) of the saturate fraction is approximately 2 (Lesueur, 2009). The average molecular weight range has been reported to vary between 470 and 880 g/mol (Lesueur, 2009). This fraction forms 5–20% of bitumen. Figure 4.2 shows two different saturate structures and the structure of cyclohexane.

4.1.2.2 Aromatics
Aromatics comprise the lowest molecular weight naphthenic aromatic compounds in bitumen, and represent the major proportion of the dispersion medium for the peptised asphaltenes. They constitute 40–65% of the total bitumen, and are dark-brown viscous liquids. The H/C ratio of the aromatic fraction could be said to range from 1.4 to 1.6. The average molecular weight range is in the region of 570–980 g/mol (Lesueur, 2009). They consist of non-polar carbon chains attached to unsaturated ring systems (aromatics) (Figure 4.3).

4.1.2.3 Resins
Resins are soluble in n-heptane. They are largely composed of hydrogen and carbon, and contain a small number of oxygen, sulfur and nitrogen atoms. They are dark brown in colour, solid or semi-solid and, being polar in nature, strongly adhesive. Resins are dispersing agents or peptisers for the asphaltenes. The proportion of resins to asphaltenes governs, to a degree, the solution (sol) or gelatinous (gel) character of the bitumen. Resins separated from bitumen are found to have molecular weights ranging from 780 to 1400 g/mol and an H/C atomic ratio of 1.4–1.7.
4.1.2.4 Asphaltenes
These are n-heptane insoluble black or brown amorphous solids containing, in addition to carbon and hydrogen, some nitrogen, sulfur and oxygen atoms. Asphaltenes are generally considered to be highly polar and complex aromatic materials of fairly high molecular weight. Different methods of determining molecular weights have led to various values, ranging widely from 800 to 3500 g/mol, based on vapour pressure osmometric measurements (a method used to measure the molecular weight of compounds based on determining the vapour pressure of solutions using Raoult’s law), and the H/C ratio of asphaltenes ranges from 0.98 to 1.6 (Lesueur, 2009). They have a particle size of 2–5 nm. The asphaltene content has a significant effect on the rheological characteristics of a bitumen. Increasing the asphaltene content produces a harder, more viscous bitumen with a lower penetration, higher softening point and, consequently, higher viscosity. Asphaltenes constitute 5–25% of the bitumen. Figure 4.4 shows a typical chemical structure of an asphaltene.

Recent studies have shown that the asphaltenes can be further separated, based on their solubility in solvents with an increasing solubility parameter,
using a preparative or chromatographic method (Schabron et al., 2010). The method separates n-heptane insoluble asphaltenes into three sub-fractions. It was seen that the relative amounts of aromatic fused-ring structures increased in the asphaltene sub-fractions as the solubility parameter of the extracting solvent increased. This method could be of interest to study further the structure and type of asphaltenes present in bitumen.

The elemental analysis of the above four groups from a bitumen with a penetration value of 100 dmm is detailed in Table 4.2.

### 4.1.3 Other classification methods

Bitumen constituents can also be divided into groups based on their polarity. This is accomplished by using an ion exchange chromatography (IEC) technique (Branthaver et al., 1993). This technique separated bitumen into five components, namely strong acid, weak acid, strong base, weak base and the neutral fraction. The structural analysis of these components was carried out by means of elemental analysis and Fourier transform infrared spectroscopy, while the molecular weight of the fractions was determined by vapour pressure osmometry. The speciation and distribution of functional groups within the fractions and the rheology of the individual fractions obtained from a bitumen sample were also investigated.

The IEC technique has been reported to be sensitive to experimental variables, and the recovery of the sample is never complete (i.e. some of the material remains permanently adsorbed in the columns). When analysing aged bitumen samples, the amount of sample permanently adsorbed on the resin was higher than that observed for fresh bitumen samples.

The separation of bitumen components based on their hydrodynamic volumes using size exclusion chromatography (SEC) was also carried out (Branthaver et al., 1993). Methods to separate bitumen into its constituents, based on molecular size or the size of associated molecules, were used. Compared with IEC, the recovery of the sample using the SEC technique was higher. Viscosity and rheological studies were carried out on the fractions obtained. Interestingly, it was stated that fractions that contained sulfoxides and ketones, which are oxidation products, were found to be

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**Table 4.2** Typical elemental analysis of the four groups of a bitumen with a penetration value of 100 dmm (Chipperfield, 1984)

<table>
<thead>
<tr>
<th>Group</th>
<th>Yield on bitumen (%)</th>
<th>Carbon (%)</th>
<th>Hydrogen (%)</th>
<th>Nitrogen (%)</th>
<th>Sulfur (%)</th>
<th>Oxygen (%)</th>
<th>Atomic ratio H/C</th>
</tr>
</thead>
<tbody>
<tr>
<td>Asphaltenes (n-heptane)</td>
<td>5.7</td>
<td>82.0</td>
<td>7.3</td>
<td>1.0</td>
<td>7.8</td>
<td>0.8</td>
<td>1.1</td>
</tr>
<tr>
<td>Resins</td>
<td>19.8</td>
<td>81.6</td>
<td>9.1</td>
<td>1.0</td>
<td>5.2</td>
<td>–</td>
<td>1.4</td>
</tr>
<tr>
<td>Aromatics</td>
<td>62.4</td>
<td>83.3</td>
<td>10.4</td>
<td>0.1</td>
<td>5.6</td>
<td>–</td>
<td>1.5</td>
</tr>
<tr>
<td>Saturates</td>
<td>9.6</td>
<td>85.6</td>
<td>13.2</td>
<td>0.05</td>
<td>0.3</td>
<td>–</td>
<td>1.8</td>
</tr>
</tbody>
</table>
present in the low molecular weight fractions. This indicated that these molecules did not readily associate with other molecules with polar groups.

4.2 Bitumen structure

4.2.1 Simple colloidal models

Bitumen is traditionally regarded as a colloidal system consisting of high molecular weight asphaltene micelles dispersed or dissolved in a lower molecular weight oily medium (maltenes) (Girdler, 1965). The micelles are considered to be asphaltenes together with an absorbed sheath of aromatic resins that act as a stabilising solvating layer. Away from the centre of the micelle, there is a gradual transition to less polar aromatic resins, these layers extending outwards to the less aromatic oily dispersion medium.

In the presence of sufficient quantities of resins and aromatics of adequate solvating power, the asphaltenes are fully peptised, and the resulting micelles have good mobility within the bitumen. These are known as ‘sol’-type bitumens, illustrated in Figure 4.5.

If the aromatic/resin fraction is not present in sufficient quantities to peptise the micelles, or has insufficient solvating power, the asphaltenes can associate together further. This can lead to an irregular, open packed structure of linked micelles in which the internal voids are filled with an intermicellar fluid of mixed constitution. These bitumens are known as ‘gel’ types, as depicted

Figure 4.5 Schematic representation of a sol-type bitumen
in Figure 4.6, the best examples being the oxidised grades used for roofing purposes. In practice, most bitumens are of an intermediate character.

The colloidal behaviour of the asphaltenes in bitumen results from their aggregation and solvation. The degree to which the asphaltenes are peptised will have a considerable influence on the resultant viscosity of the system. Such effects decrease with increasing temperature, and the gel character of certain bitumens may be lost when they are heated to high temperatures. The viscosities of the saturates, aromatics and resins depend on the molecular weight distribution and interaction between molecules. The higher the molecular weight and the greater the degree of interaction, the higher the viscosity. The viscosity of the continuous phase (i.e. the maltenes) imparts an inherent viscosity to the bitumen that is increased by the presence of the dispersed phase (i.e. the asphaltenes). The saturates fraction decreases the ability of the maltenes to solvate the asphaltenes because high saturate contents can lead to marked agglomeration of the asphaltenes. Accordingly, an increase in gel character and a lower temperature dependence for bitumen results not only from the asphaltene content but also from the saturates content.

4.2.2 More complex models
Dickie and Yen (1967) proposed a model of a colloidal system formed by asphaltenes stacking over each other. The asphaltenes in this model conform
to what is known as the ‘island’ structure (i.e. where there is a central polyaromatic nucleus with aliphatic side-chains). The polyaromatic cores of the asphaltenes result in the proposed stacking behaviour. Further studies and a refinement of the model by Mullins (2010) suggested that the asphaltene stacks form what is termed a ‘nano-aggregate’. These nano-aggregates form clusters (Figure 4.7).

It was suggested that the nano-aggregates are formed by approximately six asphaltene molecules. The polyaromatic core of the asphaltenes was reported to contain, on average, seven fused aromatic rings. The asphaltene clusters of the nano-aggregates form the colloidal portion of the bitumen.

An alternative assembly architecture dealing with asphaltene nano-aggregation was proposed by Gray et al. (2005). The asphaltenes proposed also conform to the ‘island’ configuration of the asphaltene molecule. Here, the nano-aggregation is based on hydrogen bonding, Brøndsted acid–base interactions, metal complex formation and hydrophobic group interactions as well as aromatic core stacking. The asphaltene nano-aggregate structure suggested by Gray et al. results in a more random distribution of molecules compared with the nano-aggregate structure proposed by Dickie and Yen–Mullins.

Other studies have shown that, in addition to asphaltenes that conform to the ‘island’ type of asphaltene, ‘archipelago’-type asphaltenes are also present. It has been suggested that both of these asphaltene types occur (Long et al., 2006). The ‘archipelago’ type of asphaltene consists of a few fused aromatic groups linked by aliphatic chains, as opposed to the central large fused core with aliphatic chains attached to it proposed for the ‘island’ type of asphaltene.

Regardless of the internal structure of the asphaltene nano-aggregates, the colloidal dispersion is composed of the asphaltenes as the colloidal particles dispersed in the maltene phase.
4.3 The relationship between bitumen constitution, structure and rheology

4.3.1 Influence of bitumen components on properties

The influence of bitumen components (i.e. the SARA fractions) on the properties of bitumen has been studied previously (Griffin and Miles, 1961; Griffin et al., 1959; McKay et al., 1978; Reerink, 1973). It was observed, by keeping the concentration of asphaltenes constant in a sample while varying the other components, that:

- increasing the aromatics content at a constant saturates-to-resins ratio has little effect on the rheology other than a marginal reduction in the shear susceptibility;
- maintaining a constant ratio of resins to aromatics and increasing the saturates content softens the bitumen;
- the addition of resins hardens the bitumen and reduces the penetration index and shear susceptibility, but increases the viscosity.

It has also been shown that the rheological properties of bitumen depend strongly on the asphaltene content (Lin et al., 1996). At a constant temperature, the viscosity of a bitumen increases as the concentration of the asphaltenes blended into the parent maltenes is increased. However, the increase in viscosity is substantially greater than would be expected if the asphaltenes were spherical, non-solvated entities. This suggests that the asphaltenes can interact with each other and/or the solvating medium. Even in a dilute toluene solution, the viscosity increase observed with increasing asphaltene concentrations corresponds to a concentration of non-solvated spheres some five times higher than the amount of asphaltenes used. Bitumen asphaltenes are believed to be stacks of plate-like sheets formed of aromatic/naphthenic ring structures. The viscosity of a solution, in particular a dilute solution, depends on the shape of the asphaltene particles. Size is important only if the shape changes significantly as the size increases. At high temperatures, the bonds holding the asphaltene aggregates together are broken, resulting in a change in both the size and shape of the aggregates or clusters. Consequently, the viscosity falls as the temperature increases. However, as a hot bitumen cools, associations between asphaltene nano-aggregates occur, to produce clusters.

The marked increase in non-Newtonian behaviour as bitumen cools is a consequence of the inter-molecular and intra-molecular attractions between asphaltenes and other entities. Under shear, these extended associations will deform or even dissociate in a way that is not adequately described by classical Newtonian concepts. Consequently, at ambient and intermediate temperatures, it is reasonable to conclude that the rheology of bitumen is dominated by the degree of association of asphaltene particles and the
relative amount of other species present in the system to stabilise these associations.

The question regarding the interaction between resins and asphaltenes was considered in the review by Mullins (2010). There has been some evidence to indicate that there is limited interaction between the asphaltene nano-aggregates and the resin fraction (Mullins, 2010). When considering the separation of asphaltenes from bitumen or even crude oils, it must be remembered that asphaltenes are a solubility class; that is, that fraction that is insoluble in n-heptane (although other paraffinic solvents, such as pentane, have also been used). This would mean that what would be considered as the resin fraction of the maltenes separated using n-heptane could be included in the asphaltene fraction, when pentane is used as the asphaltene precipitating solvent.

The argument put forward to support the theory that the resins do not interact with the asphaltenes to stabilise them is that the asphaltene nano-aggregates or dispersion in toluene are stable in the absence of any resin, and the nano-aggregates are similar to those observed in crude oils (Mullins, 2010). Experiments carried out where the separation of asphaltenes from crude oils showed that there was lower than expected amounts of resins present have also been cited as evidence of the limited stabilising effect of resins on asphaltene nano-aggregates.

However, other studies using microcalorimetric measurements (i.e. measurements of extremely small changes in heat content) have shown that there is indeed interaction between the resin and asphaltene phases. These measurements were seen to be comparable to those predicted by simulations. Other studies involving small angle neutron scattering measurements showed that the size of the asphaltene nano-aggregate was smaller in the presence of resins, indicating that there was an interaction between the two components.

The question regarding the interaction or extent of interaction between the resins and asphaltene components has not been addressed by the Dickie and Yen–Mullins model.

### 4.4 The relationship between broad chemical composition and physical properties during the bitumen life cycle

Atmospheric and vacuum distillation removes the lighter components from the bitumen feedstock. The loss of distillates leads to the preferential removal of saturates and the concentration of asphaltenes. Air blowing is carried out to meet certain bitumen application requirements, such as that used for roofing grades. These grades require a higher softening point than paving grade bitumen samples, at similar penetration values. Air is blown through bitumen, maintained at a temperature ranging from 240 to 320°C, until the specific
bitumen properties are obtained. Goppel and Knoterus (1955) determined that, during the bitumen blowing process, carbonyl, carboxylic acid, ester and hydroxyl groups are introduced into the bitumen. Goppel and Knoterus also stated that the molecular weight of the asphaltenes increased during the blowing process.

Air blowing of bitumen from a given vacuum residue or fluxed vacuum residue results in a considerable increase in the asphaltene content and a decrease in the aromatics content. Saturate and resin contents remain substantially of the same order as before blowing. Table 4.3 shows a comparison of the chemical composition of bitumen derived by distillation and that from blowing of the same feedstock, at similar bitumen penetration values.

### 4.4.1 Changes during bitumen processing and asphalt manufacture

Asphalts are manufactured at elevated temperatures. During this process, the properties of the bitumen change compared with those exhibited by the bitumen prior to the manufacturing process. These alterations occur because the bitumen is oxidised, a process often described as ‘ageing’. It has been established that either the rolling thin-film oven test (RTFOT) or the thin-film oven test (TFOT) simulates this ageing of bitumen (Lewis and Welborn, 1940). However, subsequent research found that the RTFOT and TFOT are not good predictors for in-service ageing (Schmidt and Santucci, 1969).

The analysis of bitumen samples after subjecting them to standard RTFOT and TFOT conditions has shown an increase in the concentrations of sulfoxide and ketone groups (Lu and Isacsson, 2002). It was also found that on oxidation, the aromatic fraction of the bitumen usually decreased and the resin and asphaltene fractions increased.

### 4.4.2 In-service changes to bitumen constitution

The oxidation of bitumen during its service life takes place at lower temperatures than those that occur during the manufacture of asphalts. These lower temperatures promote the ageing process.
temperatures have an impact on the kinetics of reactions that occur (Lu and Isacsson, 2002). The effect of temperature on bitumen oxidation kinetics has been investigated using various oxidation methods, and the results are extensively discussed in the review of bitumen ageing by Petersen (2009).

4.4.3 Chemical changes observed during the oxidation of bitumen

Bitumen contains limited concentrations of polar functional groups, which are normally present or formed during oxidation (Petersen, 2009). It has been suggested that bitumen naturally contains some functional groups: phenolic, 2-quinolone type, pyrrolic, pyridinic, sulfide and acidic moieties or functional groups. It has been further suggested that ketonic, sulfoxide and anhydride functional groups are formed due to the oxidation reactions.

Polar groups in bitumen and its fractions impact the properties of bitumen. It has been shown that the presence and concentration of polar groups (in this case the phenolic group) affects the performance of pavement (i.e. its susceptibility to cracking) (Davis and Petersen, 1967). This was also shown to be true in the case of roofing bitumen. This trend has also been shown to be valid in relation to the concentration of ketones (Branthaver et al., 1993; Martin et al., 1990; Petersen et al., 1993). The mechanism of the chemical reactions resulting in an increase in the polar content of oxidised bitumen has also been discussed (Petersen, 2009).

Studies have shown that the nature of the bitumen, as well as the void content of the asphalt pavement, plays a large role in the rate at which bitumen oxidises (Petersen, 2009). The influence of metals and aggregates was also investigated as part of SHRP (Branthaver et al., 1993). It is interesting to note that studies using anti-oxidants and additives, which have been proved to be effective when used in polymeric systems, showed that these were not universally effective when added to bitumen (Martin, 1968).

A study of the change in the composition, in relation to the SARA fractions, was carried out for bitumen while in service in asphalt pavements (Chipperfield et al., 1970) (Figure 4.8). The concept of the ‘ageing index’ has been used to study bitumen when in service: the ageing index is the ratio of the bitumen viscosity after ageing to that before ageing. It is commonly used to normalise the ageing behaviour of samples with different viscosities. The major changes in viscosity were observed to be associated with the mixing and laying process. The recovered bitumen samples were obtained from the top 3 mm of cores extracted from the test sections. Changes in the viscosity of the binder were found to be small, over time. With regard to the chemical composition, the asphaltene content increased with mixing, and showed a gradual increase with time, while the resin and aromatic content
decreased with time. Unexpectedly, the saturates content was observed to increase with time. Overall, it was observed that changes after mixing were very small, even though the initial void content of the mixtures studied was relatively high (5–8%). The ageing index is given by:

\[
\text{Ageing index} = \frac{\eta_r/\eta_0}{\eta_r/\eta_0}
\]

As stated above, bitumen is a mixture of complex compounds, and their interaction directly impacts the performance of bitumen and its products. Although a large amount of research has been carried out to study the composition of bitumen, based on solubility, molecular weight and polarity, at present no precise relationship between the composition and performance of bitumen has been established. Further research will help to elucidate the influence of bitumen components on its properties.
References


Crump GB (1981) Black but such as in esteem – the analytical chemistry of bitumen. Proceedings of the Chairman’s Retiring Address to the NW Region Analytical Division of the Royal Society of Chemistry, London, UK.


Constitution and structure of bitumens


In Europe, bitumens and polymer modified bitumens for asphalts are manufactured to three standards

- EN 12591:2009 (BSI, 2009a), covering penetration and viscosity grade bitumens
- EN 13924:2006 (BSI, 2006), covering hard paving grade bitumens
- EN 14023:2010 (BSI, 2010a), covering polymer modified bitumens (PMBs).

Bitumens for industrial uses are specified by

- EN 13304:2009 (BSI, 2009b), covering oxidised bitumens
- EN 13305:2009 (BSI, 2009c), covering hard industrial bitumens.

The key feature of all of these specifications is a combination of penetration and softening point and, in the case of the soft bitumen grades, viscosity. These properties are generally used to designate the bitumen grade required by the purchaser.

Cut-back and fluxed bitumens are bitumens blended with more or less volatile hydrocarbon components (fuels), mainly kerosene. They are characterised by a viscosity specification, EN 15322:2013 (BSI, 2013) providing the framework for such products. While such products remain popular in some parts of the world (e.g. Australia), their use has decreased markedly in the first decade of the twenty-first century in Europe.

For penetration grade bitumen, simple test methods such as those mentioned in EN 12591 are considered appropriate. Modified bitumens may require more sophisticated test methods to describe their performance adequately.
In addition to the tests specified in the standards cited above, some bitumen suppliers carry out a further range of laboratory tests to ensure that bitumens are manufactured to a consistent standard, maintaining suitability for purpose.

5.1 Penetration grade, hard penetration grade and viscosity grade bitumens

Penetration grade bitumens are specified by the penetration (BSI, 2007a) and softening point (BSI, 2007b) tests. Designation is by penetration range only: for example, 40/60 pen bitumen has a penetration that ranges from 40 to 60 inclusive. The unit of penetration is the decimillimetre (0.1 mm). This is the unit that is measured in the penetration test (discussed, along with the softening point test, in detail in Chapter 6). However, penetration grade bitumens are usually referred to without stating units. Tables 1A, 1B, 2A and 2B of EN 12591 give details of key performance parameters. They are summarised here as Tables 5.1 and 5.2.

In addition to the harder penetration grades, softer bitumens are covered by EN 12591. Because the penetration and softening point tests are not applicable to soft bitumens, the viscosity measurement at 135°C is used as the main differentiating property. Such grades are also called viscosity grade bitumens, and their specifications according to EN 12591 are summarised in Table 5.3.

Hard penetration grade bitumens represent the other end of the viscosity scale: they are generally characterised and designated by penetration range only (e.g. 10/20). Over recent years, the tendency has been to use harder bitumens in asphalts.

5.2 Polymer modified bitumens (PMBs)

During the last three decades, PMBs have become increasingly popular as a replacement for penetration grade bitumens in the upper layers of asphalt pavements. Clients usually do so because of the superior properties of PMBs. Although they represent advanced technology binders, they are still designated both by the penetration range and a minimum softening point: for example, 45/80-55 is a PMB with a penetration of 45–80 dmm and a softening point of at least 55°C. Apart from the test methods that apply to these traditional methods of nomenclature, the specification framework EN 14023:2010 (BSI, 2010a) includes in its Annex B several more sophisticated tests as key performance parameters that need to be validated:

- bending beam rheometer based on test method EN 14771:2012 (BSI, 2012a)
Table 5.1 Specifications for paving grade bitumens with penetrations from 20 × 0.1 mm to 220 × 0.1 mm: Tables 1A and 1B of EN 12591 combined, including examples of specific regional requirements (BSI, 2009a)

<table>
<thead>
<tr>
<th>Property</th>
<th>Test method</th>
<th>Unit</th>
<th>20/30</th>
<th>30/45</th>
<th>35/50</th>
<th>40/60</th>
<th>50/70</th>
<th>70/100</th>
<th>100/150</th>
<th>160/220</th>
</tr>
</thead>
<tbody>
<tr>
<td>Penetration at 25°C</td>
<td>EN 1426</td>
<td>0.1 mm</td>
<td>20–30</td>
<td>30–45</td>
<td>35–50</td>
<td>40–60</td>
<td>50–70</td>
<td>70–100</td>
<td>100–150</td>
<td>160–220</td>
</tr>
<tr>
<td>Resistance to hardening at 163°C</td>
<td>EN 12607-1</td>
<td>%</td>
<td>≥55</td>
<td>≥53</td>
<td>≥53</td>
<td>≥50</td>
<td>≥50</td>
<td>≥46</td>
<td>≥43</td>
<td>≥37</td>
</tr>
<tr>
<td>Retained penetration</td>
<td></td>
<td></td>
<td>≤0.5</td>
<td>≤0.5</td>
<td>≤0.5</td>
<td>≤0.5</td>
<td>≤0.5</td>
<td>≤0.8</td>
<td>≤0.8</td>
<td>≤1.0</td>
</tr>
<tr>
<td>Change of mass (absolute value)</td>
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<td>≤8</td>
<td>≤8</td>
<td>≤8</td>
<td>≤9</td>
<td>≤9</td>
<td>≤9</td>
<td>≤10</td>
<td>≤11</td>
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<tr>
<td>Increase in softening point – severity 1</td>
<td></td>
<td></td>
<td>≤10</td>
<td>≤11</td>
<td>≤11</td>
<td>≤11</td>
<td>≤11</td>
<td>≤11</td>
<td>≤12</td>
<td>≤12</td>
</tr>
<tr>
<td>or Increase in softening point – severity 2 a</td>
<td></td>
<td></td>
<td>≤10</td>
<td>≤11</td>
<td>≤11</td>
<td>≤11</td>
<td>≤11</td>
<td>≤11</td>
<td>≤12</td>
<td>≤12</td>
</tr>
<tr>
<td>Flash point</td>
<td>EN ISO 2592</td>
<td>°C</td>
<td>≥240</td>
<td>≥240</td>
<td>≥240</td>
<td>≥230</td>
<td>≥230</td>
<td>≥230</td>
<td>≥230</td>
<td>≥220</td>
</tr>
<tr>
<td>Solubility</td>
<td>EN 12592</td>
<td>%</td>
<td>≥99.0</td>
<td>≥99.0</td>
<td>≥99.0</td>
<td>≥99.0</td>
<td>≥99.0</td>
<td>≥99.0</td>
<td>≥99.0</td>
<td>≥99.0</td>
</tr>
<tr>
<td>Penetration index</td>
<td>Annex A b</td>
<td></td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Dynamic viscosity at 60°C</td>
<td>EN 12596</td>
<td>Pa·s</td>
<td>≥440</td>
<td>≥260</td>
<td>≥225</td>
<td>≥175</td>
<td>≥145</td>
<td>≥90</td>
<td>≥55</td>
<td>≥30</td>
</tr>
<tr>
<td>Breaking point (Fraass)</td>
<td>EN 12593</td>
<td></td>
<td>≤−5</td>
<td>≤−5</td>
<td>≤−7</td>
<td>≤−8</td>
<td>≤−10</td>
<td>≤−12</td>
<td>≤−15</td>
<td></td>
</tr>
<tr>
<td>Kinematic viscosity at 135°C</td>
<td>EN 12595</td>
<td>mm²/s</td>
<td>≥530</td>
<td>≥400</td>
<td>≥370</td>
<td>≥325</td>
<td>≥295</td>
<td>≥230</td>
<td>≥175</td>
<td>≥135</td>
</tr>
</tbody>
</table>

France  × × × × × ×
Belgium  × × × × × ×
The Netherlands  × × × × × ×
Germany  × × × × × ×
UK  × × × × × × × × × ×
Switzerland  × × × × × × × ×
Czech Republic  × × × × × × × ×
Poland  × × × × × × × ×
<table>
<thead>
<tr>
<th>Property</th>
<th>Test method</th>
<th>Unit</th>
<th>250/330</th>
<th>330/430</th>
<th>500/650</th>
<th>650/900</th>
</tr>
</thead>
<tbody>
<tr>
<td>Penetration at 25°C</td>
<td>EN 1426</td>
<td>0.1 mm</td>
<td>250–330</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>or Penetration at 15°C</td>
<td>EN 1426</td>
<td>0.1 mm</td>
<td>70–130</td>
<td>90–170</td>
<td>140–260</td>
<td>180–360</td>
</tr>
<tr>
<td>Dynamic viscosity at 60°C</td>
<td>EN 12596</td>
<td>Pa·s</td>
<td>≥18</td>
<td>≥12</td>
<td>≥7</td>
<td>≥4.5</td>
</tr>
<tr>
<td>Softening point</td>
<td>EN 1427</td>
<td>°C</td>
<td>30–38</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Resistance to hardening at 163°C</td>
<td>EN 12607-1</td>
<td>–</td>
<td>≤4.0</td>
<td>≤4.0</td>
<td>≤4.0</td>
<td>≤4.0</td>
</tr>
<tr>
<td>Ratio of viscosities at 60°C</td>
<td></td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Increase in softening point</td>
<td>°C</td>
<td>≤11</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Change of mass (absolute value)</td>
<td>%</td>
<td>≤1.0</td>
<td>≤1.0</td>
<td>≤1.5</td>
<td>≤1.5</td>
<td>≤1.5</td>
</tr>
<tr>
<td>Flash point</td>
<td>EN ISO 2719</td>
<td>°C</td>
<td>≥180</td>
<td>≥180</td>
<td>≥180</td>
<td>≥180</td>
</tr>
<tr>
<td>Solubility</td>
<td>EN 12592</td>
<td>%</td>
<td>≥99.0</td>
<td>≥99.0</td>
<td>≥99.0</td>
<td>≥99.0</td>
</tr>
<tr>
<td>Breaking point (Fraass)</td>
<td>EN 12593</td>
<td>°C</td>
<td>≤−16</td>
<td>≤−18</td>
<td>≤−20</td>
<td>≤−20</td>
</tr>
<tr>
<td>Kinematic viscosity at 135°C</td>
<td>EN 12595</td>
<td>mm²/s</td>
<td>≥100</td>
<td>≥85</td>
<td>≥65</td>
<td>≥50</td>
</tr>
</tbody>
</table>

* NR: no requirement may be used, if there is no legal or other regional requirement for this property
Specifying bitumens and checking their quality

5.3 Oxidised bitumens

Oxidised bitumens are used almost entirely for industrial applications (roofing, flooring, mastics, pipe coatings, paints, etc.). They are specified and designated by reference to both the softening point and penetration tests: for example, 85/40 is an oxidised grade bitumen with a softening point of 85 ± 5°C and a penetration of 40 ± 5 dmm. Oxidised bitumens also have

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Table 5.3 Specifications for soft paving grade bitumens with viscosity grading: Tables 3A and 3B of EN 12591 combined (BSI, 2009a)

<table>
<thead>
<tr>
<th>Property</th>
<th>Test method</th>
<th>Unit</th>
<th>V1500</th>
<th>V3000</th>
<th>V6000</th>
<th>V12000</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kinematic viscosity at 60°C</td>
<td>EN 12595</td>
<td>mm²/s</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>1000–</td>
<td>2000–</td>
<td>4000–</td>
<td>8000–</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>2000</td>
<td>4000</td>
<td>8000</td>
<td>16 000</td>
</tr>
<tr>
<td>Resistance to hardening at 120°C</td>
<td>EN 12595</td>
<td>%</td>
<td>≤2.0</td>
<td>≤1.7</td>
<td>≤1.4</td>
<td>≤1.0</td>
</tr>
<tr>
<td>Change of mass (absolute value)</td>
<td>EN 12592</td>
<td>%</td>
<td>&gt;99.0</td>
<td>&gt;99.0</td>
<td>&gt;99.0</td>
<td>&gt;99.0</td>
</tr>
<tr>
<td>Flash point</td>
<td>EN ISO 2719</td>
<td>°C</td>
<td>≥160</td>
<td>≥160</td>
<td>≥180</td>
<td>≥180</td>
</tr>
<tr>
<td>Solubility</td>
<td>EN 12592</td>
<td>%</td>
<td>≥99.0</td>
<td>≥99.0</td>
<td>≥99.0</td>
<td>≥99.0</td>
</tr>
<tr>
<td>Resistance to hardening at 120°C (thin-film oven test)</td>
<td>EN 12607-2</td>
<td>%</td>
<td>≤3.0</td>
<td>≤3.0</td>
<td>≤2.5</td>
<td>≤2.0</td>
</tr>
<tr>
<td>Ratio of viscosities at 60°C</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td></td>
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<td>or</td>
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<td></td>
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<td>NRa</td>
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</tr>
<tr>
<td></td>
<td></td>
<td>NRa</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* NR: no requirement may be used, if there is no legal or other regional requirement for this property

- dynamic shear rheometer based on test method EN 14770:2012 (BSI, 2012b)
- tensile properties by the tensile test EN 13587:2010 (BSI, 2010b)
- elastic properties by the elastic recovery test EN 13398:2010 (BSI, 2010c).

It has been shown that the softening point of PMBs has a different meaning as in the case of penetration grade bitumens, as it is strongly influenced by the type and amount of polymer added to the bitumen. This also explains why the penetration index (when determined from the penetration and softening point) does not apply to PMBs.

While styrene–butadiene copolymers were the most widely used bitumen modifiers in the first decade of the twenty-first century, in recent years other modifiers have become increasingly popular in the quest to enhance bitumen performance. Examples include paraffin, amide waxes and recycled crumb rubber. Specifications for such modified bitumens are beginning to be developed, and the same test methods used for PMBs are likely to describe the performance of such binders adequately.
to comply with solubility, loss on heating and flash point criteria (BSI, 2009b).

The softening points of oxidised grades of bitumen are considerably higher than those of the corresponding penetration grade bitumens, and therefore the temperature susceptibility (i.e. the penetration index is high) is much lower, from +2 to +8.

5.4 Hard bitumens

Hard bitumens are used solely for industrial applications (coal briquetting, paints, etc.). They are specified by reference to both the softening point and penetration tests, but are designated by a softening point range only and the prefix H: for example, H80/90 is a hard grade bitumen with a softening point between 80 and 90°C. Hard bitumens also have to comply with solubility, loss on heating and flash point criteria (BSI, 2009c).

5.5 Bitumen quality

Over many years, Shell has investigated the relationship between laboratory measured properties of penetration grade bitumens and their performance in asphalts on the road. The ability to predict the long term behaviour of asphalts becomes more important as traffic loading has increased and performance requirements have become ever more demanding. Performance on the road depends on many factors, including the design, the nature of the application and the quality of the individual components. In volumetric terms, bitumen is a relatively minor component of an asphalt, but it has a crucial role – acting both as a durable binder and conferring visco-elastic properties on the asphalt.

Essentially, satisfactory performance of a bitumen on the road can be ensured if four properties are controlled:

- rheology
- cohesion
- adhesion
- durability.

5.5.1 Rheology

Although more sophisticated test methods have been developed in the last two decades, the rheology of penetration grade bitumen at service temperatures is still adequately characterised by the values of penetration and the penetration index (Appendix A in BSI 2009a). However, for assessing the rheology of more complex modified bitumens, testing with the dynamic shear rheometer (BSI, 2012b) has found its way into the standards (BSI, 2010a). It allows users to assess bitumen stiffness and its viscous and elastic
Rheology also plays a part when it comes to assessing the tendency of the bitumen to become brittle at low temperature. While the breaking point Fraass test (BSI, 2007c) has traditionally been used for this purpose, a more advanced test, the bending beam rheometer (BBR), is now available (BSI, 2012a).

The BBR (Bahia et al., 1991) is a simple device that measures how much a beam of bitumen will deflect under a constant load at temperatures corresponding to its lowest pavement service temperature when bitumen behaves like an elastic solid. The creep load is intended to simulate the stresses that gradually increase in a pavement as the temperature falls.

Two parameters are determined in this test: the creep stiffness is a measure of the resistance of the bitumen to constant loading, and the creep rate is a measure of how the bitumen stiffness changes as loads are applied.

If the creep stiffness is too high, the asphalt will behave in a brittle manner, and cracking will be more likely. A high creep rate (sometimes denoted the m value) is desirable because, as the temperature changes and thermal stresses accumulate, the stiffness will change relatively quickly. A high value for the creep rate indicates that the bitumen will tend to disperse stresses that would otherwise accumulate to a level where low temperature cracking could occur.

Figure 5.2 shows a schematic diagram of the BBR test equipment.
5.5.2 Cohesion

The cohesive strength of bitumen is characterised either by the force ductility at low temperature (BSI, 2008) or by the tensile test (BSI, 2010b).

In the original ductility test (Figure 5.3), dumb-bell shaped specimens of bitumen are immersed in a water bath and stretched at a constant rate of 50 mm/min until fracture occurs. The distance the specimen is stretched before failure is reported as the ductility. The test temperature is adjusted, depending on the penetration of the bitumen under test (e.g. 10°C for 80–100 pen, 13°C for 60–70 pen and 17°C for 40–50 pen). Under these conditions, the test has been found to discriminate between bitumens of different cohesive strengths.

However, by recording not only the strain at fracture but also the energy required to stretch the sample during the entire test, as described in the force ductility test method (BSI, 2008), a much better discrimination of tensile and therefore cohesive behaviour of different bitumens can be obtained.
For example, the energy recording of unmodified paving grade bitumens generally only shows one maximum at low strain, which is characteristic of the stiffness of the bitumen. Thereafter, the force required to stretch the sample further drops quickly to values close to zero. On the other hand, PMBs are often characterised in this test by two maxima, the first again showing the stiffness of the bitumen compound while the second, occurring at larger strains, typically 200–400 mm, points to the effect of a polymer network in the blend, providing additional resilience. Figure 5.4 shows examples of curves obtained in this test.

### 5.5.3 Adhesion

In contrast to rheology and cohesion, the adhesion of bitumen can be measured only in combination with a substrate. Numerous laboratory tests were developed in the last century, and many of these determine the surface

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**Figure 5.3** (a) Ductility test; (b) elastic recovery of 50/70 (top) and Shell Caraphalte® 45/80-50 (bottom)
coverage of mineral aggregates with bitumen or the stability of Marshall samples before and after storage under water.

The results of these laboratory tests together with observations of performance in practice have identified key links between functional properties and the constitution of the bitumen. This work has indicated that if the molecular
weight distribution and chemical constitution of the bitumen is unbalanced, it can exhibit inhomogeneity that may adversely affect both the cohesive and adhesive properties of the bitumen (Van Gooswilligen et al., 1989).

5.5.4 Durability

Durability can be defined as the ability to maintain satisfactory rheology, cohesion and adhesion in service. The term ‘ageing’ is often used to describe the change in bitumen properties during storage, mixing, laying and in service. The following four mechanisms have been identified as the prime durability factors:

- oxidation
- evaporative hardening
- structural ageing
- exudation.

While, for penetration grade bitumens, ageing is synonymous with hardening, some PMBs may become softer under certain conditions.

5.5.4.1 Oxidation and evaporative hardening

Oxidation – reaction with oxygen from the air – is regarded as the main cause of ageing of bitumen. It is strongly accelerated by increasing temperature and increasing surface area. Also, the bitumen composition influences the rate of oxidation. It has been shown that asphaltene-rich bitumens tend to age quicker than bitumens with low asphaltene contents (Neumann, 1990). Finally, the reaction time determines the extent of oxidative ageing.

This is why the conditions during the preparation of the asphalt (i.e. at temperatures exceeding 120°C) need to be controlled, and the minimum mixing time and temperature should always be aimed at in order to minimise oxidation.

Looking at the mixture, high void contents or even interconnected voids (as in an uncompacted mix or in a porous asphalt) favour the access of oxygen. Therefore, surface courses are more prone to ageing in situ by oxidation than the lower layers of a road pavement.

It is difficult to distinguish between oxidation and evaporation in practice. Evaporation of volatile components, leaving behind a higher viscosity bitumen, can be significant at high temperatures, and may also play a role during the service life.

Tests on the bitumen

The rolling thin-film oven test (RTFOT) (BSI, 2007d) is an ageing test, and measures changes by both oxidation and evaporation. The apparatus for this test is shown in Figure 5.5. In this test, a thin film of bitumen is continuously...
Figure 5.5 The rolling thin-film oven test: (a) external view of RTFOT equipment with control panel on door; (b) internal view of RTFOT equipment with partially loaded carousel with one empty glass jar and another with binder. (Photographs courtesy of PTS International Ltd.)
rotated around the inner surface of a glass jar at 163°C for 75 min, with an injection of hot air into the jar every 3–4 s. The amount of bitumen hardening during the test (i.e. the drop in penetration and the increase in the softening point) correlates strongly with that observed during the manufacture of an asphalt. As mentioned before, PMBs may show a decrease in the softening point, which is caused by the destruction of large polymer chains – these may then react with bitumen molecules, leading to a different structure (Vonk et al., 1994).

Usually a change in mass of the sample before and after the test will be observed. A loss of mass always indicates volatile components in bitumen; however, due to reaction with oxygen, quite often the mass of the sample increases during the test.

After its introduction in the USA, the pressure ageing vessel test (PAV) has also become popular in Europe, being used to predict the changes in bitumen properties during the service life of the pavement (see Figure 5.6 and section 20.3). Again, the softening point and penetration are measured before and after the test, and changes are reported. The test is now standardised as a European Norm (BSI, 2012c).

Evaporative hardening without the interference of oxidation can be assessed by operating the RTFOT with nitrogen instead of hot air or, more directly, by measuring the volatility of the bitumen by true boiling point gas–liquid chromatography (TBP-GLC). In this test, a small sample of bitumen (150 mg) is dissolved in carbon disulfide and is separated on two chromatographic columns. The first column separates the heavy components such as asphaltenes and heavy polar aromatics from the bitumen. The hydrocarbons eluted from this column are then separated on a second column.

TBP-GLC is a rapid and accurate method of front-end volatility analysis, and is plotted on the Qualagon (described in section 5.5.5) as the percentage by mass recovered at 450 and 500°C, thus taking account of the shape of the volatility curve. The loss of 0.2% of the mass in the RTFOT correlates well with the TBP-GLC recovery limits at 450 and 500°C (Van Gooswilligen et al., 1989).

**Tests on the mixture**

The hardening due to the oxidation and evaporation of a thin film of bitumen in contact with aggregate is assessed by two mixture tests: the hot mixture storage test (Van Gooswilligen et al., 1989) and the change in the softening point of the bitumen during commercial asphalt manufacture. The hot mixture storage test simulates ageing conditions during mixing and hot storage. A prescribed mixture is manufactured in the laboratory, and a specified
quantity of this mixture is stored for 16 h at 160°C in a sealed tin. Thus, the volume of air entrained in the sample is known and is constant from test to test. The bitumen is recovered from both the mixed and stored material, and the penetration and softening points are determined from samples of bitumen recovered from these two materials. The ageing of the bitumen during mixing and storage is expressed as the difference between the softening point of the bitumen after storage and the softening point of the original bitumen.

At laboratory scale, this is a very severe test, and the change in the softening point is very much larger than that which would be found during actual bulk storage. Nevertheless, the test correlates with the hardening tendency of a mixture at high temperature when in prolonged contact with air. Again, the limited meaning of the softening point change of PMBs must be taken into account.
5.5.4.2 Structural ageing
There are hints that the structure of bitumen may slowly change during extended service life, leading to a system that is no longer a gel type but a sol. The effect is sometimes also called physical hardening, and is believed to be the result of the slow approach of the thermodynamic equilibrium of the bitumen. Asphaltenes are formed and successively no longer held in solution by the remaining maltenes. However, the results do not unambiguously prove that physical hardening is the sole cause of this process: oxidative ageing is also a suspect.

5.5.4.3 Exudative hardening
If the constitution of a bitumen is unbalanced, it may, when in contact with a porous aggregate, exude oily, less viscous components into the surface pores of the aggregate. Such constitutions can, for instance, appear when highly oxidised bitumens – either blown or strongly aged – are blended with light components, such as flux oils, to produce the desired viscosity of a fresh bitumen. This exudation process may result in a hardening of the bitumen film remaining on the surface of the aggregate. Exudation is primarily a function of the amount of low molecular weight components present in the bitumen relative to the amount and type of asphaltenes.

Shell Research has developed the exudation droplet test (Van Gooswilligen et al., 1989) to measure quantitatively the exudation tendency of a bitumen. In this test, bitumen droplets are applied to the recesses in custom-made white marble plates. The plates are stored at 60°C for 4 days under a nitrogen blanket. During this period, oily rings develop around the bitumen droplet, which can be measured under ultraviolet light using a microscope. Ring widths vary from a few tenths of a millimetre for a balanced bitumen to several millimetres for an unbalanced bitumen.

Hardening in service as a result of exudation can be substantial, and depends not only on the exudation tendency of the bitumen but also on the porosity of the aggregate. If the aggregate possesses low porosity, the quantity of exudate absorbed is negligible, irrespective of the exudation tendency of the bitumen. Similarly, if the exudation tendency of the bitumen is low, the quantity of exudate absorbed will be negligible, irrespective of the porosity of the aggregate. However, highly porous aggregate and bitumen with a strong exudation tendency may lead to quicker hardening and ensuing crack formation of the asphalt pavement.

5.5.5 Qualagon tests and test criteria
By studying the correlation between field performance and the measured properties of experimental and commercial bitumens, Shell Research has developed a set of laboratory tests to assess the quality of a bitumen. The set
of tests includes six on the bitumen and three on the asphalt. For ease of assessment, these nine results are presented in the form of a regular polygon, called the bitumen Qualagon, depicted in Figure 5.7. There are tests within the Qualagon that cover the three remaining key performance elements, apart from rheology:

- cohesion – low temperature ductility (see section 5.5.2)
- adhesion – the retained Marshall test
- durability
  - the RTFOT (oxidation stability)
  - the true boiling point – gas–liquid chromatography (volatility)
  - the exudation droplet test (homogeneity)
  - field trials (oxidation stability).

In the Qualagon, the adhesion characteristics of a bitumen are assessed by a retained Marshall test (Marshall Consulting and Testing Laboratory, 1949). In the test, eight Marshall specimens are manufactured using a prescribed aggregate, aggregate grading and bitumen content. The eight specimens are divided into two groups of four such that the average void content of the two groups is equal. One group of four is tested immediately using the
standard Marshall test, and the remaining four specimens are vacuum treated under water at a temperature of between 0 and 1°C to saturate, as far as possible, the pore volume of the mixture with water. Subsequently, the specimens are stored in a water bath at 60°C for 48 h. The Marshall stability of these four specimens is then determined. The ratio of the Marshall stability of the treated specimens to the initial Marshall stability is termed the ‘retained Marshall stability’.

Although a relatively wide scatter in absolute stability values and poor reproducibility are inherent in Marshall testing, it has been found that consideration of the percentage relative to the result from the standard procedure reduces the differences between results from different laboratories.

The oxidation stability of bitumens according to the field test is plotted on the Qualagon as the percentage retained penetration and the increase in softening point.

Many years’ experience using the Qualagon in conjunction with rheological data have confirmed that the set of criteria on which it is based provides a satisfactory quantitative measure of bitumen quality and its performance in service.

Good quality bitumens generally have properties within the ‘preferred’ area of the Qualagon. However, this does not mean that a bitumen that is partially outside the preferred area is necessarily a poor bitumen or one of low quality. The Qualagon limits are not pass/fail criteria, and the whole figure must be interpreted with care and judgement. For this reason, the Qualagon is not suitable for specification purposes, but it is an invaluable assessment tool providing an excellent guide to performance in service.

5.6 The CEN bitumen specification
Harmonisation of European standards for petroleum products was set as a target by the European Committee for Standardisation (Comité Européen de Normalisation – CEN) during the mid-1980s to eliminate barriers to trade within the member states of the EU. One of the first steps in achieving this aim was the publication of the Construction Products Directive (CPD) in December 1988. The CPD was replaced by the Construction Products Regulation (CPR) in 2011. It requires that construction products used in member states must be fit for their intended use, satisfying certain essential requirements

- mechanical resistance and stability
- safety in case of fire
- hygiene, compliance with health regulations and the environment
- safety and accessibility in use
- protection against noise
energy, economy and heat retention
- sustainable use of natural resources.

These essential requirements have to be taken into account when drafting European standards (often described as ‘ENs’ — Euro Norms). The task of producing ENs has been entrusted to the BSI and, in turn, to the BSI technical committees (TCs) and working groups (WGs). It was the BSI working group dealing with paving grade bitumens, TC227 WG1, which proposed a pan-European specification that has resulted in the publication of EN 12591:2009 (BSI, 2009a). This is based on the traditional tests that are used for characterising bitumens. The specification, shown in Tables 5.1–5.3, includes a series of mandatory tests and properties that must be adopted by all members of the EU. It also includes optional properties that can be adopted or rejected by each national body.

EN specifications are routinely resubmitted every four years, thus the next version of EN 12591 is in preparation.

5.6.1 CE marking

The European Community decided to implement harmonisation of products for construction. Accordingly, asphalt binders were considered. The process began with the original Council Directive (89/106/CEE), and has now moved to Regulation 305/2011. The effect of this harmonisation is to define the minimum specifications that will be applied throughout the EU.

The direct consequence is the mandatory CE marking certification by manufacturers of binders.

In effect, this is a legally enforceable commitment to deliver a product with declared performance, such performance being defined in the framework of associated ENs.

In the case of bitumens, CE marking is obligatory for

- paving grade bitumens complying with EN 12591:2009 (BSI, 2009a)
- PMB complying with EN 14023:2010 (BSI, 2010a).

For more detail on the CE marking of bitumens, refer to section 23.1.4.

5.7 The SHRP/Superpave bitumen specification

The Strategic Highway Research Program (SHRP), initiated in the USA in 1987, was a coordinated effort to produce rational specifications for bitumens and asphalts based on performance parameters. The motivation
was to produce pavements that performed well in service. These pavements were subsequently called ‘Superpave’ (superior performing pavements).

One of the results of this work is the ‘Superpave asphalt binder specification’, which categorises grades of bitumen according to their performance characteristics in different environmental conditions. The specification was intended to limit the potential of a bitumen to contribute to deformation, fatigue failure and low temperature cracking of asphalt pavements. The specifications do not distinguish between unmodified and modified bitumens, although some grades can only be achieved through polymer modification of bitumen.

Table 5.4 shows the complete Superpave binder specification. It is intended to control deformation, low temperature cracking and fatigue in asphalt pavements. This is achieved by controlling various physical properties measured with the equipment described in this chapter. In this specification, the physical properties remain constant for all grades but the temperature at which these properties must be achieved varies, depending on the climate in which the binder is to be used. For example, a PG 52-40 grade is designed to be used in an environment where the average 7 day maximum pavement temperature is 52°C and the minimum pavement design temperature is −40°C. Maps of the USA have been prepared showing these upper and lower temperature regimes, thus facilitating binder selection.

5.7.1 Rotational viscometer
The rotational viscometer is specified to ensure that the viscosity of the bitumen at normal application temperatures is capable of being pumped and coating the aggregate, and will enable satisfactory compaction of the asphalt.

5.7.2 Direct tension test
The dynamic shear rheometer and the bending beam rheometer (BBR) (see section 5.5.1) provide information relating to the (creep) stiffness behaviour of bitumen over a wide range of temperatures. Although stiffness can be used to estimate failure properties, for modified binders the relation between stiffness and failure is less well defined. Accordingly, an additional test to measure the strength and strain at break has been included in the Superpave specifications: the direct tension test [ASTM, 2012].

Bitumens that can be stretched long distances before failure are termed ‘ductile’, and those that break before significant stretching has occurred are termed ‘brittle’. Unfortunately, the BBR is unable to characterise fully the ability of some bitumens to stretch before failure. The direct tension test measures the ultimate tensile strain of a bitumen at low temperatures (between 0 and −36°C). A dumb-bell specimen is loaded in tension at a
### Table 5.4 The complete Superpave binder specification; where $G^\ast$, Complex Shear Modulus; $\delta$, phase angle; $S$, Stiffness modulus (Asphalt Institute, 1997)

<table>
<thead>
<tr>
<th>Performance grade</th>
<th>PG 52</th>
<th>PG 58</th>
<th>PG 64</th>
</tr>
</thead>
<tbody>
<tr>
<td>Average 7 day maximum pavement design temperature: °C</td>
<td>$&lt;52$</td>
<td>$&lt;58$</td>
<td>$&lt;64$</td>
</tr>
<tr>
<td>Minimum pavement design temperature: °C</td>
<td>$&gt; -10$</td>
<td>$&gt; -16$</td>
<td>$&gt; -22$</td>
</tr>
<tr>
<td>Minimum flash point: °C</td>
<td>230</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Viscosity, ASTM D 4402</td>
<td>135</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Maximum 3 Pa-s, test temperature: °C</td>
<td>52</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dynamic shear TP5: $G^\ast/\sin \delta$, minimum, 1.00 kPa</td>
<td>58</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Test temperature at 10 rad/s: °C</td>
<td>64</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Original binder</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mass loss, maximum, %</td>
<td>1.00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dynamic shear TP5: $G^\ast/\sin \delta$, minimum, 2.20 kPa</td>
<td>52</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Test temperature at 10 rad/s: °C</td>
<td>58</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Binder after the RTFOT</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PAV ageing temperature, °C</td>
<td>90</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dynamic shear TP5: $G^\ast/\sin \delta$, maximum, 5000 kPa</td>
<td>25</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Test temperature at 10 rad/s: °C</td>
<td>22</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Binder after the PAV test</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Physical hardening</td>
<td>16</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Creep stiffness, TP1: $S$, maximum, 300 MPa or m-value, maximum, 0.30</td>
<td>13</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Test temperature at 60 s: °C</td>
<td>10</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Direct tension, TP3: Failure strain, minimum, 1.0%</td>
<td>25</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Test temperature at 1.0 mm/min: °C</td>
<td>22</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Report</td>
<td>0</td>
<td></td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>$-6$</td>
<td>$-12$</td>
<td>$-18$</td>
</tr>
<tr>
<td>0</td>
<td>$-6$</td>
<td>$-12$</td>
<td>$-18$</td>
</tr>
</tbody>
</table>
constant rate, and the failure strain (the change in length divided by the original length) is determined. In this test, failure is defined by the stress where the load on the specimen reaches its maximum value. At this failure stress, the minimum strain at failure must be 1%.

References
Chapter 6

Routine testing and mechanical properties of bitumens

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Bitumen is a complex material. Its response to stress is equally complex. The response of a bitumen to stress is dependent on both the temperature and the loading time. Thus, the nature of any bitumen test and what it indicates about the properties of a bitumen must be interpreted in relation to the nature of the material. A wide range of tests is performed on bitumens, from specification tests to more fundamental rheological and mechanical tests.

6.1 Standard specification tests for bitumens

As a wide variety of bitumens is manufactured, it is necessary to have tests to characterise different grades. In several countries, the two tests often used to specify different grades of bitumen are the needle penetration test and the softening point test. Although they are both empirical tests, it is possible to estimate important engineering properties from the results, including high temperature viscosity and the stiffness modulus. The use of the penetration test for characterising the consistency of bitumen dates from the late nineteenth century (Bowen, 1889).

As the tests have to be carried out under well defined conditions, standard methods of testing bitumen exist, as published by the Energy Institute (IP test methods), the American Society for Testing and Materials (ASTM) and the British Standards Institution (BSI), for example. In many cases, the methods are identical. However, some methods differ in detail, for example the BS EN (European standard) and ASTM softening point methods, and in these cases a factor is provided to relate the test results obtained using the different methods.

The majority of the methods quote limits for assessing the acceptability of test results. Limits of variability for results obtained by a single operator
(repeatability) and by different operators in different laboratories (reproducibility) are specified. Thus, tolerance is given to allow for differences between operators and equipment at different locations.

**6.1.1 The penetration test**

The consistency of a penetration grade or oxidised bitumen is measured by the penetration test (e.g. according to ASTM D5 (ASTM, 2013a) or EN 1426 (BSI, 2007a)). In this test, a needle of specified dimensions is allowed to penetrate a sample of bitumen, under a known load at a fixed temperature for a known time. The test apparatus is shown in Figure 6.1.

The penetration is expressed as the distance in decimillimetres (1 dmm = 0.1 mm) that a standard needle will penetrate vertically into a sample of bitumen under specified conditions of temperature, load and load duration. Usually, the applied load is 100 g, the duration of loading 5 s and the test temperature 25°C. For each test, after specified conditioning at the test temperature, three individual measurements of penetration are taken. The average of the three values is recorded to the nearest integer. The recorded penetration is reported if the difference between the individual three measurements does not exceed a specified limit.

**Figure 6.1 The needle penetration test**
The lower the value of penetration, the harder the bitumen. Conversely, the higher the value of penetration, the softer the bitumen. This test is the basis on which penetration grade bitumens are classified into standard penetration ranges.

The specifications for paving grade bitumens and hard paving grade bitumens are provided in EN 12591 (BSI, 2009) and EN 13924 (BSI, 2006), and are discussed in detail in Chapter 5.

It is essential that the test methods are followed precisely, as even a slight variation can cause large differences in the result. The most common errors are:

- Poor sampling and sample preparation
- Badly maintained apparatus and needles
- Incorrect temperature and timing.

Temperature control is critical, to within \(\pm 0.1\,^\circ\text{C}\). Needles must be checked regularly for straightness, correctness of profile and cleanliness. Automatic timing devices are also necessary for accuracy, and these must be checked regularly. Penetration values less than 2 dmm and greater than 500 dmm cannot be determined with accuracy with this equipment. Soft bitumens require longer needles and deeper cups. Very soft bitumens are often better characterised in terms of viscosity. If the penetration value is less than 50 dmm, the repeatability for the penetration test is 2 dmm.

### 6.1.2 The softening point test

The softening point is another property commonly used to determine the consistency of a penetration grade or oxidised bitumen (e.g. according to ASTM D36 (ASTM, 2012a) or EN 1427 (BSI, 2007b) or IP 58 (EI, 2007a)). In this test, a small steel ball is placed on a sample of bitumen contained in a brass ring, and the set up is then suspended in a bath (in the form of a glass beaker) containing water or glycerine. The apparatus is shown in Figure 6.2.

Water is used for bitumen with a softening point of 80\(^\circ\text{C}\) and below, and glycerine is used for bitumen with a softening point above 80\(^\circ\text{C}\). When water is used, the initial temperature of the bath liquid is 5\(^\circ\text{C}\). The initial temperature of the bath with glycerine is 30\(^\circ\text{C}\). The bath temperature is raised at 5\(^\circ\text{C}\) per minute, and the bitumen softens and eventually deforms slowly with the ball through the ring. The softening point temperature is the temperature indicated by the thermometer at the instant the bitumen surrounding the ball touches the bottom plate 25 mm below the ring. The test is performed twice, and the average of the two measured temperatures is reported to the nearest 0.2\(^\circ\text{C}\) for softening points below or equal to 80\(^\circ\text{C}\) and 0.5\(^\circ\text{C}\) for softening points above 80\(^\circ\text{C}\).
points above 80°C. If the difference between the two results exceeds 1°C for softening points below 80°C or exceeds 2°C for softening points above 80°C, the test must be repeated. The reported temperature is designated the softening point of the bitumen. In the ASTM D36 procedure for the softening point test, the liquid in the bath is not stirred whereas in the IP 58 and EN 1427 procedure the water or glycerine is stirred. Consequently, the softening point determined by these two methods differs. The ASTM results are generally around 1.5°C higher than those obtained with the IP and EN methods (Krom, 1950).

As with the penetration test, the procedure for carrying out the softening point test must be followed precisely in order to obtain accurate results (Pfeiffer, 1950). Dimensions of the ring and the ball, the sample preparation, the rate of heating and the accuracy of temperature measurement are all critical (Pfeiffer, 1950). Automatic softening point instruments are available, and these ensure close temperature control and automatically record the result at the end of the test.

The consistency of bitumen at the softening point temperature was determined in terms of penetration value by Pfeiffer and Van Doormaal (1936). Using a specifically prepared extra long penetration needle, they found a value of 800 dmm for many, but not all, bitumens. The exact value was found to vary with the penetration index (PI) and the wax content. It has also
been demonstrated by direct measurement that the viscosity at the softening point temperature of the majority of bitumens is about 1300 Pa·s (13 000 poise) (Heukelom, 1973).

### 6.1.3 The Fraass breaking point test

The Fraass breaking point test (Fraass, 1937) is one of very few tests that can be used to describe the behaviour of bitumens at very low temperatures (as low as $-30^\circ$C). It is essentially a research tool that determines the temperature at which bitumen reaches a critical stiffness, and cracks. A number of countries with very low winter temperatures have specified maximum allowable Fraass temperatures for individual grades of bitumen.

In the Fraass test, shown in Figure 6.3, a steel plate 41 mm long, 20 mm wide and 0.15 mm thick coated with bitumen at a uniform thickness is subjected to a constant cooling rate and flexed repeatedly until the bitumen layer breaks (IP 80 (El, 2007b) and EN 12593 (BSI, 2007c)). The temperature is reduced at a rate of 1°C per minute until the bitumen reaches a critical stiffness, and cracks. The temperature at which the sample cracks is termed the Fraass breaking point, and represents an equi-stiffness temperature. It has been shown that, at fracture, the bitumen has a stiffness modulus of $2.1 \times 10^9$ Pa, which is approaching the maximum stiffness modulus of $2.7 \times 10^9$ Pa (Thenoux et al., 1987). The Fraass temperature can be predicted from the penetration and the softening point for penetration grade bitumens because it is equivalent to the temperature at which the bitumen has a penetration of 1.25 dmm (Heukelom, 1969).
6.2 Temperature susceptibility – the penetration index (PI)

All bitumens become softer when heated, and harden when cooled. One of the best known equations is that describing the temperature susceptibility of the penetration of a bitumen (Pfeiffer and Van Doormaal, 1936). If the logarithm of penetration, log pen, is plotted against temperature \( T \), a straight line is obtained such that

\[
\log \text{pen} = A T + K
\]

where \( A \) is the temperature susceptibility of the logarithm of the penetration and \( K \) is a constant.

The value of \( A \) varies from about 0.015 to 0.06, showing that there may be a considerable difference in temperature susceptibility. Pfeiffer and Van Doormaal developed an equation for the temperature susceptibility that assumes a value of about zero for road bitumens (Pfeiffer and Van Doormaal, 1936; Van der Poel, 1954). For this reason, they defined the PI as

\[
\frac{20 - \text{PI}}{10 + \text{PI}} = 50A
\]

or, explicitly,

\[
\text{PI} = \frac{20(1 - 25A)}{1 + 50A}
\]

The value of the PI ranges from around –3 for highly temperature-susceptible bitumens to around +7 for highly blown low temperature susceptible (high PI) bitumens. The PI is an unequivocal function of \( A \), and hence it may be used for the same purpose. The values of \( A \) and the PI can be derived from penetration measurements at two temperatures, \( T_1 \) and \( T_2 \), using the equation

\[
A = \frac{\log \text{pen at } T_1 - \log \text{pen at } T_2}{T_1 - T_2}
\]

The consistency at the softening point can be expressed in terms of penetration, both by linear extrapolation of the logarithm of the penetration versus temperature and by direct measurement with an extra long penetration needle at the ASTM softening point temperature. Pfeiffer and Van Doormaal found that most bitumens had a penetration of about 800 dmm at the ASTM softening point temperature (Pfeiffer and Van Doormaal, 1936). Replacing \( T_2 \) in the above equation by the ASTM softening point temperature (SP) and the penetration at \( T_2 \) by 800, they obtained the equation

\[
A = \frac{\log \text{pen at } T_1 - \log 800}{T_1 - \text{SP}}
\]
Substituting this equation in the equation for the PI and assuming a penetration test temperature of 25°C gives

\[
\text{PI} = \frac{1952 - 500 \log \text{pen} - 20\text{SP}}{50 \log \text{pen} - \text{SP} - 120}
\]

The assumption of a penetration value of 800 dmm at the softening point temperature is not valid for all bitumens. It is therefore advisable to calculate the temperature susceptibility using the penetration at two temperatures, \(T_1\) and \(T_2\).

The nomographs shown in Figures 6.4 and 6.5 enable the approximate value of the PI to be deduced from either the penetration at 25°C and the softening point temperature, or the penetration of the bitumen at two different temperatures. Due to the spread of the actual value of penetration at the softening point temperature, the value of the PI calculated from one penetration and one softening point may vary from the precise value calculated from two penetration values. However, since the penetration at 25°C and softening point are generally determined for bitumen specification control, it is normally the case that these properties are used.

One drawback of the PI system is that it uses the change in bitumen properties over a relatively small range of temperatures to characterise bitumen. Extrapolations to extremes of temperature can sometimes be misleading. The PI can be used to give a good approximation of the behaviour to be expected, but confirmation using stiffness or viscosity measurements is desirable.

### 6.3 Viscosity

Viscosity is a physical property of a fluid, and is a measure of its resistance to flow. Viscosity is defined as the ratio between the applied shear stress and the rate of shear strain. Various types of viscometers and rheometers are used to determine the flow behaviour under different conditions.

Viscometers often used for bituminous products are the cup viscometer, the capillary viscometer and the rotational viscometer. These are discussed below.

A fundamental method of measuring viscosity is using the sliding plate viscometer in which the shear stress (Pa) and the rate of strain (s\(^{-1}\)) on a thin film of bitumen between two parallel flat plates are determined. Dynamic shear rheometers are used to characterise the visco-elastic behaviour of bitumens. The fundamentals of viscosity and rheology are considered in more detail in Chapter 7.
6.3.1 Cup viscometers

The use of a cup viscometer is a simple method for determining the viscosity at a given temperature. A metal cup is filled with the material at a standard temperature, and the time is recorded in seconds for a standard volume of material to flow out through the orifice in the bottom of the cup. There are several cup viscometers available, which differ mainly in the size of opening through which the material is drained. Such viscosity tests are often used for
emulsions (EN 12846-1 (BSI, 2011a)) and cut-back bitumens (EN 12846-2 (BSI, 2011b)).

The standard tar viscometer is shown in Figure 6.6.

The dynamic (absolute) viscosity (Pa·s) is given by
\[ \eta = \text{flow time (in s)} \times \text{density} \times \text{constant} \]

The test results may also be expressed as the kinematic viscosity, \( \nu \) (mm\(^2\)/s)
\[ \nu = \text{flow time} \times \text{constant} \]
In the above equations, the value of the constant depends on the instrument used. Conversion factors for several cup viscometers are provided in Appendix 2.

6.3.2 Capillary viscometers
The flow conditions are much better defined if the opening or orifice of the cup viscometer is replaced by a long narrow tube or capillary. Capillary viscometers are essentially narrow glass tubes through which the bitumen flows. The tube has narrow and wide sections, and is provided with two or more marks to indicate a particular volume or flow, as shown in Figures 6.7 and 6.8.

The value of the kinematic viscosity is measured by timing the flow of bitumen through a glass capillary viscometer at a given temperature. Each viscometer is calibrated, and the product of the flow time and the viscometer calibration factor gives the kinematic viscosity (in mm²/s).

ASTM D2170 (ASTM, 2010a) and EN 12595 (BSI, 2007d) specify procedures to ascertain the kinematic viscosity, which is often determined at a temperature of 135°C. Measurements of kinematic viscosity at a number of different temperatures are used to obtain temperature/viscosity curves.
Figure 6.7 U tube reverse flow viscometer

Figure 6.8 Kinematic viscosity test equipment
For specifications, the dynamic viscosity at a temperature of 60°C is often required, and the test is carried out using a vacuum capillary viscometer in accordance with ASTM D2171 (ASTM, 2010b) and EN 12596 (BSI, 2007e).

6.3.3 Rotational viscometers
Rotational viscometers are based on the concept that the torque required to rotate an object in a fluid at a certain speed is a function of the viscosity of that fluid. Rotational viscometers are normally used to determine the viscosity of bitumens at application temperatures (ASTM D4402 (ASTM, 2013b) and EN 13302 (BSI, 2010)). An example of a Brookfield rotational viscometer with a Thermosel system is shown in Figure 6.9. Essentially, the device consists of a thermostatically controlled chamber containing a sample of hot bitumen. The spindle is lowered into the bitumen and rotated. The torque
required to rotate the spindle is measured, and converted into the viscosity of
the bitumen (in Pa-s).

The rotational viscosity of bitumen is usually determined at 135 or 150°C,
but with this type of apparatus the viscosity can be determined over a rela-
tively wide range of temperatures (i.e. between 120 and 180°C).

6.4 The bitumen test data chart
In the late 1960s, Heukelom developed a system that permitted penetration,
softening point, Fraass breaking point and viscosity data to be described as
a function of temperature on one chart (Heukelom, 1969). This is known as
the ‘bitumen test data chart’ (BTDC). The chart consists of one horizontal
scale for the temperature and two vertical scales for the penetration and
viscosity. The temperature scale is linear, the penetration scale is logarithmic
and the viscosity scale has been devised so that penetration grade bitumens
with ‘normal’ temperature susceptibility or penetration indices give straight
line relationships. A typical BTDC is shown in Figure 6.10.

The BTDC shows how the viscosity of a bitumen depends on temperature, but
it does not take account of the loading time. However, as the loading times
for penetration, softening point and Fraass breaking point tests are similar,
these test data can be plotted with viscosity test data on the BTDC, as shown
in Figure 6.10. As the test results on this chart form a straight line relationship,
it is possible to predict the temperature/viscosity characteristics of a
penetration grade bitumen over a wide range of temperatures using only the
penetration and softening point.

During the production and compaction of an asphalt, it is important to know
the bitumen viscosities. This is illustrated in Figure 6.11 for a dense asphalt
concrete manufactured using 200 pen bitumen. If the viscosity of the bitumen
is too high during mixing, the aggregate will not be properly coated,
whereas if the viscosity is too low, the bitumen will coat the aggregate easily
but may drain off the aggregate during storage or transportation. For satisfac-
tory coating, the viscosity should be approximately 0.2 Pa-s.

During compaction, if the viscosity is too low, the mixture will be excessively
mobile, resulting in pushing of the material in front of the roller. High viscos-
ities will significantly reduce the workability of the mixture, and little
additional compaction will be achieved. It is widely recognised that the
optimum bitumen viscosity for compaction is between 2 and 20 Pa-s.

Thus, the BTDC is a useful tool for ensuring that the correct operating temp-
eratures are selected to achieve the appropriate viscosity for any grade of
bitumen. Consideration of the viscosity requirements during asphalt pro-
duction and laying has led to the operating temperatures given in
Figure 6.10  BTDC comparing penetration grade bitumens manufactured from one crude
Figure 6.11 BTDC showing bitumen viscosities for optimal mixing and compaction of a dense asphalt concrete
Figure 6.11. Particular mixtures or circumstances may dictate other operating temperatures.

The chart can also be used for comparing the temperature/viscosity characteristics of different types of bitumen. Three classes of bitumen can be distinguished using the BTDC: class S, class B and class W bitumens.

6.4.1 Class S bitumens

The test data for a large group of bitumens can be represented by straight lines on the BTDC, within the repeatability of the test. This group, which has been designated class S (‘straight line’), comprises penetration bitumens of different origins with limited wax content. Figure 6.10 shows a chart with straight lines for a number of different penetration grade bitumens manufactured from one base crude. The lines move towards the left of the chart as the bitumens become softer. However, their slopes are equal, indicating that their temperature susceptibilities are similar. Bitumens with the same penetration at 25°C but having different origins are shown in Figure 6.12. The origin may influence the temperature susceptibility, which is reflected by the slope of the line. Accordingly, the temperature/viscosity characteristics of S-type bitumens may be determined from their penetration and softening point only.

6.4.2 Class B bitumens

The test data of class B (‘blown’) bitumens give curves on the chart as shown in Figure 6.13. The curves can be represented by two intersecting straight lines. The slope of the line in the high temperature range is about equal to that of an unblown bitumen of the same origin, but the line in the lower temperature range is less steep. Physically, there is no transition point, but it is very convenient that they are still straight lines in the penetration and viscosity regions. Each of them can be characterised with two test values. Thus, in all, four tests are required for a complete description: penetration, softening point and two high temperature viscosity measurements.

6.4.3 Class W bitumens

Class W (‘waxy’) bitumens also give curves consisting of two straight lines; however, they are different from those of blown bitumens. The two branches of the curve give slopes that are similar but are not aligned. Figure 6.13 shows an example of an S-type bitumen together with a curve for a similar bitumen with a wax content of 12%. At low temperatures, when the wax is crystalline, there is hardly any difference between the two curves. At higher temperatures, where the wax is molten, the curve for the waxy bitumen is significantly lower down on the chart. Between the two straight branches, there is a transition range in which the test data are scattered because the
Routine testing and mechanical properties of bitumens

Figure 6.12 BTDC comparing several 100 pen bitumens manufactured from different crudes
Figure 6.13  BTDC comparing class S, B and W bitumens
thermal history of the sample influences the viscosity result obtained over this range of temperatures.

6.5 Engineering properties of bitumen

The use of bitumen as such, or as a binder in diverse composite materials, is based on its mechanical properties and on the way these properties depend on the loading time and temperature. Bitumens are manufactured in a variety of grades with different mechanical properties to meet the demands of road and industrial applications. For the civil engineer and the industrial bitumen user, it is of considerable value to define the mechanical properties of bitumen in terms that are analogous to the elastic moduli of rigid materials.

For the structural design of pavements, it is essential to be able to measure and predict the fundamental mechanical characteristics of paving materials (Bonnaure et al., 1977). The stiffness modulus of an asphalt depends on the stiffness modulus of the bitumen and the volume fractions of the different mixture components (aggregates, bitumen, air). The stiffness modulus of bitumen is also a suitable parameter for the breaking properties of the bitumen itself and of the asphalts prepared with it (Heukelom, 1966).

6.5.1 The concept of stiffness modulus

A viscous material is one that is semi-fluid in nature. When stressed, it will deform or tend to deform, any deformation being permanent because it is not recovered when the loading is removed. Elastic materials also deform or tend to deform when stressed, but, when the loading is removed, any deformation is fully recovered. Bitumens are visco-elastic materials. The degree to which their behaviour is viscous and elastic is a function of both temperature and the period of loading (usually referred to as the ‘loading time’). At high temperatures or long loading times they behave as viscous liquids, whereas at very low temperatures or short loading times they behave as elastic (brittle) solids. The intermediate range of temperature and loading times, more typical of conditions in service, results in visco-elastic behaviour.

In order to define the visco-elastic properties, the concept of the stiffness modulus as a fundamental parameter to describe the mechanical properties of bitumens by analogy to the elastic modulus of solids was introduced (Van der Poel, 1954). If a tensile stress $\sigma$ is applied at a loading time $t = 0$, a strain $\varepsilon$ is instantly attained that does not increase with the loading time. The elastic modulus $E$ of the material is expressed by Hooke’s law as stress divided by strain.

In the case of visco-elastic materials such as bitumen, a tensile stress $\sigma$ applied at a loading time $t = 0$ causes a strain $\varepsilon$ that increases, but not proportionately, with the loading time. The stiffness modulus $S$, at a loading
time $t$ is defined as the ratio between the applied stress and the resulting strain at the loading time $t$. It follows from the above that the value of the stiffness modulus is dependent on the temperature and the loading time that is due to the special nature of bitumen. Consequently, it is necessary to state both the temperature $T$ and the loading time $t$ of any stiffness modulus measurement

$$S_{i,T} = \frac{\sigma}{\varepsilon_{i,T}}$$

The methods used to measure the stiffness modulus of bitumen are often based on shear deformations. The resistance to shear is expressed in terms of the shear modulus $G$, which is defined as

$$G = \frac{\text{shear stress}}{\text{shear strain}}$$

The elastic modulus and shear modulus are related by

$$E = 2(1 + \mu)G$$

where $\mu$ is Poisson’s ratio. The value of $\mu$ depends on the compressibility of the material, and may be assumed to be 0.5 for almost incompressible pure bitumens, while values of $<0.5$ have to be considered for asphalts. Thus,

$$E \sim 3G$$

In the static creep test, a constant load is applied, and the resulting deformation is then measured as a function of the loading time. The deformations at loading times from about 1 to $10^5$ s or longer can be measured.

In dynamic tests, the shear stress is usually applied as a sinusoidally varying stress of constant amplitude and fixed frequency. The deformation of the material under test also varies sinusoidally with the same frequency as the applied stress. Tests are carried out at various values of frequency, and the ratio of the stress to strain can be plotted against the inverse of the angular frequency. The degree of elasticity of the bitumen under the test conditions is given by how much the deformation response is out of phase from the applied shear stress, and this is referred to as the phase angle. For purely elastic materials the phase angle is zero. For purely viscous materials the phase angle is $90^\circ$.

In analogy to the elastic modulus $E$ and the shear modulus $G$, the stiffness modulus of bitumen can be given as (Van der Poel, 1954)

$$S_E \sim 3S_G$$

The rheology aspects and the experimental determination of stress–strain relationships for bitumen are considered in more detail in Chapter 7.
By combining creep tests with dynamic tests, a considerable range of stiffness moduli and loading times can be covered. It appears that the stiffness–loading time curves obtained at different temperatures for one grade of bitumen all have the same shape, and, if shifted along the loading time axis, would coincide.

The effect of changes in the temperature and the loading time on the stiffness modulus of three different bitumens is shown in Figures 6.14, 6.15 and 6.16.

Figure 6.14 shows a bitumen of low PI, $-2.3$. At very short loading times, the stiffness modulus is virtually constant, asymptotic towards $2.5 \times 10^9$ to $3.0 \times 10^9$ Pa, and is, in this region, largely independent of the temperature and the loading time (i.e. $S = E$). The effect of the PI is clearly illustrated by comparing Figures 6.14 and 6.15. The bitumen with the higher PI ($+5$) (Figure 6.15) is considerably stiffer at higher temperatures and longer loading times (i.e. it is less temperature susceptible).

Figure 6.16 shows the relationship for a 100 pen bitumen with a PI of $-1.0$. At a loading time of 0.02 s (which equates to a vehicle speed of around 50 km/h) the stiffness modulus is approximately $10^7$ Pa at $25^\circ$C, but falls to approximately $5 \times 10^4$ Pa at $60^\circ$C. At low temperatures, the stiffness modulus is high, and therefore permanent deformation does not occur. However, at higher temperature or longer loading times (slow moving or

Figure 6.14 The effect of the temperature and the loading time on the stiffness modulus of a low-PI bitumen
Figure 6.15 The effect of the temperature and the loading time on the stiffness modulus of a 115/15 bitumen

Figure 6.16 The effect of the temperature and the loading time on the stiffness modulus of a 100 pen bitumen
stationary traffic), the stiffness modulus is substantially reduced and, under these conditions, permanent deformation of the road surface is much more likely to occur.

In order to appreciate fully the significance of the stiffness modulus and its measurement, it is necessary to consider the deformation under stress of simple solids and liquids. The deformation behaviour of visco-elastic materials can then be derived.

6.5.2 Prediction of the stiffness modulus

If direct measurement of the stiffness modulus is not feasible, it can be predicted using the Van der Poel nomograph (Heukelom, 1973; Van der Poel, 1954). Van der Poel showed that two bitumens of the same PI at the same loading time have equal stiffness moduli at temperatures that differ from their respective softening points by the same amount. Over 40 bitumens were tested with PI values varying from +6.3 to −2.3 at many temperatures and frequencies, using both creep tests and dynamic tests. From the test data, Van der Poel produced a nomograph from which, using only penetration and the softening point, it is possible to predict the stiffness modulus of a bitumen over a wide range of conditions of temperature and loading times. The nomograph allows the prediction of the stiffness modulus within a factor of 2, which is considered of practical value in relation to the wide range of stiffness moduli. Figure 6.17 shows a Van der Poel nomograph with the stiffness modulus determined for a 40/60 pen bitumen at a loading time of 0.02 s and a test temperature of 5°C.

6.5.3 Elongation at break

In the characterisation of bitumen (and asphalt) as a construction material, essential factors are the resistance to deformation (rheology) and the permissible deformation (fracture). The stiffness modulus, relating stress and strain, was also used in explaining the breaking properties of bitumen (Heukelom, 1966). Two tests often used in this context are the Fraass breaking point test, to determine the temperature at which a bitumen obtains a certain degree of brittleness, and the ductility test, to determine the elongation at break at a fixed temperature. However, the results obtained in these tests cannot be compared directly because they are measures of different parameters (i.e. temperature and distance). The fracture of bitumen occurs after a certain deformation, the magnitude of which depends on the temperature and the loading time or rate of deformation. It was assumed that there is a relationship between fracture and rheological behaviour at the instant of failure. The rheological behaviour can be expressed in terms of the amounts of elastic strain, delayed elastic strain and viscous strain. Analysing the data from ductility tests, tensile tests and bending tests, it was shown that the
To determine the stiffness modulus of a 50 pen bitumen at the test conditions, connect 0.02 s on the loading time-scale with (53.5 + 1.5 – 5 = 50ºC) on the temperature scale. Stiffness modulus is $1.5 \times 10^8$ Pa at a PI of 0.

Conditions:
- Loading time – 0.02 s
- Temperature 5ºC
- Bitumen properties:
  - Penetration at 25ºC – 50 dmm
  - Softening point (IP) – 53.5ºC
  - PI – 0.0

**Figure 6.17** Nomograph for determining the stiffness modulus of bitumens
stiffness modulus is the main parameter, and the PI is only significant at low stiffness modulus when the viscous deformation is large. At high stiffness modulus, $S = E$. As the stiffness modulus was not measured in the ductility test, it was estimated from Van der Poel’s nomograph.

Figure 6.18 shows the elongation at break as a function of the stiffness modulus and the PI (Heukelom and Wijga, 1973).

When all the results were plotted in Van der Poel’s nomograph, a new nomograph was obtained that allows the elongation at break to be estimated, starting from the same input parameters (Heukelom 1966; Heukelom and Wijga, 1973). This nomograph is shown in Figure 6.19.

The elongation at break is closely related to the strain at break. At low levels of elongation and strain, they are equal. As the strain at break multiplied by the stiffness modulus at break is equal to the breaking strength, the latter is also a function of $S$ and the PI. The elongation being measured in tension,
To determine the elongation at break of a 50 pen bitumen at the test conditions, connect 0.02 s and 10 s on the loading time-scale with (53.5 + 1.5 – 12 = 43°C) on the temperature scale. Elongation at break at 0.02 s is 0.075 (7.5%) and at 10 s is 1.0 (100%).

Elongation = \frac{\text{increase in length}}{\text{original length}}

Conditions:
Loading time – 0.02 s and 10 s
Temperature – 12°C
Bitumen properties:
Penetration at 25°C – 50 dmm
Softening point (IP) – 53.5 °C
PI – 0.0

Figure 6.19 Nomograph for determining the elongation at break of bitumens
these are the tensile strain at break and the tensile strength. It was also found that the strain at break in tension is influenced by the presence of a hard carrier in sample testing, as is used, for example, in the Fraass test (Heukelom, 1966; Heukelom and Wijga, 1973).

6.5.4 Fatigue strength
Like many other materials, the strength of bitumen decreases as a result of repeated loading (fatigue). The fatigue phenomenon has to do with fracture produced by repeated applications of stresses smaller than the breaking strength. Fatigue tests were carried out in bending at constant stress amplitude. The fatigue strength is the stress that causes failure after \( N \) load cycles. Figure 6.20 shows the fatigue strength and the breaking strength as a function of the stiffness modulus.

Figure 6.20 The breaking strength and fatigue strength of bitumen as a function of the stiffness modulus
function of the stiffness modulus (Heukelom, 1966). The breaking strength corresponds to one cycle. The fatigue strength is reduced as the number of loading cycles increases. At higher stiffness moduli, the effect of load repetitions is considerably reduced, and there is a tendency for the lines to converge at $S = E$ (the maximum stiffness modulus of $2.7 \times 10^9$ Pa).

6.6 Cohesion

6.6.1 Ductility and force ductility

The cohesiveness of bitumen can be assessed by its ductility at a given temperature, as mentioned in Chapter 5. In this test, ASTM D113 (ASTM, 2007), a dumb bell shaped specimen of bitumen (a briquette with a specified form of the mould) is immersed in a water bath, and the two ends are pulled apart at a specified temperature and at a specified speed (usually a constant rate of 50 mm/min) until fracture occurs. The distance to which the bituminous material will elongate before breaking is reported as the ductility. The test temperature is adjusted, depending on the penetration of the bitumen in the test. In these conditions, the test has been found to discriminate between bitumens of different cohesive strengths (see Chapter 5).

Under conditions where bitumen forms a very thin thread at large values of elongation, the cohesive strength is close to zero. When bitumen is modified with elastomers, much thicker threads are formed, and the force required to stretch the material indicates a better cohesion. In the force ductility test, EN 13589 (BSI, 2008a), a moulded test specimen is extended in the ductilometer at the specified test temperature and at constant speed until fracture or an elongation of at least 1333% (400 mm) is achieved. The force–elongation curves for an unmodified bitumen and a polymer modified bitumen are presented in Figure 5.4. From the area under the curve, the deformation energy can be calculated, as described in EN 13703 (BSI, 2003), and is a measure of the cohesion.

6.6.2 Toughness and tenacity

In addition to the force ductility – making use of the ductility test set up – other tensile tests, which differ in the shape of the sample, the testing device and the applied loading, are available.

The shape of the force–elongation curve can be used not only to distinguish between modified and unmodified bitumens but also to differentiate between the level and type of modification (in particular, elastomeric components).

As is described in EN 13587 (BSI, 2010), a specimen, held by its ends between two jaws, is extended in a chamber, regulated at the test temperature, at constant speed until fracture or a given percentage elongation is
achieved. The calculations for the different parts of the area under the force–elongation curve are detailed in EN 13703 (BSI, 2003).

ASTM D5801 (ASTM, 2012b) describes a procedure to determine the toughness and tenacity of asphalts. At a specified temperature, a tension head of specified size and shape is pulled from a sample of asphalt at a specified rate, and the force is measured as a function of elongation. The toughness of the sample is defined as the work required to separate the tension head from the sample under the specified test conditions. It is calculated as the total area under the force versus elongation curve. The tenacity of the sample is defined as the work required to stretch the sample after the initial resistance has been overcome. A tangent line is placed against the force versus elongation curve as the force decreases from the initial maximum value so that the tangent line intersects the elongation axis (zero force). The area under the curve to the right of the tangent line is the tenacity of the sample.

6.6.3 Vialit cohesion

The Vialit cohesion pendulum is a test device used to assess the cohesion of bitumen for road construction. The method is described in EN 13588 (BSI, 2008b). The apparatus for carrying out the test is shown in Figure 6.21.

Figure 6.21 The Vialit pendulum test
The procedure is to place a thin film of a bitumen or a polymer modified bitumen between two cubes and to determine the energy required to remove, by pendulum impact, the upper cube. This test is of interest in situations where aggregate is placed in direct contact with traffic stresses, for example in surface dressing. It is important to use binders that have a sufficient level of cohesion for the level of traffic and the site conditions to be supported. The cohesion energy can be significantly increased by polymer modification across a wide temperature range. An example is given in Figure 6.22.

**Figure 6.22** Cohesion curves for a penetration grade bitumen and a polymer modified bitumen

The procedure is to place a thin film of a bitumen or a polymer modified bitumen between two cubes and to determine the energy required to remove, by pendulum impact, the upper cube. This test is of interest in situations where aggregate is placed in direct contact with traffic stresses, for example in surface dressing. It is important to use binders that have a sufficient level of cohesion for the level of traffic and the site conditions to be supported. The cohesion energy can be significantly increased by polymer modification across a wide temperature range. An example is given in Figure 6.22.

**References**


Rheology of bitumens

Rheology is the fundamental interdisciplinary science that is concerned with the study of the internal response of real materials to stresses. The word ‘rheology’ is derived from the Greek, and translates literally as *rheos* (stream or flow) and *logy* (science); therefore, ‘rheology’ literally means ‘the study of flow’. Bitumen rheology can thus broadly be defined as the fundamental measurements associated with the flow and deformation characteristics of bitumen.

Bitumen is a thermoplastic, visco-elastic liquid that behaves as a glass-like elastic solid at low temperatures and/or during short loading times and as a viscous fluid at high temperatures and/or during slow loading. The response of bitumen to stress is therefore dependent on both the temperature and the loading time, and consequently the rheology of bitumen is defined by its stress/strain/time/temperature response.

Traditionally, bitumen has been predominantly characterised and specified using empirical tests. A classic example is the penetration test, which has historically been the backbone for most of the specification regimes globally. The penetration test conditions neither relate to the actual stresses or strains encountered in service nor do they capture any fundamental engineering properties of the material. Instead, the procedure derives a relative measure of the ‘stiffness’ related to the material’s viscosity, and researchers have been successful in using penetration values to formulate an array of extremely useful measures that have greatly facilitated grading and predicting bitumen and asphalt performance.

This simplistic approach has worked rather well for conventional bitumen; however, the growth in use of polymer modified bitumens has necessitated a review of this thinking. Concurrently, there emerged recognition that to characterise a bituminous binder adequately over the entire performance spectrum, it is necessary to capture not only the properties at elevated temperatures but also at intermediate and low temperatures.
Early work, particularly at Shell (Bouldin et al., 1991; Collins et al., 1991), was important in establishing the importance of rheometric testing. Not only does rheometric testing provide an avenue to quantify the temperature dependence of a given property appropriately but it also provides a window to understand the rate (or frequency) dependency of the rheological properties. The US Strategic Highway Research Program (SHRP) built on these insights, and in 1993 published the initial version of the American Association of State and Highway Transportation Officials (AASHTO) performance graded (PG) binder specification, the first bitumen specification to be entirely based on fundamental material properties, derived either from rheological or strength testing (AASHTO, 2009).

This chapter provides an introduction to key rheological concepts and their relevance for bitumen, introduces some of the main test methods employed to make a rheological characterisation of bitumen, and discusses the rheological properties of unmodified and polymer modified bitumens.

7.1 Rheology, deformation and flow

Rheology is concerned with the study of the internal response of real materials to stresses. Eugene C. Bingham, a professor at Lafayette College in the USA, who was researching the properties of materials that showed both elastic properties, akin to solids, as well as flow characteristics typical of liquids, coined the term ‘rheology’ in 1929 from the Greek, as explained above. There is a body of different tests, commonly referred to as rheometry, which has been devised to quantify these properties. More often than not, they have been customised to elicit a response from a material to a specific mechanical stimulus. Rheology can be considered to be the study of materials that exhibit both solid and liquid characteristics.

At rest, a solid or a liquid retains its form; under load the material deforms, and when the load is removed this deformation can be either irrecoverable (viscous) or the deformation can be recoverable (elastic). In the simplest of terms, ideal solid materials behave in an elastic manner, and the deformation is recovered when the force or load is removed, whereas liquids behave in a viscous manner, and the load is not recovered but results in a permanent deformation of the material. Between these two ideal states, materials can exhibit what is referred to as visco-elastic behaviour, in which the response to stress is partially viscous and partially elastic.

Central to the discipline of rheology is the concept of deformation and flow. Deformation is associated with solid behaviour, and can be defined as an alteration in shape or size as a response to an applied load. Flow is associated with liquids (and gases), and is a deformation that continues for as long as the load is applied. Importantly, the effects of temperature and loading
time during the measuring process can give rise to different observations of the behaviour, and a material will exhibit a range of behaviours in response to a force, depending on the temperature and the loading time.

Although flow is typically associated with liquids, over long time-scales all materials can exhibit flow. In the Old Testament (Book of Judges, Chapter 5, Verse 5), the biblical prophetess Deborah captured this phenomenon in the phrase ‘The mountains flow before the Lord’. Implicit in this quotation is an understanding that the way a body behaves, or is perceived to behave, depends intrinsically on the time-scale of the observation and the nature of the material in question. Given sufficient time, even a mountain will flow. A famous example of long term flow behaviour is the windows of Renaissance cathedrals that have gradually become thinner at the top than at the bottom – the viscosity of glass in ambient conditions is too high to observe any cold flow during the brevity of our human lifespan, but over centuries the glass has flowed (Zanotto and Gupta, 1999).

7.1.1 Elasticity, plasticity and the behaviour of solids

The ability of a deformed material body to return to its original shape and size when the forces causing the deformation are removed is referred to as elasticity, and materials exhibiting this behaviour are referred to as elastic. Most solid materials exhibit elasticity, and the response to relatively small stresses results in a directly proportional strain: this relationship was observed by Robert Hooke in 1660, and is referred to as Hooke’s law. Hooke’s law states that the applied force \( F = kx \), where \( k \) is a constant multiplied by the displacement or change in length. The value of \( k \) depends not only on the kind of elastic material under consideration but also on its dimensions and shape. Bitumen exhibits elastic behaviour at low temperatures.

The limit of proportionality is defined as the point beyond which Hooke’s law is no longer true when deforming a material, and the elastic limit is the point beyond which the material under load becomes permanently deformed so that the material does not return to its original length when the force is removed. It marks the onset of plastic behaviour. For most brittle materials, stresses beyond the elastic limit result in fracture with almost no plastic deformation.

7.1.2 Viscosity and the behaviour of liquids

There are two basic forms of relative movement (flow) of adjacent particles of liquid: shear flow, where adjacent particles move over and past each other, and extensional flow, where particles move away from or towards each other (Barnes, 2000). Viscosity (\( \eta \)) is the measure of the resistance to flow of a liquid, and is defined as the ratio between the applied shear stress (\( \sigma \)) and the rate of shear strain (\( \gamma \)). The SI unit for viscosity is the Pascal.
second (Pa·s). The viscosity of all simple liquids, including bitumen, decreases with increasing temperature.

7.1.2.1 Newtonian behaviour and non-Newtonian behaviour

Many everyday fluids exhibit what is referred to as Newtonian behaviour (i.e. where stresses arising are linearly proportional to the local strain rate), and the viscosity is independent of the rate of deformation. The reason that the bitumen viscosity-grading system in the USA and other regions has endured for standard refined bitumens is that, at temperatures close to or exceeding 60°C and very low shear rates (which generally prevail within the standard capillary viscometers used for bitumen), bitumen tends to exhibit Newtonian behaviour.

At a high enough shear rate, all liquids will exhibit ‘non-Newtonian’ behaviour. Some liquids display what is known as shear thickening, and are termed ‘dilatant’. Shear thickening is typically associated with suspensions, and occurs as a result of restructuring of the particles in suspension as the shear rate increases. More typically, liquids display what is referred to as shear thinning, and are termed ‘pseudoplastic’, and their viscosity decreases as the shear rate increases. Most polymer modified bitumens can be considered to be shear-thinning liquids.

The behaviour of liquids under different shear conditions can be described using models. The simplest model is known as the power law model, and is widely used in engineering calculations (Hieber and Chiang, 1992)

$$\eta = m \dot{\gamma}^{n-1}$$  \hspace{1cm} (7.1)

where $\eta$ is the viscosity, $\dot{\gamma}$ is the shear rate, and $m$ and $n$ are positive constants for any given material.

In the case of $n < 1$ the viscosity decreases, and these liquids are generally referred to as pseudoplastic, when $n = 1$ they are Newtonian and when $n > 1$ the fluid is considered dilatant (i.e. the viscosity increases as a function of the shear rate). The limitation of this model is that there is no time constant (i.e. relaxation time) associated with the function, and, for example in the case of a pseudoplastic material, the viscosity monotonically increases to infinity as the shear rate decreases while tending towards zero as the shear rate increases. The most simplistic fix is known as the truncated power law model. Although this model provides a zero-shear viscosity (see section 7.2.3.1) and a characteristic time, it does not address the issue of the monotonically decreasing viscosity, and, hence, does not provide a comprehensive solution for high shear rates. In addition, this approach does not generate a realistic transition from Newtonian to non-Newtonian behaviour.
The Cross model offers, with a simple modification to the truncated power law, an approach that describes the entire range from the Newtonian region to the power law region in one equation (Cross, 1969). This model works relatively well in describing the behaviour for most polymer modified bitumens over a wide range of temperatures and shear rates.

### 7.1.3 Visco-elasticity

In the previous sections, the idealised behaviour of solids and liquids was introduced. The term visco-elasticity refers to the mechanical properties of a material that, in two limiting extremes, can result in the material behaving either as an elastic solid or a viscous fluid, depending on the temperature and the time of loading. Visco-elasticity differs from plasticity in that visco-elastic materials exhibit a time-related recovery when a load is removed: this is often referred to as a delayed elastic response. A plastic material does not return to its original form after the load is removed; plasticity is referred to as ductility in metals. Under load, a material can exhibit a mixed visco-elastic and plastic response in which a proportion of deformation occurs; this is an important characteristic of bitumen and asphalts where accumulated non-recovered deformation manifests itself in the development of ruts in asphalt pavements. Reducing this phenomenon has resulted in the adoption of ways to reduce the permanent deformation, and includes the modification of bitumen using polymers.

Bitumen can be considered a visco-elastic material where, typically, at low temperatures the elastic properties dominate, while at high temperatures the material behaves as a viscous fluid. A well known material that can be used to describe visco-elastic behaviour, particularly with respect to loading time, is Silly Putty™. If it is kneaded into a ball and thrown against the floor (a high rate of deformation), it will bounce back just like a rubber ball. If, however, it is placed over an orifice it will, over time and under the force of gravity, flow in a viscous manner. Bitumen, and in particular polymer modified bitumen, would exhibit a similar response to loading time.

The degree to which a material displays visco-elastic behaviour can be quantified. In deference to the biblical prophetess Deborah, Marcus Reiner, a professor at the Israel Institute of Technology, proposed a dimensionless number, the Deborah number \((De)\) as a measure of a material’s visco-elasticity (Reiner, 1964)

\[
De = \frac{\lambda}{t_0}
\]

where \(\lambda\) is the relaxation time and \(t_0\) is the observation time.

If the relaxation time is very long relative to the observation time, then the Deborah number is large, and the material will exhibit elastic behaviour because it cannot relax the stresses imposed on it. Conversely, if the time
of the observation is very long relative to the relaxation time, then the material will behave in a viscous manner – the energy exerted on it will be dissipated in the form of viscous flow.

As the loading periods become longer, the elastic contribution will steadily decrease, and, in terms of the Deborah number, \( De \ll 1 \). If the time that the stress is applied is much shorter than the relaxation time, then \( De \gg 1 \), and the material will behave more elastically than viscously.

In transient (steady) flow, the Weissenberg number (Wi) – rather than the Deborah number which applies to dynamic flows – is used to determine the nature of the visco-elastic response (Rohn, 1995). If the relaxation time is long and the rate of deformation high, then a material will display non-linear visco-elastic behaviour. Nevertheless, the underlying principle is the same as in the case of the Deborah number

\[
Wi = \lambda \dot{\gamma}
\]

where \( \lambda \) is the relaxation time and \( \dot{\gamma} \) is the shear rate.

An important non-linear visco-elastic effect that may accompany high Weissenberg numbers is referred to as the Weissenberg effect. This refers to a build-up of normal stresses perpendicular to the shear plane. It manifests itself in polymer modified bitumen ‘climbing’ up the shaft of a mixer during production. These effects can, particularly in asphalts, be of significant importance because they will generate confining pressures and can considerably enhance the material’s performance (Sousa et al., 1991).

### 7.1.4 Measuring visco-elasticity properties

There are two fundamentally different modes (and an infinite number of variations) of strain or stress application during tests to determine visco-elastic properties of materials: transient flow and dynamic flow.

#### 7.1.4.1 Transient flow: creep and relaxation tests

Creep and relaxation are two methods used to characterise the response of a material under transient loading. Creep is a physical phenomenon that causes non-reversible deformation of a material exposed to constant stress over a given length of time. Creep compliance \((J(t))\) is defined as the inverse of stiffness as a function of time.

For ideal elastic solids there would be an immediate response to stress during a creep test, and this would result in a constant strain. For a Newtonian liquid the same applied stress would result in an ever increasing strain, which results in a straight line if strain is plotted against time, the slope of which gives the shear rate (Barnes, 2000).
The behaviour of a visco-elastic material in creep tests varies with time, and in a generalised example consists of three regions:

- an instantaneous elastic response
- a delayed elastic response
- a steady state viscous response.

The elastic response of the bitumen dominates at short loading times and/or low temperatures, while the viscous response dominates at long loading times and/or high temperatures. The delayed elastic response dominates at intermediate loading times and temperatures. The purely viscous component and the delayed elastic component constitute the time-dependent deformation of the visco-elastic material. Although none of the viscous deformation is recovered once the load is removed, the delayed elastic deformation is recovered, but not immediately as with purely elastic deformation. Because the relative magnitudes of the three components change with loading time and temperature, both the magnitude and the shape of the creep curve will change with the loading time and the temperature (Anderson et al., 1991).

Linear visco-elastic models are used to model deformation in creep tests. A linear spring and a linear dashpot are used to represent elastic (recoverable) and viscous (irrecoverable) behaviour, respectively, and a linear spring connected in parallel to a linear dashpot is used to represent delayed (recoverable) elastic behaviour. A spring and a dashpot in series are referred to as a Maxwell element, and a spring and a dashpot in parallel are known as a Kelvin element. Combining one Maxwell element and one Kelvin element gives one of the best known models for visco-elastic behaviour: the Burgers model.

Visco-elastic creep data are typically presented in one of two ways. Total strain can be plotted as a function of time for a given temperature or temperatures. Below a critical value of applied stress, a material may exhibit linear visco-elasticity. Above this critical stress, the creep rate grows disproportionately faster. The second way of graphically presenting visco-elastic creep in a material is by plotting the creep compliance as a function of time. Below its critical stress, the visco-elastic creep modulus is independent of the stress applied. A family of curves describing the strain versus time response to various applied stresses may be represented by a single visco-elastic creep modulus versus time curve if the applied stresses are below the material's critical stress value.

If the stress is removed, then the retained ‘strain memory’ will cause the material to recover, and creep and recovery testing can therefore be used to determine how elastic a material is by measuring the recovered strain.

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Data from recovery tests can be treated and represented in the same manner as described above, and the inverse of stiffness as a function of time in recovery is known as the recovery compliance ($J_r$). The proportion recovered can be treated as either a percentage recovery or the magnitude of the creep compliance not recovered, referred to in the multiple stress creep recovery (MSCR) test for bitumen (see section 7.2.3.1) as $J_{nr}$, the non-recovered creep compliance – which has been proposed recently as a predictor for the permanent deformation characteristics of bitumen.

A relaxation test is the reverse of a creep test (Figure 7.2). In relaxation tests, the deformation is imposed suddenly and then maintained at a constant level. The total visco-elastic resistance to deformation at a constant stress level can be evaluated by measuring the total accumulated strain, after the material has had sufficient time to relax.
7.1.4.2 Dynamic flow: oscillatory tests

A dynamic shear rheometer (DSR) can be used to conduct visco-elastic analysis of a material. This is a widespread technique employed to measure bitumen properties, and is a key technique as part of the PG grading system (M 320-10 [AASHTO, 2010a]). In such tests, a sinusoidal strain is applied to a specimen, and the resulting stress is monitored as a function of frequency. This is termed ‘strain controlled testing’ and, for bitumen, is more common than the stress controlled approach in which a sinusoidal varying stress is applied and the strain response measured. If a material is exposed to a strain or stress profile as a function of time, and if the strain is sufficiently small, then it will behave in a linear visco-elastic manner, meaning that there is a linear relationship between the shear stress and the strain, as premised by Hooke’s law

\[ \sigma_{12} \propto \gamma \]  

(7.4)

where \( \sigma_{12} \) is the shear stress and \( \gamma \) is the shear strain.

The operational procedure used in DSR testing for bitumen is to impose sinusoidal strains, as an oscillatory shear, on samples of bitumen sandwiched between the parallel plates (discs) of the rheometer. The amplitude of the responding stress is measured by determining the torque transmitted through the sample in response to the applied strain (Figure 7.3). The strains that are applied during the testing must be kept small to ensure that the test remains in the linear visco-elastic region. Strain sweeps can be used to verify that testing occurs in the linear visco-elastic region. The strain must generally be less than...
0.5% at low temperatures, but can be increased at higher temperatures (Goodrich, 1988).

The loading frequency \( \omega \) is known as the angular frequency, rotational frequency or radian frequency (Eurobitume, 1996; Van der Poel, 1954), and is defined as

\[
\omega = 2\pi f
\]  

where \( \omega \) is the angular frequency (rad/s) and \( f \) is the frequency (Hz).

The response of the applied strain is the development of a stress that, for linear visco-elasticity (small strains), is sinusoidal, and the response of the material is out of phase with the applied strain, and is referred to as the phase angle \( \delta \). This is defined as the phase difference between the stress and the strain, and is also called the loss angle or the phase lag. An alternative symbol, \( \phi \), can also be used for the phase angle. For purely elastic materials, the phase angle will be zero, whereas for purely viscous materials the phase angle will be 90°. The phase angle is therefore important in describing the visco-elastic properties of a material such as bitumen.

The ratio of the resulting stress to the applied strain is called the complex shear modulus, \( G^* \), and is also referred to as the complex modulus, shear modulus or, simply, the stiffness.

The norm of the complex modulus is analogous to the magnitude of a vector (Figure 7.4), as the value is calculated from the square root of the sum of the squares of the components.

The in-phase component of \( G^* \) is called the shear storage modulus \( (G') \), or more commonly the storage modulus. The storage modulus equals the stress that is in phase with the strain divided by the strain, or

\[
G' = G^* \cos \delta
\]  

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{dynamic_shear_test.png}
\caption{Stress–strain response during oscillatory testing}
\end{figure}
The storage modulus describes the amount of energy that is stored and released elastically in each oscillation, and is therefore also known as the elastic modulus, or the elastic component of the complex modulus.

The shear loss modulus $G''$, or simply the loss modulus, is the out-of-phase component of $G^*$. The loss modulus equals the stress $90^\circ$ out of phase with the strain divided by the strain, or

$$G'' = G^* \sin \delta$$  \hspace{1cm} (7.7)

The loss modulus describes the average energy dissipation rate in the continuous steady oscillation found in the dynamic test. The loss modulus is also referred to as the viscous modulus or the viscous component of the complex modulus. The storage and loss modulus, $G'$ and $G''$, are sometimes misinterpreted as the elastic and viscous modulus, respectively. In reality, the elastic component of the response represents only part of the storage modulus, and the viscous response only part of the loss modulus. In addition, visco-elastic materials exhibit a significant amount of delayed elastic response that is time dependent but completely recoverable. The storage and loss modulus both reflect a portion of the delayed elastic response.

The loss tangent is defined as the ratio of the viscous and elastic components of the complex modulus, or simply the tangent of the phase angle

$$\tan \delta = \frac{G''}{G'}$$  \hspace{1cm} (7.8)

The complex viscosity, also termed the complex dynamic shear viscosity, is a viscosity value obtained from dynamic oscillatory tests. The complex viscosity (Pa·s) is defined as the ratio of the complex modulus and the angular frequency

$$\eta^* = \frac{G^*}{\omega}$$  \hspace{1cm} (7.9)

As the complex viscosity is a function of a complex number pair, a real part and an imaginary part of the complex viscosity can also be defined.
The real part of the complex viscosity is termed the dynamic viscosity (Pa·s), and is defined as
\[ \eta' = \frac{G''}{\omega} \] (7.10)

The imaginary part of the complex viscosity is called the out-of-phase component of \( \eta^* \) (Pa·s), and is defined as
\[ \eta'' = \frac{G'}{\omega} \] (7.11)

### 7.1.4.3 Relationship between creep recovery and oscillatory measurements
Van der Poel (1954) defined stiffness as either the inverse of the creep compliance at a loading time \( t \), or the uniaxial dynamic modulus at a loading frequency \( 1/t \). Therefore, a simple approximation between the creep compliance and the dynamic complex modulus can be defined as
\[ G^*(\omega) \approx 1/J(t) \quad \text{as} \quad t \to 1/\omega \] (7.12)

where \( G^* \) is the complex modulus, \( \omega \) is the angular frequency (rad/s) and \( J(t) \) is the creep compliance.

The reciprocal of the shear compliance \( J(t) \) obtained at time \( t_i \) is approximately equal numerically to the complex shear modulus \( (G^*(\omega)) \) at a frequency \( \omega_i \), where \( t_i = 1/\omega_i \).

The transient and dynamic visco-elastic functions can be related mathematically, although the exact relationships are complicated. However, approximate relationships exist that are frequently accurate enough for practical purposes (Anderson et al., 1994).

### 7.1.5 Representation of rheological data
In order to study the rheological characteristics of materials (including bitumens and polymer modified bitumens), the data obtained from testing need to be represented in a useful form. The most common data representation diagram and curves used for bitumen (Eurobitume, 1996) are discussed in the following sections.

#### 7.1.5.1 Simple graphical representations
An isochronal plot or isochrone is an equation, or a curve on a graph, representing the behaviour of a system at a constant frequency or loading time. In a dynamic test, such as the DSR test, curves of the complex modulus \( (G^*) \) or other visco-elastic functions versus temperature at constant frequencies are isochrones.
An isothermal plot or isotherm is an equation, or a curve on a graph representing the behaviour of a system at a constant temperature. In a dynamic test, curves of the complex modulus ($G^*$) as a function of frequency at constant temperatures are isotherms.

A Cole–Cole diagram is a graph of the loss (viscous) modulus ($G''$) as a function of the storage (elastic) modulus ($G'$). The plot provides a means of representing the visco-elastic balance of the bitumen without incorporating frequency and/or temperature as one of the axes.

A Black diagram is a graph of the magnitude (norm) of the complex modulus ($|G^*|$) versus the phase angle ($\delta$) obtained from a dynamic test. The frequency and the temperature are therefore eliminated from the plot, which allows all the dynamic data to be presented in one plot without the need to perform time–temperature superposition manipulations of the raw data (see the following section). A smooth curve in a Black diagram is a useful indicator of time–temperature equivalency, while a disjointed curve indicates the breakdown of the time–temperature superposition, and potentially the presence of a high asphaltene structured bitumen, high wax content bitumen or a highly polymer modified bitumen (Lesueur et al., 1996; Planche et al., 1996).

7.1.5.2 Master curves
To characterise fully the rheology of materials, it is important to be able to construct so-called master curves. Master curves can be constructed from both dynamic and transient loading tests. These master curves can be constructed either in the time domain or in the temperature domain. In their simplest form, master curves are produced by manually shifting modulus versus frequency plots (isotherms) at different temperatures along the logarithmic frequency axis to produce a smooth master curve (Marasteanu and Anderson, 1996).

The extended time or frequency scale used in a master curve is referred to as the reduced time or reduced frequency scale, where the reduced frequency scale is defined as

$$\log f_r = \log f + \log a(T)$$

(7.13)

where $f_r$ is the reduced frequency (Hz), $f$ is the frequency (Hz) and $a(T)$ is the shift factor.

Viscosity–temperature equations are used to characterise the temperature dependency of bitumens, and therefore to determine the shift factors needed for the time–temperature superposition principle. The time–temperature superposition principle can be expressed as (Dobson, 1969)

$$G(w, T) = G(wa(T), T_r)$$

(7.14)
where \( G \) is the modulus (and may be \( G', G'' \) or \( [G^+] \)), \( \alpha(T) \) is the shift factor, \( \omega \) is the loading frequency, \( T \) is the temperature and \( T_r \) is the reference temperature.

The shift factor \( \alpha(T) \) can be defined in several ways, depending on the mathematical equation used for its determination. Temperature dependency, as indicated by shift factors, should not be confused with temperature susceptibility, which is an empirical concept based on the change of consistency or hardness of a bitumen with temperature. The temperature dependency of the visco-elastic behaviour of bitumen is indicated by means of shift factors, and expressed as

\[
\alpha(T) = \alpha(T, T_r) \tag{7.15}
\]

and therefore depends, for a given system, only on the temperature.

The Williams, Landel and Ferry (WLF) equation (Williams et al., 1955) has been widely used to describe the relationship between the shift factors and temperature, and thereby determine the shift factors of bitumens. The equation is theoretical, based on the free volume theory (Ferry, 1971), and makes use of temperature differences, which makes it suitable for practical manipulations. The equation has also been found to be applicable to bitumen results.

The WLF equation is

\[
\log \alpha(T) = \log \frac{\eta_0(T)}{\eta_0(T_r)} = \frac{C_1(T - T_r)}{C_2 + (T - T_r)} \tag{7.16}
\]

where \( \alpha(T) \) is the shift factor at a temperature \( T \), \( \eta_0(T) \) is the Newtonian viscosity at a temperature \( T \), \( \eta_0(T_r) \) is the Newtonian viscosity at the reference temperature \( T_r \), and \( C_1 \) and \( C_2 \) are empirically determined coefficients.

The WLF equation requires three constants to be determined, namely \( C_1 \), \( C_2 \) and \( T_r \). The temperature dependency of bitumens can be described by one parameter, \( T_r \), if universal constants are used for \( C_1 \) and \( C_2 \) in the WLF equation. Williams et al. (1955) proposed that if \( T_r \) is suitably chosen for each material, then \( C_1 \) and \( C_2 \) could be allotted universal values of 8.86 and 101.6, respectively. Brodnyan et al. (1960) showed that, for bitumens, the universal parameters fitted the data for \( T - T_r > -20^\circ \text{C} \), but at lower temperatures the predicted shift factors were too great.

Anderson et al. (1991) have found that for aged and unaged bitumen the constants in the WLF equation are all essentially the same value, with \( C_1 = 19 \) and \( C_2 = 92 \) based on a defining temperature, \( T_d \), which is bitumen specific. These values have also been obtained by Jongepier and Kuilman (1969).
Unfortunately, $T_r$ is difficult to determine. Brodnyan et al. (1960) suggested that $T_r$ is very similar to the softening point. Williams et al. (1955) proposed that $T_r$ was related to the glass transition temperature ($T_g$) by the relationship

$$T_r - T_g = 50^\circ C \quad (7.17)$$

An alternative equation that can be used to describe the relationship between the shift factors and temperature is the Arrhenius equation:

$$\log a(T) = -\frac{E_a}{2.303R}\left( \frac{1}{T} - \frac{1}{T_r} \right) \quad (7.18)$$

where $a(T)$ is the horizontal shift factor, $E_a$ is the activation energy (typically 250 kJ/mol), $R$ is the universal gas constant (8.314 J/K mol), $T$ is the temperature (K) and $T_r$ is the reference temperature (K).

The Arrhenius expression or function requires only one constant to be determined, namely the activation energy. The reference temperature ($T_r$) can be arbitrarily chosen (Nielsen, 1995).

Both the Arrhenius and WLF equations are based on theoretical considerations, and therefore their parameters provide some insight into the molecular structure of bitumen (Marasteanu and Anderson, 1996).

Master curves are only valid for the reference temperature used in the master curve plot. Time–temperature superposition must be applied to calculate the rheological properties at other temperatures. It must be remembered that, in interpreting the master curve, both the time dependency, as indicated by the master curve, and the temperature dependency, as indicated by the shift factors, must be considered in evaluating the rheological properties of the visco-elastic material. For this reason, isochronal plots are probably more informative and more easily interpreted when characterising the rheological properties of bitumen (Anderson et al., 1992).

The shape of the master curve of the complex modulus as a function of the reduced frequency, on a log–log scale, resembles the shape of a hyperbola. The curve has a horizontal asymptote (glassy modulus) at high frequencies and an asymptote at an angle of $45^\circ$ (viscous part) at low frequencies, with a transition range in between.

Modulus curves at low temperatures crowd together at high frequency values, and at very high frequencies they nearly all coincide with one horizontal asymptote. The limiting elastic behaviour is therefore not only independent of frequency but also nearly independent of temperature. The elastic modulus corresponding with this asymptote is called the glassy modulus ($G_g$). However, under viscous conditions, there is no convergence to a single viscous asymptote, as viscosity strongly depends on temperature,
and therefore each temperature gives rise to a separate viscous flow asymptote. The phase angle can also be shifted together with its modulus value, to obtain a curve of the phase angle versus the logarithm of the reduced frequency.

As with Black diagrams, breaks in the smoothness of the master curve indicate the presence of structural changes with temperature within the bitumen, as would be found for waxy bitumens, highly structured ‘gel’-type bitumens and polymer modified bitumens.

A plot of \( \log a(T) \) versus temperature with respect to the reference temperature curve is generally prepared in conjunction with a master curve. These values can be considered to be viscosity changes with respect to the viscosity at the reference temperature, and give a visual indication of how the properties of a viscoelastic material change with temperature. The viscoelastic behaviour of a bitumen is therefore represented by two curves, namely viscosity as a function of temperature (the shift factor versus temperature) and modulus as a function of frequency at a fixed temperature (the master curve).

### 7.2 The rheology of bitumen

The behaviour of bitumen can be discussed in three regions of differing behaviour, namely

- a low temperature linear elastic region
- a high temperature viscous region
- an intermediate temperature visco-elastic region.

Linear behaviour is fulfilled at low temperatures and short loading times (high frequencies), where the bitumen behaves as an elastic solid. The linearity is also maintained (for unmodified bitumens) at high temperatures and long loading times (low frequencies), where the material behaves almost entirely as a Newtonian fluid. The area where non-linearity is prominent is therefore in the range of moderate temperatures and moderate loading times (Van der Poel, 1954). This range of temperatures and loading times corresponds to the conditions experienced in the field.

#### 7.2.1 Low temperature linear elastic region

The bending beam rheometer (BBR) is the most widely used test device for determining the stiffness of bitumen at low temperatures. In principle, the BBR is a constant stress extensional rheometer that yields a Young’s modulus \( E \). However, generally the stiffness is referred to in the bitumen industry as a flexural stiffness \( S(t) \). The test method has been adopted as a standard test in AASHTO’s ‘Standard specification for performance-graded asphalt binder’ (AASHTO M 320-10) (AASHTO, 2010a). There is also a detailed

During this testing, a beam of bitumen is subjected to a constant stress by means of a loaded piston (100 g) in a three-point bending machine. This beam is suspended in an approximately equi-dense cooling fluid, and the travel of the piston measured as a function of time (Lee, 1997). The relationship between the relevant parameters is

\[ S(t) = \frac{pL^3}{4\delta(t)bh^3} \]  

(7.19)

where \( L \) is the beam length, \( h \) is the beam height, \( p \) is the applied constant load, \( b \) is the beam width and \( \delta(t) \) is the displacement at time \( t \).

For specification purposes, \( S(t) \) and the slope of the curve, commonly referred to as the \( m \) value, are both determined at a load time of 60 s. Alternatively, the stiffness curves can be obtained at different temperatures, and flexural stiffness master curves constructed. The data can also be treated in exactly the same way as creep test data, to compute compliance information. To convert the results from the extensional mode to the shear mode, it is imperative to take Poisson’s ratio (\( \nu \)) into consideration. Poisson’s ratio accounts for the fact that, when extending, an object will compress in the perpendicular direction and expand in the axial direction (in which it is being pulled). For the ideal case of an incompressible, isotropic material, \( \nu \) should be equal to 0.5. However in practice, the value of Poisson’s ratio can be significantly smaller.

7.2.2 High temperature viscous region

At very low shear rates or very low stress levels, almost all bitumens exhibit Newtonian behaviour. Non-Newtonian behaviour appears gradually as the shear rate or stress level increases. Generally, unmodified bitumens tend to exhibit Newtonian behaviour at temperatures greater than approximately 60°C. However, polymer modified bitumens tend to be shear susceptible at temperatures above 60°C and sometimes at mixing and compaction temperatures, and therefore exhibit non-Newtonian behaviour at higher temperatures than those associated with unmodified bitumens (Anderson et al., 1991).

Historically, and in the USA particularly, the Asphalt Institute vacuum capillary viscometer has been the standard equipment used to determine the viscosity of conventional bitumens, and more recently of polymer modified bitumens. A detailed description of the equipment is given in The Asphalt Handbook (AI, 2007). Generally, the viscosity is measured with this device at 60°C or 135°C.
The drawback of capillary viscometers is that a particular shear rate cannot be selected because the value depends on the viscosity of the material. The lower the viscosity, the faster it flows through the tube and the higher the shear rate. This does not present difficulties for conventional grades of bitumen, which tend to behave in a Newtonian manner at the rates observed in a vacuum capillary viscometer. However, the non-Newtonian behaviour of polymer modified bitumens gives rise to limitations in this approach.

Rotational viscometers consist of one cylinder rotating coaxially inside a second (static) cylinder containing the bitumen sample. The material between the inner cylinder and the outer cylinder is therefore analogous to the thin bitumen film found in the sliding plate viscometer. The torque on the rotating cylinder or spindle is used to measure the relative resistance to rotation of the bitumen at a particular temperature. The torque value is then altered by means of calibration factors to yield the viscosity of the bitumen. Rotational viscometers are also commonly used as a practical means of determining the viscosity and/or the shear rate dependency of bitumen, because such instruments allow the testing of a wide range of bitumens over a wide range of temperatures, more so than most other viscosity measurement systems (Zacharias and Emery, 1994).

The shear susceptibility, or shear rate dependency, of bitumen can be defined as the change in rheological properties of the bitumen as a function of the loading time. The shear susceptibility of bitumen can be represented by modelling the flow properties using a power law model in which the logarithms of the shear stress and shear strain rate are linearly related (Traxler et al., 1944)

\[ \log \sigma = c \log \dot{\gamma} + B \]  

(7.20)

where \( \sigma \) is the shear stress, \( \dot{\gamma} \) is the \( d\gamma(t)/dt \) shear strain rate, \( c \) is the degree of complex flow and \( B \) is a constant.

A more common representation of the shear rate dependency is the following function

\[ \eta = \frac{\sigma}{\dot{\gamma}^c} \]  

(7.21)

where \( \eta \) is the apparent viscosity.

Viscosity testing, although a more fundamental method of determining the rheological performance of bitumen, does not provide information on the time dependence of bitumen. These measurements are consequently not adequate for describing the visco-elastic properties needed for the complete rheological evaluation of bitumen.
7.2.3 Intermediate temperatures and the visco-elastic region

At in-service pavement temperatures, bitumen has properties that are in the visco-elastic region. At these temperatures, the bitumen therefore exhibits both viscous and elastic behaviour and displays a time-dependent relationship between the applied stress or strain and the resultant strain or stress (Goodrich, 1988). The conventional methods of characterising the rheological properties of bitumen cannot therefore completely describe the visco-elastic properties that are needed to relate fundamental physical binder properties to performance. The penetration and softening point tests, although conducted around the temperatures in question, are almost completely empirical, and hence not useful for characterising the visco-elastic behaviour of bitumen.

The visco-elastic characteristics of bitumen are typically determined by means of time-dependent tests. The two most common means of determining the visco-elastic properties of bitumen are by creep and recovery measurements (transient loading) and dynamic oscillatory measurements (alternating stress and strain of constant amplitude and frequency).

7.2.3.1 Creep and recovery tests

Steady state creep measurements can be used to calculate an apparent viscosity at intermediate pavement service temperatures, ranging from 0 to 25°C (Griffen et al., 1956; Moavenzadeh and Stander, 1967; Romberg and Traxler, 1947). To conduct such measurements, it is necessary to apply a shear stress to the bitumen until the strain rate becomes constant. At low temperatures, longer times are required for delayed elasticity to be expended and steady state flow to occur. This results in the bitumen being subjected to very large strains, causing geometric non-linearity. The delayed elasticity and geometric non-linearity can be overcome by assuming that the steady state strain rate has been attained at a series of shear stress levels, and extrapolating the calculated apparent viscosities to a zero shear rate. This apparent viscosity at a zero shear rate is known as the zero shear viscosity ($\eta_0$), and is equivalent to the maximum Newtonian viscosity of the bitumen at the test temperature.

The time–temperature superposition principle can be applied to creep test data to produce a master curve from the creep data. The horizontal shift factor ($a(T)$) needed for the production of the master curve can be produced for each creep curve determined at a particular test temperature. These shift factors can be plotted against temperature to form, together with the master curve, a complete characterisation of the linear stress–strain–time–temperature response of the bitumen. The time dependency of the bitumen is reflected in the master curve whereas the temperature dependency is reflected in the temperature shift factors ($\log a(T)$).
Recently, creep and recovery testing has been proposed as a standard means of more accurately predicting the rutting behaviour of bitumen and asphalts (AASHTO TP 70-09 (AASHTO, 2009) and AASHTO MP 19-10 (AASHTO, 2010b)). In this approach, a sample of bitumen is subjected to a 1 s load and a recovery time of 9 s using a DSR. The test starts with the application of a low stress (0.1 kPa) for ten creep/recovery cycles, after which the stress is increased to 3.2 kPa and repeated for an additional ten cycles. It is considered that the higher levels of stress applied during the test are more representative of the conditions experienced in the pavement (Federal Highways Administration, 2011). Results from the multiple stress creep recovery (MSCR) test are represented as a percentage recovery and non-recovered creep compliance ($J_{nr}$).

### 7.2.3.2 Oscillatory tests: dynamic shear rheometry

The rheology of bitumen is typically measured by using oscillatory type testing, generally conducted within the region of linear visco-elastic response. This approach allows the viscous and elastic (visco-elastic) nature of the bitumen to be determined over a wide range of temperatures and loading times (Anderson et al., 1994; Goodrich 1988; Petersen et al., 1994a). It is important that strain sweeps are carried out to ensure that the dynamic tests are conducted in the linear visco-elastic range so that the material functions, such as the complex modulus ($G^{\ast}$) and the complex viscosity ($\eta^{\ast}$), are independent of the applied strain levels.

Sliding plate rheometers were one of the first devices developed to characterise the rheology of bitumens. These devices can be used for transient or stress and strain controlled experiments, but they are also appropriate for performing oscillatory shear experiments. The strain amplitude of the oscillations needs to be kept sufficiently small to ensure linearity and angular frequencies, and gaps are adjusted to minimise inertial effects.

The DSR using plate–plate geometry is the most common example of a rheometer (Figure 7.5). In such tests, a sample of bitumen, which is sandwiched between two parallel discs or plates, is subjected to alternating shear stresses and strains. The test can be either stress controlled or strain controlled, depending on which of these variables is controlled by the test apparatus. The test condition usually used to determine the dynamic rheological properties of the bitumen is typically the controlled strain condition (Goodrich 1988; Petersen et al., 1994b). The use of the controlled strain condition ensures that the strains are kept small and therefore within the linear visco-elastic region.

The principle that is used to relate the equivalency between time and temperature and thereby produce the master curve is known as the time–temperature superposition principle or the method of reduced variables (Ferry,
1971). Because bitumens are linear visco-elastic in a wide range of their applications and because they are found to be ‘thermo-rheologically simple’, the time–temperature superposition principle can be used to determine master curves and shift factors. Typically, dynamic data are first collected over a range of temperatures and frequencies, a standard reference temperature is selected that is usually between 0 and 25°C, and the data at all other temperatures are then shifted with respect to time until the curves merge into a single smooth function.

The shifting may be carried out based on any of the visco-elastic functions, such as the complex modulus \( G^* \), and if the time–temperature superposition principle is valid, the other visco-elastic functions will all form continuous functions after shifting. This continuous curve represents the binder behaviour at a given temperature for a wide range of frequencies. The complex modulus increases with decreasing penetration values for a given frequency, and over the reduced frequency scale of the master curve the complex modulus of the

**Figure 7.5** A dynamic shear rheometer typically used in bitumen testing
bitumen varies by over six orders of magnitude (i.e. powers of $10$). Master curves of the complex modulus for a range of bitumens are given in Figure 7.6.

Over the last two decades, DSR testing of bitumen has become much more widespread, primarily as a result of the US bitumen research programme of the Strategic Highway Research Program (SHRP). SHRP developed and implemented a number of binder rheological and fracture (failure) tests that measure the fundamental properties of bitumen. These new or improved test methods were used to develop a performance related specification as part of Project A-002A (Anderson and Kennedy, 1993), which introduced DSR testing and rheology into bitumen specifications. Properties, such as the complex modulus ($G^*$) and the phase angle ($\delta$), are measured across a range of frequencies and temperatures.

Although the rutting of asphalts is influenced primarily by mixture properties, the properties of the binder are also important. This is particularly true for polymer modified bitumens, which are claimed to enhance the rutting resistance of asphalt pavements. A measurement of the non-recoverable deformation of bitumen at high temperatures using the DSR was established as
a suitable rutting parameter for bitumen. A loading time of 0.1 s was chosen to represent the loading time within the pavement, which can be attributed to a truck tyre travelling at 80 km/h (Petersen et al., 1994a). This 0.1 s loading time is equivalent to a sinusoidal loading at a frequency of 1.59 Hz. The specification criterion for rutting was taken as the inverse of the loss compliance, \(1/J''\), which is numerically equal to the complex modulus divided by the sine of the phase angle, \(G^*/\sin \delta\) (Anderson and Kennedy, 1993). The SHRP specification states that, at the maximum pavement design temperature, \(G^*/\sin \delta\) for an unaged bitumen must be greater than 1.0 kPa, and for a bitumen aged by rolling thin-film oven test it must be greater than 2.2 kPa (Anderson and Kennedy, 1993; Petersen et al., 1994a).

Fatigue cracking generally occurs late in the life of the pavement, and therefore the bitumen needs to be tested after appropriate long term ageing. The fatigue parameter was chosen to reflect the energy dissipated per load cycle, which can be calculated as \(G^* \sin \delta\) in a dynamic shear test (Ferry, 1971). The SHRP specification requires that, at the intermediate pavement design temperature, the value of \(G^* \sin \delta\) at a frequency of 1.59 Hz must be less than 5.0 MPa (Anderson and Kennedy, 1993; Petersen et al., 1994b).

The selection of the basically empirical parameters of the complex modulus divided by the sine of the phase angle (\(G^*/\sin \delta\)) and the complex modulus multiplied by the sine of the phase angle (\(G^* \sin \delta\)) to represent the binder characteristics from the available DSR rheological data to describe the performance of bitumens presents some limitations. This is particularly relevant for modified binders, such as polymer modified bitumens, which, due to their complex rheological behaviour, require greater quantities of data for their complete characterisation. Additionally, the small strain measurements associated with the DSR are not considered to be wholly representative of pavement conditions, and this has given rise to alternative approaches such as the MSCR test outlined in section 7.2.3.1.

### 7.2.4 Relationships between fundamental and empirical bitumen testing

Many studies link between index properties taken from rheological test data and empirical tests such as penetration and the softening point (Eurobitume, 2012; Nicholls et al., 1999). In general, unmodified bitumen relationships can be successfully developed between fundamental and empirical tests. For example, the penetration value of bitumen can be estimated using the following formula (Gershkoff, 1995)

\[
\log(G^*|_{25^\circ C, f=0.4 \text{ Hz}}) = 2.923 - 1.9 \log(\text{Pen})
\]  

\[\text{(7.22)}\]
where $G^*_{25^\circ C, f=0.4 \text{ Hz}}$ is the complex modulus at $25^\circ C$ and 0.4 Hz and Pen is the penetration.

Such approaches work reasonably well for unmodified bitumens, but when relating the rheological properties of polymer modified bitumens to empirical tests, the more complex rheology of the latter gives rise to difficulties, and a more detailed approach is often required to characterise them.

### 7.2.5 Rheology of polymer modified bitumen

The use of synthetic polymers to modify the performance of conventional bituminous binders dates back to the early 1970s (Taylor and Airey, 2008), with these binders subsequently having decreased temperature susceptibility, increased cohesion and modified rheological characteristics. Polymers have been primarily used to improve the high temperature properties of bitumen and provide asphalts with a better resistance to deformation (Collins et al., 1991; Goodrich, 1988; Isacsson and Lu, 1995; King et al., 1993). In addition, the polymer modified bitumen should possess improved resistance to thermal and fatigue cracking (Brown et al., 1990).

A polymer modified bitumen extends the temperature range, sometimes referred to as the plasticity interval. In simple terms, the plasticity interval can be defined as the temperature range between the measure of high temperature performance (e.g. the softening point or criteria based on the complex modulus in the SHRP PG grading approach) and the low temperature measure of performance (e.g. a brittleness point or limiting stiffness value determined by the BBR). In rheological terms, the plasticity interval broadly describes the temperature range between the bitumen behaving as an elastic solid and as a viscous liquid. The extent to which polymer modification extends the plasticity interval is dependent on the degree of modification, the polymer properties, the polymer content and the nature of the base bitumen.

Plastomers modify bitumen by forming a tough, rigid, three dimensional network to resist deformation, while elastomers have a characteristically high elastic response and, therefore, resist deformation by stretching and recovering their initial shape. Elastomers, such as styrene–butadiene–styrene (SBS) copolymers, derive their strength and elasticity from the physical cross-linking of the molecules into a three dimensional network within the bitumen. The polystyrene end-blocks impart strength to the polymer, while the polybutadiene rubbery matrix mid-blocks give the material its elasticity. The effectiveness of these cross-links diminishes rapidly above the glass transition temperature of polystyrene (approximately $100^\circ C$), although the polystyrene domains will reform on cooling, restoring the strength and elasticity of the copolymer (Isacsson and Lu, 1995).
Many polymer modified bitumens tend to behave more as a polymer than as a bitumen. For a cross-linked SBS (or styrene–butadiene) polymer blend, four distinct regions are found:

- A flow region, where the properties are primarily driven by the viscous response.
- A plateau region, where the storage modulus is more or less independent of the frequency (or, conversely, temperature).
- A transition region at intermediate temperatures.
- A glassy region, where the properties are entirely controlled by the physico-chemical make-up of the bitumen at low temperatures.

Conventional bitumens are characterised by a flow region that transitions into the glassy region (i.e. no truly visco-elastic region can be identified). Figure 7.7 shows schematically how cross-linking greatly enhances the elastic response relative to a conventional bitumen.

The effect of SBS modification on the rheological parameters (the complex modulus and the phase angle) has been combined in the form of Black space diagrams, an example of which is shown in Figure 7.8. The morphology, and therefore the rheological characteristics of the polymer modified bitumens, are functions of the mutual effect of the polymer and the bitumen and, consequently, are influenced by the bitumen composition and the polymer nature and content. At low polymer contents (up to 3%), the behaviour of the modified binder remains closer to that of the base bitumen. At
higher polymer contents, the polymer network in the SBS polymer modified bitumens results in a continued reduction in the phase angle (increased elastic response) at low complex modulus values.

Isochronal plots of the complex modulus ($G^*$) and the phase angle ($\delta$) versus the temperature at 0.02 Hz for a base bitumen with an increasing quantity of SBS are shown in Figures 7.9 and 7.10. Although there are only minor increases in $G^*$ at low temperatures due to SBS modification, there is

---

**Figure 7.8** Black space diagram showing the effect on the rheology of bitumen with increasing SBS addition

**Figure 7.9** Isochronal plots of the complex modulus at 0.02 Hz for SBS modified bitumens
considerable evidence of extreme polymeric modification at high temperatures, with the establishment of a plateau region indicative of a dominant polymer network.

The phase angle isochrones in Figure 7.10 illustrate the improved elastic response (reduced phase angles) of the modified binders compared with their respective base bitumen. Whereas the phase angles of the base bitumen approaches 90°, indicating predominantly viscous behaviour, with increasing temperature the SBS polymer significantly improves the elastic response of the modified binders. This increase in elastic response at high temperatures can be attributed to the viscosity of the base bitumen being low enough to allow the elastic network of the polymer to influence the mechanical properties of the modified binders.

References
AASHTO (2010a) AASHTO M 320-10. Standard specification for performance-graded asphalt binder. AASHTO, Washington, DC, USA.
AASHTO (2010b) AASHTO MP 19-10. Standard specification for performance-graded asphalt binder using multiple stress creep recovery (MSCR) test. AASHTO, Washington, DC, USA.


Society for Testing and Materials, West Conshohocken, PA, USA. ASTM STP 212.


On the majority of roads, conventional bitumen grades possess satisfactory adhesion and mechanical properties for use in asphalt pavements for the vast majority of traffic and climatic conditions that are encountered. However, demands made on roads increase year by year and, in some cases, the limits of performance of conventional bitumens have been reached mainly due to

- increases in rainfall and temperature variations
- increases in axle weight, tyre pressures and freight movement, leading to a higher maintenance requirement and consequent increased costs for road owners
- a tendency to use thinner layers in pavements
- an ongoing demand for financial efficiency and cost savings to reduce the frequency of maintenance required in order to minimise disruption to traffic flow, and to provide an increased service life.

Thus, road failures happen earlier than expected, and maintenance downtime becomes more expensive for road owners. One means of overcoming these challenges is to modify bitumen with polymers or other additives to provide improved pavement performance. Modification technology depends on the specific applications and techniques (surface dressings, thin surface course systems, porous asphalt, surface courses for heavily trafficked roads, bases for heavily trafficked roads, anti-cracking membranes, bridge waterproofing layers, etc.). The degree of improvement required, and hence the cost, will depend on the particular needs of the site where treatment is proposed.

In this chapter, the main additives used for bitumen modification for road applications (Brown et al., 1990) are described. The polymers most commonly used are elastomers, plastomers, reclaimed tyre rubbers and, to a lesser extent, viscosity modifiers and reactive polymers. Special asphalts for road and industrial applications, developed through particular processes such as
the use of clear pigmentable binders, multigrade bitumens, cold application bituminous adhesives and a special bitumen additive for odour neutralisation are also discussed. Finally, basic properties of polymer modified bitumens (PMBs) and the related test methods, the current industry practice for manufacturing and handling PMBs, and some general information on the performance of PMBs for use on road applications are considered.

8.1 The role of bitumen modifiers in asphalt pavements

Traditional bitumens, derived from the distillation and/or blowing of crude oils, possess adequate performance characteristics, but increasing the high temperature performance can sometimes lessen the low temperature performance properties of the bitumen. The development of steps to counter this effect has been the incentive that has driven the early stages of the development of PMBs for use in roads. Modified binders are binders whose properties have been changed by the use of a chemical agent that, when added to the original bitumen, alters its chemical structure and physical and/or mechanical properties. As the bituminous binder is responsible for the visco-elastic behaviour of asphalts, it plays a large part in determining many aspects of road performance, particularly resistance to deformation and cracking, the two most common reasons for the structural failure of thicker pavements. In general, the proportion of any induced strain in asphalt that is attributable to viscous flow (i.e. non-recoverable flow) increases with both the loading time and the temperature. One of the prime roles of many bitumen modifiers is to increase the resistance of the asphalt to deformation at high road temperatures without adversely affecting the properties of the bitumen or asphalt at other temperatures. This is achieved by one of the two following methods, both of which result in a reduction in the permanent strain. The first approach is to stiffen the bitumen so that the total visco-elastic response of the asphalt is reduced. The second approach is to increase the elastic component of the bitumen, thereby reducing the viscous component. Increasing the stiffness of the bitumen is also likely to increase the dynamic stiffness of the asphalt. This will improve the load spreading capability of the material, increase the structural strength and lengthen the expected service life of the pavement. Alternatively, it may be possible to achieve the same structural strength but with a thinner layer. Increasing the elastic component of the bitumen will improve the flexibility of the asphalt. This is important where high tensile strains are induced.

For many years, researchers and chemists have experimented with modified bitumens, mainly for industrial uses, adding asbestos, special fillers, mineral fibres, plastomers, thermoplastic elastomers and rubbers. In the 1970s, the development of PMBs (Jiqing et al., 2014; Taylor and Airey, 2008; Yetkin, 2007) and the use of additives for bitumen by controlled industrial
processes in Europe was very much linked to the development of new mixture designs for asphalts such as thin surface course systems, which provide improved performance. For instance, the highly gap graded asphalts containing modified binders or additives, particularly fibres (asbestos, organic or mineral fibres), made it possible to increase the maximum binder content without the danger of the asphalt fatting up (the migration of binder to the surface of the asphalt causing a reduction in surface texture), providing the asphalt with the required level of cohesion and waterproofing the substrate. The addition of fibres helped in the development of stone mastic asphalt in Germany, subsequently adopted in many other countries. Another application of modified binders is in the anti-cracking membranes and waterproofing layers on steel and concrete bridges, because of their ability to withstand the considerable movements that occur in the surfacing of this type of construction. Modified binders have also been considered for structural layers because they improve fatigue strength. In the case of hot rolled asphalt, it is recognised that some polymers can improve the embedment of precoated chippings into the asphalt, particularly when laying takes place in less favourable weather conditions. In the case of open textured asphalts (essentially porous asphalt), the most important objectives for the modified binder are to improve its cohesive strength, which assists the resistance to weathering and its adhesion to aggregate particles, and in turn reduces the risks of the binder being stripped by water. Furthermore, the use of modified binders or fibres (the two additives are often combined for porous asphalts with very high void contents of over 25%) improves the resistance to binder drainage during transportation and laying, by increasing the binder viscosity. It allows higher binder contents to be used, increasing the thickness of the binder film, thus making the asphalt more durable by delaying binder ageing.

The most common modifying agents are polymers. These are macromolecules in which the same group of atoms is repeated many times. These repeated groups can be formed from one or several different molecules (monomers). Most of the additives assessed for modifying bitumen are shown in Table 8.1. Bitumens modified by polymers often prove to be the most cost-effective alternative to conventional bitumen because they improve targeted aspects of the performance of roads, and the polymers employed to modify bitumens are readily available at reasonable cost. This has led to the development of a wide range of proprietary asphalts made with PMBs and a range of PMBs that can be added to generic asphalts. The advantages of using these products have been proved in service for decades, while complying with product standards in different countries.

For the modifier (polymer, additives) to be effective and for its use to be both practicable and economic, it must:
Table 8.1 Examples of additives used to modify bitumen (Jiqing et al., 2014; Taylor and Airey, 2008; Yetkin, 2007)

<table>
<thead>
<tr>
<th>Type of modifier</th>
<th>Examples</th>
<th>Abbreviation</th>
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<tbody>
<tr>
<td>Thermoplastic elastomers</td>
<td>Styrene–butadiene elastomer</td>
<td>SBE</td>
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<tr>
<td></td>
<td>Styrene–butadiene–styrene elastomer (linear or radial)</td>
<td>SBS</td>
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<tr>
<td></td>
<td>Styrene–butadiene rubber</td>
<td>SBR</td>
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<td></td>
<td>Styrene–isoprene–styrene elastomer</td>
<td>SIS</td>
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<td></td>
<td>Styrene–ethylene–butadiene–styrene elastomer</td>
<td>SEBS</td>
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<td></td>
<td>Ethylene–propylene–diene terpolymer</td>
<td>EPDM</td>
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<td></td>
<td>Isobutene–isoprene random copolymer</td>
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<td></td>
<td>Polyisobutene</td>
<td>PIB</td>
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<td></td>
<td>Polybutadiene</td>
<td>PBD</td>
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<td></td>
<td>Polyisoprene</td>
<td>PI</td>
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<tr>
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<td>Natural rubber</td>
<td>NR</td>
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<tr>
<td>Thermoplastic polymers</td>
<td>Ethylene–vinyl acetate</td>
<td>EVA</td>
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<tr>
<td></td>
<td>Ethylene–methyl acrylate</td>
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<tr>
<td></td>
<td>Ethylene–butyl acrylate</td>
<td>EBA</td>
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<td>Atactic polypropylene</td>
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<td>Acrylic resin</td>
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<td>Phenolic resin</td>
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<td>Sulfonic acid, sulfuric acid</td>
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<td>Carboxylic anhydrides or acid esters</td>
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<td>Dibenzoyl peroxide</td>
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<td>Silanes</td>
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<td>Organic or inorganic sulfides</td>
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<td>Alumino-magnesium silicate</td>
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<td>Organic amines</td>
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<td>Anti-oxidants</td>
<td>Amides</td>
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<td>Natural asphalts</td>
<td>Phenols</td>
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<td>Organo-zinc or organo-lead compounds</td>
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<td>Trinidad Lake Asphalt</td>
<td>TLA</td>
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<td>Gilsonite</td>
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<td>Rock asphalt</td>
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be readily available
be cost effective
blend with bitumen
resist degradation at asphalt mixing temperatures
improve resistance to flow at high road temperatures without making the bitumen too viscous at mixing and laying temperatures or too stiff or brittle at low road temperatures
improve binder cohesion or adhesion properties.

In addition, the modifier, when blended with bitumen, should
be capable of being processed using conventional equipment
maintain its premium properties during storage, transportation, application and in service
be physically and chemically stable during storage, transportation, application and in service
achieve a coating or spraying viscosity at normal application temperatures.

### 8.2 The modification of bitumen

#### 8.2.1 The modification of bitumen using thermoplastic elastomers

Thermoplastic elastomers include polyurethane, polyether–polyester copolymers and olefinic or styrenic block copolymers, but it is the styrenic block copolymers that have proved to have the greatest potential when blended with bitumen (Bonemazzi et al., 1996; Bull and Vonk, 1984; Lewandowski, 1994). Styrene block copolymers, also termed styrene elastomers, thermoplastic rubbers or thermoplastic elastomers, may be produced by a sequential operation of successive polymerisation of styrene and butadiene (resulting in styrene–butadiene elastomers, SBEs) or styrene and isoprene (resulting in styrene–isoprene elastomers). The linear copolymer (styrene–butadiene–styrene, SBS) consists of a central rubbery backbone chain joined at each
end to a polystyrene chain. Alternatively, a di-block precursor can be produced by successive polymerisation of styrene and a mid-block monomer, followed by reaction with a coupling agent. Thus, not only linear copolymers but also multi-armed SBEs can be produced. These polymers are often referred to as radial, star or branched copolymers. When added to bitumen, a radial SBE may result in a much higher viscosity and softening point when compared with a bitumen to which has been added the same level of linear copolymer (e.g. linear SBS or SB). This may make it more difficult to achieve good blending and dispersion of the radial polymer in a bitumen compared with a bitumen made with the linear block copolymers.

In general, SBEs derive their strength and elasticity from a physical cross-linking or entanglements of molecules: a cross-linking tri-dimensional network can be achieved by the association of the polystyrene end-blocks into separate domains to form semi-crystalline zones, providing the physical cross-links with the polybutadiene chains. Cross-links are formed at close to ambient temperatures but are broken at elevated temperatures, only to reform again on cooling, being then a three dimensional reversible network (Figure 8.1). SBEs are characterised by a glass transition point ($T_g$), and, as elastomers, exhibit increased tensile strength with elongation. They have the ability to revert to their initial condition after an applied load has been removed. It is the polystyrene end-blocks that impart strength to the polymer, and the rubber mid-block that gives the material its exceptional elasticity (Vonk and Gooswilligen, 1989). At temperatures above the glass transition point of polystyrene ($100^\circ$C), the polystyrene softens as the domains weaken, and will even dissociate under stress, thus allowing easy processing. On cooling, the domains re-associate, and the strength and elasticity are restored (i.e. the material is thermoplastic). Once this has occurred, the level of shear exerted on the swollen particles is critical if a satisfactory dispersion is to be achieved within a realistic blending time at the manufacturing plant.

Ultraviolet (UV) fluorescence microscopy is a technique relying on the use of UV light to cause polymer molecules (but not the bitumen) to fluoresce at a wavelength that can be detected by the eye. Thus, the polymer molecules appear as bright objects against a dark (bitumen) background. Figures 8.2 and 8.3 show examples of typical morphologies of SBE blended into a bituminous matrix (Oliver and Khoo, 2012), the yellow nodules being the SBE particles dispersed in the black bitumen matrix. The addition of a thermoplastic polymer to bitumen makes a significant change to its physical properties, even at fairly low concentrations (<10%). Medium or, preferably, high shear mixers are required to disperse thermoplastic elastomers adequately into the bitumen matrix. The mixing and manufacturing of PMB will be discussed later in this chapter. When the SBE is added to the hot bitumen, the
bitumen constituents immediately start to penetrate the SBE particles, causing the styrene domains of the polymer to become solvated (the styrene domain of the SBS polymer becomes soluble with specific bitumen species). For instance, SBE, while absorbing the oily maltenes in the bitumen, may swell
by as much as nine times its volume when used at high concentrations (above 4%), and, from a morphology viewpoint, is forming a ‘continuous polymer-rich phase’ (Figure 8.2) in the bitumen. At lower concentrations (around 2–3%), the SBE will be dispersed in the bitumen matrix, giving a more ‘bitumen-rich phase’ (Figure 8.3).

The modification of bitumen morphology results in an increase in the softening point of the blend with the incremental addition of SBE, leading to a

**Figure 8.2** UV fluorescence microscopy of 4.5% SBS blended with bitumen

**Figure 8.3** UV fluorescence microscopy of 3% SBS blended with bitumen
typical ‘S-shape’ curve as shown in Figure 8.4. The rate of increase in the softening point of the modified binder also depends on the bitumen type, the SBE content, the SBE chemical structure and the grades of the SBS polymer.

As discussed in Chapter 4, bitumens are complex systems that can be subdivided into groups of molecules, saturates, aromatics, resins and asphaltenes. Saturates and aromatics can be viewed as carriers for the ‘polar’ aromatics (i.e. the resins and asphaltenes). The polar aromatics are responsible for the visco-elastic properties of the bitumen at ambient temperatures. This is due to the association of the polar molecules that leads to large structures, which in some cases may result in the generation of three dimensional networks (i.e. ‘gel’-type bitumen). The degree to which this association takes place depends on the temperature, the molecular weight distribution, the concentration of the polar aromatics, and on the solvency power of the saturates and aromatics in the maltene phase. If the concentration and molecular weight of the asphaltenes is relatively low, the result will be a ‘sol’-type bitumen. The addition of thermoplastic elastomers with a molecular weight similar to or higher than asphaltenes disturbs the phase equilibrium: the polymer and the asphaltene will then ‘compete’ for the solvency power of the maltene phase, and, if insufficient maltenes are available, phase separation (also sometimes called de-mixing) between the polymer and the bitumen may occur. There are other parameters that have an effect on the compatibility of polymers with bitumen. These parameters may contribute

Figure 8.4 Typical ‘S-shape’ curve of SBS modified bitumen blends (softening point with increasing content of linear and radial SBSs)
to the instability of the system, depending on the handling conditions (e.g. the storage conditions of the PMBs such as time, temperature, efficiency of mixing system). The quality of the polymer dispersion in the bitumen phase is influenced by a number of factors:

- the difference in solubility parameters of the polymers and the maltene phase of the bitumen
- the amount and type of asphaltenes present in the bitumen
- the type and concentration of the polymer
- the manufacturing processes and handling conditions of the SBE modified binder (Whiteoak, 1990).

The complexities of the inter-relationships between these factors are shown in Figure 8.5. It can be seen that very high aromaticity levels of the bitumen constituents lead to a weakening of the polystyrene domains (of the thermoplastic rubber), causing low softening points and low flow resistance properties (Collins et al., 1991). At low aromaticity levels, however, insufficient polymer will be incorporated into the bitumen, which also leads to low flow resistance properties.

In this context, the molecular weight of the polymer directly relates to the solubility parameters (Nielsen and Charles, 2005; Redelius, 2000). It is thus easier to blend polymers of low molecular weight than those of high molecular weight. Solubility parameters can vary according to the SBS type and within different commercial grades of SBS. This will depend on the ratio

**Figure 8.5** Effect of aromaticity and asphaltene content on the stability of thermoplastic rubber (TR) bituminous blends (Whiteoak, 1990)
of polymer segments in the case of copolymer structure and on the chemical structure of the polymer (linear or branched, molecular weight and nature of the double bonds inside the butadiene chains). Suppliers of polymers can provide different grades of SBEs for example, giving different properties to the bitumen (Bulatovic et al., 2013; Jianga and Zhang, 2011; Zhang et al., 2014).

The stability of SBE modified bitumens during handling (storage, transport and delivery to the customer site) can be enhanced by adding chemical co-modifiers such as peroxides, polyphosphoric acid, carboxylic anhydrides or esters, sulfur or any cross-linking materials. Sulfur vulcanisation, as used commonly in the rubber industry since 1839, is currently the predominant technology to improve the stabilisation of SBE modified bitumen products in a variety of handling conditions. The mechanism of vulcanisation by sulfur is a fairly complex chemical reaction (Akiba and Hashim, 1997; Krejsa et al., 1937; Weng et al., 2002). It is strongly dependent on the bitumen type, the polymer structure, the polymer concentration and the processing conditions at the plant. The SBE particles, which are sheared and shattered to a very small size, are homogeneously dispersed in the bitumen matrix. There are competing kinetic reactions involving sulfur compounds, SBE chains and bitumen species. From the literature, it can be concluded that a reaction between bitumen and sulfur (S) occurs predominantly within the aromatic and naphthenic centres (Ar) of the bitumen species, leading to higher asphaltene/resin ratios and to the formation of mono or polysulfide bonding such as Ar–S–Ar, Ar–SS–Ar, Ar–SSS–Ar and Ar–SSSSS–Ar (Weng et al., 2001). Fine SBE particles are wrapped by some aromatic components in the bitumen through sulfide or polysulfide links (Gawel, 2000), creating polymer–(S)ₓ–bitumen chains links (Figure 8.6).

These sulfur bonds contribute to the enhancement of polymer compatibility with bitumen, and may also influence the rheological properties of the binder, depending on the cross-link density and bonding lengths. Sulfur reacts with the carbon atoms next to the carbon–carbon double bonds of polybutadiene chains in SBEs by substituting allylic hydrogen atoms with sulfur by a free radical mechanism (Akiba and Hashim, 1997; Gawel, 2000; Krejsa et al., 1937; Weng et al., 2001, 2002). Thus, storage-stable sulfur cross-linked SBE modified bitumen is generally obtained by creating an irreversible three dimensional network, although the reactions between sulfur and SBE modified bitumens have not been studied extensively in the literature.

8.2.2 The modification of bitumen using plastomers

Some plastomers are used in the bitumen industry as an alternative to elastomers for road paving applications. Polyethylene, polypropylene, polyvinyl
chloride, polystyrene and ethylene–vinyl acetate (EVA) copolymer are the main non-rubber thermoplastic polymers that have been examined in recent decades. As thermoplastic polymers, they are characterised by softening on heating and hardening on cooling. These polymers tend to influence the penetration more than the softening point properties when added to bitumen, which is the opposite tendency of thermoplastic elastomers. Polyolefins have been used for the modification of bitumen due to their relatively low cost and the benefits that they induce. The light components of bitumen are usually absorbed by the polyolefins. A bi-phasic morphological structure is formed (i.e. the polyolefin is dispersed in the bitumen matrix). As the concentration of the polymer increases, phase inversion phenomena occur, leading to a

![Figure 8.6](image_url) Representation of SBS–S–SBS bonding in bitumen
polymer matrix that can be detected in higher concentration polyolefin modified bitumen formulations. In addition to this, the regular and long chains of polyolefin materials have a tendency to pack closely and crystallise, which leads to a lack of interaction between the bitumen and the polyolefin and causes instability of the modified bitumen. The compatibility of polyolefins with bitumen is usually found to be very poor because of the non-polar nature of the polyolefin materials (Fawcett and McNally, 2000; Yousefi, 2003). Adding polyolefins to bitumen does not significantly increase its elastic properties, and, when heated, the polyolefin can separate, which may give rise to a coarse dispersion on cooling. PMBs obtained with olefinic polymers are used, but the storage stability of the system remains an unresolved issue. A possible solution is the use of polyethylene based copolymers, in which the co-monomer is polar, either inert or reactive with respect to the bitumen species. Typical examples are EVA and ethylene–butyl acrylate random copolymers (Gonzalez et al., 2004; Isacsson and Lu, 1999), while maleic anhydride and glycidyl methacrylate groups being grafted to polyethylene chains or being incorporated by a copolymerisation process to polyethylene could interact with the reactive functions of bitumen species, leading to better compatibility of the polyolefin with the bitumen.

EVA copolymers, as thermoplastic materials, are commonly used for modifying bitumen. EVA has a random structure produced by the copolymerisation of ethylene and vinyl acetate. EVA copolymers with a low vinyl acetate content possess properties similar to low density polyethylene. As the level of vinyl acetate increases, the properties of the copolymer alter, which may induce changes in the properties of the bitumen. The properties of EVA copolymers are classified by molecular weight and vinyl acetate content as follows.

- **Molecular weight.** Standard practice for EVAs is to measure the melt flow index (MFI), a viscosity test that is inversely related to the polymer molecular weight: the higher the MFI, the lower the molecular weight and viscosity. This is analogous to the penetration test for bitumen: the higher the penetration, the lower the average molecular weight and viscosity of the bitumen.

- **Vinyl acetate content.** Regular polyethylene segments of the EVA chain pack closely together and form so-called ‘crystalline’ regions, and can be represented graphically, showing the main effects of vinyl acetate species on the properties of a bitumen (Figure 8.7). At the same time, the bulky vinyl acetate groups disrupt this closely packed arrangement to give ‘non-crystalline’ or ‘amorphous rubbery’ regions. The crystalline regions are relatively stiff, and have a considerable reinforcing effect whereas the amorphous regions are rubbery. Obviously, the more vinyl
acetate groups present (or the higher the vinyl acetate content), the higher the proportion of rubbery regions will be, and, conversely, the lower the proportion of crystalline regions.

A wide range of EVA copolymers is available, specified by both the MFI and the vinyl acetate content. EVA copolymers are easily dispersed and have good compatibility with bitumen. They are thermally stable at the temperatures at which asphalt is normally mixed. However, during static storage, some separation between the polymer and bitumen may occur, and it is therefore recommended that the blended product should be thoroughly mixed before use.

**8.2.3 The modification of bitumen by the addition of rubbers**

Polybutadiene, polyisoprene, natural rubber, butyl rubber, chloroprene and random styrene–butadiene rubber, among other rubbers, have all been used to modify bitumen: their effect is mainly to increase its viscosity. Rubbers are also used in a vulcanised (cross-linked) state. For example, particles of crumb rubber (CR) reclaimed from old scrap tyres, are used to modify and improve the performance of conventional bitumen while providing an outlet for the disposal of waste tyres (Gillen, 2007; Oliver, 1999; WRAP, 2008). This technology began in the 1960s, and there are two main types of process, described as ‘wet’ and ‘dry’. In the ‘wet’ process, the CR modifier is added to the bitumen while heating to 180–210°C, prior to the manufacture of the hot asphalt. In the ‘dry’ process, the CR modifier is mixed directly into the
conventionally prepared asphalt at the paddle mixer. The CR modifier acts as an ‘elastic mineral aggregate’, and replaces part of the aggregate compound. The description of the processes for manufacturing CR modified bitumen (CRMB) and its applications are detailed in section 22.5.

Typical concentrations of CR particles in bitumen are between 8 and 15%. High CR particle content (above 20%) allows the formation of a physical network that partially prevents the sedimentation of the particles. In all cases, when dispersed in bitumen, these crumb particles strongly influence its rheology and increase its viscosity, which may sometimes be detrimental to its workability. A high temperature is necessary (180–195°C), which unfortunately results in the generation of odours and emissions. In addition, storing the CRMB at high temperature may accelerate the process of particle sedimentation to the bottom of the tank or truck, which results in inconsistent supply quality. Accordingly, it is usually recommended that the CRMB blends are continuously agitated during storage to maintain the stability of the product, or even mixed immediately with aggregates. The preparation of these blends directly at the asphalt plant is also said to be advantageous. Chemical compatibility and processing conditions, such as time, temperature and shear, are important in obtaining the desired properties and stable binder product as per specification (Pérez-Lepe et al., 2003). Successful developments were made and patent applications on this were filed by Shell Bitumen (Shell Mexphalte RM®) to improve the homogeneity and the stability of CRMB during transportation to and storage at the asphalt plant. Other advances have resulted in limiting the maximum mixing and laying temperature to 170°C, thus avoiding the generation of odours and emissions from the rubber.

8.2.4 The modification of bitumen by the addition of viscosity modifiers

As bitumen is responsible for the visco-elastic behaviour characteristic of asphalt, it plays a large part in determining many aspects of road performance, particularly resistance to deformation (Edwards, 2005) and resistance to cracking (Prowell, 2005; Suleiman, 2011), the two most common structural failure modes in roads. (Structural failure is usually interpreted as requiring the replacement of the surface course, binder course and the base of the pavement.)

A wax additive is used to modify the binder, which itself is used subsequently to lower the mixing and compaction temperatures in asphalts. Several performance properties of asphalts can be improved by adding low molecular weight polyethylene, synthetic wax obtained from a Fischer–Tropsch synthesis or paraffin wax to bitumen. Fischer–Tropsch synthetic waxes have a microcrystalline structure, and consist of particles with a large
number of carbon atoms, up to 100. Due to the morphology of waxes, they can be used to lower the viscosity of conventional bitumen at higher temperatures (typically above 120°C) and increase the softening point. In relation to the modification of viscosity, the following points should be noted.

- At temperatures above the melting point of the wax additive, the viscosity of the bitumen is significantly reduced. This, in turn, allows for a reduction in asphalt mixing and laying temperatures by up to 30°C, thereby saving energy and reducing fume emissions. Also, critical applications, where hand laying may be required or material has to be laid at low ambient temperatures, may be facilitated as a result of the longer time period available between asphalt mixing and the time when compaction is no longer possible.

- At the crystallisation temperature of the additive, the viscosity of the modified bitumen rises sharply, which, in turn, may lead to higher asphalt stiffness, thereby reducing the permanent deformation characteristics of the carriageway. However, the low temperature properties may be adversely affected.

Figure 8.8 shows typical viscosity–temperature relationships for an unmodified penetration grade bitumen and the same bitumen modified with a selected paraffin wax content. The magnitude and nature of the effect on bitumen rheology depends on the bitumen itself as well as on the type and quantity of additive. Bitumen composition is of critical importance, but there are many additives that can change the visco-elastic properties of bitumen: examples are natural bitumens such as rock asphalt, which harden bitumen, and paraffinic flux oils or aromatics, which soften bitumen.

![Figure 8.8](image-url)
8.2.5 The modification of bitumen by reactive chemistry

In addition to physical blends of bitumens and polymers, another way to improve the binder properties is through chemical modification: reactive ethylene terpolymers, comprising ethylene, ester groups of methyl, ethyl or butyl acrylate and glycidyl methacrylate groups (also known as reactive epoxy functions), can chemically react with bitumen species (carboxylic groups present in asphaltenes) and can be used to enhance asphalt performance and increase the compatibility between the polymer backbone and the bitumen, keeping the product stable during storage and transportation (Dupont, 2014; Kanabar, 2010; Keyf et al., 2007). There are many other reactive polymeric materials described in the literature or in patent publications that can be used to enhance the properties of bitumen, examples being grafted maleic anhydride styrene block copolymers, polymers with high vinyl content, and hydroxyl, carboxylic or silane-grafted polyolefins (Chaverot et al., 2012; Cong et al., 2011; Crews and Kalinowski, 1954). As this involves various chemical reactions between the reactive functions of the polymer and bitumen species, in certain instances some catalysts such as phosphoric acid (PA) can be used. However, in such circumstances, processing conditions (mixing time, temperature, polymer content) should be carefully defined to avoid uncontrolled kinetic reactions and gelling of the reacted product. In the case of reactive ethylene terpolymers, PA can be used to catalyse the reactivity of glycidyl methacrylate (epoxy) groups with bitumens that have carboxylic reactive functions.

Among the different acids that can be used in bitumen, polyphosphoric acid (PPA) added at a low content (<2%) to improve asphalt binder properties through chemical modification occurs in numerous patented technologies described in articles and patent publications that appeared in the USA in the 1990s and 2000s. In the main, these lead to increased asphalt stiffness without deterioration of its low temperature properties (Baumgardner et al., 2005; Masson, 2008), greater adhesion to aggregates and an increase in the performance grade range by one to two classification grades (e.g. an increase in the grade of the binder from PG70 to PG76 for the high temperature range) to meet Superpave – see also section 5.7. (Superpave specifications are commonly used in the USA – see section 12.4.5). PPA is different from orthophosphoric acid, with no free water, and is defined as an inorganic polymer modifier having different distributions of chain lengths, with the number of repeating units varying from one chain to another. The modification of bitumen with PPA appears to be a complex physico-chemical process, and the resulting properties may be strongly dependent on the nature of the bitumen. The investigation of the reactions occurring between the bitumen species and PPA is difficult due to the large number of molecules with different chemical structures and their possible interactions. Many
mechanisms have been proposed in the literature to explain the increase in bitumen stiffness by PPA addition, for instance a model is proposed based on the reactivity of asphaltenes with PPA molecules, through an acid–base reaction mechanism and esterification reactions between those molecules (Orange et al., 2004). These reactions may cause a de-agglomeration of the aggregates in the asphaltenes, which improves bitumen rheology. In combination with a polymer or a rubber, a positive effect also seems to occur: adding very small amounts of PPA jointly with SBS or CR to bitumen with a suitable formulation is used to improve the handling and performance of PMBs (TRB, 2009; Zhang and Hu, 2013), achieving higher Superpave performance grades while improving the mixing and compaction characteristics. (Superpave performance grades are commonly used in the USA. They apply to the performance of asphalts, and are based on the proposition that the properties of a bitumen should be related to the conditions under which it is to be used.)

8.3 Other modified products used for road and industrial applications

8.3.1 Multigrade bitumens

If it was possible to create the ideal pavement binder, it would have a uniform designed ‘stiffness’ and ‘flow’ behaviour across the operational temperature range to combat both softness/deformation and fatigue/brittleness. Multigrade bitumens are a step in this direction, as they are designed to be less temperature susceptible than penetration grade bitumens, resulting in improved performance at high temperatures and better low temperature characteristics than hard grades, making them ideal for use in surface or binder courses. Multigrade bitumens are generally manufactured through a unique processing route that involves catalytic oxidation using PA. This process makes the product perform as both

- a soft bitumen at low temperatures
- a hard, stiffer bitumen at higher temperatures, but not so stiff that premature cracking occurs at low temperatures.

Multigrade bitumens are specified in national/international bitumen standards such as the Australian AS 2008 standard (AS, 2008) and the European EN 13924-2 standard (BSI, 2014), to provide a framework for specifying the properties and relevant test methods for multigrade bituminous binders.

8.3.1.1 Shell Multiphalte®

In the 1990s, Shell developed and introduced to the market a range of proprietary multigrade bitumens called Multiphalte after more than 10 years of laboratory research and field trial experience (Koole et al., 1991).
These products are part of a well established technology with an improved viscosity–temperature relationship designed to enhance stiffness and viscosity at high service temperatures, with the aim of substantially improving deformation resistance. At low temperatures they have lower stiffness, and are therefore less likely to crack.

This can be seen, for example, when a Multiphalte bitumen is compared with a conventional penetration grade bitumen on the bitumen test data chart (Figure 8.9). The top left corner of the chart shows that at low temperatures Multiphalte is less susceptible to low temperature cracking than conventional bitumen, meaning that as a component in an asphalt it is less brittle at low winter temperatures. The bottom right corner shows that, for the same high temperature, Multiphalte has a higher viscosity than conventional bitumen, meaning that it improves the resistance of pavements to deformation and reduces stripping during high summer road temperatures.

Shell Multiphalte is produced and used in asphalt applications (sealing, surface or binder course applications) worldwide (Europe, Argentina, Australia, etc.), typically in the pen range 20/30, 35/50 and 45/60, although some grades can be tailor-made to suit specific market requirements and countries. In France, the high modulus asphalt version (Multiphalte HM ‘High Modulus’ with a 20/30 penetration range) is available to satisfy certain French asphalt specifications. There are two Multiphalte grades in Australia: the M1000 grade is designed to resist rutting, and is often used on airport runways, while the M500 grade is more commonly employed in sprayed sealing applications.

Figure 8.9 Bitumen test data chart illustrating the properties of a Multiphalte bitumen compared with a conventional bitumen.
Figure 8.10 illustrates the concept of Shell’s Multiphalte bitumen, showing the improved resistance to deformation of the M1000 grade compared with reference asphalts made with C320 bitumen (40/60 pen) and a highly modified PMB (A15E), using the wheel track tester. The wheel tracking machine measured the rut depth generated by a loaded wheel at 60°C over 10,000 passes, following the Australian reference standard AG:PT/T231 (AS, 2006).

Asphalts manufactured with Multiphalte can be mixed, transported, laid and rolled with minimal changes to traditional and well established procedures. The most critical difference is that it is somewhat more viscous than conventional bitumen, and consequently needs to be mixed and laid at a temperature 10°C higher than normal. All other paving operations remain within the normal range.

8.3.2 Synthetic pigmentable binders

The different methods of achieving a coloured asphalt surfacing are described in the literature (Pierard et al., 2013). One important way of producing a coloured surface is to use a pigmentable binder. The most common pigmented asphalt is red, which is produced by substituting ferrous oxide for the same proportion of fines (usually 0.5–6%, depending on the purity of the ferrous oxide), which has a size <0.063 mm. Appropriately coloured aggregates can be used to ensure that the overall appearance of the material is
maintained when the aggregate is exposed after trafficking. The main drawbacks to colouring mixtures using conventional bitumens are that

- the only acceptable colour that can be achieved is a fairly dark red
- the quantity of the expensive ferrous oxide required to achieve an acceptable red is fairly high, which significantly increases the cost of the mixture.

There are many applications for coloured asphalts, as illustrated in Figures 8.11–8.13, including architectural applications and use in historical monuments, parks and squares, sports venues and stadiums, boardwalks and promenades (bicycle and pedestrian lanes), office buildings, car parks, bus lanes, speed regulation, crossing paths in roundabouts, school exits, tunnels, multifunctional lanes (Genardini, 1994; Gustafsson, 1988; Lohan, 1999; O’Connor, 1999).

8.3.2.1 Shell Mexphalte C®

To enable asphalt to be pigmented in colours other than red, Shell developed a range of synthetic clear binders (Mexphalte C) for various applications that contain no asphaltenes (Le Coroller and Herment, 1989; Schellekens and Korenstra, 1987). These can be pigmented as required, to produce a broad spectrum of coloured asphalts enabling pavements to be matched to the local environment. Mexphalte C products possess rheological and

Figure 8.11 Tunnel de Lorentweiler, Luxembourg
Figure 8.12 Bus lanes in Tours, France

Figure 8.13 Parking area in Orange, France
mechanical properties similar to penetration grade bitumen, and the temperature at which the synthetic binder is mixed with aggregates is usually lower than the corresponding penetration grades. Coloured asphalts manufactured using Mexphalte C products require around 0.5–1% of pigment to achieve a satisfactory colour, whereas only red can be achieved with penetration grade bitumen using up to 6% ferrous oxide in the mixture. Inorganic pigments can be mixed alone or in combination with a synthetic binder, to give a range of colours, examples being:

- ferrous oxide for red, yellow, brown and black
- titanium dioxide for white
- chromium oxide for green
- cobalt oxide or cobalt aluminate for blue.

A range of Mexphalte C products is available for multiple applications.

- Mexphalte C 70/100 is a clear synthetic binder. A clear cationic synthetic emulsion modified with polymers can also be formulated and used as a slurry seal and for surface dressing applications (light traffic only).
- Synthetic binder modified with polymers: Mexphalte C LT requires lower asphalt mixing and paving temperatures, Mexphalte C P2 is used for heavy traffic and Mexphalte C 35/50 is used for special applications, but note that it has to be handled at high temperatures.
- Mexphalte C P3j is a mastic joint solution that combines the durability of a PMB and a clear synthetic binder.

**8.3.3 Modified binders for industrial applications**

For industrial applications, Shell has developed a wide range of high performance PMB products used in Europe and Asia, under the brand names of Shell Tixophalte®, Shell Caritop® and Shell Flintkote®.

**8.3.3.1 Shell Tixophalte®**

The first development of a cold applicable bituminous adhesive with improved wet or tack properties under the brand name of Tixophalte was undertaken in Shell’s R&D laboratory in The Netherlands. Tixophalte was first trialled in 1981, and this product has since been reformulated several times to comply with health and safety regulations, with new variants being developed for specific applications – revisions include the bitumen, the polymer, additives, the mineral filler, and use of a non-chlorinated and non-toxic solvent (Deygout and Seive, 2004).

The current formulation of Tixophalte is a ready to use bituminous mastic that can be applied as a filler, joint sealer or adhesive for any job, from large
scale industrial applications to domestic situations. Its waterproofing properties mean it can be used underwater. It is easy to apply, and offers long lasting protection and waterproofing. It is capable of withstanding movement, weathering, chemicals and other environmental factors.

The main uses for this material are

- in building applications (sealing leaks and making emergency repairs even on wet surfaces, gluing isolation panels onto metal or concrete surfaces, fixing roofing felt overlaps and many other roofing uses, plumbing and waterproofing tasks, sealing joints/connections in chimneys, guttering, skylights and water evacuations)
- in infrastructure applications (sealing joints and drainage systems in concrete, drainage channels and concrete bridge decks, sealing induction loops, sealing between tram rails and asphalt)
- in hydraulic applications (for waterproofing cracks and joints in waterways, water canals, reservoirs, dams and bridges).

8.3.3.2 Shell Caritop®

Caritop is a range of polymer modified binders designed and manufactured by Shell specifically for use in the industrial market. Examples of its use include

- the manufacture of carpet tiles (lower mixing temperature during the manufacturing process and an increased filler content or a wider use of filler types, and improved mechanical performance of the finished tile)
- the production of roofing felts (improved resistance to high and low temperatures, improved cold bend and adhesion properties, and reduced risk of damage during installation)
- the production of mastic flooring (improved high temperature stability and low temperature flexibility, increased tolerance to building movement, better durability and increased fatigue resistance).

8.3.3.3 Shell Flintkote®

From its conception over 50 years ago as a cost effective and reliable waterproof coating, the Flintkote product range has expanded to create an integrated portfolio of products (Shell Flintkote Colourflex for exposed waterproofing and decoration) that are designed to work together, offering a seamless barrier to keep water out. Their wide range of uses includes waterproofing wet areas (e.g. bathrooms, kitchens and toilets), waterproofing metal roofs, waterproofing blockwork and concrete structures, waterproofing mastic flooring, waterproofing flat concrete roofs and mastic asphalt roofs, the treatment of water tanks and the protection of steelwork.
Bitumen and asphalt operations are facing increasing public concern about odour emissions, especially for production facilities located in urban areas. Bitumen and asphalt producers are seeking appropriate solutions in order to resolve disputes, maintain their image and ensure the long term viability of their production sites. These complaints can necessitate significant investment to treat expelled gases, and may even threaten the continued operation of the asphalt plant. Although treatments of expelled gases are often effective in reducing concentrations of volatile organic compounds, these systems require modification of the facilities, making their use rather costly. It is for this reason that Shell undertook intensive research in early 2006 in collaboration with a partner that specialised in the treatment of odours. This led to the joint development of the Shell Bitufresh additive that can be introduced into hot bitumen with ease and is designed to minimise bitumen odour conditions effectively for both workers during asphalt mixing and laying operations, and also to reduce odour nuisance for local residents in areas where bitumen is being used (SIA, 2008). The mechanism employed by this product is fundamentally different from other bitumen additives that work by masking the smell. Bitufresh has proven to be effective at low dosage levels because of its well balanced mixture of neutralising components (around 40–60 ppm in bitumen, as depicted in Figure 8.14). This has been supported by both laboratory and field trials undertaken with external partners. After treatment, the characteristics of the bitumen remain unchanged and bitumen odour neutralisation continues for a period of at least 2 weeks. A patent application for this technology was filed in 2007.

Figure 8.14 Effectiveness of Bitufresh in reducing bitumen fume odour. (From an external evaluation performed by Science Industry Australia (SIA))
8.4 Properties of PMBs and related test methods

PMBs used in road applications are normally tested using traditional bitumen tests, investigating, for example, penetration, softening point, Fraass breaking point and viscosity, and properties after ageing in a rolling thin-film oven and a pressure ageing vessel. In addition, there are often specific tests, such as for storage stability, elastic recovery, force ductility, and more complex rheological characterisation using the dynamic shear rheometer and the bending beam rheometer. The methods used for such tests may follow EN (for Europe) or ASTM (for USA, Asia) standards, depending on the country in which the PMBs are produced and sold.

Phase separation or incompatibility between the polymer and bitumen components can be demonstrated by a simple hot storage test in which a sample of the PMB is placed in a cylindrical tube and stored vertically at elevated temperature, usually between 160 and 180°C, in an oven, typically for 2 or 3 days or longer, depending on the test method or specification. At the end of the storage period, the top and bottom of the sample are separated and tested. Incompatibility is usually assessed by the difference in softening point between the top and bottom samples – if the difference is less than a range of 2–5°C, the binder is usually considered to be storage stable, depending on local specifications. The storage stability test is usually carried out as per ASTM D7173 (ASTM, 2011) or EN 13399 (BSI, 2011).

The main factors influencing storage stability and binder properties are:

- the amount and molecular weight of the asphaltenes and the aromaticity of the maltene phase
- the amount of polymer present and its molecular weight and structure
- the storage conditions (temperature, time, mixing systems).

Shell Bitumen has developed a number of compatible and stable SBE-modified binders, obtained from sulfur cross-linking technology or reactive chemistry, for a variety of surfacing applications, including deformation resistant hot rolled asphalt, stone mastic asphalt and thin surface course systems. The philosophy of producing a compatible system is based on the practicalities and user friendliness of the bitumen to facilitate the medium and long term storage that is often required. The effect of compatibility on storage stability can be seen in Figure 8.15, which shows the numerical difference in the softening point of the top and bottom samples of two bitumens containing 7% SBS after 1, 3, 5 and 7 days’ storage at 140°C. The results clearly show that the compatible system is extremely stable whereas the incompatible system has separated dramatically after 7 days’ storage. The implications of this for storage are obvious. In practice, it is possible to use PMBs that are not storage stable. However, they must be handled with great care, and
The elastic recovery test is a useful method for assessing whether a polymeric material that has been added to a bitumen provides a significant elastomeric characteristic. In this test, the PMB specimen is pulled to a specified distance at a specified speed and at a specified temperature (e.g. 25 or 10°C), then cut into two parts as seen in Figure 8.16, and the percentage of recoverable strain from the elongated specimen is measured after a fixed time period. The test methods used are ASTM D6084 (ASTM, 2013) and EN 13398 (BSI, 2010a).

The property of cohesion allows PMBs to be differentiated from unmodified paving grade bitumens by several test methods: force ductility, the tensile test and the Vialit pendulum reflect different characteristics of PMBs, and are not considered to be equivalent. It is intended that the most appropriate cohesion test method should be used in each case, at the discretion of the supplier. Ductility is that property of bitumen that permits it to undergo substantial deformation or elongation. The test method describes the procedure for determining the ductility of an asphalt measured by the distance by which it will

**Figure 8.15** Effect of the compatibility of the polymer in the bituminous matrix on the storage stability test result

stored in tanks with stirrers or extended circulation to prevent separation of the polymer.
elongate before breaking, as seen in Figure 8.17. Unless otherwise specified, the test can be undertaken at ambient (25°C) or lower temperatures (5°C) and at a speed of 5 cm/min. At other temperatures, the speed should be specified. The testing can be carried out as per ASTM D113 (ASTM,
2007), EN 13589 (BSI, 2008), EN 13703 (BSI, 2003) or a method specified in a particular country. Meeting the ductility specification is one of the major criteria for PMBs in Asia and China, and force ductility is mostly applied in Europe as per the PMB product specification EN 14023 (BSI, 2010b) using various temperatures and traction speeds. To quantify this effect, the force required to stretch the asphalt sample is recorded during the measurement phase. The area below the resulting force–distance function can be calculated, and represents the cohesive energy.

8.5 Manufacture, storage and handling of PMBs

The manufacture of PMBs can be undertaken in either fixed or mobile plants. The basic scheme, however, remains the same, although the utilities available may vary according to what is available locally. Mobile plants are more dependent on associated sites for utilities, which may include hot oil, steam and electricity. These plants are mainly used at large construction sites for short periods. Fixed plants, however, are independent depots or associated with refineries, and are intended to provide continuous long term production; utilities are provided on a permanent basis at such plants. Several companies specialise in constructing complete PMB plants.

PMBs generally consist of a blend of a bitumen and a polymer, usually an SBE. In some cases, other ingredients are added to the base bitumen, to assist in blending and achieving particular properties. In addition to the normal safety, quality and environmental considerations involved in handling hot bitumen, the manufacture of PMBs requires:

- accurate blending of materials and control of manufacture, to ensure that the products conform to target specifications
- monitoring and control of the temperature at all phases of manufacture, storage, transport and field usage, to avoid premature deterioration of polymers at high temperatures
- maintaining the homogeneity of some PMB mixtures that may segregate in storage
- avoiding contamination with other products that may alter the performance characteristics of the end product.

A PMB manufacturing processes can be described in four main steps:

1. base bitumen is heated to the required temperature of 180–190°C and fed to the tank
2. fluxing, where a suitable low viscosity non-volatile component is added to the tank (this stage is optional, and may be required for correction of the properties of the final product, or improving the compatibility between the polymer and the bitumen)
3 polymer addition and dissolution into bitumen with a high shear mill (using polymer pellets/beads) with simple or multiple passes, and/or in low shear mixing (using the polymer in a powdered form) within the temperature range 180–190°C.

4 conditioning of the PMB blend under low shear mixing for a few hours, thus allowing the polymer to swell into the bitumen and ensuring a homogeneous mixture.

Batch or continuous processes can be used for the manufacture of PMBs. Prior to the mill, there is almost always a pre-mix tank to permit charging of the polymer into the bitumen, after which the combination can be passed through the mill system to the conditioning tank. Polymer pellets are sheared in the mill and mixed into the base bitumen. This mixture is then permitted to swell for a certain period of time, thus allowing the polymer to disperse fully. When the PMB product is homogeneous, it is ready for use.

The master batch process involves the production of a PMB concentrate using a high dosage of polymer. Usually this takes longer to manufacture, but the PMB concentrate can be diluted with conventional bitumen to produce the end grade required with a lower polymer content.

There are several considerations relating to the storage of PMBs. It is important to remember that both the handling of the material and the storage temperature are critical if the quality of the product is to be maintained. The tanks for storage of the finished PMB are designed to minimise deterioration of the product in storage, including strict control of temperature, a minimal surface area to reduce the potential for oxidation, and mixing or circulating equipment to ensure that the product remains homogeneous during the storage period. Shell Bitumen has handling guidelines for PMBs, with different temperature requirements depending on the length of storage for the Shell Cariphalte® PMBs. Storage recommendations for Cariphalte are usually as follows:

- for short term storage (3–4 days), the recommended temperature is around 165–175°C
- for long term storage (up to 2 weeks), the recommended temperature is below 140–160°C
- for very long term storage (more than 2 weeks), the recommended temperature is below 100°C.

For truck loading of the PMB in the loading gantry, all supply lines throughout the PMB plant, including the loading equipment, are designed and procedures established so as to avoid any contamination during a change in product or cleaning of the supply lines. The use of oils such as kerosene,
diesel or gas oil for flushing lines must be avoided. Where lines need to be flushed, it is usually done with hot bitumen or a finished product.

8.6 Performance of PMBs for road application: Shell Cariphalte® example

PMBs provide improved balance between low and high temperature properties, such as high temperature stiffness (Figure 8.18) and low temperature flexibility. For instance, a pavement needs to be resistant to deformation at high temperatures during summer and to cracking during winter when the temperature of the pavement can fall to negative values. So, it needs both elastic and viscous components, and each needs to be dominant in the different temperature domains. In addition, increasingly aggressive traffic conditions require higher performance binders, economic considerations such as the desire for thinner pavements are the impetus for the production of premium binders, and quality issues are behind the desire for improved cohesion and adhesion properties in all pavements.

Shell PMB products are sold to customers under the brand name Cariphalte, providing solutions for different types of roads and meeting different customer needs and applications. A few examples of the uses of Cariphalte PMBs are given below:

![Rut depth vs. Number of passes](image)

**Figure 8.18** Anti-rutting performance results from the Hamburg wheel tracking test performed under warm water at 50°C (stone mastic asphalt 0/11 S granular curve using 35/50 bitumen and 2.5% and 7% SBS modified bitumens)
- in surface courses such as thin surface course systems and those made with stone mastic asphalt
- in drainage (porous) asphalt applications
- for an increased level of fuel resistance (Cariphalte Fuelsafe, specially designed to make asphalt surfaces last longer when exposed to the potential damage caused by spilt and leaking fuel)
- for motor racing circuits and test tracks (Cariphalte Racetrack®)
- for asphalts designed to possess high resistance to deformation and high flexibility, especially in binder courses
- for high performance road applications in combination with reclaimed asphalt pavement (Cariphalte RC)
- as the absorbing layer in pavements subject to reflection crack stresses (Cariphalte SAMI).

References


Kanabar N (2010) Comparison of Ethylene Terpolymer, Styrene Butadiene, and Polyphosphoric Acid Type Modifiers for Asphalt Cement. MA thesis, Department of Chemistry, Queen’s University, Kingston, ON, Canada.


Polymer modified bitumens and other modified binders


Whatever the end use, application conditions usually require bitumen to behave as a mobile liquid. In principle, there are three ways to make a highly viscous bitumen into a low viscosity liquid:

- heat it
- dissolve it in solvents
- emulsify it.

Bitumen emulsions provide a convenient and environmentally friendly option in which the bitumen is liquefied in small particles by dispersing it in water.

Currently, bitumen emulsions are used for road maintenance and repairs (e.g. surface dressing and slurry seals) and for parts of structural pavements (tack coats, bond coats, prime coat, cold mixes, cold recycling and gravel emulsion).

A phase in chemistry is defined as ‘a mechanically separate, homogeneous part of a heterogeneous system’ (Infoplease, 2014). Bitumen is not mixed into water, it is held in suspension, so bitumen and water are part of a two phase system. They work together in the emulsion (the heterogeneous system) in a special way that maintains the homogeneous nature of each ingredient.

An emulsion is defined by the International Union of Pure and Applied Chemistry as:

A fluid colloidal system in which liquid droplets are dispersed in a liquid. An emulsion is denoted by the symbol O/W if the continuous phase is an aqueous solution and W/O if the continuous phase is an organic liquid (an ‘oil’). More complicated emulsions such as O/W/O are also possible.
In the particular context of bituminous binders, an emulsion is further defined in EN 12597 (BSI, 2014) as ‘an emulsion in which the dispersed phase is bituminous’ (i.e. O/W) and which is ‘thermodynamically meta-stable’.

The inherent thermodynamic instability of all emulsions, which is caused by a relatively high interfacial tension between the two immiscible oil and water phases, leads to a requirement for energy input to form an increased interfacial area between them (i.e. to form dispersed-phase droplets). This energy requirement can be reduced by the addition of surface-active agents (surfactants) to the system that adsorbs at the phase interface and lower interfacial tension. Surfactants, steric stabilisers (some polymers) or certain clay minerals can also kinetically stabilise an emulsion against re-coalescence.

In bitumen emulsions, the surfactants and any other additives required are usually solubilised prior to the manufacture of the emulsion in the water phase. In general, surfactants that are relatively more soluble in water (hydrophilic) rather than oil (lipophilic) tend to promote the formation of O/W rather than W/O emulsions. In the case of the former, the water phase may then also be referred to synonymously as the aqueous or continuous phase of the emulsion.

By far the most common surfactants used in bitumen emulsion production are cationic in nature such as alkyl diamines, amidoimidazolines and quaternary ammonium chlorides, although anionic, non-ionic and zwitterionic surfactants and clay minerals are used for certain specialised applications. The typical bitumen droplet diameter size range of bitumen emulsions is of the order of tenths of a micrometre to a few tens of micrometres, and the droplet size distribution is more or less poly-dispersed, depending on the specific formulation and manufacturing conditions used.

The most recent worldwide update of bitumen emulsion usage was carried out for the 2010 World Emulsion Congress through an enquiry in more than 100 countries. This showed that the total production in 2009 was roughly 8 million tonnes, a similar volume to that used in the previous 5 years (Le Bouteiller, 2012). This represents between 10% and 12% of the total global road bitumen demand. Consumption by type of application around the world is shown in Figure 9.1.

9.1 A brief history of the development of bitumen emulsions

In 1906, the first patent covering the application of dispersions of bitumen in water for road building was taken out (Van Westrum, 1906). Initially, efforts were made to form emulsions by purely mechanical means. However, it rapidly became apparent that mechanical action alone was insufficient, and, since these pioneering days, emulsifiers have been used in the process (Albert and Berend, 1916).
At first, the naturally occurring organic acids in the bitumen were utilised by adding sodium hydroxide or potassium hydroxide to the aqueous phase. The subsequent reaction formed an anionic soap that stabilised the dispersion (Bradshaw, 1960).

A great variety of acidic chemicals has been used to promote the stability of anionic bitumen emulsions. These include residues from fatty acid distillation, rosin acids, hydroxystearic acid and lignin sulfonates, all of which may be blended with the bitumen prior to emulsification or dissolved in the alkaline aqueous phase.

Since the early 1950s, cationic emulsifiers have become increasingly popular because of their affinity for many solid surfaces. This is an important property in road construction because the good adhesion of bitumen to different types of mineral aggregate is essential. The most widely used cationic emulsifiers are amines, amidoamines and imidazolines (Schwitzer, 1972).

In the mid-1980s, following the development of polymer modified binders, modified bitumen emulsions based on different manufacturing processes were produced. The residues of these bitumen emulsions had superior properties compared with traditional emulsions, and exhibited improved performance in specific applications (discussed further in section 9.6).

### 9.2 The components of emulsions

Bitumen emulsions are heterogeneous, two phase fluid systems consisting of two immiscible liquids, bitumen and water, stabilised by an emulsifier. The bitumen is dispersed throughout the continuous aqueous phase in the form of discrete particles, typically 1.0–10 µm in diameter, that are held in suspension by electrostatic charges imparted to the bitumen particles by an emulsifier (REA, 2013a). Components that are typically used to manufacture bitumen emulsions are shown in Table 9.1.
Bitumen emulsions can be divided into four classes

- Cationic emulsions
- Anionic emulsions
- Non-ionic emulsions
- Clay-stabilised emulsions.

The first two are, by far, the most widely used.

The terms ‘anionic’ and ‘cationic’ stem from the electrical charges on the bitumen globules. This identification system originates from one of the fundamental laws of electricity – like charges repel, opposite charges attract. If an electrical potential is applied between two electrodes immersed in an emulsion containing negatively charged particles of bitumen, they will migrate to the anode. In that case, the emulsion is described as ‘anionic’. Conversely, in a system containing positively charged particles of bitumen, they will move to the cathode, and the emulsion is described as ‘cationic’. The bitumen particles in a non-ionic emulsion are neutral and, therefore, will not migrate to either pole. These types of emulsion are rarely used on highways.

Clay-stabilised emulsions are used for industrial applications rather than for road applications. In these materials, the emulsifiers are fine powders, often natural or processed clays and bentonites, with a particle size much less than

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**Table 9.1 Emulsion components and their functions**

<table>
<thead>
<tr>
<th>Component</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Bitumen</strong></td>
<td>Conventional (commonly 180/200 pen grade; heavier grades such as 15/25 for special tack coats or 50/100 for microsurfacing or slurry seals are used) Modified (discussed in detail in section 9.6.3 of this chapter)</td>
</tr>
<tr>
<td><strong>Solvent</strong></td>
<td>Solvents may be included in the bitumen to improve emulsification to reduce settlement, the curing rate at low temperatures or to provide the right binder viscosity after curing. In addition, solvents can be used to produce primer emulsions, which, in some cases, are formulated with low bitumen content</td>
</tr>
<tr>
<td><strong>Water</strong></td>
<td>Water used to manufacture emulsions may come from various sources: municipal systems, wells, etc. Whatever the source of the water, it must contain a minimum amount of mineral and organic impurities</td>
</tr>
<tr>
<td><strong>Adhesion promoters</strong></td>
<td>Water resistance is an important property of cold asphalt mixtures and seals. The cured film from some anionic emulsions and occasionally also cationic emulsions may not have sufficient adhesion to aggregates, in which case adhesion promoters based on surface-active amine compounds are added</td>
</tr>
<tr>
<td><strong>Calcium chlorides and sodium chlorides</strong></td>
<td>Bitumen contains a small amount of salt, which can lead to an osmotic swelling of the droplets in an emulsion, as water is drawn into the droplet. This results in an increase in emulsion viscosity. Calcium chloride or sodium chloride (anionic emulsions) is included (0.1–0.2%) to reduce the osmosis of water into the bitumen and to minimise changes in the viscosity</td>
</tr>
</tbody>
</table>

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that of the bitumen particles in the emulsion. Although the bitumen particles may carry a weak electrical charge, the prime mechanism that inhibits their agglomeration is the mechanical protection of the surface of the bitumen by the powder together with the thixotropic structure of the emulsion, which hinders movement of the bitumen particles.

9.2.1 Functions of emulsifiers
Emulsifiers perform several functions within a bitumen emulsion. They

- make emulsification easier by reducing the interfacial tension between the bitumen and water
- determine whether the emulsion formed is the water-in-oil or oil-in-water type
- stabilise the emulsion by preventing the coalescence of droplets
- dictate the performance characteristics of the emulsion such as the setting rate and adhesion.

The emulsifier is the single most important constituent of any bitumen-in-water emulsion. In order to be effective, the emulsifier must be water soluble and possess the correct balance between hydrophilic (having an affinity for water) and lipophilic (having an affinity with lipids (organic compounds that are insoluble in water but soluble in organic solvents or fats)) properties. Emulsifiers can be used singly or in combination to provide special properties.

In the emulsion, the ionic portion of the emulsifier is located at the surface of the bitumen droplet while the hydrocarbon chain orientates itself on the surface of the bitumen, and is firmly bound to it. This is illustrated in Figure 9.2.

Emulsions may also contain unbound emulsifier that can influence the final properties of the emulsion, particularly their breaking and adhesion performance. The ionic portion of the emulsifier imparts a charge to the droplets themselves, and counter-ions such as sodium or chloride diffuse into the water phase.

Emulsifying agents are large organic molecules that have two distinct parts to them. These parts are called the ‘head’ and the ‘tail’. The head portion consists of a group of atoms that chemically have positive and negative charge areas. These two charged areas give rise to the head becoming polar (as in the poles of a magnet). Because of this polarity, and the nature of some of the atoms in this polar head, the head is soluble in water. The tail consists of a long-chain organic group that is not soluble in water but is soluble in other organic substances such as oils (bitumen). Thus, an emulsifying agent is a molecule with both water-soluble and oil-soluble portions. This unique characteristic gives the chemical its emulsifying ability.
In an anionic emulsion there are many billions of bitumen droplets with the emulsifying agent at the water–bitumen interface. The tail portion of the emulsifying agent aligns itself in the bitumen while the positive portion of the head floats around in the water, leaving the rest of the head negatively charged and at the surface of the droplet. This imparts a negative charge to all the droplets. An example can be represented by

\[ R-\text{COONa}^+ \]

**Figure 9.2** Schematic diagram of charges on bitumen droplets
The cationic emulsifying agent functions similarly to the anionic emulsifying agent: the negative portion of the head floats around in the water leaving a positively charged head. This imparts a positive charge to all the droplets. Because positives repel each other, all the droplets repel each other and remain as distinct bitumen drops in suspension (Bitumina, 2014). An example can be represented by

\[ R-\text{NH}_3^+\text{Cl}^- \]

In non-ionic emulsifiers, the hydrophilic head group is covalent (indicating a chemical bond that involves the sharing of electron pairs between atoms), polar and dissolves without ionisation (the process by which an atom or a molecule acquires a negative or positive charge by gaining or losing electrons). Any charge on the bitumen emulsion droplets is derived from ionic species in the bitumen itself. An example can be represented by

\[ R-\text{COO(}CH_2CH_2O)\text{xH} \]

In simple emulsifiers with the above chemical structures, R represents the hydrophobic (i.e. repelled by water) portion of the emulsifier, and is usually a long-chain hydrocarbon consisting of 8–22 carbon atoms derived from natural fats and oils such as tallow or from petroleum such as alkylbenzenes. The hydrophilic head group can variously contain amines, sulfonates, carboxylates, ether and alcohol groups. Emulsifiers with poly-functional head groups containing more than one of these types are widely used.

Complex wood-derived emulsifiers include Vinsol Resin, tannins and lignosulfonates, which contain polycyclic hydrophobic portions and several hydrophilic centres. Proteinaceous materials such as blood and casein have also been used in bitumen emulsifiers. In general, the structure of these molecules is complex, as can be seen by the structure for lignosulfonate illustrated in Figure 9.3.

Cationic emulsions constitute the largest volume of emulsions manufactured worldwide. They are produced from the following types or mixtures: monoamines, diamines, quaternary ammonium compounds, alkoxylated amines and amidoamines, and can be represented structurally as shown in Table 9.2.

Most of these are supplied in neutral basic form, and need to be reacted with an acid to become water soluble and cationic in nature. Therefore, cationic emulsions are generally acidic with a pH < 7. Hydrochloric acid is normally used, which reacts with the nitrogen atom to form an ammonium ion. This reaction can be represented as follows

\[ \text{amine } + \text{ acid } \rightarrow \text{alkylammonium chloride} \]
Figure 9.3 The structure of lignosulfonate

Table 9.2 Chemical components of cationic emulsions (R=C_8–22)

<table>
<thead>
<tr>
<th>Example of chemical structure</th>
<th>Chemical type</th>
</tr>
</thead>
<tbody>
<tr>
<td>R–NH₂</td>
<td>Monoamine</td>
</tr>
<tr>
<td>R–NH₂CH₂CH₂CH₂NH₃⁺2Cl⁻</td>
<td>Diamine</td>
</tr>
<tr>
<td>RCONHCH₂CH₂CH₂NH(CH₃)₂Cl⁻</td>
<td>Amidoamine</td>
</tr>
<tr>
<td>CH₃</td>
<td>Quaternary ammonium compound</td>
</tr>
<tr>
<td>R–N–CH₃Cl⁻</td>
<td>Alkoxylated amine</td>
</tr>
<tr>
<td>CH₃</td>
<td></td>
</tr>
<tr>
<td>(CH₂CH₂O)ₓH</td>
<td></td>
</tr>
<tr>
<td>R–N–(CH₂CH₂O)ₓH</td>
<td></td>
</tr>
</tbody>
</table>
Quaternary ammonium compounds do not need to be treated with acids as they are already salts and water soluble, but the water-phase pH can be adjusted with acid if required, to modify the performance of the emulsion.

Anionic emulsifiers constitute the second largest volume of emulsion produced worldwide and are usually stabilised with fatty acids or sulfonate emulsifiers.

Fatty acids are insoluble in water, and are made soluble by reacting with an alkali, normally sodium or potassium hydroxide, so that anionic emulsions are alkaline with a pH $> 7$. (Cationic emulsions usually have a pH in the range 2–3, with anionic emulsions having a pH in the range 10–11. A liquid having a pH value of 7 is neutral, being neither acidic nor alkaline.)

Sulfonates are usually supplied as water-soluble sodium salts. Further neutralisation is not required, but an excess of sodium hydroxide is used in order to keep the pH of the emulsion higher than 7 and also to neutralise the natural acids contained in the bitumen.

Non-ionic emulsifiers are not produced in significant quantities, and are normally only used to modify both anionic and cationic emulsions. Typical non-ionic emulsifiers include nonylphenolethoxylates and ethoxylated fatty acids.

9.3 The manufacture of bitumen emulsions

Most bitumen emulsions are manufactured by a continuous process using a colloid mill (a machine used to reduce the particle size of a solid in suspension in a liquid or to reduce the droplet size of a liquid suspended in another liquid). This equipment consists of a high speed rotor that revolves at 1000–6000 revs/min in a stator. The clearance between the rotor and the stator is typically 0.25–0.50 mm and is usually adjustable.

Hot bitumen and emulsifier solutions are fed separately but simultaneously into the colloid mill, the temperatures of the two components being critical to the process. The viscosity of the bitumen entering the colloid mill should not exceed 0.2 Pa·s (2 poise). Bitumen temperatures in the range 100–140°C are used in order to achieve this viscosity with the penetration grade bitumens that are normally used in emulsions. To avoid boiling the water, the temperature of the water phase is adjusted so that the temperature of the resultant emulsion is less than 90°C. As the bitumen and emulsifier solutions enter the colloid mill, they are subjected to intense shearing forces that cause the bitumen to break into small globules. The individual globules become coated with the emulsifier, which gives the surface of the droplets an electrical charge. The resulting electrostatic forces prevent the globules from coalescing.
When the bitumen is not a soft penetration grade bitumen or when a polymer modified binder is used, the process is more difficult. Higher temperatures are needed to allow the bitumen to be pumped to and dispersed in the mill, and dispersion of the bitumen requires more power input to the mill, which further increases the product temperature. Pressurised mills are used with bitumens having a high viscosity at normal emulsification temperatures and to allow higher throughput with normal bitumens. Emulsions with temperatures up to 130°C are produced under high pressure, and the emulsion output must be cooled below 100°C before being discharged into normal storage tanks.

As an alternative to a colloid mill, a static mixer may be used. This contains no moving parts. The high shear necessary to produce an emulsion is generated by pumping the input materials at high speed through a series of baffles designed to produce highly turbulent flow. The benefits of having no moving parts and no shaft seals are obvious; additional benefits claimed are closer control of the bitumen particle size in the emulsion produced, with consequent closer control of critical emulsion properties (discussed in section 9.5).

A batch process can be used for the production of small volumes of emulsion. The type of mixer that is used is chosen to suit the consistency of the end product: it may be a high speed propeller for low viscosity road emulsions or a slow Z-blade mixer for paste-like industrial emulsions.

Schematic diagrams of continuous and batch emulsion manufacturing facilities are shown in Figure 9.4.

9.4 Storage and handling of bitumen emulsions

The production rate of emulsion plants is generally higher than the quantity necessary to meet the current demand. Storage facilities make it possible to have longer production runs, thus improving plant productivity. Modern emulsions can be stored for up to several months without major changes in their physical properties. It is advisable to use small diameter vertical storage tanks with a minimum horizontal cross section with a dip tube filling pipe that extends to a point at or near the bottom of the storage tank. Vertical tanks are superior to horizontal tanks because they minimise the surface area exposed to air. Horizontal tanks are acceptable but not recommended because they expose the emulsion to air at its surface, which builds a crust on the product. The crust will not liquefy nor will it pump or spray. Accordingly, if a storage facility has to use horizontal tanks, keeping them full to limit exposure to air is recommended.

Emulsions of different ionic types should never be mixed, and tanks should be thoroughly cleaned before refilling with a different ionic type. Provision
Figure 9.4 Schematic diagrams of (a) a continuous emulsion plant and (b) a batch emulsion plant.
should also be made to ensure proper agitation of stored emulsion to prevent settling, decantation or creaming. Mixers may be used in the tank. They may be simple propeller mixers that revolve slowly or larger anchor sweep mixers. It is important not to mix the emulsion too frequently. It should require mixing once a week or for a short time before being transported for use. Bitumen emulsions are sensitive to frost, which can cause irreversible breaking, so appropriate measures should be put in place to prevent stored emulsion from freezing (Baumgardner, 2006).

9.5 Properties of bitumen emulsions

Surface dressing applications are, by far, the largest use for bitumen emulsions. Hence, the emulsion properties discussed below relate particularly to surface dressing requirements. Notwithstanding, the principles have general application. The most important properties of bitumen emulsions are:

- stability
- viscosity (or, more accurately, rheology)
- breaking
- adhesivity.

There are conflicting requirements for the properties of bitumen emulsions. The ideal emulsion would be stable under storage, transport and application conditions but would break rapidly very soon after application, leaving a binder having the properties of the original bitumen adhering strongly to the road and the aggregates. It would have a low viscosity for ease of handling and application and would flow to minimise irregular spraying but would not flow due to road irregularities, cambers or gradients.

It is generally assumed that the bitumen produced when an emulsion breaks is the same as the bitumen that was used to produce the emulsion, but there are exceptions. The emulsifier used may modify the recovered bitumen, particularly in relation to rheological properties that are surface dependent such as adhesion. Clay-stabilised emulsions (such as Shell Flintkote) can be formulated to produce bitumen films that have non-flow properties even at very high temperatures. These are useful for roofing and insulating applications.

9.5.1 Emulsion stability

9.5.1.1 Settlement

Emulsions, particularly those having a low bitumen content and low viscosity, are prone to settlement.

At ambient temperatures, the grades of bitumen normally used in emulsions have a density that is slightly higher than that of the aqueous phase of the emulsion. Consequently, the bitumen particles tend to fall through the aqueous phase, resulting in a bitumen-rich lower layer and a bitumen-deficient upper
layer. The velocity of the downward movement of the particles can be estimated using Stokes’ law (Stokes, 1851)

$$v = \frac{2}{9} gr^2 \frac{\varsigma_1 - \varsigma_2}{\eta}$$

where $g$ is the gravitational force, $r$ is the particle radius, $\varsigma_1$ is the specific gravity of the dispersed phase, $\varsigma_2$ is the specific gravity of the aqueous phase and $\eta$ is the viscosity of the aqueous phase.

However, Stokes’ law applies to particles that are free to move and have no inter-particular forces, conditions that are frequently not met in a bitumen emulsion.

Settlement can be reduced by equalising the densities of the two phases. One way of achieving this is to add calcium chloride to the aqueous phase. However, because the coefficients of thermal expansion of bitumen and the aqueous phase are not the same, their densities can be made equal only at one specific temperature. As large particles settle more rapidly than do small ones, settlement can be abated by reducing either the mean particle size or the range of particle sizes that are present. Increasing the viscosity of the aqueous phase will also reduce the rate of settlement. Indeed, if the aqueous phase behaviour can be made non-Newtonian by introducing a yield value, settlement can be eliminated completely. In addition to gravity, there are repulsive forces between the bitumen droplets caused by the layers of emulsifier on the droplets that impede or accelerate settlement.

Coalescence follows settlement in two stages. First, bitumen droplets agglomerate into clumps; this reversible phenomenon is called flocculation. Second, the resultant flocks fuse together to form larger globules; this irreversible process is called coalescence. This can be spontaneous or it can be induced by mechanical action.

9.5.1.2 Stability during pumping, heating and transportation
Two bitumen particles in an emulsion will coalesce if they come into contact. Contact is prevented by electric charge repulsion and the mechanical protection offered by the emulsifier. Any effect that overcomes these forces will induce flocculation and coalescence. Flow of the emulsion, caused by pumping, heating (convection currents) or transportation is one such effect. Some emulsifiers have a tendency to foam, which is itself a potential cause of coalescence because bitumen particles in the thin film of a bubble are subjected to the forces of surface tension.

9.5.2 Emulsion viscosity
As surface dressing emulsions are almost always applied by spray, their viscosity under spraying conditions is of prime importance.
The viscosity of emulsions is normally measured as the time of effluence from a flow cup with a standard orifice at different temperatures.

Saybolt Furol cups at 25 and 50°C are used according to the American Society for Testing and Materials (ASTM) standards in the USA (ASTM D244:2009 (ASTM, 2009)), and standard tar viscometers at 40 and 50°C are used in Europe (EN 12846-1:2011 (BSI, 2011)). Dynamic viscosity is used in Europe (EN 13302:2010 (BSI, 2010a)) to characterise the behaviour of the emulsion after spraying (e.g. in surface dressing or tack coat applications) or during mixing with aggregates (e.g. in coating applications).

Emulsions having a high concentration of the dispersed phase (bitumen) rarely have Newtonian viscosity characteristics (e.g. the apparent viscosity changes with the shear rate at which the viscosity is measured). In addition, the rate of change of viscosity with temperature is not the same for different emulsions. When comparing two different emulsions, it is possible for one to have a lower viscosity and better spray distribution at the spraying temperature of 85°C while also showing higher viscosity and less run off from the road at a road temperature of 30°C compared with the other emulsion.

Single point viscosity measurements can, therefore, be misleading, although more data are difficult to obtain.

In principle, there are four methods of increasing the viscosity of an emulsion:

- by increasing the concentration of the dispersed phase (bitumen)
- by increasing the viscosity of the dispersed phase
- by increasing the viscosity of the continuous phase (the aqueous phase)
- by reducing the particle size distribution range.

The opposite changes will decrease the viscosity of an emulsion. As stated, emulsion viscosity is almost independent of the viscosity of the dispersed phase (bitumen). It is possible to produce emulsions based on hard bitumen (<10 pen) that are readily pourable at 10°C.

9.5.2.1 Increasing the bitumen content
At low bitumen contents, the effect is small. At high bitumen contents, a small increase in concentration can induce a dramatic change in the viscosity that may be uncontrollable.

9.5.2.2 Modification of the aqueous phase
The viscosity of a bitumen emulsion is highly dependent on the aqueous phase composition. It has been shown that, in the case of conventional cationic road emulsions, the viscosity can be increased by decreasing the acid content or increasing the emulsifier content. Additives intended specifically as viscosity modifiers can also be used.
Latex addition in the aqueous phase for modified bitumen emulsion production is another example (discussed in section 9.6.3).

9.5.2.3 Increasing the flow rate through the mill

By increasing the flow rate through the mill, the particle size distribution of the emulsion will be changed. At bitumen contents of less than 65%, the viscosity of the emulsion is virtually independent of the flow rate. However, at bitumen contents greater than 65% when the globules of bitumen are packed relatively close together, inducing a change in the particle size distribution by changing the flow rate has a marked effect on the viscosity, as shown in Figure 9.5.

Figure 9.5 Emulsion viscosity (in degrees Engler) as a function of the flow rate for different bitumen contents
9.5.2.4 Decreasing the viscosity of the bitumen in the mill
If the viscosity of the bitumen entering the colloid mill is reduced, the particle size of the emulsion will be reduced, which, in turn, tends to increase the viscosity of the emulsion.

9.5.3 Breaking of emulsions
It is important to know when an emulsion has ‘broken’. ‘Breaking’ is the loss of water from the emulsion. In the process of breaking, an emulsion changes from a liquid to a continuous film of bitumen. Determining whether an emulsion has broken is very easy: the colour turns from brown to black. Some specifications prohibit an emulsion to be covered by the succeeding layer unless the emulsion has broken. A truck reversing into a paver over an emulsion tack coat, or bond coat that has not broken will lift the emulsion on its tyres, thus reducing or negating the effect of the tack coat or bond coat.

Once in contact with a solid surface such as a layer of asphalt or aggregate, the emulsion is able to coalesce (set), and the water either runs off or evaporates. By careful control of the emulsion chemistry, its rate of coalescence, or setting rate, can be adjusted to be slow, medium or rapid setting in order to have optimum versatility.

According to the European standard EN 13808:2013 (BSI, 2013), there are different methods for determining the breaking behaviour. It is compulsory to declare a performance class according to one of these methods (for more details, see section 9.7.2).

European standards can be used for medium setting and fast setting emulsions, in which the breaking value is a dimensionless number corresponding to the quantity of reference filler, in grams, needed to coagulate 100 g of bitumen emulsion (EN 13075-1:2012 (BSI, 2012)). For slow setting and over stabilised emulsions, a European standard specifies a method for the determination of the mixing stability of the emulsion with cement (EN 12848:2009 (BSI, 2009)).

There are, in principle, six parameters that can be used to change the breaking properties of emulsions

- the bitumen content
- the aqueous phase composition (type and content of emulsifier, pH value)
- the particle size distribution
- environmental conditions
- aggregates
- the use of breaking agents.

How each of these parameters changes the breaking properties is set out as follows.
9.5.3.1 Bitumen content
At high bitumen contents, the bitumen particles are more likely to come into contact with each other, resulting in an increase in the breaking rate.

9.5.3.2 Aqueous phase composition
The breaking rate of a bitumen emulsion has been shown to be increased by reducing the acid content, increasing the emulsifier content or by decreasing the ratio between the acid and emulsifier contents.

9.5.3.3 Particle size distribution
The smaller the size of the bitumen particles, the finer will be the dispersion, resulting in a slower breaking rate of the emulsion.

9.5.3.4 Environmental conditions
The evaporation of water is influenced by the incident wind velocity, humidity and temperature in that order. Temperature and humidity are related: as the air temperature falls, the relative humidity increases. Working at night with emulsions can therefore be difficult at low ambient temperatures and the relative humidity reaching 100%, causing the loss of water to cease entirely.

At higher ambient temperatures, the bitumen particles in the emulsion are more mobile, and the bitumen is softer. In such circumstances, particles are more likely to come into contact and, therefore, more likely to coalesce.

9.5.3.5 Aggregates
As stated above, the spraying conditions for fast setting bitumen grades initiate the emulsion breaking process. Accordingly, it is imperative that aggregates are applied very soon after the emulsion has been applied to the surface of the road. This is necessary to ensure that the emulsion is still capable of wetting the aggregates. When the aggregates are applied, breaking is accelerated by the absorption of emulsifier onto the aggregate and the evaporation of water. The former can be completely inhibited by the use of coated aggregates. Conversely, dust can cause rapid breaking of the emulsion onto the dust with no adhesion of bitumen to the aggregates. Within these extremes, the geometry of the aggregates (i.e. their size and shape) has a considerable influence on the breaking rate of the emulsion.

The emulsion contains emulsifier molecules in both the water and on the surface of the droplets. Some of the emulsifier ions form micelles (particles of colloidal dimensions) and, in a stable emulsion, equilibrium exists as shown in Figure 9.6. If some of the emulsifier ions are removed from the solution, the balance is restored by ions from the micelles and the surface of the droplets. This occurs when an emulsion comes into contact with a
mineral aggregate. The negatively charged aggregate surface rapidly absorbs some of the ions from the solution, weakening the charge on the surface of the droplets. This initiates the breaking process, as shown in Figure 9.7. A point is reached where the charge on the surface of the droplets is so depleted that rapid coalescence takes place. The aggregate is now covered in hydrocarbon chains, and, as a result, the liberated bitumen adheres strongly to its surface.

Figure 9.6 Emulsifier ions forming micelles in a stable solution

Figure 9.7 Schematic diagram of an emulsion breaking
9.5.3.6 The use of breaking agents

The use of breaking agents can accelerate the breaking of an emulsion. For surface dressing emulsions, it is possible to spray a chemical breaking agent either simultaneously with the emulsion or just after the emulsion has been applied to the road. Care is required in the use of breaking agents. Applying too little will have no effect, but applying too much may cause the emulsion to break and adversely affect its adhesivity. Poor distribution of a breaking agent can have similar effects.

9.5.4 Emulsion adhesivity

It is very important in all applications where bitumen is used as an adhesive between solid surfaces that the bitumen ‘wets’ the surface to create the maximum effective contact area. With dry substrates, the ‘critical surface tension of wetting’ of the aggregate must be high enough to ensure that the bitumen spreads easily over the surface. The resultant adhesion generally exceeds the cohesion of the bitumen. However, when the surface of an aggregate is covered with water, the wetting of the aggregate becomes a three phase phenomenon that can only occur if the balance of the interfacial energies favours wetting by the bitumen. Cationic emulsifiers are particularly efficient at reducing the free surface energy of a polar aggregate, forming a thermodynamically stable condition of minimum surface energy resulting from the emulsifier being attracted to the aggregate surface (Heukelom and Wijga, 1973; HRB, 1968).

Most cationic emulsifiers are also anti-stripping agents. Consequently, initial bonding is assured. However, the quality of the bond between the bitumen and the aggregate depends on a number of factors:

- the type and amount of emulsifier
- the bitumen grade and constitution
- the pH of the emulsifier solution
- the particle size distribution of the emulsion
- the nature of the aggregate.

9.6 Modification of bitumen emulsion properties

9.6.1 Particle size distribution

The particle size distribution of bitumen emulsions influences many of the emulsion properties that are critical to achieving success in application and service. Automatic equipment is available that can measure both the mean particle size and the particle size distribution of emulsions, but bitumen emulsions can present difficulties because they usually contain a few very large particles and may contain some that are very small.

The strong influence of the particle size distribution on the properties of bitumen emulsions is due to the surface area of a spherical particle being
proportional to the square of its diameter, and its mass is a function of the cube of its diameter. Many performance properties of an emulsion are influenced by the amount of ‘free’ emulsifier in the aqueous phase (i.e. the amount of emulsifier that has not been absorbed onto the bitumen particles). The amount of emulsifier that is absorbed onto the bitumen particles depends on the total surface area of those particles. Even a small proportion of bitumen present as sub-micrometre particles can create a large surface area.

The distribution of the emulsion droplet size is dependent on the interfacial tension between the bitumen and the aqueous phase (the lower the interfacial tension, the easier the bitumen disperses) and on the energy used in dispersing the bitumen. For a given mechanical energy input, harder bitumens will produce coarser emulsions, and high penetration or cut-back bitumens will produce finer emulsions. It is possible to influence the particle size and distribution by modifying the materials and process used to make an emulsion.

9.6.1.1 The addition of acid to the bitumen
The addition of naphthenic acids to a non-acidic bitumen is important for the production of anionic emulsions. The acids react with the alkaline aqueous phase to form soaps that are surface active and that stabilise the dispersion. The addition of naphthenic acids causes a decrease in the mean particle size of the emulsion without changing its size distribution.

9.6.1.2 Manufacturing conditions
Manufacturing conditions have a substantial influence on the resulting particle size distribution of the emulsion.

- Temperature. Increasing the temperature of either the aqueous phase or the bitumen normally decreases the mean particle size of the emulsion.
- Bitumen content. Increasing the bitumen content increases the mean particle size and tends to reduce the range of particle sizes.
- Composition of the aqueous phase. For cationic emulsions manufactured using hydrochloric acid and an amine emulsifier, the particle size can be decreased by increasing either the acid or the emulsifier content: if the ratio of acid to amine is kept constant, the particle size can also be reduced by increasing the amine/acid content; the size distribution does not appear to be related to the concentration of these two components.
- Operating conditions of the colloid mill. The gap and rotational speed of the colloid mill strongly influence the particle size and distribution of the emulsion: a small gap will result in a small particle size with a relatively narrow range of sizes; high rotational speed will produce a small particle size.
Decreasing the viscosity of the bitumen. If the viscosity of the bitumen entering the colloid mill is lowered, the particle size of the emulsion will be reduced, which will tend to increase the viscosity of the emulsion.

9.6.2 Effects of bitumen properties

9.6.2.1 Influence of the ionic content of bitumen in aqueous suspension

Bitumens usually contain a small amount of ionic material, principally sodium chloride (an ion is an atom or molecule, or a group of either, that has lost or gained one or more electrons and, as a result, possesses an electrical charge). Typical concentrations are less than 0.1%, which is in the low hundreds parts per million (ppm). This small amount of ionic material in conjunction with the aqueous medium in the emulsion can exert considerable influence on the mixture’s viscosity. At concentrations up to 20 ppm, the ionic material has little or no effect. As the concentration increases, the viscosity of emulsions produced from the bitumen rises (for a given bitumen content) to a maximum. When the concentration exceeds approximately 300 ppm, the emulsion viscosity suddenly falls. These characteristics can be attributed to osmosis (the natural movement of water from low ionic concentrations to a higher concentration on the other side of a semi-permeable membrane to achieve an ionic concentration balance; this process can generate physical pressure). If a particle of bitumen has an ionic content it may be subject to osmosis. If this occurs, water will enter the particle, and the particle will expand under the osmotic pressure. If the emulsion has a substantial proportion of its particles susceptible to osmosis, then there will be a substantial increase in the viscosity. If the difference in ionic concentration between the particle and the surrounding emulsion medium is high enough, the particle will continue to expand to a point where it ruptures. If this point of failure is reached, it can trigger coagulation between other particles, which is undesirable. When creating an emulsion, its osmotic characteristics can be controlled. If the bitumen used to manufacture the emulsion has a low ionic content, this can be modified by adding sodium chloride to the bitumen before the process is started. If the ionic content is high, running the risk of osmotic failure, ionic material (e.g. calcium chloride) can be added to the aqueous component of the emulsion. This control of ionic concentration between the emulsion medium and the content of an osmotic particle will manage the particle size and, therefore, control the influence this parameter has on the emulsion viscosity.

9.6.2.2 Bitumen density

High bitumen density can cause rapid settlement in emulsions, leading to coagulation during static storage. The problem can be alleviated by adding a high boiling point solvent (e.g. kerosene) to bitumen that is a grade harder than would normally be used, or by increasing the specific gravity of the
aqueous phase (e.g. by adding calcium chloride or by using an emulsifier that imparts a yield value to the aqueous phase).

9.6.2.3 Acid value
The presence of natural naphthenic acids in the bitumen is beneficial to the production of most anionic emulsions. The acids react with the excess of alkali in the aqueous phase during emulsification, acting as an efficient natural emulsifier. However, some industrial emulsions cannot be made with acidic bitumens: the naphthenic acids take precedence at the particle surface over the emulsifiers in the aqueous phase, and give an emulsion that is either unstable or has insufficient stability.

9.6.3 Polymer modified bitumen emulsions
9.6.3.1 Definition
Modified emulsions are those that produce a residue of modified binder. A polymer can be mixed in bitumen to obtain a polymer modified bitumen that can then be emulsified (called single phase modified emulsions). Ethylene–vinyl acetate and styrene–butadiene–styrene are most commonly used in paving grade emulsions.

Polymers can be also be added during the aqueous phase as latex (dual-phase modified emulsions). Latex is a water-based polymer, and comes in anionic, non-ionic and cationic forms. It is important that the latex type should be compatible with the emulsion. Styrene–butadiene rubber, polychlorophene and natural rubber latex are commonly used in road emulsions (Figure 9.8).

9.6.3.2 Properties of residual binder
The properties of the binder after the breaking of a bi-phasic modified emulsion depend on the type of bitumen, the type of modifier, their compatibility and the relative modifier/bitumen concentrations. The action caused by the modifier results in improvements in the original binder similar to those obtained with elastomeric modified bitumens, as follows

- increased cohesion at medium and high service temperatures
- improved performance at low temperatures
- increased plasticity interval (= ring and ball temperature – Fraass point)
- better ageing behaviour
- improved rheological behaviour.

In the case of single phase modified emulsions, the performance of the residual binder should be very similar to that of the modified bitumen used for its production, although the colloid mill mechanical action as well as the emulsifier physico-chemical action can modify the performance slightly.
With dual phase modified emulsions, there is a concern as to how to recover material that is truly representative of what is applied in the field. The different methods of obtaining the residual binder of emulsions in the laboratory (distillation, oven stabilisation, vacuum, etc.) do not satisfactorily reproduce the breaking mechanism of these emulsions, which varies with the type of application in either surface dressing mixtures or microsurfacing. For this reason, it is preferable to check the performance of the different types of dual phase modified emulsions by an indirect method by means of tests carried out on the final product.

9.6.3.3 Manufacture
Modified emulsions are manufactured following the same basic method as conventional emulsions except for the addition of the modifying agent. As single phase modified emulsions are manufactured from previously modified bitumen, its high viscosity requires that the emulsion manufacturing temperature is higher than in the conventional emulsion manufacturing process. Accordingly, the final thermal balance leads to an emulsion temperature at the outlet of the mill in excess of 100°C. In these conditions, it is necessary to modify the conventional manufacturing equipment to make it capable of working under pressures ranging from 1.5 to 2 bar and to use a cooling system at the outlet of the mill (generally a heat exchanger). As mentioned above, in the dual phase modified emulsion case, the modifier is typically

![Diagram showing single phase and dual phase modified emulsions](image)

**Figure 9.8** (a) Single phase and dual phase (b) modified emulsions: differences

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added as a latex dispersion. It can be incorporated either in the aqueous phase or the bitumen or even added subsequently to the emulsion. Dispersion in the aqueous phase is the most commonly used method. It allows for accurate regulation of the emulsification parameters, especially pH, as well as achieving good homogeneity in the dispersion (World Road Association, 1999).

9.6.3.4 Method of use
Emulsions containing polymer modified bitumen are generally used in exactly the same way as their unmodified equivalents. It must not be assumed that the use of polymer modified products provides a safety margin for poor contracting practices. On the contrary, the modified products usually require greater care in their use and are less tolerant of unsuitable site and weather conditions. Their benefits are in the performance levels that can be obtained under high traffic stress conditions.

9.6.4 Manufacturing variables and emulsion properties
Previous sections have provided an overview of the factors that influence the properties of emulsions, and a number of alternative approaches are available to the emulsion manufacturer to adjust emulsion properties. However, it is virtually impossible to adjust one property of the emulsion without influencing others. This interdependence is illustrated in Figure 9.9.

9.7 Classification and specification of bitumen emulsions
9.7.1 US specifications
Paving grade emulsions are classified according to the sign of the charge on the droplets in anionic and cationic emulsions while the test methods are defined in ASTM specification D244 (ASTM, 2009). In accordance with an ASTM publication (ASTM, 2002), cationic emulsions are classified by an alphanumeric designation: the first part indicates the breaking rate (RS or rapid setting, QS or quick setting, MS or medium setting, and SS or slow setting) followed by numbers and text indicating the emulsion viscosity and residue properties. For example, CRS-2 would be a reactive (rapid setting) cationic emulsion of high viscosity. CSS-1h would be a non-reactive (slow setting) cationic emulsion with a hard bitumen residue.

9.7.2 European specifications
The European standard EN 13808 (BSI, 2013) specifies the requirement for the performance characteristics of conventional and modified cationic emulsions. They are defined as an expression in letters and numbers that describes the different characteristics of emulsions: their polarity, binder content, binder type, type and amount of flux (if any) and breaking value. The European standard provides a framework of specifications and classes
for the properties of cationic bituminous emulsions and is based on the use of four tables

- Table 1 defines the denomination of the abbreviated terms
- Table 2 describes 12 classes according to properties of the emulsion (breaking behaviour, viscosity and storage stability among other properties)
- Tables 3 and 4 define the properties and performance classes applicable to the residual binders obtained after distillation, recovery, stabilisation and ageing procedures.

Some examples of abbreviated terms according to the European specification are shown in Table 9.3.

**9.8 Uses of bitumen emulsions**

**9.8.1 Road uses**

The vast majority of bitumen emulsions are used in surface dressing, slurry seals or microsurfacing applications, and these are discussed in detail in Chapter 21.

However, their versatility makes them suitable for a wide variety of applications, which are briefly described below.
9.8.1.1 Tack coats
Tack coating is the application of a conventional bitumen emulsion to facilitate adhesion between layers in an asphalt pavement. The function of the tack coat is to minimise the effect of residual surface dust on the existing surface and to provide an adhesive surface for the overlay.

There is considerable variation in the type of emulsion used for tack coats worldwide. In many countries, unmodified slow setting anionic or cationic emulsions are used, which may be diluted with water, but Europe and South America use unmodified rapid setting cationic emulsions. It is necessary for the tack coat to wet out any dust on the surface of the lower layer, and this favours emulsions of small particle size and some solvent content.

9.8.1.2 Bond coats
A bond coat is a proprietary polymer modified bitumen emulsion used to promote adhesion between layers in the construction of a paved area, or to bond a new surface course to an existing road surface when carrying out road maintenance or construction.

Modern road surface courses such as stone mastic asphalts, thin surface course systems and porous asphalts may be laid thinner than is the case with traditional surface courses. Such layers place greater reliance on the inter-surface bonding to reduce the risk of slippage between the two courses. Some modern surface courses are more open in texture and consequently more porous. In such cases, the bond coat has an important function in helping to waterproof and inhibit the ingress of water to the structural layers (REA, 2013b)

New developments include bond coats based on very hard binders that cure rapidly and avoid sticking to the tyres of traffic or construction equipment.

The best equipment for applying bond coats is an integral paver. This machine is equipped with an emulsion spray system that places the emulsion on the pavement surface immediately in front of the hot asphalt.

![Table 9.3](image)

<table>
<thead>
<tr>
<th>Type according to EN 13808</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>C 69 B 2</td>
<td>Cationic, nominal binder 69%, produced from bitumen, Class 2 breaking value</td>
</tr>
<tr>
<td>C 65 BP 3</td>
<td>Cationic, nominal binder 65%, produced from bitumen, containing polymers, Class 3 breaking value</td>
</tr>
<tr>
<td>C 69 BF 3 70/100</td>
<td>Cationic, nominal binder 69%, produced from bitumen, containing more than 3% (m/m) flux, Class 3 breaking value and 70/100 pen grade bitumen</td>
</tr>
</tbody>
</table>
Specifications may require that, after application, the emulsion must be allowed to break before the asphalt is laid, unless it is applied by an integral paver. As pointed out in section 9.5.3, verification of an emulsion having broken is simple: it turns from brown to black in colour.

More information on bond coats can be found in section 5.5 of the UK’s asphalt transport, laying and compaction standard (BSI, 2010b).

9.8.1.3 Fog seals
Fog seals are a method of adding diluted emulsions to an existing pavement surface to improve sealing or waterproofing. This may reduce or prevent further stone loss by holding the aggregate in place, or simply improve the surface appearance. However, inappropriate use can result in slick pavements and tracking of excess material.

The Asphalt Emulsion Manufacturers Association defines a fog seal as ‘a light spray application of dilute asphalt emulsion used primarily to seal an existing asphalt surface to reduce ravelling and enrich dry and weathered surfaces’. Others refer to fog seals as enrichment treatments, because they add fresh asphalt to an aged surface and extend the life of the surface of the pavement. Fog seals are also useful in surface dressing (what is described in the USA as a ‘chip seal’) applications to hold chips in place in fresh seal coats (Caltrans, 2003).

9.8.1.4 Prime coats
Emulsion prime coats are applied to unbound subbases in order to seal the surface before the application of the asphalt layers. The primer seal prevents the ingress of water into the layer and the loss of fines from wind or water erosion, and ideally allows construction vehicles to drive over the surface without particles on the surface being picked up by tyres.

A few millimetres of penetration are readily achievable if the compacted material is not too dense, but may be very difficult in practice with fine graded and highly compacted bases. Penetration can be achieved using very slow setting cationic or anionic emulsions containing a low bitumen and solvent content, but, in some cases, deep penetration can be very difficult.

However, current thinking suggests that deep penetration may be unnecessary, as dense and highly compacted bases are already very robust and merely need to be sealed from the intrusion of water. This can be achieved using a very thin primer application with minimal penetration.

9.8.1.5 Soil stabilisation
Cationic slow setting emulsions can be used for the stabilisation of uncrushed naturally occurring gravels and sandy soils. Generally, soils with a sand
equivalence value of more than 25 (a measure of the clay content) can be
treated with some degree of success for use as a base material for hot overlay
or for minor roads where a seal coat may be sufficient. Materials of even
lower sand equivalence can, in some cases, be treated successfully using a
combination of emulsion and a hydraulic binder such as lime or cement.

9.8.1.6 Cold in-place recycling
Surfaces courses or even the full depth of the pavement can be recycled in
place either by a specially built mobile plant or by simple equipment.
Cold recycling uses bitumen emulsions either alone or in combination with
cement or lime. Typically, a cationic slow setting emulsion is used. Structural
materials can be produced from emulsion and crushed aggregates or
reclaimed asphalt pavement, and such mixtures can meet acceptable per-
formance criteria comparison favourably with hot mix asphalts. Depending
on the aggregate grading, medium or slow setting emulsions can be used.
Cold mixtures that combine bitumen emulsion with cement can give much
improved bearing capacities (AkzoNobel, 2014). Rejuvenation agents may
be included, depending on either the penetration or viscosity grade of the
bitumen in the existing pavement.

9.8.1.7 Cold plant mixtures
Growing concern about energy conservation and the desire for environ-
mentally friendly processes have generated an interest in the production of
cold asphalts.
Although the idea of wide-scale production of cold materials is relatively
recent in countries such as the UK, elsewhere the manufacture and utilisation
of ‘emulsified asphalts’ has been commonplace for a number of years
(Bradshaw, 1974).
France has been using cold materials since the 1960s for strengthening and
reprofiling lightly trafficked roads, and collaborative work between contrac-
tors and road authorities led to the development of a material called ‘grave
e´mulsion’. Although traditionally a continuously graded 20 mm material,
modern grave e´mulsion typically comprises a 14 mm grading with a bitumen
content of 4–4.5%. Grave e´mulsion can be stockpiled, laid using conven-
tional paving equipment or by a blade grader, and compacted at ambient
temperatures. Although the material was originally used for minor mainten-
ance works, more recently it has been applied in structural layers of
moderately trafficked roads. A national standard for grave e´mulsion was
introduced in France in 1993 (AFNOR, 1993).
Similar developments have taken place in the USA, where environmental
issues and the remoteness of some sites from asphalt plants provided the

Cold mixture technology presents a new set of challenges to engineers who have traditionally specified hot asphalts. Whereas hot mixtures rely on the visco-elastic properties of bitumen, emulsion mixtures introduce a new series of conditions that must be met in order that such materials can be successfully produced and laid. The surface chemistry of the aggregate begins to have an important role, and emulsions must be tailored to the mineralogy of different rock types.

The classic concept of how cold mixtures work is that the emulsion breaks, either during mixing or compaction, coating the aggregate, after which there is an increase in strength over time. The strength development is a result of the expulsion of water from the aggregate matrix and the coalescence and subsequent cohesion of the bitumen particles. However, the characteristics of initial workability or being able to stockpile the material and the subsequent development of mechanical strength in situ form conflicting requirements. By tailoring the emulsion to produce a mixture that will remain workable for days or weeks, the development of cohesion in the compacted mixture and, hence, the strength gain of the matrix will also be retarded.

If the emulsion is tailored to produce a rapid break, then the mixture will quickly exhibit developing stiffness and, hence, will only have a brief workability window. Accordingly, there exists the potential for innovative technology (such as ‘half-warm mixtures’, discussed in section 9.9) to address such problems and develop cold mixtures that perform in a manner similar to that of hot mixed products.

Finally, cold porous asphalt mixtures based on modified emulsions can be used for hot porous asphalt repairs or maintenance.

### 9.8.2 Miscellaneous uses of bitumen emulsions

Details of miscellaneous applications for emulsions have been published by the Road Emulsion Association (REA, 2013c). Bitumen emulsions are also used in other civil engineering works and in horticultural and agricultural applications. Some examples are given below.

#### 9.8.2.1 Slip layers and concrete curing

Bitumen emulsions are used to create a membrane between layers of concrete, the objective being to retain the strength of the upper layer by preventing water seepage into the lower layers. This avoids rigid adhesion between
layers of different ages and strengths and helps to produce a stronger upper layer by preventing water absorption into the lower layers. Bitumen emulsion is also sprayed onto the top surface of freshly laid concrete to prevent the evaporation of water.

9.8.2.2 Protective coats
Bitumen emulsions are used for protecting buried concrete, pipelines and ironwork. To enhance the adhesive and cohesive characteristics of the cured binder film, a polymer modified emulsion is normally used.

9.9 Trends and new developments in the use of bitumen emulsions
Although bitumen emulsion based techniques are known to be useful for all classes of roads, their use may also be possible in the following applications:

- asphalts for heavily trafficked roads (based on the use of polymer modified emulsions)
- warm emulsion based mixtures, to improve both their maturation time and mechanical properties
- half-warm technology, in which aggregates are heated up to 100°C, producing mixtures with similar properties to those of hot asphalts
- high performance surface dressing (e.g. glass fibre reinforced chip seals, scrub seals with modified emulsified binder and special microsurfacings) on heavily trafficked roads.

9.10 Further information on bitumen emulsions
For further information on the subject of bitumen emulsions, see Bitumen Emulsions – General Information and Applications (SFÉRB, 1991), which is available in both French and English.

References


World Road Association (Technical Committee Flexible Roads [C8]) (1999) Use of Modified Bituminous Binders, Special Bitumens and Bitumens with Additives in Road Pavements. World Road Association–PIARC, Paris, France.
10.1 Introduction

Aggregate is the largest constituent in asphalts, typically 92–96% by mass; the type of aggregate, its mineralogy, and physical and chemical properties will have a significant impact on asphalt performance.

Suitable aggregates, and their properties, for use in conforming to European asphalt mixtures can be specified by using EN 13043 (BSI, 2002). This standard defines aggregate as a ‘granular material used in construction’, and separates this into one of three types – natural, manufactured or recycled – defined as follows:

- Natural aggregate: ‘aggregate from mineral sources that has been subjected to nothing more than mechanical processing’ (e.g. crushed rock, sands and gravel, often referred to as primary aggregate).
- Manufactured aggregate: ‘aggregate of mineral origin resulting from an industrial process involving thermal or other modification’ (e.g. blast furnace slag).
- Recycled aggregate: ‘aggregate resulting from the processing of inorganic or mineral material previously used in construction’ (e.g. reclaimed asphalt).

Further categorisation of aggregates is given by the description for particle size:

- coarse aggregate: substantially retained on a 2 mm test sieve
- fine aggregate: substantially passing a 2 mm test sieve
- all-in aggregate: a combination of coarse and fine aggregates
- filler aggregate: substantially passing a 0.063 mm test sieve.

In the UK, the Specification for Highway Works (SHW), Volume 1, Clause 901.3 (Highways Agency et al., 2008), states that natural, recycled unbound and manufactured (artificial) aggregates for use in asphalts shall be...
clean, hard and durable and shall comply with BS EN 13043 [BSI, 2002]. Where recycled coarse aggregate or recycled concrete aggregate is used in bituminous mixtures, it shall have been tested in accordance with SHW Clause 710 and the content of other materials (Class X) including wood, plastic and metal shall not exceed 1% by mass.

The majority of asphalts are produced with natural crushed rock aggregate, although the trend is for ever greater use of recycled aggregate as a replacement, usually for economic or environmental reasons.

In the UK, the demand for construction aggregate from primary sources in 2011 was 145.9 million tonnes (Mt), comprising about 91 Mt of crushed rock and 55 Mt of sand and gravel (Bide et al., 2013). Of this, about 19.3 Mt of crushed rock (one-fifth of all the crushed rock produced) and 1.9 Mt of sand and gravel were coated with bitumen to produce asphalt. Figure 10.1 shows the relative proportions of crushed rock used for all construction activities in 2011, with asphalt referred to as ‘roadstone coated’.

10.2 Origin and type
The Earth is thought to have formed about 4.6 billion years ago, when the solar system was young. At that time the surface was very hot, but eventually, after cooling, the Earth’s crust formed; today the Earth’s crust is typically 7–70 km thick.

![Figure 10.1 UK production (90.9 Mt) of crushed rock by end use 2011 (Bide et al., 2013)](image-url)
The Earth’s crust is composed of ‘plates’ that, due to forces within the Earth, move relative to each other, resulting in the formation of mountains, ocean trenches and ridges, in a process known as ‘plate tectonics’. In addition to plate tectonics, processes of erosion, weathering, and chemical and biological action have been operating over the history of the Earth, resulting in the rocks encountered today.

The formation of rocks by these processes is known as the rock cycle, and it is summarised in Figure 10.2. Overall, these processes form igneous, metamorphic and sedimentary rocks.

### 10.2.1 Rock types

There are three main types of rock: igneous, sedimentary and metamorphic.

#### 10.2.1.1 Igneous rock

Igneous rock results from the solidification of the molten magma at or beneath the Earth’s surface. The magma is either extruded onto the surface (volcanic, extrusive) through lines of weakness in the crust as lava, or it may solidify at depth to form large igneous bodies underground (plutonic). Those formed beneath but close to the Earth’s surface, usually as smaller deposits, are known as hypabyssal.

Lavas extruded at the surface cool quickly and are characterised by their small crystalline structure (e.g. basalt, rhyolite), while intrusive (hypabyssal and plutonic) lavas cool much more slowly, and are characterised by larger crystal structures (e.g. granite, gabbro).
Igneous rocks can be defined by their mineralogical composition (Figure 10.3) and crystal size. Rhyolite and granite, for example, have the same composition (acidic) but very different crystal sizes, and therefore have different names, as is the case also for basalt and gabbro (basic).

The silica ($\text{SiO}_2$) content of igneous rocks is sometimes used to categorise rocks further as being acidic, intermediate, basic or ultrabasic on the basis of the following percentages:

- $>65\%$ by mass – acidic
- $52–65\%$ by mass – intermediate
- $45–52\%$ by mass – basic
- $<45\%$ by mass – ultrabasic.

(Note: the terms acidic and basic used here are geology expressions and do not indicate a pH level.)

Sometimes, the slowly cooling igneous mass deep down is moved by Earth movements higher into the crust where it cools more rapidly (i.e. two or more stages of crystallisation), resulting in a porphyritic texture – large crystals surrounded by smaller ones. Porphyritic rocks are frequently used as aggregates.

Major sources of igneous rock for the production of asphalt include

- basalt
- dolerite

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**Figure 10.3** Rock type according to mineralogy

<table>
<thead>
<tr>
<th>Volcanic:</th>
<th>Plutonic:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rhyolite</td>
<td>Granite</td>
</tr>
<tr>
<td>Dacite</td>
<td>Granodiorite</td>
</tr>
<tr>
<td>Andesite</td>
<td>Diorite</td>
</tr>
<tr>
<td>Basalt</td>
<td>Gabbro</td>
</tr>
<tr>
<td>Komatite</td>
<td>Peridotte</td>
</tr>
</tbody>
</table>

**Silica ($\text{SiO}_2$): % by mass**

- Orthoclase
- Quartz
- Plagioclase
- Ca-rich
- Na-rich
- Pyroxene
- Olivine
- Amphibole
- Muscovite
- Biotite

**Minerals: vol. %**

- Orthoclase
- Quartz
- Plagioclase
- Na-rich
- Ca-rich
- Pyroxene
- Olivine
- Amphibole
- Muscovite
- Biotite

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10.2.1.2 Sedimentary rock

Sediments evolve from

- the weathering and transport of particles (in air or water) and resultant sedimentation
- the precipitation of minerals from water
- the accumulation of biogenic matter (fossils).

The resultant sediments are then lithified or stuck together, in the lengthy process of diagenesis, in which the vast accumulations of sediments are transformed into rock. This can often be seen as a series of ‘beds’ or layers, which are characteristics of sedimentary rocks.

Sediments can now be seen in rivers, shallow seas and beaches. Glacial sand and gravel deposits can be found around many parts of the UK. These are associated with old river terraces formed from glaciers melting and depositing sediments during and at the ends of the ice ages. These sediments are unconsolidated (not stuck together), and can be easily extracted as sands and gravels.

Once sediments have been lithified and consolidated they become rocks (e.g. sands become sandstone, and sands and gravels become conglomerate).

For example, sandstone is formed from the cementation of accumulated sand that has been transported long distances and predominantly deposited in shallow seas, beaches and rivers. In the UK, the term ‘gritstone’ is used to describe a coarse sandstone in which the grain size is typically greater than 0.5 mm. Conglomerate is a rock formed from the cementation of sand and gravel that occurred in rivers and on beaches. Limestone is complicated, as it may be formed from a combination of fossil fragments and precipitated calcium carbonate, but always in marine conditions. Limestone can be very pure, consisting of a very high percentage of calcium carbonate, but, after formation, some of the calcium ions may be replaced by magnesium ions, which may be dissolved in the groundwater, thus transforming the limestone into a dolomite.

Major sources of sedimentary rock for the production of asphalt include

- limestone
- dolomite
- sandstone (gritstone).
10.2.1.3 Metamorphic rock

Metamorphic rock results from existing rock (sedimentary, igneous or metamorphic) being subjected to increased heat and pressure. The appearance and properties of these rocks are dependent on the varying degrees of heat, pressure and fluid activity to which they have been subjected. Small amounts of heat and pressure lead to minor textural changes (e.g. slate), and greater amounts result in substantial changes (e.g. gneiss). It should be noted that metamorphic rocks have not been melted, otherwise they would be igneous rocks. Rather, they are changed in the ‘solid state’; that is, the overall chemical composition of the rock stays the same but the contents are rearranged, resulting in textural changes and growth of minerals that are stable at set temperatures and pressures. Metamorphic rocks often tend to be layered or fissile, which often makes them unsuitable for aggregate in asphalt.

Major sources of metamorphic rock for the production of asphalt include meta-quartzite.

10.2.2 Petrographic examination and classification of rocks

A basic procedure for the petrographic examination and classification of rocks for use as aggregates is given in EN 932-3 (BSI, 1997). Table 10.1 is derived from this standard, and lists some common rock types classified according to their origin. Table 10.2 shows a simplified petrographic report of a plutonic igneous rock using this standard.

10.2.3 Manufactured aggregates

Some asphalt aggregates are manufactured, typically as a by-product of other industrial processes (e.g. steel slag and blast furnace slag). Some of these aggregates have properties that are very beneficial for asphalts (e.g. skid resistance from calcined bauxite) and have a good history of use, while others, particularly those with little or no history of use, will require

<table>
<thead>
<tr>
<th>Table 10.1 Simplified petrographic terms for aggregate from EN 932-3</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Igneous</strong></td>
</tr>
<tr>
<td>Extrusive/hypabyssal</td>
</tr>
<tr>
<td>Dacite</td>
</tr>
<tr>
<td>Basalt</td>
</tr>
</tbody>
</table>

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greater testing and need careful consideration in asphalt design and its application.

### 10.2.4 Recycled aggregates

Aggregate can be formed using recycled materials from other sources: for example, the use of bottom ash from municipal incinerators, foundry sand, recycled rail ballast, construction and demolition waste and milled asphalt from roads (the main source of recycled aggregate used in the production of asphalts).

The properties of these aggregates need to be carefully considered for inclusion in asphalt, not only with regard to their physico-chemical properties but also their suitability for use and their impact on the natural environment. The use of reclaimed asphalt (RA, but sometimes referred to as RAP from ‘reclaimed asphalt pavement or planings’) as aggregate poses the least potential risk because when it was originally manufactured it was almost certainly identified as suitable for incorporation in asphalt, and its use within European-compliant mixtures is controlled by EN 13108-8:2005 (BSI, 2005a).

The move towards performance specifications in European standards for aggregate and asphalt and the production of recycled aggregates following national quality protocols will minimise concerns and allow recycled aggregates and aggregates from primary sources to compete equally. Such an example is the quality protocol Aggregates from Inert Waste: End of Waste Criteria for the Production of Aggregates from Inert Waste.
applicable in Wales, Northern Ireland and England, which has the stated aim ‘to provide increased market confidence in the quality of products made from waste and so encourage greater recovery and recycling’.

10.3 Aggregate processing

It is a combination of the nature of the raw material and the requirements of the market that determine the type of processing required. The essential elements of aggregate processing are breaking the rock into smaller sizes or fractions, and then separating them into the different sizes (aggregate) required by the market. The useable sizes can range from a maximum dimension of about 1 m, such as large stone blocks (armourstone) for coastal defences – which do not go through the crushing process – to fractions of a millimetre, where the rock may have gone through a multi-stage crushing process to produce the correct size and shape of aggregate.

The stages from extraction through to processing determine the aggregate shapes and sizes, but can also affect aggregate quality and integrity. Establishing a stable, controlled production process reduces the variability of products and can be achieved through the application and operation of a formal quality system.

10.3.1 Extraction

The extraction process to recover the material will depend on the characteristics of the deposit. Very often, it will be covered with soils and other material (overburden) that is not suitable for use as aggregate. This has to be removed and stored for later use (e.g. in the quarry restoration process once extraction has finished).

Sand and gravel deposits (Figure 10.4) are usually the result of ancient glacial river systems, and often occur as thin strips or sheets in lowland or offshore areas. Extraction of land and marine based deposits can be over quite large areas, but are usually not more than a few metres deep. Marine deposits (approximately 8% of Britain’s primary aggregate production) are mainly recovered by suction dredging, whereas the terrestrial deposits can be recovered using hydraulic excavators or draglines. The deposit can either be worked wet (recovered from under the water) or dry (using pumps to keep the extraction area dry).

Hard rock deposits (Figure 10.5) are usually won using conventional drill and blast methods using, typically, 115–150 mm diameter holes filled with bulk ammonium nitrate fuel oil explosives, although some weaker rocks can be excavated directly into dump trucks or broken using a ‘ripper’ attachment on an excavator or on the back of a bulldozer. The objective is to fragment
Figure 10.4  Aggregate Industries’ Newbold Quarry: a sand and gravel deposit. (Courtesy of Aggregate Industries)

Figure 10.5  Aggregate Industries’ Duntilland Quarry Scotland: a large igneous rock deposit. (Courtesy of Aggregate Industries)
the rock to a size suitable for loading and primary crushing (Figure 10.6),
and then transporting to the plant for further processing, all using methods
that are both safe and minimise environmental impacts.

Once the material has been won, it is transported to the processing plant by
way of a conveyor belt or dump truck. The increased usage of modular,
mobile processing equipment has meant that some or all of the required
processing and loading of trucks for delivery may now take place within the
extraction area itself. This equipment is able to follow the extraction area as it
develops within a quarry, but can also be moved from quarry to quarry fairly
easily. However, many quarries, particularly the larger ones, will have static
fixed plant, where the raw feed material is processed to produce the
products required for the market, remote from the extraction area and
conveniently located for road transport or rail access.

10.3.2 Crushing
10.3.2.1 Crushing process
Crushing is usually the first step of processing after extraction of rock, and
reduces large rock, mechanically, to more useful conforming and saleable
sizes, typically 75 mm and less. Crushing is usually a sequence and can
entail one, two, three or more crushing phases, known as primary, secondary and tertiary crushing.

The first stage – primary crushing – is the first reduction in stone size. This is achieved using either a compression crusher (jaw or gyratory crusher types, see Figure 10.9) or impact crusher (see Figure 10.10). The primary crusher output is fed to a secondary crusher (e.g. a cone crusher) for further reduction in stone size. This is followed by screening its output into appropriate aggregate sizes or reprocessing with further crushing cycles (tertiary) using cone crushers to achieve the desired shape and sizes. Figure 10.7 shows a primary crusher being fed with blast rock.

Some operations provide a stage either immediately prior to or after primary crushing known as ‘scalping’.

**Scalping**

Scalping is a coarse screening to remove the 40 mm and down fraction, and is often carried out prior to primary crushing. This removes weak rock, clays and silts, to ensure a cleaner, good quality rock feed and to improve primary crusher performance. However, on large primary crushing installations it is more common to carry out the scalping process once the rock has undergone primary crushing.
On some sites, this scalped material is blended with a range of different sized aggregates, to produce a lower grade material, sometimes referred to as crusher runs. In very dirty rock deposits, the product derived from the scalping operation will often be diverted as a waste material.

**Simplified crushing process flow**

Figure 10.8 is a simplified process flow through the crushing stages before final screening and storage into the various product sizes needed by the quarry. In this process, raw material is first screened through a vibrating deck called a ‘vibrating grizzly’, to remove small pieces of rock along with any clay and soils followed by primary, secondary and tertiary crushings that result in the desired aggregate sizes of appropriate grade and quality, the types of crusher being dependent on the rock type.

10.3.2.2 Crusher types

**Compression crushers**

Compression crushers apply pressure to the rock between fixed and moving parts in a repeated squeezing action, reducing the rock in size until it is small enough to pass through a crushing chamber. There are two types of compression crusher (Figure 10.9).

- **Jaw crushers** work by squeezing rock in the crushing chamber between two jaws, one of which is fixed while the other is angled and moves backwards and forwards applying the crushing force. The distance between the jaws reduces as the crushed rock travels down the chamber.
- **Gyratory crushers** work by having a gyrating crushing mantle set within a bowl, and provide a continuous crushing action. Rock is crushed and reduced in size as it travels down the crushing chamber.
Figure 10.9 (a) Jaw and (b) gyratory crushers. (Courtesy of Matthew Allen)
Compression crushers, although relatively economical, can have a tendency to create poor particle shapes for some rock sources. In these instances, impact crushers may be used for primary crushing, but at a slightly higher cost per tonne.

**Impact crusher**

In impact crushers, as the name suggests, the breaking down of the rock is achieved when the high speed rotating blow bars or swinging metal hammers impact on the cascade of rock discharged into the crusher. Large rocks are either broken down by this primary action or by the secondary action of rock against rock as particles are impelled at high speed from the process. Figure 10.10 shows the simplified layout of a horizontal impact crusher.

**Cone crusher**

Cone crushers (Figure 10.11) are sometimes called gyratory cone crushers, and although they can be used as a primary crusher they are more often used as secondary or tertiary crushers. In operation as a secondary or tertiary crusher, pre-crushed aggregate is fed through the top of a cone crusher between two conical surfaces, called the mantle and the concave liners, into the crushing chamber.

![Horizontal action impact crusher](image-url)
The mantle rotates eccentrically within the crushing bowl, and this gyrating action causes the gap between the mantle and liners to change dimension and perform the actual crushing as the rock falls through.

10.3.3 Screening

Once scalped and crushed, the multi-sized rock is passed over a number of different screens that are typically double or triple decked, to sort the aggregate into individual aggregate sizes. These sizes will reflect the aggregate product range, and are typically 40, 28, 20, 14, 10 and 6.3 mm, with the finest fraction, less than 4 mm, being referred to as crushed rock fines.

Screen decks are modular units usually manufactured from rubber or polyurethane, but perforated metal plate and woven wire screen decks can be used to produce a high grade, low flake product for use as surface dressing or high specification asphalt surface course aggregates.

10.4 Coarse and fine aggregates

The performance of an asphalt is largely predetermined by the characteristics of its components: bitumen, aggregates and air voids. However, for this marriage to be successful, the properties of the aggregates need to be known, fully understood and adequately specified.

The properties can be considered within a group of five broad classifications, namely

- geometrical properties (e.g. size, shape and particle packing)
mechanical properties (e.g. strength and hardness)
physical properties (e.g. particle density and water absorption)
chemical properties (e.g. adhesion)
durability, weathering properties (e.g. freeze–thaw resistance).

Some of these properties (i.e. those for geometric requirements, e.g. grading, size, shape and cleanliness (clay and silt content)) are partly controlled, and are a function of the aggregate processing, while others (e.g. physical, mechanical and chemical properties) are an inherent characteristic of the aggregate unaltered by quarry processing.

In Europe, the standard EN 13043 (BSI, 2002) provides a range of categories for aggregate properties and the associated test methods: this allows purchasers and designers of asphalts the means to select appropriate limiting values for the wide range of aggregates used in asphalts. It is usually the responsibility of the purchaser to determine and define appropriate EN 13043 categories for properties that are relevant to the particular end-use of an aggregate. Where a property is not relevant to both the mixture or its application, the specifier should identify this using a ‘no requirement’ description.

Aggregate properties referred to in EN 13043 and their applicable test methods are outlined in the following sections: note that this standard includes mechanical properties as a part of physical properties, and the following sections have been ordered in line with EN 13043.

10.4.1 Properties and testing of coarse, fine and all-in aggregate

10.4.1.1 Geometrical properties

Aggregate size
This is described using the designation \(d/D\), in terms of lower (\(d\)) and upper (\(D\)) sieve sizes: for example, 4/10 mm single size.

Grading (particle size distribution)
Aggregate gradings are determined in accordance with EN 933-1 (BSI, 2012a) and specified as appropriate to the aggregate sizes \(d/D\) from tables within the standard. Grading categories are used and expressed as follows: \(G_C\) for coarse aggregate, \(G_F\) for fine aggregate and \(G_A\) for all-in aggregate. In addition, for coarse aggregates the numerical specification limits for the minimum percentage by mass passing the sieve size represented by \(D\) and the maximum amount passing the sieve size represented by \(d\) are stated. For fine (\(G_F\)) and all-in (\(G_A\)) aggregates, only the minimum percentage appropriate to the sieve size \(D\) is stated.

For example, when specifying aggregate size it is sufficient to quote the aggregate size and grading category (e.g. 10/20, \(G_C\) 85/35). From this
example, it can be determined that the product is a coarse aggregate \((G_C)\) and will have between 0 and 15\% by mass of aggregate larger than a 20 mm sieve size \((D)\), this fraction being known as the ‘oversize’. Also, between 0 and 35\% by mass of aggregate is less than the 10 mm sieve \((d)\), this fraction being known as the ‘undersize’.

**Fines content**
The fines content \((f)\) specified in EN 13043 relates to the percentage by mass passing a 0.063 mm sieve for coarse and fine aggregates, and is expressed as \(f_x\), where \(x\) is the maximum percentage passing the 0.063 mm sieve. It is determined as part of the washing and sieving test in accordance with EN 933-1.

**Fines quality**
Some fines such as clay minerals are harmful to asphalt and can occur in the fines fraction of some aggregate sources. The presence of clays leads to high rates of water absorption, swelling and increased moisture sensitivity of the mixture, resulting in stripping.

The methylene blue test specified in EN 933-9, Annex A (BSI, 2009a), is used to assess the quantity of potentially harmful fines in fine aggregate for the 0/0.125 mm fraction. The test is carried out by adding an aqueous solution of methylene blue dye to a fine aggregate sample held in suspension with water, and measuring the quantity of added dye at the point where adsorption of the dye has stopped. The test exploits the fact that clay minerals absorb basic dyes from aqueous solutions, and the higher the quantity of dye absorbed, the greater is the quantity of potentially harmful fines present in the fine aggregate.

Where the fines content of either fine or all-in aggregate \((D \leq 8 \text{ mm})\) is greater than 3\%, the amount of harmful fines \((\text{e.g. swelling clays})\) needs to be considered. A measure of such presence is determined using the methylene blue test according to EN 933-9. This method describes separate tests for two size fractions

- 0/2 mm
- 0/0.125 mm (Annex A).

The results are reported with the units of g/kg, and expressed according to the appropriate category as \(\text{MB}_v\) value for the 0/2 mm fraction, and \(\text{MB}_F\) value for the 0/0.125 mm fraction. EN 13043 (BSI, 2002) specifies the requirement for the 0/0.125 mm fraction to be used for the assessment of ‘fines quality’ for fine and all-in aggregates.
The French publication *The Use of Standards for Hot Mixes* (Sétra, 2008) recommends a limiting value of $MB_{F10} (10 \, g/kg)$ for the ‘quality of fines and particles $< 0.125 \, mm$ in sands and graded aggregate’.

**Shape of coarse aggregate**

The shape of aggregate particles is a very important property, and has a large bearing on asphalt performance, more cubical shaped particles providing better aggregate interlock and denser particle packing. The flakiness index (FI) and the shape index (SI) are the specified characteristics determined using the method in EN 933-3 (BSI, 2012b) and EN 933-4 (BSI, 2008a), respectively.

The flakiness index is the percentage of particles of a coarse aggregate that have a thickness of less than one-half of the nominal size ($D$) using specific bar sieves, and is expressed as a percentage of the mass of the test portion as $F_{I \text{value}}$. The greater this value, the flakier is the aggregate. Good cubical shaped aggregates would have an FI of 20 or less.

The shape index is a measure of individual particles in a sample of aggregates classified on the basis of the ratio of their length to thickness using a measuring gauge. The shape index is the mass of particles when their length divided by their thickness exceeds 3, expressed as a percentage of the total mass of sample. Figure 10.12 shows cubical, flaky and elongated particles of an igneous aggregate.

**Percentage of crushed and broken surfaces in coarse aggregates**

This property is not considered for crushed rock particles, but is particularly relevant to natural gravels, which may contain either a proportion or be totally composed of rounded particles. It is usually specified as a control to ensure that a good mechanical interlock is possible in the asphalts into which they are to be incorporated, where this property is most usually necessary.

![Figure 10.12 Aggregate particle shapes. (Courtesy of Benjamin Allen)](image-url)
The content of crushed or broken particles, including totally crushed particles and the percentage of totally rounded particles, is determined using the method in EN 933-5 (BSI, 1998a), and declared in accordance with the relevant category and expressed as $C_{x/y}$, where $x$ denotes the percentage of totally crushed or broken and partially crushed or broken particles by mass and $y$ denotes the percentage of totally rounded particles by mass.

**Angularity of fine aggregate**

The angularity of fine aggregate is determined in accordance with EN 933-6 (BSI, 2014a). It is an indirect method of measuring fine aggregate angularity using a flow test, and records the time needed for a given amount of fine aggregate to flow out of a particular shaped vessel: the result is expressed as a flow coefficient with the symbol $E_{cs}$ value. Rougher surfaces or a more angular particle shape will result in longer flow times: this can be interpreted in terms of asphalt mixture design as needing more compactive energy to achieve full compaction but likely to prove more stable in use.

**Current areas of research**

Test methods for the characterisation of geometrical properties rely predominantly on dimensional measurements using sieves, gauges and visual judgement (crushed and uncrushed). They are well tried and proved within the limits of acceptability. However, advanced image analysis techniques have been developed in the past decade that more precisely characterise the shape, angularity and texture of aggregates.

These techniques involve the acquisition, processing and analysis of images of aggregate particles. Some of the techniques used to capture two dimensional images include optical microscopy, digital imaging and laser scanning. Aggregate geometrical characteristics determined by these techniques have been related to their mechanical properties and those of the resultant asphalt. They have also been employed to measure aggregate surface texture and show how this compares with friction and skid resistance (Dunford, 2013).

More recently, X-ray computed tomography techniques have been used to measure the internal structure of asphalt. This non-destructive technique is used to obtain three dimensional images of asphalt specimens, and is based on the attenuation of X-ray radiation as a function of the material density. The technique allows the visualisation of the internal structure consisting of aggregates, bitumen and air voids. Aggregate degradation as a result of traffic (loading) has also been monitored (Kutay et al., 2010).

It is clear that over the next decade many of these technological advancements for measuring aggregate properties will become more widely used and possibly be incorporated into aggregate specifications.
10.4.1.2 Physical properties

Resistance to fragmentation of coarse aggregate

Aggregates must be hard and tough enough to resist fragmentation (degradation and disintegration) when being stockpiled, transported and mixed in production processes and also to withstand the stresses from their in-service applications.

The Los Angeles (LA) test to EN 1097-2 (BSI, 2010a) determines an aggregate’s resistance to fragmentation by degrading a sample placed in a rotating steel drum with 11 large (400–445 g) steel balls (45–49 mm in diameter). As the drum rotates, the aggregate particles are broken down due to impacts with the steel balls and the other aggregate particles in the drum.

The result is reported as LA-value, calculated from the mass of aggregate that has broken down to smaller sizes expressed as a percentage of the total mass of aggregate. The lower the value, the tougher and more fragmentation resistant is the aggregate. A value of LA30 or less is needed to make a good roadstone aggregate.

Figure 10.13 shows the degree of fragmentation following the Los Angeles test on a 10/14 mm igneous aggregate.

Resistance to polishing of coarse aggregate for surface courses

The action of tyres on the road surface results in polishing of the exposed surface aggregate, and the state of polish is a key factor affecting the road
surface’s skid resistance. An aggregate’s resistance to polishing is therefore a most important characteristic, and aggregates with a high resistance to polishing sell at a premium relative to those with less resistance. Its resistance depends on its micro-texture characteristics, which are determined using the polished stone test in EN 1097-8 (BSI, 2009b), under standardised conditions similar to those occurring at the surface of the road.

The apparatus is illustrated in Figure 10.14: four test specimens, each consisting of between 36 and 46 (passing a 10 mm sieve but retained on a 7.2 mm flaky sieve) aggregate particles, bound in a rigid resin matrix (Figure 10.15) are clamped around the periphery of a large road wheel with which a second, smaller, solid rubber-tyred wheel has contact. Both wheels rotate, and a corn emery abrasive, similar to that used on a coarse emery paper, is fed into the interface between the two wheels, and the aggregate samples are polished for 3 h. The first cycle is followed by a second cycle of 3 h of polishing with fine emery flour instead of the corn emery. The samples are removed from the apparatus, and their degree of polishing measured using the Transport Research Laboratory portable skid resistance tester.

The results are expressed from the mean of the four test specimens as the aggregate polished stone value (PSV), in the form PSV\textsubscript{value}. The higher the number, the greater the resistance to polishing, and an aggregate having a PSV of 60 or more is generally regarded as providing a good level of

![Figure 10.14](image)

**Figure 10.14** Simple diagram of polished stone value (PSV) machine. (Courtesy of Matthew Allen)
polishing resistance. The in-service life of a higher PSV aggregate (i.e. the length of time from first trafficking to the time when the road surface has polished to the extent that it no longer has its anticipated degree of skid resistance) will be longer than that of a lower PSV aggregate.

Table 10.3 is an extract from the Highways Agency’s Interim Advice Note (IAN) 156/12 (Department for Transport, 2012). IAN 156/12 amended HD 36/06 (Highways Agency et al., 2006) for the aggregate PSV after the incorporation of the results of recent research. It shows how the Agency specifies requirements for the PSV of aggregates used in ‘hot applied thin surface course’ systems on the UK’s trunk road network (which includes motorways) as a function of

- the traffic frequency (commercial vehicles/lane/day)
- the site (site description in terms of different degrees of risk to road users)
- the investigatory level (IL): a factor (dictated by the skid resistance policy for the road) to maintain an adequate in-service performance in relation to the skid resistance of the surface course.

Resistance to surface abrasion
For aggregates used in surface course mixtures, it is also important to ensure that the aggregate will not be worn away too quickly in service by specifying
<table>
<thead>
<tr>
<th>Site category</th>
<th>Site description</th>
<th>Investigatory level (IL)</th>
<th>Minimum PSV required for a given IL, traffic level and type of site</th>
</tr>
</thead>
<tbody>
<tr>
<td>A1</td>
<td>Motorways where traffic is generally free flowing on a relatively straight line</td>
<td>0.30</td>
<td>50</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.35</td>
<td>50</td>
</tr>
<tr>
<td>A2</td>
<td>Motorways where some braking regularly occurs</td>
<td>0.35</td>
<td>50</td>
</tr>
<tr>
<td>B1</td>
<td>Dual carriageways where traffic is generally free flowing on a relatively straight line</td>
<td>0.30</td>
<td>50</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.35</td>
<td>50</td>
</tr>
<tr>
<td>B2</td>
<td>Dual carriageways where some braking regularly occurs</td>
<td>0.35</td>
<td>50</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.40</td>
<td>55</td>
</tr>
<tr>
<td>C</td>
<td>Single carriageways where traffic is generally free flowing on a relatively straight line</td>
<td>0.35</td>
<td>50</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.40</td>
<td>50</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.45</td>
<td>53</td>
</tr>
<tr>
<td>G1/G2</td>
<td>Gradients &gt;5% longer than 50 m as per the Highways Agency’s guidance HD 28 (‘skid resistance’)</td>
<td>0.45</td>
<td>55</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.50</td>
<td>60</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.55</td>
<td>68+</td>
</tr>
<tr>
<td>K</td>
<td>Approaches to pedestrian crossings and other high-risk situations</td>
<td>0.50</td>
<td>65</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.55</td>
<td>68+</td>
</tr>
<tr>
<td>Q</td>
<td>Approaches to major and minor junctions on dual carriageways and single carriageways where frequent or sudden braking occurs but in a generally straight line</td>
<td>0.45</td>
<td>60</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.50</td>
<td>65</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.55</td>
<td>68+</td>
</tr>
<tr>
<td>R</td>
<td>Roundabout circulation areas</td>
<td>0.45</td>
<td>50</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.50</td>
<td>68+</td>
</tr>
<tr>
<td>S1/S2</td>
<td>Bends (radius &lt;500 m) on all types of road, including motorway link roads; other hazards that require combined braking and cornering</td>
<td>0.45</td>
<td>50</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.50</td>
<td>68+</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.55</td>
<td>HFS</td>
</tr>
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</table>

Department for Transport (2012)
a maximum aggregate abrasion value (AAV) determined in accordance with EN 1097-8:1999, Annex A [BSI, 2009c].

In this test, a sample is prepared using specimens of 6.3/14 mm aggregate particles, retained on a 10.2 mm grid sieve, bound in a rigid resin matrix. This is weighed then mounted face down against a large circular rotating steel disc. Coarse abrasive sand is fed between the disc and the faces of the resin-bound aggregates as the wheel rotates for 500 revolutions, causing the sand to abrade the aggregate specimens. At the end of the test, the sample is removed from the machine, cleaned and re-weighed. The percentage of mass lost is expressed as AAV\textsubscript{value}. The lower the value, the more abrasion resistant is the aggregate. Figure 10.16 shows two gritstone AAV specimens, before and after testing for the AAV, and illustrates the abrasive effects of the test on the aggregate particles.

For a particular application, the selection of an appropriate aggregate AAV will be dependent on the amount of traffic, the design life and the mixture type. Table 10.4 illustrates this with an example from the UK’s Highways Agency Design Manual for Roads and Bridges, HD 36/06 (Table 3.2), and shows the specification of appropriate maximum AAV levels for a number of asphalt types.

**Resistance to wear of coarse aggregate**

The wear of aggregates at particle points and edges due to breakdown as a result of attrition is an important aggregate property, particularly for mixtures such as porous or open graded asphalt with many point-to-point aggregate contacts.

For this aggregate property, the micro-Deval test to EN 1097-1 [BSI, 2011a] is used as a control measure, and its result calculated and reported as a micro-Deval coefficient, \( M_{\text{DE value}} \).
The test method uses a rotating steel vessel containing an aggregate sample of predetermined mass, a 5 kg charge of small (9.5–10.5 mm dia.) steel balls and a volume (2.5 litres) of clean water. After 1200 revolutions, about 2 h of rotation, the now worn aggregate sample is discharged from the vessel and reweighed, and the percentage loss of mass expressed as $M_{DE}$ value. The lower the numerical value, the more resistant to wear through attrition is the aggregate.

Figure 10.17 shows the effect and degree of ‘particle rounding’ that occurs as a result of this test on an igneous aggregate.

| Traffic at the design life: commercial vehicles/lane/day | 
| <250 | 251–1000 | 1001–1750 | 1751–2500 | 2501–3250 | >3250 |
| 14 | 12 | 12 | 10 | 10 | 10 |
| Maximum AAV for aggregates for hot rolled asphalt and surface dressing, and for aggregate in slurry and microsurfacing systems |
| 16 | 16 | 14 | 14 | 12 | 12 |
| Maximum AAV for aggregate in thin surface course systems, exposed aggregate concrete surfacing and coated macadam surface course |

Highways Agency et al. (2006)
Resistance to abrasion from studded tyres of coarse aggregates to be used for surface courses

In countries where vehicles use studded road tyres in winter, this results in accelerated wear of the asphalt at the road surface. As a consequence, it becomes an important factor in the specification and selection of aggregates used in surface course mixtures.

Where required, the resistance to abrasion from studded tyres for the coarse aggregate fraction in asphalt is determined in accordance with EN 1097-9 (BSI, 2014b).

The test is performed on coarse aggregate between 11.2 and 16 mm that is rotated (90 ± 3 rev/min) in a drum with steel abrasive balls in the presence of water.

After a specified number of revolutions, the abrasion loss rate of the aggregate is calculated and reported as a Nordic abrasion value, AN_value. The lower the numerical value, the more resistant to abrasion from studded tyres is the aggregate.

Particle density and water absorption

Particle density and water absorption are determined in accordance with EN 1097-6 (BSI, 2013a), Clause 7, 8 or 9, depending on the size of the aggregate.

- Clause 7: wire basket method for aggregate particles passing the 63 mm test sieve and retained on the 31.5 mm test sieve.
- Clause 8: pyknometer method for aggregate particles passing the 31.5 mm test sieve and retained on the 4 mm test sieve.
- Clause 9: pyknometer method for aggregate particles passing the 4 mm test sieve and retained on the 0.063 mm test sieve.

The tests give a measure for particle density expressed in the units Mg/m³ for

- the apparent particle density
- the oven-dried particle density
- the saturated and surface-dried particle density.

The aggregate’s water absorption after immersion for 24 h is expressed as WA_{24} value.

Bulk density

The bulk density of an aggregate is determined in accordance with EN 1097-3 (BSI, 1998b), and is expressed in Mg/m³.

A physical property of an aggregate not considered in EN 13043 is its thermal expansion and contraction (i.e. its thermal coefficient, α). As
aggregates represent about 85% of the total volume of a typical asphalt, the thermal coefficient of asphalt is greatly influenced by that of the aggregate. The thermal coefficient of any natural aggregate depends on its mineralogy. In general, siliceous aggregates with a high quartz content exhibit a high thermal coefficient \((10.01 \times 10^{-6} \text{ to } 13.00 \times 10^{-6} \text{ mm/mm/} ^\circ \text{C})\), whereas some pure limestones that consist mainly of calcite exhibit a lower value \((5.58 \times 10^{-6} \text{ to } 6.10 \times 10^{-6} \text{ mm/mm/} ^\circ \text{C})\) (Mukhopadhyay et al., 2007). The thermal coefficient of a bituminous binder is 10–20 times higher than that of the aggregate.

Thus, thermal stresses can develop in the binder film surrounding an aggregate particle as a result of differential thermal contraction. Under extreme low temperatures and/or when stiff binders are used, these thermal stresses may cause localised cracking in the asphalt. These cracks can then facilitate moisture entry into the asphalt, causing damage and a subsequent reduction in durability (El Hussein et al., 1998).

10.4.1.3 Durability
Water absorption value as a screening test for freeze–thaw resistance
The water absorption value is used as an indicator of an aggregate’s resistance to freezing and thawing action. If its water absorption is 2% or less, then an aggregate is considered as suitably freeze–thaw resistant without further testing.

Aggregates with water absorption values greater than 2% can be very resistant to freeze–thaw action, but need to be tested for this property more directly. Examples of this category include

- Jurassic limestone and sandstone
- blast furnace slags
- Permian limestone
- dolomite
- carboniferous sandstone.

Resistance to freezing and thawing
The vulnerability of an aggregate to damage from periods of freezing and thawing will depend on the climate, with the severity of any damage being related to the frequency and severity of these freeze–thaw cycles. This risk increases with the degree of water saturation, and is significantly increased with sea water.

EN 13043 gives guidance on the selection of an appropriate level of resistance to freezing and thawing based on climate, which can be

- Mediterranean
Also, the possible environmental conditions applying to the product in service are considered:

- frost free or dry situation
- partial saturation, no salt
- saturated, no salt
- salt (sea water or road surfaces)
- airfield surfacings.

The ability of an aggregate to withstand changes in volume as a result of conditions under freeze–thaw cycling will depend on its petrographic type and the size distribution of the pores within the aggregate.

EN 13043 gives two methods for determining an aggregate’s ability to resist freezing and thawing, often referred to as its soundness: EN 1367-1 (BSI, 2007) and EN 1367-2 (BSI, 2009d).

In the UK, the magnesium sulfate test is usually specified (EN 1367-2). In this test, particles of aggregate of size 10–14 mm are tested for soundness by subjecting pre-weighed samples to soaking and drying cycles in a magnesium sulfate solution, and measuring the mass of aggregate degraded to finer than that which passes a 10 mm test sieve, due to salt crystallisation and its associated cracking. This degradation is reported as a percentage, in the form MS\text{value}.

Aggregates for UK conditions are considered satisfactory for general purpose use if their magnesium sulfate soundness category is MS\text{25} or better (i.e. a lower value). Other categories of magnesium sulfate soundness may be more appropriate for climatic conditions outside the UK.

Vulnerable aggregates tend to be derived from highly weathered rocks and some conglomerates and breccias, and can include, for example,

- schist
- mica schist
- phyllite
- shale
- chalk.

10.4.1.4 Resistance to thermal shock

Aggregates are subjected to heating and drying as a part of the production of asphalt, and undergo an element of thermal shock as a consequence. EN 13043 provides a method, EN 1367-5 (BSI, 2011b), for determining the extent to which aggregates degrade by heating during the drying
process, and knowledge of this property can be useful when assessing new and potential resources with no previous history of use in asphalt, particularly recycled and manufactured aggregates.

The test involves the preparation of two aggregate samples. One sample is tested for resistance to fragmentation using the Los Angeles test (described in section 10.4.1.2), and the other sample is tested for fragmentation after exposure to thermal shock. Thermal shock involves soaking a sample of the aggregate in water and subjecting it to thermal shock by heating it to 700 °C for 3 min.

The difference between the LA coefficient before and after thermal shock is defined as ‘the loss in strength due to thermal shock’.

10.4.1.5 Affinity of coarse aggregates for bituminous binders

Good coating and adhesion of bitumen to aggregate is essential, but the loss of bond between aggregate and bitumen in service due to the presence of water, known as ‘stripping’, has a major detrimental effect on the integrity of an asphalt pavement. Although many factors affect the affinity of the aggregate for bitumen, it is accepted that the mineralogy and chemical composition of the aggregate are important contributory factors.

Generally, aggregates with high silica contents (acid) have a good affinity for water (hydrophilic) but not for bitumen (Figure 10.18). They are more likely to suffer from stripping after exposure to water than those with lower silica contents (basic) that have an affinity for bitumen rather than water (hydrophobic). Aggregate binder affinity for highly siliceous aggregates can be sufficiently improved with the addition of adhesion promoters.

Figure 10.18 Typical range of silica content (SiO₂) for some common rocks
EN 13043 requires the affinity of coarse aggregates to bituminous binders to be determined in accordance with EN 12697-11 (BSI, 2012c). The latter standard gives three methods:

A. A rolling bottle containing a loose bitumen-coated sample of aggregate, a glass rod device providing a stirring action, and water are used. After rolling for defined periods of time, the sample is removed from the roller bottle and visually examined for bitumen stripping.

B. A sample of bitumen-coated aggregate is immersed in distilled water for 48 h, and the number of particles that are no longer completely coated is assessed.

C. An aggregate sample is mixed with bitumen to give complete coverage of the aggregate. This prepared sample is first subjected to stripping in boiling water for 10 min. The proportion of exposed aggregate following this is determined by placing the sample in contact, for 5 min, with either hydrochloric acid (for calcareous aggregates) or hydrofluoric acid (for siliceous aggregates) of given concentrations. A titration method is used to determine the volume of hydrochloric acid or hydrofluoric acid consumed by the reaction of these reagents with the free surface of the aggregate. Defined calibration curves are used to estimate the proportion of the exposed aggregate surface from the volume of reagent consumed.

In the UK, the Ministry of Defence, Defence Infrastructure Organisation (DIO) Specification 13 (Ministry of Defence, 2009a) is specified in Table 3.1 using EN 12697-11 Method B, and the requirement is ‘not greater than 6 particles from a 150 particle test sample’.

There are other tests or means of evaluation. In the UK, the Highways Agency’s Specification for Highway Works, Series 900, Clause 953 (Highways Agency et al., 2008), describes the saturation ageing tensile stiffness (SATS) test, which is used to assess the durability of asphalts. In this test, cylindrical specimens of the asphalt are conditioned in a pressure vessel, stiffness levels being determined before and after conditioning, to establish a percentage of retained stiffness. It is essentially a measure of the adhesion between the aggregate and the bitumen where deterioration in asphalts due to a gradual loss of adhesion results in a reduction in mixture stiffness.

10.4.1.6 ‘Sonnenbrand’ of basalt
‘Sonnenbrand’ is a German word that translates into English as ‘sunburn’. It is used in connection with basalt to describe a particular type of rock decay that is present in some basalt that develops over time under certain atmospheric conditions. This visual observation can take place within months of
extraction or extend over many years. Affected rocks first show staining (grey/white-coloured spots) and efflorescence, and later develop hairline cracks that reduce the strength of the aggregate, in some cases resulting in total breakdown. When an aggregate is known or suspected of being afflicted with ‘sonnenbrand’, then the aggregate’s loss of mass and resistance to fragmentation is determined using the EN 1367-3 (BSI, 2001) and EN 1097-2 (BSI, 2010a) test methods.

10.4.1.7 Chemical
The chemical requirements for coarse and fine aggregates within EN 13043 are determined in accordance with EN 932-3 (BSI, 1997). This method employs a visual examination of the aggregate to determine the constituent rock or mineral types (see Table 10.2). Where a more quantitative study is required, then an analytical approach is needed using methods such as X-ray fluorescent techniques, principally to identify an aggregate’s oxide composition (see Figure 10.19).

Other chemical requirements are stated within EN 13043 essentially for slag (manufactured) aggregate and recycled aggregate. These test methods are based on EN 1744-1 (BSI, 2009e) procedures, falling under the following categories.

Coarse lightweight contaminators
The term ‘coarse lightweight contaminators’ refers to the presence of organic contaminants larger than 2 mm. These are not likely to occur in natural rocks, and are usually specified where manufactured or recycled aggregates are used.

Testing is carried out in accordance with Clause 14.2 of EN 1744-1, and the result expresses the amount of organic contaminants as a percentage of the coarse aggregate.

Constituents that affect the volume stability of blast furnace and steel slags
In the cases of blast furnace and steel slags, there is a specific requirement for establishing and measuring the volume stability of these materials

- air-cooled blast furnace slag aggregate shall be free from dicalcium silicate disintegration when tested in accordance with EN 1744-1
- air-cooled blast furnace slag aggregate shall be free from iron disintegration when tested in accordance with EN 1744-1
- the volume stability of steel slag aggregate shall be determined in accordance with EN 1744-1.
### AGGREGATE PROPERTIES SUMMARY DATA SHEET

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* All tests carried out are UKAS accredited unless otherwise denoted by *

Comments:

* Full report available upon request.

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**Figure 10.19** Summary of physical and chemical properties for a granite aggregate, part 1. (Courtesy of Aggregate Industries)
10.4.1.8 Summary of properties and testing: example of a producer’s property data sheet

Figures 10.19 and 10.20 give an example of a suite of tests carried out by a UK aggregates supplier on a granite aggregate for the UK market. This example includes not only physical and chemical properties applicable to the production of asphalts, and described in this chapter, but also includes properties relating to unbound aggregate mixtures, such as subbase for pavement foundations (described as Type 1, in Figure 10.19).

Figure 10.20 Summary of physical and chemical properties for a granite aggregate, part 2. (Courtesy of Aggregate Industries)
10.5 Filler aggregate

The main function of filler is to act as mastic when it is combined with the binder. It provides added stiffness and stability to the mixture, as well as acting as a packing mineral to control the void structure of the mixture.

Filler aggregate is defined in EN 13043 as ‘aggregate, most of which passes a 0.063 mm sieve, which can be added to construction materials to provide certain properties’, and is derived according to EN 13043 as being

- added filler, ‘filler aggregate of mineral origin, which has been produced separately’
- mixed filler, ‘filler aggregate of mineral origin, which has been mixed with calcium hydroxide’ or
- fines, ‘the particle size fraction of an aggregate which passes the 0.063 mm sieve’ (e.g. baghouse fines from the dust collection devices at asphalt mixing).

The requirements for filler aggregate in EN 13043 with regard to geometrical, physical and chemical properties and consistency of production apply to mixed filler, added filler and fines from the fine aggregate when this content exceeds 10% by mass of the fine aggregate.

Sometimes added filler with requirements for mineral type or mixed fillers are specified for the purpose of improving a mixture’s adhesion property and durability. These are described below as ‘beneficial fillers’. In the UK, guidance on the use and specification of asphalts is given in PD 6691:2010 (BSI, 2010b), and recommends that, for BS EN 13108-4 (BSI, 2006) hot rolled asphalt mixtures, any ‘added filler’ shall ‘consist of limestone, hydrated lime or cement’, and the UK’s Ministry of Defence DIO Specification 40 (Ministry of Defence, 2009b), Porous Friction Course for Airfields, requires that

between 1.5% and 2.0% by mass of the combined aggregate/filler aggregate ... shall be CL 90-S lime to BS EN 459-1 which shall be added to the mixture as part of the fraction passing the 0.063 mm sieve. If additional material of this grading is required, it shall be crushed limestone.

EN 459-1 (BSI, 2010c) is the European standard for building lime in which lime is defined as

calcium oxide and/or hydroxide, and calcium-magnesium oxide and/or hydroxide by the thermal decomposition (calcination) of naturally occurring calcium carbonate (for example limestone, chalk, shells) or naturally occurring calcium magnesium carbonate (for example dolomitic limestone, dolomite).
Included within the scope of EN 459-1 is the use of lime for civil engineering applications, including asphalts. This standard gives definitions for the different types of lime, and their classification, and chemical and physical properties.

Using EN 459-1, the CL 90-S lime specified in the DIO Specification 40 has the following derivation

- CL: a calcium lime, mainly either calcium oxide and/or calcium hydroxide that has no hydraulic or pozzolanic activity
- 90: a characteristic content of calcium oxide and magnesium oxide equal to or greater than 90%
- S: the product is hydrated lime (Ca(OH)_2) and not quicklime (CaO), which is designated with the letter Q instead of an S, and that it is in powder form rather than being either a putty (S PL), slurry or milk of lime (S ML).

Other chemical and physical properties are required by EN 459-1 appropriate to this classification.

Recognised beneficial fillers for use in asphalts include

- hydrated lime
- cement
- limestone (with high calcium carbonate content)
- mixed filler (with high calcium hydroxide content).

While limestone filler is the most commonly used because of its lower cost and wide availability, both hydrated lime and cement have greater beneficial effect.

When beneficial filler is used instead of, or as a supplement to, reclaimed (baghouse) filler, the binding properties of the mastic are improved, resulting in better adhesion between the aggregate and the mastic. In the UK, the durability of adhesion for base and binder course mixtures that have been designed according to the requirements of the Highways Agency’s Specification for Highway Works, Series 900, Clause 929 (Highways Agency et al., 2008), are required to be tested and shown to achieve a SATS durability index above 80%. Mixtures that include 2% of CL 90-S hydrated lime filler are deemed to satisfy the durability requirements of this specification without the need for SATS testing.

In general, aggregates with high carbonate content such as limestone are easier to coat with binder than aggregates with high silica content. This is due to the fact that siliceous aggregates contain high concentrations of hydroxyl groups with greater affinity for carboxylic acid and water. The carboxylic acid components that are present in the binder are adsorbed by the
surface of these aggregates, generating a binder–aggregate bond. However, these bonds with carboxylic acids are also prone to displacement in the presence of water (Petersen and Pancher, 1998). Such displacement is reduced when hydrated lime is used in the mixture and also, but to a lesser degree, when limestone filler is used.

### 10.5.1 Properties and testing

Within EN 13043 the requirements for testing filler properties is treated separately from other aggregates. The properties required through specification should be limited to those relating to the particular filler type, end use application or source of the coarse and fine aggregate.

#### 10.5.1.1 Geometrical requirements

**Grading**

Gradings are carried out in accordance with EN 933-10 (BSI, 2009f), which describes the reference method used for filler grading, for type test purposes and, in cases of dispute, both for natural or manufactured filler using air jet sieving.

At least 70% filler must pass the 0.063 mm sieve, and 100% must pass the 2 mm sieve size.

**Harmful fines**

It is usually considered that added fillers from beneficial sources are not likely to contain harmful fines, but when harmful fines in fillers are considered likely, their presence is assessed using the methylene blue test in accordance with EN 933-9, Annex A (BSI, 2009a) (and reported as MBFvalue: a description of this method is given in section 10.4.1.1 under the heading ‘Fines quality’ for fine and all-in aggregate).

#### 10.5.1.2 Physical requirements

**Water content**

The water content of added filler is determined in accordance with EN 1097-5 (BSI, 2008b), which limits it to a maximum of 1% by mass.

**Particle density**

Particle density is determined in accordance with EN 1097-7 (BSI, 2008c), and the results declared.

**Stiffening properties**

The addition of filler to form mastic with binder will stiffen an asphalt and affect a mixture’s workability and deformation resistance. The degree of stiffening is a function of many properties, with those relating to filler within
EN 13043 including the voids in dry compacted filler (Rigden) and the ‘delta ring and ball’.

**Voids of dry compacted filler (Rigden)**

The void content of dry compacted filler is known as the ‘fractional voids’, and is often referred to as the ‘Rigden voids’ after P. J. Rigden, who introduced the concept and a method for measuring them in 1947 (Rigden, 1947).

In EN 13043, the determination of the fractional voids is based on Rigden’s original method, and is specified in accordance with EN 1097-4 (BSI, 2008d). The result is reported within an EN 13043 class as $V_{xx/yy}$, where $xx$ and $yy$ define the overall range for individual results. The method involves the compaction of a dry filler test portion (10 g), using a standard compactive effort, into a small metal mould, to establish a reference bulk density. Using this bulk density and the particle density of the filler, the volume and percentage of fractional voids can be calculated.

After Rigden, it is considered that the bitumen required to fill the voids in the dry compacted sample is the ‘fixed’ bitumen (forming the mastic phase with the filler), and the bitumen in excess of this is the ‘free’ bitumen – the fraction that coats the coarse aggregate and lubricates the mixture. Higher Rigden void contents lead to a greater stiffening effect, and knowledge of the amount of voids can be important in both asphalt mixture design and quality control.

French guidance on the specification for Rigden void content is given in *The Use of Standards for Hot Mixtures* (Sétra, 2008), and for the characterisation of fillers used in asphalt this recommendation is $V_{28/45}$. When specified to EN 13043, all results are expected to be within this specification range, with a further requirement that at least 90% of the last 20 values be within 4 units of each other.

**‘Delta ring and ball’ of filler aggregate for asphalts**

The ‘delta ring and ball’ ($\Delta_{R&B}$) value determined in accordance with EN 13179-1 (BSI, 2013b) measures the increase in the softening point of a laboratory-prepared bitumen/filler mastic sample, respectively 62.5/37.5 by volume, relative to the softening point of the bitumen used in the test. The bitumen is 70/100 grade to EN 12591 (BSI, 2009g). The resulting increase in the softening point is reported as the $\Delta_{R&B}$ value (to the nearest 0.5°C), and results are declared within a relevant specified category, which is appropriate to the particular application.

In the UK, the stiffening property of added fillers used in EME2 base and binder course mixtures are recommended – according to PD 6691 (BSI,
2010b) – to be category $\Delta_{R&B} 8/16$, with the note that ‘Fillers with delta ring and ball values above 16 but not greater than 20 may be used where there is a history of satisfactory use in asphalt.’

**10.5.1.3 Chemical requirements**

When required, the following chemical requirements of filler in EN 13043 are tested.

- Water solubility, determined in accordance with EN 1744-1, with results declared as a percentage of solubility by mass, in the form $WS_{value}$ for categorisation purposes.
- Water susceptibility, determined in accordance with EN 1744-4 (BSI, 2005b).
- Calcium carbonate content of limestone filler aggregate, determined in accordance with EN 196-21 (replaced by EN 196-2:2013 (BSI, 2013c)), with results declared as the calcium carbonate content, percentage by mass, in the form $CC_{value}$ for categorisation purposes.
- Calcium hydroxide content of mixed filler, determined in accordance with EN 459-2 (BSI, 2010d), with results declared as the calcium hydroxide content, percentage by mass, in the form $K_{avalue}$ for categorisation purposes.

**10.5.2 Requirements for consistency of filler production**

EN 13043 requires at least one of the following properties to be measured as a means of establishing and measuring the consistency of filler production.

- The ‘bitumen number’ of the added filler. This is the amount of water, in ml, needed to be added to 100 g of filler to reach a reference consistency defined by a penetration value of 5–7 mm, in accordance with EN 13179-2 (BSI, 2000). The results are declared as $BN_{value}$, and for consistency purposes the values are required to be within the specified range, with the additional requirement that, on the basis of the last 20 values, 90% of the results are required to be within 6 units of each other.
- The loss on ignition of coal fly ash. This is determined in accordance with EN 1744-1, Clause 17, with the additional requirement that a producer’s declared range shall not be greater than 6% by mass. Also, when aggregates contain non-volatile oxidisable constituents, then the loss on ignition is corrected in accordance with EN 196-2.
- The particle density of the added filler. This is determined in accordance with EN 1097-7, with the units of Mg/m$^3$, and where this value is declared as a range, then this is not to exceed 0.2 Mg/m$^3$. 

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The loose bulk density in kerosene. An apparent density determined using a specified measuring flask with 10 g of filler in 25 ml of kerosene. The density is derived by measuring the height of the filler sediment in the kerosene after 6 h in accordance with EN 1097-3, Annex B [BSI, 1998c]. The producer’s range of values is required to be between 0.5 and 0.9 Mg/m³.

The Blaine test. This measures the specific surface of filler, in units of m²/kg, as specified in EN 196-6 [BSI, 2010e], with the requirement that a producer’s declared range shall be not greater than 140 m²/kg.

10.6 Assuring aggregate quality

The preceding sections describe the characterisation and categorisation of aggregate, and how it is measured and specified. However, this is of little value if the producer is unable to supply consistently to the declared grade.

Consistency is achieved through the application of a suitable and independently verifiable quality system that underpins the appropriate level of processing and inspection needed to control production. The establishment of aggregate declared properties is achieved by the application of a statistically based sampling and testing procedure, which is not always demanded by standards or specifications. This provides a more reliable measure of an aggregate’s true performance level than non-statistically based methods.

All aggregate used in the production of asphalt will have undergone some processing with subsequent assessment and testing, to deliver a final product that can be shown to meet a product specifier’s demands. It is the specifier’s responsibility to ensure that their specification can deliver a product ‘fit for its intended purpose’, and it is the aggregate producer’s responsibility to deliver a consistent supply of aggregate that meets their claimed performance levels.

With effect from 1 July 2013 in all European member states, the Regulation 2011 (EC, 2011) made it mandatory that all products are produced to harmonised European standards – this includes aggregates used in asphalts to EN 13043, which are required to be CE marked.

CE marking ensures that aggregates used within construction products not only satisfy the requirements of the relevant European standards but are also safe to use. The adherence to recognised systems of factory production control (FPC) gives the assurance that both of these requirements are being fulfilled.

Annex B (Normative) of the aggregate standard EN 13043:2002 deals with the FPC requirements, and specifies a system for aggregate production
that ensures they conform to the relevant requirements of the standard. The essential elements of this are

- organisation
  - responsibility and authority
  - management representative for factory production control
  - management review
- control procedures
  - document and data control
  - sub-contract services
  - knowledge of the raw material
- management of production
- inspection and test
  - general
  - equipment
  - frequency and location of inspection, sampling and tests
- records
- control of non-conforming products
- handling, storage and conditioning in production areas
- transport and packaging
  - transport
  - packaging
- training of personnel.

A new European standard is currently being developed that will specify requirements for both the initial type testing (ITT) and the FPC of aggregates. The FPC system will describe methods of controlling the sourcing and processing of aggregate, and will be combined with sampling and testing routines to provide ongoing assurances that aggregate products consistently conform to those properties that were determined in the ITT.

References


Aggregates in asphalts


Mukhopadhyay AK, Neekhra S and Zollinger DG (2007) Preliminary Characterization of Aggregate Coefficient of Thermal Expansion and Gradation for Paving Concrete. Texas Transportation Institute, Texas A&M University, College Station, TX, USA, Report FHWA/TX-05/0-1700-5.


Choosing asphalts for use in flexible pavement layers

There are three phases to the process of designing an asphalt pavement:

- choosing the particular asphalt to be used in each layer of the pavement
- design of the asphalt mixture
- design of the layers in a flexible pavement.

Design of the asphalt mixture and design of the pavement layer thicknesses are described in Chapters 12 and 13, respectively. This chapter discusses the asphalts that are used for particular roles in a flexible pavement in different regions of the world.

11.1 Asphalts used in flexible pavements in Europe

11.1.1 Surface courses

Different asphalts are available for the top layer of a road constructed with a flexible pavement. The top layer is normally called the surface course, and the asphalt chosen for that role depends on the requirements for the road surface.

The surface course constitutes the top layer of the pavement, and should be able to withstand high traffic and environment-induced stresses without exhibiting unacceptable levels of cracking and rutting. It should also have an even profile for the comfort of the user (in the UK, surface regularity) and possess a suitable texture to ensure adequate skid resistance. Depending on the applicable specification in any particular region, functional characteristics such as skid resistance, noise reduction and durability are often required for surface courses. In some cases, rapid drainage of surface water is achieved by having a porous structure, while in other cases the surface course should be impermeable in order to keep water out of the pavement structure. As indicated, the surface layer is important for the pavement performance but no single material can provide all the desired characteristics.
(e.g. being porous and impermeable at the same time). A wide range of surface layer products can, therefore, be considered appropriate, depending on specific requirements

- asphalt concrete (AC)
- asphalt concrete for very thin layers (AC-TL)
- asphalt for ultra-thin layers (AUTL)
- stone mastic asphalt (SMA)
- hot rolled asphalt (HRA)
- porous asphalt (PA)
- double layered porous asphalt (2L-PA)
- mastic asphalt (MA)
- soft asphalt (SA).

11.1.1.1 Asphalt concrete
AC is asphalt in which the aggregate particles are continuously graded or gap graded to form an interlocking structure. Dense asphalt concretes are often used in the base layers in flexible pavements.

11.1.1.2 Asphalt concrete for very thin layers
AC-TL is asphalt for surface courses with a thickness of 20–30 mm in which the aggregate particles are generally gap graded to form stone to stone contact and to provide an open surface texture. This mixture is often used in France where it is called béton bitumineux très mince (BBTM).

11.1.1.3 Asphalt for ultra-thin layers
AUTL is a surface course that is laid on a bond coat (a proprietary tack coat made with polymer modified bitumen) at a nominal thickness of 10–20 mm with properties suitable for the intended use. The method of bonding is an essential part of the process, and the final product is a combination of the bonding system and the asphalt. AUTL is an asphalt in which the aggregate particles are generally gap graded to form stone to stone contact with an open surface texture. Several varieties of this layer are often used to provide a good, new, low noise running surface.

11.1.1.4 Stone mastic asphalt
SMA is a gap graded asphalt with bitumen as a binder, composed of a coarse crushed aggregate skeleton bound with a mastic mortar. This mixture is often used as a surface layer in cases where high stability is needed. The surface structure generates low noise levels.

11.1.1.5 Hot rolled asphalt (HRA)
HRA is a dense, gap graded asphalt in which the mortar of fine aggregate, filler and high viscosity binder are major contributors to the performance
of the laid material. Coated chippings (nominally single size aggregate particles with a high resistance to polishing, which are lightly coated with high viscosity binder) are always rolled into and form part of an HRA surface course. This durable surface layer is often used as a surface layer in the UK.

11.1.1.6 Porous asphalt
PA is an asphalt with a very high level of interconnected voids (18–25%), which allow the passage of water. It generates low noise levels and very little spray.

11.1.1.7 Double layered porous asphalt
The top layer of porous asphalt typically uses an aggregate with nominal size of 4–8 mm and is laid about 25 mm thick. The second/bottom layer is a thicker layer of porous asphalt with a larger aggregate nominal size (11–16 mm). As a result of the finer texture at the top (resulting in fewer tyre vibrations), this material is designed to produce lower noise levels than a porous asphalt laid in a single layer.

11.1.1.8 Mastic asphalt
MA is designed to be of low void content. The binder content is adjusted so that the voids are completely filled. MA is pourable and can be spread in its working temperature condition. It requires no compaction on site. This mixture is very durable and has often been used as a surface layer in certain countries.

11.1.1.9 Soft asphalt
SA is a flexible mixture of aggregate and soft bitumen grades and is used in the Nordic countries for secondary roads.

The selection of the surface course is a matter of identifying the most appropriate material during the design process. The functional requirements can conflict. For example, low noise generation may require the use of a double layered porous asphalt but that would conflict with the requirement for the surface to be very durable. The durability of surface layers can be improved by using higher quality materials. The higher costs of these components are recouped, at least in part, by the lower costs of traffic measures and user costs.

11.1.2 Bases and binder courses
Throughout Europe, in fully flexible pavements, the most dominant type of asphalt used in these layers is asphalt concrete (AC). Different variations of AC are used and further details are given in this chapter.
11.1.3 European asphalt standards

On 1 January 2008, the European Committee for Standardisation (CEN) adopted a unified approach to mixture design, manufacture and CE marking of asphalts.

The ultimate aim is to specify asphalts in terms of fundamental properties based on performance. However, given the differences in knowledge and experience with the basic specifications for this type of mixture in Europe, it is not currently possible to adopt a fundamental approach.

Eight different asphalts are covered by the various parts of BS EN 13108 (BSI, 2005, 2006a–g) and are shown along with the type testing and factory production control standards (BSI, 2006h–i) in Table 11.1.

11.2 UK pavements

Layers in UK flexible pavements are shown in Figure 11.1.

Pressures are applied at the surface of a road by the tyres of vehicles running over the pavement. The structural wear caused by lighter traffic (i.e. cars and lighter goods vehicles) is considered to be negligible according to paragraph 2.6 of the UK’s traffic assessment standard (Highways Agency et al., 2006a). It is only heavier commercial vehicles (i.e. those having a gross vehicle weight in excess of 3.5 tonnes) [also defined in paragraph 2.6 of the UK’s traffic assessment standard that are considered to cause structural damage to a pavement]. The highest pressure will be exerted by heavy goods vehicles having six or more axles, and the pressure will typically be of the order of 8.5 bar, and may well be higher for particular loads. The function of a road pavement is to dissipate such pressures down through the pavement, the foundation and the subgrade such that none of the materials in each of these layers deteriorates as a result of the effects of traffic.

The role played by each layer in a road with a flexible pavement, together with the effects that the location has on the choice of asphalt, is discussed below.
11.2.1 Subgrade and foundation

The subgrade will play a significant role in determining the service life of a pavement (see Table 13.2). In addition, the foundation also plays a key role in the performance of the road. However, foundations are not discussed in this book in any detail. Advice on foundations can be found in Interim Advice Note 73/06 (Highways Agency et al., 2009) and Chapter 32 of the ICE Manual of Highway Design and Management (Payne and Walsh, 2011).

11.2.2 Base

The primary function of the base is to act as the main structural layer in the pavement. Although UK pavement specifications allow a wide range of materials to be chosen for use as a base, including many cementitious mixtures, the material found in the vast majority of bases are asphalts. Properly designed asphalts provide the stiffness necessary to dissipate the pressures imposed by traffic without causing distress in the pavement.

Figure 11.1 UK road constructed with a fully flexible pavement: PA, porous asphalt; TSCS, thin surface course system
layers, the foundation or subgrade. Mixtures formulated for bases need to have an adequate stiffness modulus and the required degree of resistance to fatigue. The formulation of these mixtures has evolved over the years, through a reduction in the size of the aggregates, and the use of harder bitumens.

The thickness of the base in a heavily trafficked pavement will typically be of the order of 200 mm or more. Although UK specifications no longer place a limit on the maximum thickness of asphalt layer that can be placed in a single lift, it would be wise to lay in two layers if the thickness of the base exceeds 120 mm. The reason for laying a thickness no greater than this is the difficulty of achieving the required void contents (the usual UK specification has an upper limit of void content) and the surface regularity specification, with the latter being the crucial parameter. The thicker a layer is, the more difficult it is to achieve the surface regularity tolerance. Thus, bases on pavements carrying heavy traffic will usually be constructed in two layers, with the first being described as the lower base and the second being described as the upper base, as depicted in Figure 11.1.

11.2.3 Binder course

The surface regularity of a finished road is a key determinant of its quality. It is a measure of the number of undulations in the surface. The fewer the undulations or irregularities, as the normal UK specification describes them in Clause 702 of the Specification for Highway Works (Highways Agency et al., 2008), the more comfortable is the ride for those travelling on the road. UK surface regularity specifications require that the surface course is within a specified tolerance of the specified finished level. The most common tolerance is $\pm 6$ mm of the specified finished level. However, a tolerance of +6 to 0 mm is not unusual (on racing circuits the tolerance may be even smaller). Asphalts are placed in layers using specialist items of equipment called ‘pavers’. These are discussed in detail in Chapter 15. Every time a layer of asphalt is laid by a paver, the action of the paver reduces any irregularities in the laid mat. Thus, the irregularities at the top of the upper base are less than those in the lower base. Similarly, the irregularities on top of the binder course are less than those in the upper base. Finally, the irregularities in the surface course are less than those in the binder course. Tolerances apply at each layer, with the tolerance becoming tighter closer to the surface. Compliance after each layer has been placed is a specification requirement but achieving regularity limitations is most important on top of the surface course. It is for this reason that a binder course is necessary. Without a fourth layer, asphalt contractors may find it difficult to achieve compliance with specified regularity requirements.

Secondary, albeit important, benefits of having a binder course are
the contribution made by the binder course in dissipating pressure from traffic (this property is described as the stiffness of the asphalt, i.e. a measure of its load spreading ability; this is discussed further in Chapter 13)

- preventing water reaching the base and foundation layers and the subgrade.

### 11.2.4 Surface course

The surface course is the visible portion of the road structure. It is this layer that suffers most from the effects of climate aggression and vehicle loads.

The formulation of the surface course asphalt will be the factor that is key in determining the characteristics of the surface layer of the pavement. These attributes include

- durability
- surface regularity
- skid resistance
- noise reduction
- resistance to rutting
- resistance to cracking
- water drainage
- appearance.

The appearance of the finished surface is rarely, if ever, given any thought. However, one of the many benefits of a thin surface course system that has been properly designed and skilfully laid is that it looks so much more attractive than does a chipped hot rolled asphalt surface course.

Investment in research and development of surface course asphalt has brought major innovations over the last 25 years. Examples of these initiatives are improved methods of choosing aggregates and the use of special bitumens, including, most importantly, polymer modified bitumens, which have expanded the range of types of surface course. Transport Scotland’s work on surface courses is worthy of particular note (Transport Scotland, 2010).

### 11.3 Asphalts used in flexible pavements in the UK

Referring to Figure 11.1, each of the layers in the pavement has to perform specific functions as discussed below.

#### 11.3.1 Base

The base is the main structural layer in the pavement, although the other layers will also contribute to the structural strength of the finished pavement. Many materials are available to act as a base in a modern pavement.

It is permissible to construct bases using a group of materials described as ‘hydraulically bound mixtures’ (HBMs). Many of these mixtures are cementitious.
However, the bulk of bases, as specified and as constructed, are asphaltic. The asphalts that are permitted for use as bases on UK trunk roads (the major arterial routes in the UK) are:

- DBM125
- DBM50/HDM50
- EME2
- HRA50.

The names used for particular asphalts in the UK have arisen over many years. Current descriptions continue to be used, with dense bitumen macadam (DBM) and heavy duty macadam (HDM) (discussed below) being examples. DBM and HDM will become ‘dense bituminous material’ and ‘heavy duty material’ in 2015. The family of standards governing asphalts is the BS EN 13108 series (see Table 11.1).

Generically, DBM is an asphalt concrete, so production of this material is governed by BS EN 13108-1 [BSI, 2006a]. The word ‘macadam’ comes from John Loudon McAdam (1756–1836) – the spelling of the surname is correct. McAdam is credited with inventing the system of placing increasingly smaller aggregate sizes and finally blinding the surface with dust. (In fact, the Romans used a similar system 2000 years ago (Mitchell, 2011)). The provisions of BS EN 13108-1 (like all the other parts of BS EN 13108) apply throughout Europe. However, the UK has a document that does not have the status of a European standard but is the core of any asphalt production specification in the UK. It is designated PD 6691 [BSI, 2010b]. PD stands for ‘published document’ and the role of this document is explained in the second paragraph of its Introduction as: ‘This Published Document gives the recommended choices for the mixtures most commonly used in the UK’.

HDM is also an asphalt concrete, so its production is governed by BS EN 13108-1 [BSI, 2006a].

Two options in bases are DBM50 and HDM50. The ‘50’ in the description indicates that these asphalts are manufactured with 40/60 pen bitumen as per BS EN 12591 [BSI, 2009]. DBM50 and HDM50 can be treated as identical materials. These materials account for the majority of bases in heavily trafficked UK roads. Table 11.2 gives typical declared gradings for a DBM50 and an HDM50. (The standards that apply in European member states set limits that apply to particular mixtures but it is for individual producers to ‘declare’ gradings within those limits, hence the caption for Table 11.2.)

As can be seen from Table 11.2, the mixture details for DBM50 and HDM50 are identical, and this will be the case with the DBM50 and HDM50 asphalts offered by many asphalt manufacturers. So why bother
having two different asphalts available when they are in fact identical? The reason is that these mixtures have been available historically and customers expect them to continue to be available. Over time, one or both designations may disappear following a rationalisation exercise within PD 6691 (BSI, 2010b).

DBM125 is a dense bitumen macadam, like DBM50, but is made with a 100/150 pen bitumen.

EME2 stands for enrobé à module éleveé class 2. EME2 is treated as an asphalt concrete, and thus its production falls under BS EN 13108-1 (BSI, 2006a). This asphalt contains very hard bitumen (15–20 pen is the target, which is not a range supported in the hard grade bitumen standard in BS EN 13924 (BSI, 2006j), in which the relevant classes are 10/20 and 15/25 pen). EME2 was developed following studies by the Highways Agency in France. French highway engineers have been using mixtures made with very hard bitumens since 1985 (Williams, 2007), and so have amassed a great deal of valuable experience in the behaviour of asphalts made with very hard bitumens. More information on EME2 can be found in Sanders and Nunn (2005) and Williams (2007).

HRA stands for hot rolled asphalt. Generically, HRA is a ‘hot rolled asphalt’ so its production is governed by BS EN 13108-4 (BSI, 2006d). The ‘50’ in the description HRA50 indicates that these asphalts are manufactured using 40/60 pen bitumen as per BS EN 12591 (BSI, 2009). HRA has been in use in the UK for many years, the first standard for the material being published in January 1935 (BSI, 1935). Traditionally, in the UK there were two categories of asphalt, described as gap graded and continuously graded. Gradings, of course, describe the proportion of aggregate passing a particular sieve size; an example of this is shown in Table 11.3. Where a mixture included a proportion of all or most of the sieve sizes, the material was

### Table 11.2  Examples of declared mixture details for DBM50 and HDM50

<table>
<thead>
<tr>
<th>Grading (passing):</th>
<th>DBM50</th>
<th>HDM50</th>
</tr>
</thead>
<tbody>
<tr>
<td>40 mm sieve</td>
<td>100%</td>
<td>100%</td>
</tr>
<tr>
<td>31.5 mm sieve</td>
<td>99%</td>
<td>99%</td>
</tr>
<tr>
<td>20 mm sieve</td>
<td>86%</td>
<td>86%</td>
</tr>
<tr>
<td>6.3 mm sieve</td>
<td>52%</td>
<td>52%</td>
</tr>
<tr>
<td>2 mm sieve</td>
<td>30%</td>
<td>30%</td>
</tr>
<tr>
<td>0.25 mm sieve</td>
<td>13%</td>
<td>13%</td>
</tr>
<tr>
<td>0.063 mm sieve</td>
<td>6.0%</td>
<td>6.0%</td>
</tr>
<tr>
<td>Binder content</td>
<td>$B_{\min}4.0$ (4.0%)</td>
<td>$B_{\min}4.0$ (4.0%)</td>
</tr>
<tr>
<td>Temperature of the mixture</td>
<td>150–190°C</td>
<td>150–190°C</td>
</tr>
</tbody>
</table>
categorised as ‘continuously graded’ (i.e. asphalt concretes (previously called ‘bitumen macadams’)). Where the grading was discontinuous, the material was categorised as ‘gap graded’ (i.e. hot rolled asphalts). An example of both categories is shown in Figure 11.2, which is drawn from the data presented in Table 11.3. However, the distinction based on the gaps or otherwise in gradings is not easy to perceive and this method of classification, while of interest historically, is of little practical assistance in categorising asphalts.

The designer, therefore, has, in effect, a choice of four materials to use in the

**Figure 11.2** Grading envelopes for a continuously graded asphalt (AC32 DBM50 base) and a gap graded asphalt (HRA 30/14 surf)

**Table 11.3** Gradings for a continuously graded asphalt (AC32 DBM50 base) and a gap graded asphalt (HRA 30/14 surf)

<table>
<thead>
<tr>
<th></th>
<th>AC32 DBM50 base</th>
<th>HRA 30/14 surf</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Target</td>
<td>Min.</td>
</tr>
<tr>
<td>40</td>
<td>100</td>
<td>98</td>
</tr>
<tr>
<td>31.5</td>
<td>99</td>
<td>90</td>
</tr>
<tr>
<td>20</td>
<td>86</td>
<td>77</td>
</tr>
<tr>
<td>14</td>
<td>66</td>
<td>61</td>
</tr>
<tr>
<td>10</td>
<td>61</td>
<td>57</td>
</tr>
<tr>
<td>6.3</td>
<td>52</td>
<td>45</td>
</tr>
<tr>
<td>30</td>
<td>8</td>
<td>18</td>
</tr>
<tr>
<td>0.5</td>
<td>8</td>
<td>11</td>
</tr>
<tr>
<td>0.25</td>
<td>3</td>
<td>9</td>
</tr>
<tr>
<td>0.063</td>
<td>6</td>
<td>3</td>
</tr>
</tbody>
</table>
Choosing asphalts for use in flexible pavement layers

### Table 11.4 Advantages and disadvantages of available base materials

<table>
<thead>
<tr>
<th>Asphalt</th>
<th>Advantages</th>
<th>Disadvantages</th>
</tr>
</thead>
<tbody>
<tr>
<td>DBM125</td>
<td>Good mechanical interlock, low bitumen content, reasonable crack and rut resistance</td>
<td>Soft bitumen, low stiffness, relatively high cost</td>
</tr>
<tr>
<td>DBM50/HDM50</td>
<td>Good mechanical interlock, relatively low cost, medium stiffness, high deformation resistance</td>
<td></td>
</tr>
<tr>
<td>EME2</td>
<td>High stiffness, impermeability</td>
<td>Relatively high cost</td>
</tr>
<tr>
<td>HRA50</td>
<td>Impermeability</td>
<td>Relatively high cost, low deformation resistance</td>
</tr>
</tbody>
</table>

Base. Some designers offer all options to contractors pricing tenders and accept whichever option the winning bidder offers, which is the approach required by HD 26/06 (Highways Agency et al., 2006b). Other designers may decide to be more selective by taking account of the advantages and disadvantages of each option. Table 11.4 summarises the advantages and disadvantages of each available option.

Note that the low deformation resistance of HRA50 is not relevant in a base, as deformation in thicker pavements is restricted to the surface course and binder course (Nunn, et al., 1997). However, the binder course is usually made of the same generic material as the base, and an HRA50 as a binder course would carry with it significant concerns about deformation reaching a critical level (i.e. where replacement is warranted) much earlier than would be the case with the other options.

#### 11.3.2 Binder course

As explained above, the binder course is present primarily to allow the asphalt contractor to achieve the very tight tolerances applied to the finished levels of the surface course. In addition, however, binder courses usually also have a higher quantity of bitumen in the asphalts used for their construction. This increases the durability of the layer. It also improves the waterproofing characteristic of the layer. section 4.2.1 of RN42 (Nicholls et al., 2008) states that

There is a need for the properties required for the binder course, particularly impermeability to limit any flow of water from the surface course downwards and/or from the base upwards. With surfacing materials being made thinner and more open and with base layers often being designed with relatively low binder contents, the need for an impermeable binder layer becomes vital.

The types of asphalt used for binder courses are much the same as those used for bases, with one interesting addition

- DBM125
- DBM50/HDM50
EME2
HRA50
SMA.

It is important to note that DBM125 is available as a base or a binder course. The difference between these two mixtures is that the binder course variant has approximately 0.5% more bitumen than the base variant. The same is true of DBM50 and HDM50. In contrast, for any particular source, both EME2 and HRA50 mixtures used in base and binder courses are the same (i.e. they will have the same proportion of bitumen in the asphalt).

SMA stands for stone mastic asphalt, an asphalt that was developed in Germany in the 1960s. At that time in Germany, studded tyres were widely used in winter, and the Germans had a choice of two running surfaces – asphalt concrete and gussasphalt. As has been explained, asphalt concrete is continuously graded, and has a relatively low binder content. It was found to wear rapidly under studded tyres, failing by way of a mechanism called ravelling, which means the asphalt is abraded away. Attempts to counter this tendency by increasing the binder content simply resulted in the material failing in deformation in the wheel tracks. The performance of gussasphalt, however, was quite different. Being gap graded, it has a relatively high binder content but, because of this, it was expensive to produce and lay.

Thus, in the late 1960s, split mastic asphalt appeared, split being the German for ‘stone’. Stone mastic asphalt was a proprietary gap graded material, with a high stone content and a high binder content. The mixture is durable and, being negatively textured, generates lower levels of noise and spray. A positively textured surface is one where some constituents of the asphalt project upwards from the surface. A negatively textured surface is one that has voids in the surface. Negatively textured surface courses generate significantly lower noise levels, and result in lower noise levels than is the case with positively textured surface courses. An example of a positively textured surface course would be a chipped hot rolled asphalt surface course. An example of a negatively textured surface course would be an asphalt concrete close graded surface course. Studded tyres were banned in Germany in 1975 but the benefits of SMAs were, by that time, apparent. SMA did not find its way into the German national specification until 1984. The use of SMA spread to other countries in Europe throughout the 1980s. Sweden, France, Belgium, Denmark and other countries all used variations of this material. In the early 1990s, formulations based on SMA gradings appeared in the UK, and some are in use today as thin surface course systems (TSCS) complying with Clause 942 of the standard UK highway specification (Highways Agency et al., 2008).

SMA also exists as binder courses and regulating courses in the UK. The very
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useful advantage of SMA binder courses is that they can be laid as thin as 30 mm. In the design of pavements, the total asphalt thickness is derived using procedures that are described in HD 26/06 (Highways Agency et al., 2006b). Sometimes it is very helpful in the process to be able to specify a thin binder course. Most asphalt binder courses have to be at least 50 mm thick, as per Table 1A of BS 5949-87:2010 (BSI, 2010a). Table 1A applies to asphalt concrete binder courses; hot rolled asphalt binder courses can be as thin as 45 mm but are not recommended because of their propensity to rut relatively easily.

Generally, the binder course is made of the same generic material as the base, albeit that the binder course material may have more bitumen than that of the base. So, for example, the base would be HDM50 and the binder course would be HDM50, or the base would be EME2 and the binder course would be EME2.

11.3.3 Surface course
On heavily trafficked UK roads, two types of surface course predominate

- chipped HRA surface course
- thin surface course systems.

Chipped HRA was the only surface course on major roads for many years, until the introduction of thin surface course systems in the 1990s.

11.3.3.1 Chipped hot rolled asphalt surface course
Chipped HRA consists of a layer of HRA into which precoated chippings are rolled. The most common type of HRA surface course consists of approximately

- 30% of 14 mm single-sized aggregate
- 55% sand
- 9% very fine aggregate (called filler)
- 6.5% bitumen binder.

After the HRA surface course has been placed and before it has been rolled, precoated single-sized chippings (usually 20 mm) are spread onto the surface. The chippings are then pushed into the surface as part of the compaction process. The chippings are coated with bitumen of the same type as was used in the manufacture of the asphalt itself. After being coated, the chippings are allowed to cool, and are then transported to the site to be stored until they are added to the surface course at a specified rate of spread. The chippings are necessary to give the surface adequate skid resistance, thus allowing vehicles to brake in the shortest possible length. Good frictional characteristics are essential, and will reduce injuries and fatalities. It is most important for those involved in highways to be familiar with the factors that affect skid resistance.
HD 28/04 (Highways Agency et al., 2004) is entitled Skid Resistance, and provides essential information on this very important topic.

A chipped HRA surface course using the above formulation is laid such that, when the chippings have been added and the material has been rolled (more meaningfully, compacted), it is 40 mm thick, in accordance with Table 1B of BS 594987:2010 (BSI, 2010a).

11.3.3.2 Thin surface course systems
Thin surface course systems are proprietary materials that can be laid as the running surface of a carriageway. They can usually be laid at thicknesses from 40 mm down to a few millimetres.

This class of materials first appeared in late 1992, when two UK companies imported French technology (the products were called Safepave, which was produced by Associated Asphalt, and UL-M (ultra-mince being French for ‘very thin’), which was introduced by Alfred McAlpine). Other companies followed, with Tarmac, for example, undertaking trials in 1992 and 1993 of its Masterpave. A major difficulty in the usage of these asphalts on trunk roads was that the national specification did not recognise the possibility that such materials may appear.

The emergence of the proprietary Safepave and UL-M mixtures and others, the effects of harmonisation activities in relation to standards, combined with stories of this wonderful stone mastic asphalt material from Germany stimulated significant interest in a new breed of what were generically referred to in the industry as ‘stone mastic asphalts’.

Indeed, the level of interest generated was such that the Road Engineering and Environmental Division of the Highways Agency commissioned a demonstration trial at the TRL. The resultant report (Project Report 65, commonly referred to as PR 65), was published in 1994 (Nunn, 1994). This report was extremely successful in that it served to highlight the availability of alternative mixtures to the industry.

However, the difficulty remained that the national specification did not recognise these mixtures, and thus their use was limited to non-trunk road locations. As a result of the pressure for usage from producers, external customers and Highways Agency and TRL staff who had realised the benefits of these materials, the Highways Agency set about considering how best to allow the use of such materials on trunk roads. The solution was to put in place a system that allowed the use of proprietary mixtures, which had been rigorously tested in every single respect relevant to the life and performance of surface courses. This started with close control over the ingredients through to the establishment of controls over the way the material is laid. This system is known as HAPAS (Highway Authority
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Table 11.5 Permitted pavement surface course materials on UK trunk roads

<table>
<thead>
<tr>
<th>Country</th>
<th>Default surface course</th>
</tr>
</thead>
<tbody>
<tr>
<td>England</td>
<td>Thin surface course system</td>
</tr>
<tr>
<td>Northern Ireland</td>
<td>Thin surface course system Chipped hot rolled asphalt Dense macadam</td>
</tr>
<tr>
<td>Scotland</td>
<td>Chipped hot rolled asphalt</td>
</tr>
<tr>
<td>Wales</td>
<td>Thin surface course system</td>
</tr>
</tbody>
</table>

Notes
1. In Northern Ireland, asphalt concrete surface courses can only be used on roads that are not high speed.
2. In Scotland, surface dressing may be permissible on roads that are not high speed.
3. All countries may permit other treatments subject to a departure from standard.

Product Approval Scheme) or the BBA HAPAS (British Board of Agrément), the UK body that provides reassurance to manufacturers, users, specifiers, insurers and regulators of construction products and systems. More information on HAPAS can be found in section 23.3.3.

The Specification for Highway Works now accepts the use of thin surface course systems, and these materials are addressed in Clause 942 (Highways Agency et al., 2008). It should be noted that this clause requires the contractor to guarantee the integrity of the surface course and the workmanship for a period of 5 years from the date of opening to traffic, but it is very important to be aware that this applies to the trunk road network only. The 5 year guarantee includes defects such as fretting, ravelling, stripping and loss of chippings.

Table 11.5 summarises the choice of surface course on trunks roads in the different countries of the UK as per Tables 2.2E, 2.2W, 2.2S and 2.2NI in HD 36/06 (Highways Agency et al., 2006c).

So, on trunk roads, the default surface course is, in most cases, a thin surface course system. However, some authorities continue to specify chipped HRA. Primarily, this is because of a belief that the lifespan of HRA surface courses is significantly better than that of TSCS. A detailed study undertaken by TRL (Nicholls et al., 2010) states in section 2.5.1.1 of the report that a TSCS will last for, on average, 12–13 years (for thin asphalt concretes with 0/14 and 0/10 mm gradings, generally with polymer modified bitumen). A chipped HRA surface course is expected to have a lifespan of around 18 years according to the ICE Manual of Highway Design and Management, which states a range of 14–24 years (Payne and Walsh, 2011: Table 2 of Chapter 50). The surface regularity on TSCSs is superior to that of chipped HRA. TSCSs are also superior in relation to traffic noise, which may well be approximately two-thirds of that on chipped HRA. TSCSs also exhibit reduced spray as compared with chipped HRA. TSCSs are
superior in aesthetic terms to a chipped HRA surface course. However, they do use a higher proportion of high polished stone value (PSV) aggregate compared with chipped HRA surface courses. Thus, the only disadvantages of a TSCS compared with a chipped HRA surface course are the lower lifespan and the greater usage of premium aggregate.

11.3.3.3 Other asphalt surface courses
A number of other asphalts are available for use as surface courses

- asphalt concrete
- porous asphalt
- generic stone mastic asphalt.

Asphalt concrete surface courses
Asphalt concrete surface courses can be used on roads having low traffic levels. There are five types of asphalt concrete surface course available

- dense
- close graded
- medium graded
- fine graded
- open graded.

Generally speaking, asphalt concrete surface courses are used on roads that carry light traffic. Open graded mixtures were widely used in the UK, particularly in England, prior to the emergence of heavier commercial vehicles. Having an open texture means that they are accessible to water and are likely to degrade rapidly where there are frequent freeze–thaw weather cycles. Their durability is a function of the nature of the pavement itself and the type of traffic to which the road is subject. The use of a surface dressing may be necessary as a sealant. Close graded, dense and medium graded surface courses are less susceptible to intrusion and damage by surface water. However, they are only suitable for light traffic or applications such as footways, car parks, playgrounds etc. Fine graded surface course was previously described as fine cold asphalt, and it is only suitable for footways etc.

Porous asphalt
Porous asphalt is a high void asphalt concrete surface course. The void content can be between 18 and 25% (most asphalts have void contents between 4% and 8%). The high voidage in porous asphalt results in the material being pervious, so rainwater passes right through the layer. This is illustrated in Figure 11.3.

The rainwater passing through the porous asphalt surface course runs along the top of the impervious asphalt concrete binder course and into a drain.
placed below the channel to catch this water. Another feature of the material is that it has exceptionally low traffic noise characteristics, generating around half the noise of a chipped hot rolled asphalt surface course.

The disadvantage of porous asphalt is its lifespan, which is approximately 8 years. A number of trials have been carried out in the UK but the material has performed disappointingly and, as a consequence, it appears unlikely that it will enjoy widespread usage in the UK in the foreseeable future.

**High stone content hot rolled asphalt**
A number of specifiers favour this material, which is also known as medium temperature asphalt (MTA). The asphalt in a chipped HRA surface course often has 30% of 14 mm aggregate (sometimes represented as 30/14). High stone content asphalts are either 55% of 10 mm or 14 mm aggregate (i.e. 55/10 or 55/14). It is generally easier to lay than a traditional chipped HRA surface course. It has no precoated chippings as it is textured when laid properly.

**Generic stone mastic asphalt**
This material would not have the frictional characteristics favoured in the UK. Accordingly, it is not used in the UK.

### 11.4 Asphalts used in flexible pavements in France
#### 11.4.1 The evolution of asphalts in France
Most roads in France are flexible pavements and are constructed using asphalt in the base, binder course and surface courses. France is considered to have a very high quality road network as a result of knowledge and experience of asphalt and pavement performance acquired over many years by both highway administrators and road contractors. Several of the paving materials developed in France, such as high modulus asphalt and thin
surface course layers, have been adopted in other countries. The most frequently used paving grade bitumens are 35/50 and 50/70 pen bitumens. In addition, since the 1980s, special harder grades of bitumen (e.g. 10/20 or 20/30 pen) have been used in specific circumstances to meet the demands of heavy traffic loading or climatic conditions. Polymer modified bitumens are used extensively in France.

In common with other European countries, during the 1950s most asphalts used in France were generic ‘recipe’ type mixtures, and the limitations of such an approach for coping with increasing traffic levels was beginning to manifest itself in the form of pavement damage through rutting, surface cracking and premature ageing of bitumen. This was exacerbated because many of the recipe mixtures at that time were based on relatively small nominal size aggregates (16 mm maximum size) and were manufactured using natural sand and crushed gravel. In addition, the bitumens commonly used were relatively soft, with 80/100 and 180/220 pen grades being used to construct pavements at that time.

By the mid-1960s, French roads were exhibiting unacceptable levels of rutting due to increased levels of loading from commercial vehicles. The monitoring of badly rutted roads and subsequent laboratory studies resulted in a different approach to the manufacture of asphalts, which dictated the direction taken by asphalt technology in France. Instead of using natural sands and gravel, asphalts were to be manufactured with crushed rock. In addition, harder penetration bitumen grades of 40/50 and 60/70 pen were employed. These designs formed the basis for the semi-coarse asphalt concretes (bétons bitumineux semi-grenus (BBSG)) used today.

For surface courses, the invention of polymer modified bitumen in the 1970s led to the development of mixture designs suitable for use in very thin layers designed to ensure the highest levels of grip and durability. This began with 40 mm nominal thickness surface course and, in the 1980s, to the development of very thin layer asphalt layers known as béton bitumineux très mince (BBTM). These asphalts are now widely used on many parts of the road network, in layer thicknesses of 15–20 mm. In the early 1980s, béton bitumineux drainant (porous asphalt) was introduced to improve driving visibility and skid resistance by reducing surface water. A further adaptation of thin asphalt and porous asphalt has led to the development of small nominal size aggregate asphalts (0/6 mm), known as béton bitumineux mince, for the reduction of traffic noise.

For base and binder course materials, in the 1980s, high modulus asphalts, enrobé à module élevé (EME), were first introduced with the aim of reducing thicknesses and optimising aggregate resources. These mixtures have a nominal aggregate size of 10–20 mm, the most commonly used size being 14 mm. A
Choosing asphalts for use in flexible pavement layers

A hard bitumen grade of 10/20 or 20/30 pen is used with relatively high binder contents of 5.5–6.0% by mass of aggregate in the mixture, to ensure good durability and protect against fatigue damage and water sensitivity, while producing high stiffness modulus mixtures capable of withstanding high traffic loading.

BBME (bétons bitumineux à module élevé – high modulus asphalt) first appeared in the mid-1990s, using gradings with nominal sizes of 0/10 or 0/14 mm. It is designed using 20/30 pen bitumen. This asphalt is another mixture type used in binder course or surface course to overcome the phenomenon of rutting.

### 11.4.2 Current practice for asphalt pavements in France

All asphalts in France are required to meet European (CEN) specifications. Table 11.6 summarises the majority of asphalts used today in France.

The CEN standards allow two different approaches to asphalt mixture design. The first is an empirical approach, in which the final composition is based on the characteristics of the mixture (e.g. the nominal size of aggregate, the air void content, the particle size distribution and the target binder content). Alternatively, the fundamental approach measures a number of material properties of the mixture (e.g. complex stiffness modulus, fatigue and resistance to rutting), and includes requirements for water sensitivity of the mixture and the percentage of air voids within the mixture. The latter is the approach that is most often used in France, and a mixture design procedure has been developed over many years to assess the properties of asphalt mixtures. In this approach, the target binder content and the grading envelope are not predetermined but are designed to achieve the performance criteria. Furthermore, the measured mechanical performance of the mixture is given a performance class, ranging from 1 to 4, with 4 being the highest levels of loading and/or most severe climatic conditions. The pavement design, and hence selection of the type and thickness of binder course and surface course, will reflect the performance class and thickness of base layer proposed.

Although EME and BBME are in widespread use in France, in particular on the major road networks, the most common material used for the asphalt base layers is grave bitume. This asphalt has a grading of nominal size 0/14 or 0/20 mm, and typically contains 35/50 pen bitumen with a binder content between 3% and 5%. Variation of the bitumen content and the grading curve is permitted to achieve varying defined performance classes (from 2 to 4).

Béton bitumineux semi-grenus (BBSG) (semi-coarse asphalt concrete) remains the most widely used asphalt in France. These mixtures use a continuous grading curve with a nominal size of 0/10 or 0/14 mm, reconstituted from at least three granular fractions combined to meet the specification. The bitumen content of paving grade (typically 35/50 or 50/70 pen) is 5–5.8% by mass of
## Table 11.6 Asphalts in France

<table>
<thead>
<tr>
<th>Norms EN</th>
<th>Nom</th>
<th>Grading Class</th>
<th>Thickness</th>
<th>Disc.</th>
<th>% content ppc</th>
<th>PCG test</th>
<th>Duriez</th>
<th>Rutting test</th>
<th>% Modulus (15°C, 10 Hz)</th>
<th>Defor rage 3 m</th>
<th>Test in situ</th>
</tr>
</thead>
<tbody>
<tr>
<td>0/10</td>
<td>BBSG EB</td>
<td>1</td>
<td>4</td>
<td>5-7</td>
<td>3.4</td>
<td>5.7</td>
<td>&gt;11</td>
<td>5-10</td>
<td>&gt;0.70</td>
<td>10 cm</td>
<td>5-8</td>
</tr>
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<td>BBSG EB</td>
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<td>2</td>
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<td>5.4</td>
<td>6-9</td>
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<td>&gt;0.70</td>
<td>10 cm</td>
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<td>5-7</td>
<td>3.2</td>
<td>5.4</td>
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<td>10 cm</td>
<td>3-8</td>
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<td>4</td>
<td>6-7</td>
<td>3.6</td>
<td>5.7</td>
<td>&gt;11</td>
<td>5-10</td>
<td>&gt;0.70</td>
<td>10 cm</td>
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<td>3.5</td>
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<td>3.5</td>
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<td>&gt;0.70</td>
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<td>3.5</td>
<td>3.2</td>
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<td>10 cm</td>
<td>5-8</td>
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<td>10 cm</td>
<td>3-8</td>
<td>≤10</td>
</tr>
<tr>
<td>Nom Grading Class</td>
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<td>k</td>
<td>% content ppc</td>
<td>PCG test</td>
<td>Duriez</td>
<td>Buiting test %</td>
<td>Modulus (15°C, 10 Hz) MPa</td>
<td>Fatigue (10°C, 25 Hz): m def</td>
<td>Difor angle 3 m</td>
<td>Test In situ</td>
</tr>
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</tr>
<tr>
<td>880r 0/6</td>
<td>1 2 3.4 4.9 20-26</td>
<td>≤10</td>
<td>11</td>
<td>≤20</td>
<td>≤0.80</td>
<td>1 cm</td>
<td>350</td>
<td>0.6</td>
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<tr>
<td>20/60</td>
<td>2 4.6</td>
<td>≥0.80</td>
<td>1 cm</td>
<td>350</td>
<td>0.6</td>
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</tr>
<tr>
<td>0/10</td>
<td>1 2 4.5</td>
<td>20-26</td>
<td>≤20</td>
<td>≤0.80</td>
<td>2 cm</td>
<td>250</td>
<td>0.8</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>2 6.9</td>
<td>2 8.9</td>
<td>10-12</td>
<td>≤20</td>
<td>≤0.75</td>
<td>3 cm</td>
<td>4-9</td>
<td></td>
<td></td>
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<tr>
<td>881 0/6</td>
<td>1 2 3.2 5.1 20-26</td>
<td>≤10</td>
<td>11</td>
<td>≤20</td>
<td>≤0.80</td>
<td>1 cm</td>
<td>350</td>
<td>0.6</td>
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<tr>
<td>2 4.6</td>
<td>2 6.9</td>
<td>10-12</td>
<td>≤20</td>
<td>≤0.75</td>
<td>2 cm</td>
<td>250</td>
<td>0.8</td>
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<tr>
<td>3 8.9</td>
<td>3 10-12</td>
<td>10-24</td>
<td>≤20</td>
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<td>3 cm</td>
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<tr>
<td>881M 0/6</td>
<td>1 2 3.5 5.1 20-26</td>
<td>≤10</td>
<td>11</td>
<td>≤20</td>
<td>≤0.80</td>
<td>1 cm</td>
<td>350</td>
<td>0.6</td>
<td></td>
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<tr>
<td>2 4.6</td>
<td>2 6.9</td>
<td>10-12</td>
<td>≤20</td>
<td>≤0.75</td>
<td>2 cm</td>
<td>250</td>
<td>0.8</td>
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<tr>
<td>3 8.9</td>
<td>3 10-12</td>
<td>10-24</td>
<td>≤20</td>
<td>≤0.75</td>
<td>3 cm</td>
<td>4-9</td>
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</tr>
<tr>
<td>881O 0/6</td>
<td>1 2 3.7 4.9 20-26</td>
<td>≤10</td>
<td>11</td>
<td>≤20</td>
<td>≤0.80</td>
<td>1 cm</td>
<td>350</td>
<td>0.6</td>
<td></td>
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<tr>
<td>2 4.6</td>
<td>2 6.9</td>
<td>10-12</td>
<td>≤20</td>
<td>≤0.75</td>
<td>2 cm</td>
<td>250</td>
<td>0.8</td>
<td></td>
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<tr>
<td>3 8.9</td>
<td>3 10-12</td>
<td>10-24</td>
<td>≤20</td>
<td>≤0.75</td>
<td>3 cm</td>
<td>4-9</td>
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<tr>
<td>882 0/6</td>
<td>1 2 3.5 5.1 20-26</td>
<td>≤10</td>
<td>11</td>
<td>≤20</td>
<td>≤0.80</td>
<td>1 cm</td>
<td>350</td>
<td>0.6</td>
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<tr>
<td>2 4.6</td>
<td>2 6.9</td>
<td>10-12</td>
<td>≤20</td>
<td>≤0.75</td>
<td>2 cm</td>
<td>250</td>
<td>0.8</td>
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<tr>
<td>3 8.9</td>
<td>3 10-12</td>
<td>10-24</td>
<td>≤20</td>
<td>≤0.75</td>
<td>3 cm</td>
<td>4-9</td>
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<tr>
<td>882M 0/6</td>
<td>1 2 3.7 4.9 20-26</td>
<td>≤10</td>
<td>11</td>
<td>≤20</td>
<td>≤0.80</td>
<td>1 cm</td>
<td>350</td>
<td>0.6</td>
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<tr>
<td>2 4.6</td>
<td>2 6.9</td>
<td>10-12</td>
<td>≤20</td>
<td>≤0.75</td>
<td>2 cm</td>
<td>250</td>
<td>0.8</td>
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<tr>
<td>3 8.9</td>
<td>3 10-12</td>
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<tr>
<td>883 0/6</td>
<td>1 2 3.7 4.9 20-26</td>
<td>≤10</td>
<td>11</td>
<td>≤20</td>
<td>≤0.80</td>
<td>1 cm</td>
<td>350</td>
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<tr>
<td>2 4.6</td>
<td>2 6.9</td>
<td>10-12</td>
<td>≤20</td>
<td>≤0.75</td>
<td>2 cm</td>
<td>250</td>
<td>0.8</td>
<td></td>
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<tr>
<td>3 8.9</td>
<td>3 10-12</td>
<td>10-24</td>
<td>≤20</td>
<td>≤0.75</td>
<td>3 cm</td>
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</tr>
<tr>
<td>883M 0/6</td>
<td>1 2 3.7 4.9 20-26</td>
<td>≤10</td>
<td>11</td>
<td>≤20</td>
<td>≤0.80</td>
<td>1 cm</td>
<td>350</td>
<td>0.6</td>
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<tr>
<td>2 4.6</td>
<td>2 6.9</td>
<td>10-12</td>
<td>≤20</td>
<td>≤0.75</td>
<td>2 cm</td>
<td>250</td>
<td>0.8</td>
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<tr>
<td>3 8.9</td>
<td>3 10-12</td>
<td>10-24</td>
<td>≤20</td>
<td>≤0.75</td>
<td>3 cm</td>
<td>4-9</td>
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</table>

Choosing asphalts for use in flexible pavement layers...
aggregate. The mixtures are traditionally laid between 50 and 80 mm thick and are designed to accommodate the requirements of each pavement layer.

Following many years of the increasing use of harder grades of bitumen in all layers of the pavement, some softer grades are now being used more commonly in France. This is, in part, due to the increasing incorporation of reclaimed asphalt pavement (RAP) in asphalts. Concurrently, there has been a rise in the use of warm asphalt mixtures in France, with lower temperature versions of the common mixtures outlined in this section being regularly produced and used.

11.5 Asphalts used in flexible pavements in Germany

In Germany, an asphalt pavement is constructed in a traditional manner (i.e. a base, a binder course and a surface course). All asphalts must comply with BS EN 13108 (BSI, 2005, 2006a–g). Implementation of the requirements of BS EN 13108 in Germany is specified in the German national standard TL Asphalt-StB (FSGV, 2013a). Layers in a flexible pavement in Germany are shown in Figure 11.4 (see Table 11.7).

The base is the lowest layer in a German asphalt pavement. It is placed on a foundation that is typically formed of an unbound material.

For the base and binder courses, asphalt concretes are normally used. The composition of the asphalt concretes in the base and binder courses differ significantly in terms of their void content, binder content and grading curve. To distinguish between base and binder course asphalts the base is called

- **AC T** (asphalt concrete Tragschicht (base))

and the binder course is called

- **AC B** (asphalt concrete Binderschicht (binder course)).

For the surface course, a range of asphalts is available

- **AC D** (asphalt concrete Deckschicht (surface course))
- **SMA** (stone mastic asphalt)

Surface course (asphalt concrete (AC D), stone mastic asphalt (SMA), mastic asphalt (MA) or porous asphalt (PA))

Binder course (AC B) (for heavily trafficked roads only)

Base (AC T)
PA (porous asphalt)
MA (mastic asphalt).

Materials used in Germany for particular layers in flexible pavements and their thickness ranges are shown in Table 11.7.

The thickness of these layers depends on climatic conditions and the applicable traffic load

- surface course – up to 40 mm
- binder course – 60–80 mm (for heavily trafficked roads only)
- base – up to 220 mm.

All asphalt layers are classified as L, N or S types, depending on the anticipated loading

- L – low loading
- N – normal loading
- S – severe loading.

11.5.1 Stone mastic asphalt
SMA was developed in Germany in the 1960s. It was originally designed to resist studded tyres. Today, SMA is used in Europe, Australia, the USA and Canada to reduce deformation in heavy duty asphalt pavements.
Typically, SMA is a gap graded mixture and normally contains cellulose fibres to prevent binder drainage (the fibres hold the bitumen in the mixture). As SMA is mainly used for heavily trafficked asphalt pavements, the use of polymer modified bitumen should be given careful consideration.

11.6 Asphalts used in flexible pavements in the USA

Over 93% of paved surfaces in the USA are asphalts (usually described in the USA and areas that have adopted US practice as hot mix asphalts (HMA)). As in Europe, the surface transportation system is mature, which places the emphasis on the maintenance, repair, rehabilitation or reconstruction of existing pavements, rather than new pavement construction.

In the USA, there is no national specification used throughout the country for road construction. The Federal Highway Administration (FHWA) and the American Association of State Highway and Transportation Officials (AASHTO) publish specifications, test methods and guidelines for a multitude of activities, but a state department of transportation (DOT) typically administers highway projects and publishes its own specifications. Local governments and other specifiers generally follow the lead of the state DOT specification, or use it outright for their projects. For airfield pavement construction, the Federal Aviation Administration publishes a standard specification that is modified for use on local projects.

Asphalt selection includes identification of the type, classification and specific aggregate and bitumen requirements that may be appropriate for a particular project. For this discussion, mixture type refers to general grading descriptions (dense, gap or open graded), while mixture classification refers to the nominal maximum aggregate size (NMAS) of a mixture, which is defined as ‘one sieve size larger than the first sieve to retain more than 10 per cent’ in the AASHTO standard M 323-13 (AASHTO, 2013), which uses the standard sieve nest for characterising the grading of aggregates used in asphalt in the USA.

As there is such a wide range of conditions in the USA, no single mixture type or classification can provide requirements for component materials that apply across the entire country. Consequently, specifiers need to select the appropriate bitumen grade and any aggregate quality requirements appropriate for the particular use of a particular asphalt. Tools such as LTPPBind, Version 3.1 (FHWA, 2003) are publicly available and provide a means of selecting the proper grade of bitumen according to the anticipated range of pavement temperatures, traffic characteristics and desired reliability for projects throughout North America. When selecting the bitumen grade, consideration should also be given to the proportion and nature of any reclaimd asphalt permitted for use in the mixture design.
Aggregate quality requirements often vary depending on the location of the asphalt within the pavement structure, the asphalt type and regional considerations reflecting the availability of particular materials. Test procedures and criteria for aggregate quality are described in Chapter 10.

### 11.6.1 Dense graded mixtures

As in other regions, dense graded asphalts have been widely used for paving in the USA since early in the twentieth century. Mixture design systems have evolved over time in a quest to provide a means of comprehensively analysing mixture qualities that are related to pavement performance.

Superpave (superior performing asphalt pavement) was a product of the Strategic Highway Research Program, a programme of focused research into long term pavement performance studies, cement and concrete, maintenance and bitumen that took place between 1987 and 1992. It includes a system for grading and selecting the correct bitumen for a particular asphalt, and a volumetric mixture design procedure that uses a gyratory compactor for moulding laboratory specimens. The performance graded (PG) bitumen specification was adopted first, with most state DOTs including either the AASHTO M 320-10 (AASHTO, 2010a) specification or a modification thereof. The Superpave mixture design procedure came into use more gradually but is currently the most commonly used method in the USA, although the Marshall method continues to be used, especially for airfield pavements. Superpave is discussed in detail in section 12.4.5.

Although no single standard specification is used throughout the USA, recommendations for mixture type selection have been jointly published by the National Asphalt Pavement Association (NAPA) and the FHWA in a document entitled *HMA Pavement Mixture Type Selection Guide* (NAPA and FHWA, 2001). This guide references the Superpave mixture design system for dense graded asphalt, which identifies mixture classifications according to nominal maximum particle size. Cross sections through US flexible pavements are shown in Figure 11.5.

Although there are significant differences in how Superpave has been implemented from one agency to another, the use of NMAS (nominal maximum aggregate size – nominal stone size (NSS) in the UK) to classify dense graded mixtures is commonplace, and is an important consideration for mixture selection, as lift thickness recommendations are directly related to the NMAS. Table 11.8 lists typical applications and compacted lift thicknesses commonly used for NMAS designations used with dense graded mixtures. Not all of these mixture classifications are used throughout the USA.

Dense graded mixtures may also be classified according to whether or not they are used as a surface course or according to the anticipated traffic level.
Surface course mixtures used where there is high speed traffic often require more stringent criteria for coarse aggregates in order to resist polishing and the loss of surface friction in wet conditions.

Performance related tests for resistance to rutting and various forms of cracking are increasingly used in state DOT specifications. Examples of these tests are listed in Table 11.9.

### 11.6.2 Speciality asphalts
Open graded mixture and SMA (stone matrix asphalts in the USA, stone mastic asphalts in the UK) have gained acceptance in parts of the USA for the

#### Figure 11.5 Layers in flexible pavements in the USA (NAPA and FHWA, 2001)

Surface course mixtures used where there is high speed traffic often require more stringent criteria for coarse aggregates in order to resist polishing and the loss of surface friction in wet conditions.

### Table 11.8 Superpave mixture classifications

<table>
<thead>
<tr>
<th>NMAS: mm</th>
<th>Typical application(s)</th>
<th>Typical range of compacted lift thickness</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.75</td>
<td>Thin surface course</td>
<td>19–38</td>
</tr>
<tr>
<td>9.5</td>
<td>Surface course</td>
<td>38–63</td>
</tr>
<tr>
<td>12.5</td>
<td>Surface course, binder course</td>
<td>50–100</td>
</tr>
<tr>
<td>19.0</td>
<td>Binder/base course</td>
<td>75–125</td>
</tr>
<tr>
<td>25.0</td>
<td>Binder/base course</td>
<td>100–150</td>
</tr>
<tr>
<td>37.5</td>
<td>Base course</td>
<td>125–175</td>
</tr>
</tbody>
</table>
same reasons as described for European countries. In 1990, industry, government and research organisations organised a tour of six western European countries to determine what could be learned from their asphalt paving practices that could be readily applied to the USA. The findings from this initiative led to the introduction of SMA, and the revision and reintroduction of permeable surface layer mixtures, borrowing and adapting the requirements for each mixture to conditions in the USA. As with dense graded mixtures, individual state DOTs have established their own specifications and requirements for these mixtures suited to the local climate and available materials.

There has been a great deal of interest in developing thin asphalts, particularly for use in pavement preservation. Examples include a thin overlay mixture (TOM) classification, developed by the Texas Department of Transportation, and thin, bonded overlay mixtures that usually require placement using an integral paver. Integral (or integrated) pavers are equipped with a bar that sprays a tack coat on the surface immediately before the asphalt is laid. While there are no standard definitions or accepted nomenclature for these asphalts, there is a trend towards their increased use due to the need to preserve and improve the functional characteristics of existing pavements using an approach that is less expensive and time consuming than traditional cold milling and asphalt inlay, and more durable under high traffic conditions than surface treatments.

### 11.7 Asphalts used in flexible pavements in India
A flexible pavement in India consists of asphalt layers, granular layers and the subgrade layer. A typical cross section of a flexible pavement in India for a national highway pavement is shown in Figure 11.6.

<table>
<thead>
<tr>
<th>Description</th>
<th>Test method</th>
<th>Distress mode(s)</th>
<th>Requirement(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hamburg wheel track testing of compacted hot mix asphalt (HMA)</td>
<td>AASHTO T 324 (AASHTO, 2014)</td>
<td>Rutting, sensitivity to moisture damage</td>
<td>Maximum deformation after a prescribed number of passes – the minimum number of passes to a defined inflection point. Usually performed at 50°C.</td>
</tr>
<tr>
<td>Asphalt Pavement Analyzer</td>
<td>AASHTO T 340 (AASHTO, 2010b)</td>
<td>Rutting</td>
<td>Maximum deformation after a prescribed number of passes at the high pavement temperature used for bitumen selection.</td>
</tr>
<tr>
<td>Indirect tensile strength</td>
<td>AASHTO T 322-07 (AASHTO, 2007)</td>
<td>Cracking</td>
<td>Minimum, or range of indirect tensile strengths measured at a prescribed temperature.</td>
</tr>
<tr>
<td>Texas overlay test</td>
<td>Tex-248-F (TxDOT, 2014)</td>
<td>Cracking</td>
<td>Minimum cycles to failure.</td>
</tr>
</tbody>
</table>
For national highways and expressways, the thickness of the asphalt layers is in the range 100–200 mm. The asphalt course is often constructed in three lifts: an upper layer, a middle layer and a lower layer. For other carriage-ways, such as class 2 highways, provincial highways, and urban or municipal roads, there are typically two asphalt layers: an upper layer and a lower layer. The upper layer, commonly 40–50 mm thick, is described as the surface course. The middle layer and lower layer, commonly 50–100 mm thick, are considered as acting as a binder course.

The most commonly used asphalts in India are specified in a standard published by the India Roads Congress (IRC, 2009)

- dense graded bituminous concrete (BC) surface course
- dense graded dense bituminous macadam (DBM) binder course.

Dense bituminous macadam (DBM) is mainly used as a base/binder course and profile corrective courses. The work consists of single or multiple layers of DBM on a previously prepared base or subbase. The thickness of individual layers is in the range 50–100 mm.

Bituminous concrete for use in surface courses can be a single layer of bituminous concrete on a previously prepared bituminous bound surface. A single layer can be 25, 40 or 50 mm thick.

The asphalt mixture design uses the Marshall method as specified in the Ministry of Road Transport and Highways (MoRT&H, 2013) specifications for roads and bridges in India. The Marshall method is discussed in detail in section 12.4.3. Aggregate gradings for DBM and BC mixtures are listed in Table 11.10.

In India, unmodified bitumen is graded by viscosity, and the commonly used grades are VG 30 and VG 40. The commonly used modified bitumens are

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### Table 11.10 Grading limits for DBM and BC asphalts in India (MoRT&H, 2013)

<table>
<thead>
<tr>
<th>Asphalt</th>
<th>Percentage passing sieve size: mm, listed below: %</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>45.0</td>
</tr>
</tbody>
</table>

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polymer modified bitumens (PMBs) and crumb rubber modified bitumens (CRMBs). The mechanical requirements for asphalts using unmodified and modified binders are given in IRC:111-2009 (IRC, 2009).

IRC:111-2009 recommends that when the traffic exceeds 2000 commercial vehicles per day (CVPD) per lane and the highest daily mean temperature exceeds 40°C, VG 40 or modified bitumen is used in BC and top layers of DBM.

11.8 Asphalts used in flexible pavements in China

Roads in China commonly consist of a flexible pavement with a semi-rigid base constructed with aggregate that is stabilised with lime or concrete. A typical flexible pavement in China is shown in Figure 11.7.

For major highways, including expressways, the thickness of the asphalt course is in the range 120–180 mm. The asphalt course is often constructed in three lifts: an upper layer, a middle layer and a lower layer. For other carriageways, such as class 2 highways, provincial highways, and urban or municipal roads, there are typically two layers, an upper layer and a lower layer. The upper layer, commonly 30–50 mm, is regarded as a surface course. The middle layer, commonly 40–60 mm thick, and the lower layer, 50–100 mm thick, are considered as acting as a binder course.

The asphalts commonly used in China are AC (asphalt concrete, a dense graded asphalt), ATB (asphalt treated base, a dense graded bitumen stabilised aggregate), ATPB (asphalt treated permeable base, an open graded asphalt), AM (asphalt macadam, semi-open graded mixture), SMA (stone mastic asphalt) and OGFC (open graded friction course). In some circumstances, Superpave mixtures (see section 12.4.5) are employed. For semi-rigid flexible pavements, the asphalts for individual layers are generally selected from those shown in Table 11.11.

**Table 11.11** Asphalts used in flexible pavements in China, as per JTJ F40-2004 (Research Institute of Highways, 2004)

<table>
<thead>
<tr>
<th>Asphalt course</th>
<th>Asphalt</th>
</tr>
</thead>
<tbody>
<tr>
<td>Upper layer</td>
<td>AC-13, AC-16, SMA-13, SMA-16, SUP-13</td>
</tr>
<tr>
<td>Middle layer</td>
<td>AC-16, AC-20, SUP-20</td>
</tr>
<tr>
<td>Lower layer</td>
<td>AC-25, AM-25, SUP-20, ATB-25, ATB-30</td>
</tr>
</tbody>
</table>
Design of individual asphalts is usually based on the Marshall method, as required by the national specification JTJ F40-2004 (Research Institute of Highways, 2004). Aggregate gradings for commonly used asphalts are listed in Table 11.12.

The most commonly used bitumens are heavy traffic bitumens AH-70 and AH-90 (penetration grade), used in the lower and middle layers, and sometimes in the upper layer. The commonly used polymer modified bitumens (PMBs) use SBS and SBR modifiers. In China, these modified binders are generally used in the upper layer for high class highways, such as expressways or class 1 or 2 highways. If the expressway has heavy duty traffic, PMB is also used in the middle layer.

Table 11.12 Grading limits for commonly used asphalts in China, as per JTJ F40-2004 (Research Institute of Highways, 2004)

<table>
<thead>
<tr>
<th>Asphalt mixture</th>
<th>Percentage passing sieve size: mm, listed below: %</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>31.5 26.5 19.0 16.0 13.2 9.5 4.75 2.36 1.18 0.6 0.3 0.15 0.075</td>
</tr>
<tr>
<td>SMA-13</td>
<td>100 90–100 50–75 20–34 15–26 14–24 12–20 10–16 9–15 8–12</td>
</tr>
</tbody>
</table>

References


AASHTO (2010a) AASHTO M 320-10. Standard specification for performance-graded asphalt binder. AASHTO, Washington, DC, USA.


Choosing asphalts for use in flexible pavement layers

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MoRT&H (Ministry of Road Transport and Highways) (2013) Specifications for Road & Bridge Works, 5th revision. IRC, New Delhi, India.

NAPA (National Asphalt Pavement Association) and FHWA (Federal Highway Administration) (2001) HMA Pavement Mixture Type Selection Guide. NAPA/FHWA, Lanham, MD/Washington, DC, USA.


Choosing asphalts for use in flexible pavement layers

Payne IR and Walsh ID (2011) ICE Manual of Highway Design and Management (Walsh ID et al. [eds]). Institution of Civil Engineers, London, UK.


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In many countries, asphalts that have traditionally been found to perform satisfactorily for acceptable periods have evolved over the years. Indeed, these empirical mixtures continue to be used world over. However, the best approach in terms of maximising the functionality and minimising the environmental effects is to design the asphalt for a particular purpose. Both European and US engineers are moving in that direction. This chapter discusses the methods used to design asphalt mixtures themselves. It begins with a very brief look at the history of the development of asphalt mixture design methods, considers the very important concepts of volumetrics and grading, and then describes the major asphalt design methods.

US practice is to use the word ‘asphalt’ for bitumen. In this chapter, the word ‘bitumen’ has been used except where the word ‘asphalt’ features in a quote, the title of an organisation, a process or a reference. In the sections on volumetrics and grading, US symbols have been employed. Note that Appendix 8 contains a comparison of US and EU symbols used in matters associated with mixture volumetrics.

12.1 A very brief history of asphalt mixture design methods

In the early days of asphalt pavements, the mixture employed would either have been one that the contractor chose or recommended or it may have been a proprietary mixture. In 1905, Clifford Richardson published his book *The Modern Asphalt Pavement*, with a second edition in 1908 (Richardson, 1908). It is clear that Richardson carried out tests in order to settle on the constituents for a particular asphalt.

In acknowledgement of the debt he owed to other scientists, Sir Isaac Newton wrote in 1676, ‘if I have seen further it is by standing on the shoulders of giants’. It is apt to begin here by acknowledging that today’s
achievements in the world of asphalt mixture design would not have been possible without the discoveries of the past.

In the mid-1920s, Charles Hubbard and Frederick Field in conjunction with the newly created Asphalt Association (which later became the Asphalt Institute) developed a system for designing asphalts called the Hubbard Field method of design. It was used by US state highway departments in the 1920s and 1930s, and continued in some states until the 1960s.

In 1927, Francis Hveem, a resident engineer in the California Division of Highways, began development of a method of assessing the bitumen content based on a calculation of the surface area of the constituent aggregate blend, something he achieved by 1932.

Bruce Marshall, an employee of the Mississippi Department of Highways, is probably the most well known pioneer associated with asphalt mixture design. He devised the Marshall method in the 1930s. Indeed, versions of his method remain in use today in areas throughout the world.

In 1987, the American Association of State Highway and Transportation Officials (AASHTO) member states began the 5-year research programme entitled the Strategic Highway Research Program (SHRP). It began because of an increasing number of premature pavement failures. In an attempt to deal with this problem, the states initiated the development of a coordinated, well funded, national research effort to derive improved specifications for bitumens and, ultimately, asphalts. It consisted of four areas of activity:

- bitumen (i.e. asphalt in the USA)
- concrete and structures
- highway operations
- long term pavement performance.

One of the outcomes of the bitumen study area was the Superpave system, the final product of the bitumen and asphalt mixture research programme. Superpave is a method of designing asphalt mixtures, and is used throughout the USA and areas that adopt US asphalt technology practices.

On a worldwide basis, the most widely implemented asphalt mixture design methods are as follows:

- Hubbard Field
- Hveem or California
- Marshall or Corps of Engineers
- Superpave.

In order to understand the various methods of designing asphalt mixtures, users must be familiar with the concepts of both mixture volumetrics and aggregate grading (often described as gradation). This chapter begins with
sections on these important subject areas and then considers the development history, an outline and usage of a number of different asphalt design mixtures.

Much has been published about the development of asphalts and asphalt mixture design methods (Hveem, 1970; White, 1985; Goetz, 1989; Leahy and McGennis, 1999).

12.2 Mixture volumetrics

As has been explained, volumetric analysis is a key element of asphalt mixture design. Accordingly, a sound understanding of this topic is necessary in order to design high quality, functional asphalts.

Pavement engineers often consider asphalts in terms of the proportion by weight of bitumen and/or aggregate. However, in relation to asphalt mixture design, it is important to consider an asphalt in terms of its three components

- aggregate
- bitumen
- air.

The complex interplay of these three components is the key to understanding the behaviour of asphalts both in the laboratory and in service.

The composition of an asphalt can be expressed either in terms of weight or volume. However, as air has no mass, if all three components are to be considered, then constituents need to be expressed in volumetric terms. While a typical asphalt may contain 5% bitumen and 95% aggregate by weight, the scenario changes completely when composition is considered volumetrically. Figure 12.1 denotes the composition by volume of three types of asphalt (Advanced Asphalt Technologies, 2011), stone matrix asphalt would be described as stone mastic asphalt in the UK.

![Figure 12.1 Typical composition by volume of different asphalts (Advanced Asphalt Technologies, 2011)](image-url)
The simplified layout of the component diagram shown in Figure 12.2 (sometimes described as a phase diagram) helps users to visualise the volumetric and mass relationships that are used in the analysis of an asphalt.

There now follows an explanation of some of the abbreviations used in Figure 12.2.

12.2.1 Air voids (VIM/Va)

The air void content is a measure of the total volume of air expressed as a percentage of the bulk volume of the compacted mixture. The air is distributed throughout a compacted asphalt, mostly between the coated aggregate particles. The air void content does not include pockets of air within individual aggregate particles or air contained in microscopic surface voids or capillaries on the surface of the aggregate.

Air voids define the performance of an asphalt. Air voids are a key consideration in designing the asphalt, and will have a profound effect on the life of the asphalt in service. If the air void content is too high, the resulting mixture may exhibit additional permeability to air and water, resulting in moisture damage.

**Figure 12.2 Symbols for constituents in an asphalt**

- \( V_{ma} \) = volume of voids in mineral aggregate
- \( V_{mb} \) = bulk volume of compact mix
- \( V_{mm} \) = voidless volume of paving mix
- \( V_{fa} \) = volume of voids filled with asphalt
- \( V_a \) = volume of air voids
- \( V_b \) = volume of asphalt binder
- \( V_{ba} \) = volume of absorbed asphalt binder
- \( V_{se} \) = volume of mineral aggregate (by bulk specific gravity)
- \( V_{sb} \) = volume of mineral aggregate (by effective specific gravity)

- \( M_{mix} \) = total mass of asphalt mixture
- \( M_b \) = mass of asphalt binder
- \( M_{be} \) = mass of effective asphalt binder
- \( M_{agg} \) = mass of aggregate
- \( M_{air} \) = mass of air = 0

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and age hardening. When air void content is too low, the binder content is likely to be high, resulting in an asphalt that is prone to bleeding and plastic deformation.

Determination of the air voids is one of the prime objectives of volumetric analysis, but there is no direct means of ascertaining its value. Air voids are determined by comparing the specific gravity (or relative density – specific gravity and relative density are the same property here but note that ‘relative density’ is the preferred term now) and is the ratio of the density (mass of a unit volume) of a substance to the density of a given reference material (in this case water, specific gravity usually means relative density with respect to water) of a compacted specimen with the maximum theoretical density of the mixture used to make that specimen. For example, if the compacted density of an asphalt specimen is 95.3% of the theoretical maximum specific gravity, the air void content is

$$VIM = 100 \left( \frac{G_{mm} - G_{mb}}{G_{mm}} \right)$$

where VIM is the air voids in the compacted mixture expressed as a percentage of the total volume, $G_{mm}$ is the maximum specific gravity of the mixture and $G_{mb}$ is the bulk specific gravity of the compacted mixture.

### 12.2.2 Bitumen content ($P_b$)

The bitumen content of an asphalt is one of its most important characteristics. Using a bitumen content lower than required may result in a dry, stiff mixture that is difficult to place and compact and will be prone to cracking and other durability problems while possessing lower resistance to fatigue cracking. Too much bitumen is likely to make the asphalt uneconomic because of the relatively high cost of bitumen and, in performance terms, it is likely to make the asphalt prone to bleeding and deformation.

$P_b$ is expressed and specified as a percentage of the total weight of the asphalt. However, expressing bitumen content by weight has two major drawbacks (Advanced Asphalt Technologies, 2011)

- it is the bitumen content by volume and not by weight that dictates performance
- most aggregates tend to absorb some bitumen, and that bitumen does not significantly contribute to the durability of a mixture.

### 12.2.3 Bitumen absorption ($P_{ba}$)

Any bitumen that is absorbed into the aggregate particles [$P_{ba}$] does not influence the performance characteristics of an asphalt but has the effect of
changing the specific gravity of the aggregate. Bitumen absorption is expressed as a percentage proportion by weight of the aggregate (not the asphalt!), and is calculated using the following equation

$$P_{ba} = \frac{100(G_{se} - G_{sb})G_b}{G_{se}G_{sb}}$$  \hspace{1cm} (12.2)

where $P_{ba}$ is the absorbed bitumen expressed as the percentage proportion of the total weight of the aggregate, $G_{se}$ is the effective specific gravity of the aggregate, $G_{sb}$ is the bulk specific gravity of the aggregate and $G_b$ is the specific gravity of bitumen.

### 12.2.4 Effective bitumen content ($P_{be}$)

The term effective binder content ($P_{be}$) is used to describe the amount of bitumen in a mixture excluding that portion of the bitumen that is absorbed by the aggregate. It is the effective binder content that is used in mixture volumetrics, and it is the portion of the total bitumen content that remains as a coating on the outside of the aggregate particles. It is this bitumen content on which the service performance of an asphalt depends. It is calculated using the following equation

$$P_{be} = P_b - \frac{P_{ba}P_s}{100}$$  \hspace{1cm} (12.3)

where $P_{be}$ is the effective bitumen content expressed as the percentage proportion of the total weight of the mixture, $P_b$ is the bitumen content expressed as the percentage proportion of the total weight of the mixture, $P_{ba}$ is the absorbed bitumen expressed as the percentage proportion of the total weight of the aggregate and $P_s$ is the aggregate content expressed as the percentage proportion of the total weight of the mixture.

### 12.2.5 Voids in the mineral aggregate (VMA)

The voids in the mineral aggregate (VMA) is the volume of void space between the aggregate particles of a compacted asphalt. VMA is numerically equal to the air void content plus the effective binder content by volume, and can be calculated using the following equation

$$VMA = 100 - \frac{G_{mb}P_s}{G_{sb}}$$  \hspace{1cm} (12.4)

where VMA is the voids in the mineral aggregate, $G_{mb}$ is the bulk specific gravity of the compacted mixture, $G_{sb}$ is the bulk specific gravity of the aggregate and $P_s$ is the aggregate content, expressed as the percentage proportion of the total weight of the mixture.

Establishing a single design air void content (such as the value of 4.0% used in Superpave mixtures) and then controlling VMA is the same as controlling the effective binder content. For example, a Superpave 12.5 mm mixture
designed at 4.0% air voids with 14.0% minimum VMA has a minimum effective binder content of $14.0 - 4.0 = 10.0\%$ by volume.

12.2.6 Voids filled with bitumen (VFB)

Voids filled with bitumen (VFB) is the percentage of VMA filled with bitumen, the balance being air voids (VIM). The bitumen content in the expression of VFB is the effective bitumen content. VFB can be calculated using the following equation

$$VFB = 100 \left( \frac{VMA - VIM}{VMA} \right)$$

(12.5)

where VFB is the voids filled with bitumen expressed as a percentage of the VMA, VMA is the voids in the mineral aggregate expressed as a percentage proportion of the bulk volume and VIM is the air voids in the compacted mixture expressed as a percentage proportion of the total volume.

In designing asphalts, VFB is closely related to both VMA and $P_{be}$. This occurs because the designed air voids is constant at about 4.0%, and, as VMA increases, both $P_{be}$ and VFB increase. Thus, in most cases, VFB can be considered to be simply an indicator of the richness of the mixture, as can VMA or $P_{be}$ (Advanced Asphalt Technologies, 2011).

It is not entirely clear what aspects of performance are related to VFB that are not also strongly related to other volumetric factors, especially $V_{be}$. Some engineers have suggested that fatigue resistance increases with increasing VFB. However, VFB and $P_{be}$ are closely related. Recent research strongly suggests that $P_{be}$ is a better overall indicator of resistance to fatigue cracking in asphalts. Therefore, in order to control or evaluate resistance to fatigue cracking, engineers should either use $P_{be}$ or VMA at a constant design air voids content. There is then little need to specify VFB independently (Advanced Asphalt Technologies, 2011).

12.2.7 Bulk specific gravity ($G_{sb}$) of an aggregate

The bulk specific gravity is the ratio of the weight in air of a unit volume of a permeable aggregate (including both permeable (to water and bitumen) and impermeable voids within the aggregate particles) at a particular temperature to the weight in air of an equal volume of gas free distilled water at the same temperature. It can be calculated using the following equation

$$G_{sb} = \frac{\text{weight of a unit volume of permeable aggregate in air}}{\text{weight of a unit volume of permeable aggregate in gas free distilled water}}$$

(12.6)

As an asphalt contains a mixture of coarse aggregate, fine aggregate and filler, the bulk specific gravity of the mixture can be determined using the
following equation

\[ G_{sb} = \frac{P_1 + P_2 + \cdots + P_n}{P_1/G_1 + P_2/G_2 + \cdots + P_n/G_n} \]  \hspace{1cm} \text{(12.7)}

where \( G_{sb} \) is the bulk specific gravity for the total aggregate; \( P_1, P_2, \ldots, P_n \) are individual percentages by weight of aggregates, where \( P_1 \) is the percentage by weight for aggregate fraction 1 and so on; and \( G_1, G_2, \ldots, G_n \) are individual bulk specific gravities of the aggregates, where \( G_1 \) is the bulk specific gravity of the aggregate in aggregate fraction 1 and so on.

**12.2.8 Apparent specific gravity (\( G_{sa} \)) of an aggregate**

The apparent specific gravity of an aggregate is the ratio of the weight in air of a unit volume of an impermeable aggregate at a particular temperature to the weight in gas free distilled water of a unit volume of the same impermeable aggregate at the same temperature. It can be calculated using the following equation

\[ G_{sa} = \frac{\text{weight of a unit volume of impermeable aggregate in air}}{\text{weight of a unit volume of impermeable aggregate in gas free distilled water}} \]  \hspace{1cm} \text{(12.8)}

**12.2.9 Effective specific gravity (\( G_{se} \)) of an aggregate**

The effective specific gravity of aggregate is the ratio of the weight in air of a unit volume of a permeable aggregate (excluding voids permeable to bitumen) at a fixed temperature to the weight in air of an equal volume of gas free distilled water at a stated temperature. It can be calculated using the following equation

\[ G_{se} = \frac{100 - P_b}{100/G_{mm} - P_b/G_b} \]  \hspace{1cm} \text{(12.9)}

where \( G_{se} \) is the effective specific gravity of the aggregate, \( G_{mm} \) is the maximum specific gravity of mixed material (no air voids), \( P_b \) is the bitumen content expressed as the percentage proportion of the total weight of the mixture at which the ASTM D2041 test \( (G_{mm}) \) \text{ (ASTM, 2011)} was performed and \( G_b \) is the specific gravity of the bitumen.

It is clear from the above that, for a given aggregate, the apparent specific gravity will have the highest value followed by the effective specific gravity and then the bulk specific gravity. Mathematically, it may be represented as

\[ G_{sa} > G_{se} > G_{sb} \]  \hspace{1cm} \text{(12.10)}

If the results of testing do not follow this relationship, then those results should be rejected and the material resampled and/or retested.
12.2.10 Specific gravity of a bitumen ($G_b$)
The specific gravity of a bitumen is the ratio of the weight in air of a unit volume of bitumen at a particular temperature to the weight in air of an equal volume of gas free distilled water at the same temperature. The methodology of determining the specific gravity of bitumen is given in AASHTO T 228 (AASHTO, 2009) or ASTM D70 (ASTM, 2009).

12.2.11 Bulk specific gravity of an asphalt ($G_{mb}$)
The bulk specific gravity of an asphalt is the ratio of the weight of a particular volume of asphalt including the permeable and impermeable voids in the specimen to the weight of the same volume of gas free distilled water. The standard procedure for determining the bulk specific gravity of a compacted asphalt involves weighing the specimen in air and in water. If the water absorption of the aggregates is less than 2.0%, saturated surface dry specimens are used (the procedure is outlined in AASHTO T 166) (AASHTO, 2013a). For specimens having high water absorption values (>2%), paraffin coated specimens should be used in the specific gravity determination (the procedure is given in AASHTO T 275) (AASHTO, 2007a).

$G_{mb}$ can be calculated using the following equation

$$G_{mb} = \frac{A}{B - C}$$  \hspace{1cm} (12.11)

where $G_{mb}$ is the bulk specific gravity of the compacted specimen, $A$ is the mass of the dry specimen in air (g), $B$ is the mass of the saturated surface dry specimen in air (g) and $C$ is the mass of the specimen in water (g).

The set up for determining $G_{mb}$ is shown in Figure 12.3.
12.2.12 Maximum specific gravity of the loose mixed material 
\((G_{\text{mm}})\)

The theoretical maximum specific gravity of an asphalt \((G_{\text{mm}})\) is the specific gravity of the mixture at zero air voids content. It is one of the most important tests to be performed in the design of an asphalt mixture, and high levels of repeatability are required for the mixture designer to draw accurate conclusions and make appropriate adjustments to the mixture design. Like bulk specific gravity, the theoretical maximum specific gravity does not affect the performance of an asphalt. However, it is essential in determining volumetric factors that are good indicators of performance, such as the air void content and VMA (Advanced Asphalt Technologies, 2011). The maximum specific gravity is determined by measuring the specific gravity of the loose asphalt having removed all of the air trapped in the mixture by subjecting it to a partial vacuum (vacuum saturation). The complete procedure is outlined in ASTM D2041 (ASTM, 2011). Following this procedure, the maximum specific gravity of a loose asphalt specimen is calculated using the following formula

\[
G_{\text{mm}} = \frac{A}{A + D - E} \quad (12.12)
\]

where \(G_{\text{mm}}\) is the theoretical maximum specific gravity of the loose mixture, \(A\) is the mass of the oven dry specimen in air (g), \(D\) is the mass of container filled with water at 25°C to the calibration mark (g) and \(E\) is the mass of the container with the specimen filled with water at 25°C to the calibration mark (g).

\(G_{\text{mm}}\) can also be calculated using the following equation

\[
G_{\text{mm}} = \frac{100}{P_s/G_{\text{se}} + P_b/G_b} \quad (12.13)
\]

where \(G_{\text{mm}}\) is the maximum specific gravity of the mixture at no air voids, \(P_s\) is the aggregate content expressed as the percentage proportion of the total weight of the mix, \(P_b\) is the bitumen content expressed as the percentage proportion of the total weight of the mix, \(G_{\text{se}}\) is the effective specific gravity of aggregate and \(G_b\) is the specific gravity of bitumen.

12.3 Grading
12.3.1 Historical development of packing theory

Aggregates typically constitute some 92–96% of an asphalt by mass. Clearly, the nature and grading of the aggregate in a particular asphalt will have a profound effect on the performance properties of that asphalt. Kandhal and Parker (1997) suggested that aspects of the aggregate can be related to particular asphalt defects. Table 12.1 is based on Kandhal and Parker’s findings.

Aggregate properties are considered in detail in Chapter 10 except for aggregate grading, which constitutes the subject area of this section.
Aggregate grading (often described as gradation) is the distribution of particle sizes in a batch of aggregates. It is one of the most important mixture design properties. A change in the size distribution of the aggregates can result in a different load distribution over the surface of the aggregate particles. The best grading for an asphalt concrete is that which gives the densest particle packing, thus increasing stability by having more inter-particle contacts and reducing the air voids. (This is obviously not the case for asphalt concretes that are designed to be permeable, e.g. porous asphalt or open graded friction course.) In dense mixtures, however, there must be sufficient air voids to permit the bitumen to be incorporated and assure durability without filling all the space to avoid bleeding and/or rutting (Roberts et al., 1996) (a condition sometimes described as voids over-filled with binder).

Arranging aggregates in an asphalt mixture is akin to packing matter into a confined space. Packing of spheres into a confined space and measuring its density has intrigued scientists and mathematicians since Isaac Newton (1643–1727). The German mathematician Johannes Kepler (1571–1630) worked on sphere packing in three dimensional Euclidian space, and proposed an arrangement known as close packing (either face centred cubic or hexagonal close packing, both of which have average densities of \( \pi/(3\sqrt{2}) \) approximately 0.740480489...). This is known as the Kepler conjecture, and states that no other arrangement of spheres has a higher density.

Fuller and Thompson (1907) studied different combinations of stones and sand with the aim of producing the densest concrete. The main objective of their study was to improve the quality of the mixture and decrease its cost by using the optimum quantity of cement. As a guide to obtaining the best concrete with constant cement content, the authors concluded that the stones should be evenly graded from fine to coarse, as an excessive amount of fine or middle sizes is very harmful to strength. They also concluded that the diameter of the largest grain of sand should not exceed one tenth of the diameter of the largest stone, and that the coarser the stone used, the coarser

### Table 12.1 Aggregate properties related to asphalt performance

<table>
<thead>
<tr>
<th>Aggregate property/test</th>
<th>Performance measure</th>
</tr>
</thead>
<tbody>
<tr>
<td>Grading and size</td>
<td>Permanent deformation and fatigue cracking</td>
</tr>
<tr>
<td>Uncompacted void content of coarse aggregate</td>
<td>Permanent deformation and fatigue cracking</td>
</tr>
<tr>
<td>Flat or elongated particles in coarse aggregate</td>
<td>Permanent deformation and fatigue cracking</td>
</tr>
<tr>
<td>Uncompacted void content of fine aggregate</td>
<td>Permanent deformation</td>
</tr>
<tr>
<td>Methylene blue test of fine aggregate (passing the 0.075 mm sieve)</td>
<td>Permanent deformation resulting from loading and environmental effects (i.e. asphalt stripping)</td>
</tr>
<tr>
<td>Micro-Deval [magnesium sulfate soundness]</td>
<td>Ravelling, potholes</td>
</tr>
</tbody>
</table>

Kandhal and Parker (1997)
the sand must be – an approach that provided a more dense and watertight concrete mixture.

Following this work, Talbot and Richart (1923) and Weymouth (1938) evaluated the size distribution of spheres that maximised density. They determined that when plotting on a semi-log basis (i.e. a linear scale on one axis and a log scale on the other axis) with the percentage passing a particular sieve (plotted on the linear y axis) versus the particle size (plotted on the logarithmic x axis) the maximum packing was obtained when a straight line on the graph had a slope of 0.5. This can be represented by the following equation

\[
P = 100 \left( \frac{d}{D} \right)^n
\]  

(12.14)

where \( P \) is the percentage of material by weight passing a given sieve with an opening of size \( d \), \( D \) is the maximum particle size in a given aggregate blend and \( n \) is the an exponent that affects the coarseness or fineness of the grading given by the slope.

Nijboer (1948) investigated the effect of particle size, taking shape into account, for asphalts and, as determined previously, he found that the densest packing was produced by a straight line in a semi-log plot of the percentage by weight passing a sieve size versus the size of the sieve. Empirically, it was found that for asphalts and aggregates as rough, shaped material this line had a slope of 0.45. Goode and Lufsey (1962) published a validation of Nijboer’s work, and described the numerical procedure for drawing a semi-power chart for gradings. This is the chart commonly referred as the 0.45 power chart (Meininger, 1992). The 0.45 power chart has become a cornerstone, as its use has been widely accepted worldwide including in the Superpave design method discussed later in this chapter.

12.3.2 Aggregate grading determination methodologies using packing theory

Although the development of semi-log graphs of the percentage of material passing each sieve versus sieve size is still used for determining aggregate gradings, an alternative method was proposed by Robert Bailey (formerly of the Illinois Department of Transportation) using particle packing concepts. This approach is discussed below.

It is a fact that granular particles such as aggregates cannot be placed to achieve 100% packing. Thus, there will always be voids between the aggregate particles. Vavrik et al. (2001) summarised the factors on which the degree of packing of aggregates depend:

- **Type and amount of compactive energy.** Several types of compactive force can be used, including static pressure, impact (e.g. Marshall
hammer) or shearing (e.g. gyratory shear compactor or California kneading compactor). Higher density can be achieved by increasing the compactive effort: for example, higher static pressure, more blows of the hammer, or more tamps or gyrations.

- **Shape of the particles.** Irregular elongated particles tend to resist packing in a dense configuration. Regular, blocky particles tend to arrange in dense configurations.

- **Surface texture of the particles.** Particles with smooth textures will more easily re-orient into denser configurations. Particles with rough, textured surfaces will resist sliding against one another, leading to lower density configurations.

- **Size distribution (gradation) of the particles.** Single sized particles will not pack as densely as a mixture of particle sizes.

### 12.3.2.1 Bailey’s method

In the early 1980s, Robert Bailey developed a method of grading analysis that takes into account the packing characteristics of individual aggregates, providing a quantified criterion that can be used to control mixture properties such as workability, segregation and compactibility (Vavrik et al., 2001). In a mixture design with a given compactive effort, three aggregate properties control the packing characteristics:

- **grading**
- **surface texture**
- **shape.**

Changing the grading of a mixture will influence the amount of void space in the aggregate skeleton.

The Bailey method marks a significant departure by defining coarse and fine aggregates as follows:

- **coarse aggregate** – large aggregate particles that when placed in a unit volume create voids
- **fine aggregate** – aggregate particles that can fill the voids created by the coarse aggregate in the mixture.

The Bailey method determines the coarse fraction as those particles that create voids, while the fine fraction is those particles that fit into the voids created by the coarse aggregate. It is this premise on which the method is based. It is also clear from the above that more than a single aggregate size is needed to define the descriptions coarse or fine. Figure 12.4 shows how the Bailey method makes use of a break point to define coarse and fine aggregates. The break point that determines the sieve size separating the coarse and fine aggregates is designated as the primary control sieve.
(PCS), which is determined using the following equation

$$\text{PCS} = \text{NMPS} \times 0.22$$

where PCS is the primary control sieve for the overall blend and NMPS is the nominal maximum particle size for the overall blend being one sieve size larger than the first sieve to retain more than 10% of the aggregate.

In addition, the term NMAS (nominal maximum aggregate size) is defined (AASHTO, 2014a) as the sieve size that is one size larger than the largest sieve that retains more than 10% aggregate. The significance of the NMAS in the selection and performance of a layer is examined later in this chapter.

Vavrik et al. (2001) explain that the value of 0.22 in the above equation was determined empirically. Various particle shape combinations and the resultant void sizes were studied. The analysis revealed the spacing as shown in Figure 12.5 and Table 12.2.
The void sizes vary from 0.15$d$ to 0.29$d$, where $d$ is the NMPS of the aggregate. Four cases were studied, considering all three particles to be round and replacing each round particle with a flat particle until all three particles are flat. Aschenbrener (2002) states that these figures were a result of a two dimensional analysis that gave particle diameter ratios of 0.155 (all round) to 0.289 (all flat). The void sizes are listed in Table 12.2.

A value of 0.22 was chosen, because it is the average of all four scenarios above, and the primary control sieves for Superpave gradings (the Superpave mixture design method is discussed below) are as shown in Table 12.3.

Table 12.3 clearly demonstrates that the coarse aggregate void size is a function of particle size and shape, and therefore the average size of the coarse aggregate voids in, say, a 9.5 mm NMAS mixture is smaller than the voids in a 37.5 mm NMAS mixture. Because of this change in average void size, a different size of material will be required to fit into the voids created. The primary control sieve determines not only which aggregates are considered as coarse and fine individually but also what portion of each individual aggregate is considered as coarse and fine for use in a given blend. The sieve found to be closest to the calculated value serves as the primary control sieve for a given blend. The amount of material above and below this specific sieve directly determines the volumes of coarse aggregate and fine aggregate (Vavrik et al., 2001).

### Table 12.2 Void sizes for various scenarios

<table>
<thead>
<tr>
<th>Details of particles</th>
<th>Void size</th>
</tr>
</thead>
<tbody>
<tr>
<td>All round</td>
<td>0.15$d$</td>
</tr>
<tr>
<td>2 Round/1 flat</td>
<td>0.20$d$</td>
</tr>
<tr>
<td>1 Round/2 flat</td>
<td>0.24$d$</td>
</tr>
<tr>
<td>All flat</td>
<td>0.29$d$</td>
</tr>
</tbody>
</table>

### Table 12.3 Primary control sieve for each corresponding nominal maximum particle size in the Superpave method

<table>
<thead>
<tr>
<th>Mixture NMPS: mm</th>
<th>NMPS x 0.22</th>
<th>PCS: mm</th>
</tr>
</thead>
<tbody>
<tr>
<td>37.5</td>
<td>8.250</td>
<td>9.5</td>
</tr>
<tr>
<td>25</td>
<td>5.500</td>
<td>4.75</td>
</tr>
<tr>
<td>19</td>
<td>4.180</td>
<td>4.75</td>
</tr>
<tr>
<td>12.5</td>
<td>2.750</td>
<td>2.36</td>
</tr>
<tr>
<td>9.5</td>
<td>2.090</td>
<td>2.36</td>
</tr>
<tr>
<td>4.75</td>
<td>1.045</td>
<td>1.18</td>
</tr>
</tbody>
</table>
12.3.3 Impact of aggregate grading on mixture volumetrics

This section considers how a compliant grading can result in the alteration of the mixture volumetrics of an asphalt (Advanced Asphalt Technologies, 2011). As an example, a grading is classified as coarse or fine depending on whether it passes below or above the maximum density line, respectively. In reality, both the gradings are dense, but one is coarser than the other in comparison with the maximum density gradation.

A concept based on packing theory and maximum density grading is the continuous maximum density (CMD) grading, which can be calculated using the following equation

\[ P_{\text{CMD}}(d_2) = P(d_1) \left( \frac{d_1}{d_2} \right)^{0.45} \]

where \( P_{\text{CMD}}(d_2) \) is the percentage passing the continuous maximum density grading for sieve size \( d_2 \), \( d_1 \) is one sieve size larger than \( d_2 \) and \( P(d_1) \) is the percentage passing sieve \( d_1 \).

The usefulness of the CMD grading is that it allows a comparatively detailed analysis of how closely a grading follows the maximum density grading calculated using Equation 12.16. Figure 12.6(a) shows a 9.5 mm grading compared with the standard maximum density grading as calculated using Equation 12.14, while Figure 12.6(b) shows the deviation using Equation 12.16. While both graphs show deviation, the graph in Figure 12.6(b) is much clearer in the way the deviation is shown. As an example, the aggregate grading actually follows the maximum density grading below the 1.18 mm sieve (as is evident from Figure 12.6(b)) while the upper graph (Figure 12.6(a)) shows a different picture.

This is further exemplified in Figure 12.7, where several gradings are considered and corresponding CMD plots are developed.

Figures 12.6 and 12.7, demonstrate that while there is a significant difference in their gradings, the fine aggregate fractions follow the maximum density grading very closely. The importance of the CMD plot is exemplified by the following explanation. Consider the dense/coarse grading plot in Figure 12.7 – the chart in Figure 12.7(a) indicates that the fine aggregate portion deviates significantly from the maximum density grading, and changing this portion may help to bring the VMA within limits, whereas the chart in Figure 12.7(b) shows that this portion already follows the maximum density grading and that the portion from 2.36 to 9.5 mm needs to be changed. This concept may be extended to the discussion on the modification of existing asphalt designs in which the changes will mostly consist of altering the air voids or the binder content, which are, primarily, modifications in VMA. The CMD plots can be used as a tool by the mixture designer to alter...
that portion of the grading that will have an impact on the VMA, as shown in Figure 12.8.

12.4 Mixture design methods
This section briefly considers a number of mixture design methods. The evolution, the method itself and an indication of usage is given for each of the following asphalt mixture design methods

- Hubbard Field
- Hveem or California
- Marshall (US Army Corps of Engineers)
- Smith or Asphalt Institute
- Superpave.

Figure 12.6 (a) A 9.5 mm aggregate grading compared with the maximum density grading; (b) deviation from the continuous maximum density grading for the same 9.5 mm grading (Advanced Asphalt Technologies, 2011)
While American design methods are highly regarded worldwide, it should be noted that other methods exist that take a different approach, most notably the French mixture design approach whose ‘methodology is based on component characteristics, water-sensitivity testing, void content assessments using gyratory compaction, resistance to permanent deformation, stiffness and fatigue resistance’ (Laboratoire Central des Ponts et Chaussées, 2007). It has not been possible to include all of the various methods within this chapter.

**12.4.1 Hubbard Field method**

By the early 1920s, the principles of soil compaction and the effect of water on density had been quantified and the Proctor method of evaluating soil compaction properties had been developed. Prevost Hubbard and Fredrick Field of the Asphalt Association (later to become the Asphalt Institute) developed a
method of mixture design in which asphalt samples were compacted using a Proctor hammer (Hubbard and Field, 1935) and by direct compression. The Hubbard Field mixture design method was developed for sheet asphalt and sand base mixtures with at least 65% of the aggregate passing the 2 mm sieve and 100% of the aggregate passing the 5 mm sieve. Sheet asphalts were pre-mixed asphalts made with bitumen and clean, angular sand dosed with mineral filler. Sheet asphalts were used as surface courses in the late 19th and early 20th century and consisted of a pre-mixed layer of graded aggregate, mineral filler and binder. They were laid in thicknesses varying from 40 to 55 mm and provided a smooth riding surface. Hubbard and Field demonstrated remarkable engineering wisdom in developing the traffic simulation test for the selection of the asphalt mixture. As the traffic at that time was dominated by wagons with steel wheels, Hubbard and Field used a punching shear test to simulate the punching effect of the steel wheels on the sheet asphalt mixtures. The punching test was conducted at 60°C, which was considered to be the highest pavement service temperature. The maximum load required to force a 50 mm diameter by 25 mm high compacted sample through a 43.75 mm diameter orifice under conditions of this test was reported as the stability value. It is interesting to note that some of the systems that were adopted in the Hubbard Field method have been followed in subsequent mixture design methodologies. Some of those methodologies are as follows

1. Measure the bulk density of the specimens.
2. Develop the following plots
   a. bulk density versus bitumen content
   b. stability versus bitumen content
   c. percentage voids in the total mixture versus bitumen content
   d. percentage voids in the aggregate mass versus bitumen content.
3. These plots were then used to select the design bitumen content by comparing them against design criteria.

4. The criteria, which included limits on stability values and percentage voids in the total mixture, were established by comparing the test values with the pavement performance.

5. The bitumen content would be selected based on the mid-point of the voids range: for example, if the voids requirement is from 3 to 5%, the bitumen content corresponding to a void content of 4% would be chosen. If the stability criteria are met in that mixture, it was adopted as the bitumen content.

6. The plot of bitumen content versus voids in the aggregate mass was used to adjust the grading when this was necessary to meet the voids criteria (Goetz, 1989).

Although the shortcomings of the punching shear type of test became apparent when applied to coarser mixtures, the principles of a design procedure incorporating stability and durability criteria in a systematic, quantitative way became well established. It has been reported that the Hubbard Field method was still in use by several states in the USA in the 1970s (Prowell and Ray Brown, 2006).

12.4.2 Hveem or California method

Francis Hveem is credited with the development of the Hveem or California method of asphalt mixture design. Hveem joined the California Division of Highways in 1917, and developed the stabilometer test during the late 1920s and early 1930s. Around 1927, Hveem was given the task of developing a method to determine the appropriate amount of ‘oil’ (bitumen) to add to aggregates in order to produce a ‘hard and smooth’ road surface that would not deform under traffic (Vallerga and Lovering, 1985). While working with oil–aggregate mixtures, Hveem observed that coarser gradings tended to require less oil than finer gradings, and made the connection that the surface area of the aggregate varied with the grading. He went on to identify a method for estimating the surface area of aggregate developed by a Canadian engineer, Captain L. N. Edwards, for concrete mixtures (Hveem, 1942, 1970).

The usage of ‘oil–aggregate mixtures’ to study the determination of design bitumen content using surface area concepts was first developed in the western USA, with the highway departments of California and Nebraska developing formulae based on the surface area concept (Goetz and Wood, 1960; Vallerga, 1953). The California Division of Highways (later the California Department of Transportation) carried forward work on the surface area concept, and used the Hveem method for the development of its highway network.
Hveem realised that, in addition to the surface area, the optimum bitumen content, or at least the point where the optimum bitumen content was exceeded and the stability decreased, was affected by the surface texture of the aggregate. Accordingly, a surface factor was used by Hveem in combination with the calculated surface area to determine the optimum bitumen content. Hveem observed that a smaller bitumen film thickness was required for smaller particles than for larger particles. Hveem continued to work on the surface area concept of aggregates in his quest for a rational mixture design method, and developed some protocols to measure the surface area factors accurately for fine and coarse aggregates. Hveem used the centrifuge kerosene equivalent (CKE) test to determine the surface area of the fine aggregates, taking into account the surface area, aggregate absorption and adjustment for surface texture. For coarse aggregates, the surface area is determined by the amount of SAE 10 oil that is retained on the aggregate. Hveem stated that the CKE method indicated the optimum bitumen content in 95% of cases (Hveem, 1970; War Department, 1948).

Hveem also evaluated the stability of asphalts. He hypothesised that, depending on the roughness and angularity of the aggregate, the film thickness at which the particles would become overly lubricated by the bitumen and therefore unstable would vary (War Department, 1948). Hveem developed his first stabilometer in 1930. The Hveem stabilometer evolved into a hydraulic device into which a compacted sample of asphalt was placed. The sample was loaded vertically on its flat surface, and the radial force transmitted to the surrounding hydraulic cell was measured. The stability value was calculated using the following equation

$$S = \frac{22.2}{P_v D_2 / (P_v - P_h)} + 0.222$$

(12.17)

where $P_v$ is the vertical pressure (\(\cong 400\) psi), $P_h$ is the horizontal pressure at a vertical pressure of $400$ psi and $D_2$ is the displacement of the sample (\(\cong \) number of turns of the handle).

The assumptions made by Hveem in developing the test (Stanton and Hveem, 1934) were that

- asphalt is a plastic solid, and a measure of its resistance to plastic deformation should be a measure of its stability
- the degree of distortion of an asphalt concrete pavement ‘is the accumulated results of a large number of quick shoves or pushes, each lasting a small fraction of time’.

The optimum bitumen content using the Hveem method was determined as follows. First, the bitumen contents for which moderate to heavy bleeding was observed on the surface of the compacted sample were eliminated. Next, any
bitumen contents that failed the minimum stability value were eliminated. Finally, the highest bitumen content that had at least 4% air voids was selected as the optimum. A swell test was also conducted to determine the resistance of the specimens to water induced damage (Vallerga and Lovering, 1985).

Vallerga and Lovering (1985) quote Hveem's own summary of his mixture design philosophy in 1937 as follows:

For the best stability, a harsh, crushed stone with some gradation, mixed with only sufficient asphalt to permit high compaction with the means available. For greatest resistance to abrasion, raveling, aging and deterioration, and imperviousness to water, a high asphalt content, broadly speaking, the richer the better. For impermeability, a uniformly graded mixture with a sufficient quantity of fine sand (fine sand is more important than filler dust). For non-skid surfaces, a large quantity of the maximum sized aggregate within the size limits used. For workability and freedom from segregation, a uniformly graded aggregate. To reduce the above factors to as simple a consideration as possible, it seems to be the best rule to use a dense, uniformly graded mixture without an excess of dust and to add as much oil or asphalt as the mixture will tolerate without losing stability.

In this quotation, the word ‘asphalt’ is that used in the USA for bitumen.

Much more detail of the development of the Hveem mixture design method can be found in documents published by Caltrans (1983) and Vallerga and Lovering (1985). For a step by step explanation of the Hveem mixture design method, a chapter in Manual Series No. 2 by the Asphalt Institute (1997) provides that information.

The Hveem mixture design method was used by some authorities prior to the implementation of the Superpave method. A study by Kandhal and Koehler carried out in 1984 in the USA indicated that, at that time, ten US states were using the Hveem mixture design method (Kandhal and Koehler, 1985). The performance of the Hveem method has been summarised by Vallerga and Lovering (1985)

It can be concluded that the Hveem method has served the state of California well over the past 50 years in the design and construction of a paved State highway system second to none.

**12.4.3 Marshall or Corps of Engineers method**

The basic concepts of the Marshall mixture design method were developed by Bruce Marshall around 1939 while he was an employee of the Mississippi State Highway Department. Marshall used many of the principles found in the Hubbard Field method. Instead of using a Proctor hammer, which covered
only part of the specimen face, the hammer size was increased to match the diameter of the specimen. The weight and drop of the hammer were increased to maintain a similar amount of compactive energy. Air voids, and later the VMA, were calculated for the compacted specimens. Marshall changed the orientation of his mechanical property test by turning the specimen on its side, placing it between two nearly semi-circular loading heads and loading it until it failed. The peak load sustained by the specimen was known as the Marshall stability. White (1985) has published an outstanding account of the development of the Marshall mixture design method.

During World War II, the Corps of Engineers was charged with selecting a method of asphalt mixture design to cope with the increasing tyre pressures found on military aircraft as aircraft weights began increasing during that period. The first research efforts addressing mixture designs for airfields within the US War Department were conducted at the US Army Tulsa District (War Department, 1943). Four mixture design methods were evaluated for their suitability, and the following conclusions were drawn:

The Hubbard field test is considered the most satisfactory method for general utility. The Texas punching method is similar but apparently over-emphasizes gradation and is not reproducible with different personnel. The Hveem stability test should be good for research but is not self-sufficient for routine testing since it measures chiefly internal friction. Experience has indicated that it is a good method of determining the most satisfactory bitumen content for a given aggregate. The Skidmore test is also not self-sufficient and it is believed to measure chiefly cohesion and interlock.

A continuation of the above work was assigned to the Waterways Experiment Station (WES) in September 1943. The first step was to develop ‘a simple apparatus suitable for use with the present California bearing ratio (CBR) equipment to design and control asphalt paving mixtures’ (War Department, 1944). The Marshall method at the time of the WES study (1943) had been used in southern US states by several highway agencies for about 4 years (War Department, 1944).

In the first phase of the study begun in 1943 (White, 1985), comparisons were made between the Hubbard Field and Marshall mixture design methods using a wide range of asphalts. From this study it was concluded that the Marshall stability test gave results that were comparable to the Hubbard Field stability test. Furthermore, the Hubbard Field test was not readily adaptable to the field CBR equipment, and the Marshall apparatus was also more portable. Therefore, the Marshall method was selected for additional study to evaluate some additional objectives. Some of the reasons given for recommending the adoption of the Marshall procedure for the design and field control of asphalt paving mixtures (War Department, 1944) were...
a. The Marshall stability test stresses and fails the entire specimen. Definite shear planes divide the specimen into four separate pieces. The stability value is a measure of the resistance of the specimen to the development of internal shear planes. This resistance is considered directly to the degree to which masses of particles are bonded together mechanically or with bituminous material. The Hubbard Field method produces a shear around the perimeter at the base of the specimen. Therefore stability is measured by a small fraction of the specimen. Several rock fragments may be contained within the specimen and would reflect increased stability only if they are in close proximity to the lower edge. Erratic stability values between identical specimens are possible as the laws of chance place more or less rock particles within the shear section. A number of check tests are therefore required for accurate details.

b. Tests can be made more rapidly with less effort by the Marshall stability method.

c. The equipment of the Marshall stability method weighs less and is more compact.

d. The density secured by the Marshall method appears to duplicate that obtained in the field more closely than does the Hubbard Field method. Therefore the amount of asphalt required in a mix is more accurately determined by the Marshall test.

Marshall design stability and flow criteria have evolved over the years and an example of the criteria used in the late 1970s is shown in Table 12.4. For the present day user, the Marshall criteria have evolved significantly to provide criteria based on the traffic that the pavement will carry, and are presented in Table 12.5.

A brief description of the Marshall mixture design method (the test procedure can be found in Manual Series No. 2 by the Asphalt Institute (1997)) is as follows

1. Aggregate selection. The aggregate selection consists of conducting the grading and other tests such as the aggregate specific gravity and so on to determine the suitability of an aggregate for asphalt mixture design.

| Table 12.4 Typical Marshall design criteria (Asphalt Institute, 1979) |
|---------------------------------|-----------------|-----------------|-----------------|
| Mix criteria | Light traffic ($<10^4$ ESALs) | Medium traffic ($10^4$–$10^5$ ESALs) | Heavy traffic ($>10^5$ ESALs) |
| Compaction (number of blows on each end of the sample) | 35 | 50 | 75 |
| Stability (minimum) | 224 N (500 lbs) | 336 N (750 lbs) | 6672 N (1500 lbs) |
| Flow (0.25 mm (0.01 inch)) | 8 | 20 | 8 | 18 | 8 | 16 |
| Percentage air voids | 5 | 3 | 5 | 3 | 5 |
Aggregates are chosen based on compliance with local specifications. The traditional Marshall mixture design requires aggregates to have a maximum size of 25 mm. For aggregate sizes larger than 25 mm and up to 38 mm, the modified Marshall method should be followed. This is outlined in section 5.16 of MS-2 (Asphalt Institute, 1997).

2. Binder selection. The binder is selected based on the local specifications, which may require the binder to be selected based on penetration, absolute viscosity, or the Superpave performance grading.

3. Preparing laboratory specimens. The Marshall method requires that cylindrical samples of approximately 64 mm in height and 102 mm in diameter are prepared. The mixture designer may start the sample preparation with about 1.2 kg of aggregates per mould. If the height is more than the permissible 64 mm, the following equation can be used to adjust the height of the specimen

\[
\text{adjusted mass of aggregate} = 63.5 \times \frac{\text{mass of aggregate used}}{\text{specimen height (mm) obtained}}
\]

(12.18)
In order to determine the optimum binder content, specimens at six different binder contents around the expected binder content are prepared. For example, if the optimum binder content is expected to be around 4.5%, specimens are prepared with binder contents of 3.0%, 4.0%, 4.5%, 5.0%, 5.5% and 6.0%. Three samples are prepared for each binder content chosen. As a starting point, the binder content can be estimated from the following equation (Asphalt Institute, 1997)

\[
DBC = 0.035a + 0.04b + Kc + F
\]  

(12.19)

where DBC is the approximate design bitumen content, as the percentage by total weight of the mixture; \(a\) is the percentage by weight of the mineral aggregate retained on a 2.36 mm sieve; \(b\) is the percentage by weight of the mineral aggregate passing the 2.36 mm sieve and retained on a 0.075 mm sieve; \(c\) is the percentage by weight of the mineral aggregate passing the 0.075 mm sieve; \(K\) = 0.15 for 11–15% passing the 0.075 mm sieve, 0.18 for 6–10% passing the 0.075 mm sieve and 0.20 for 5% or less passing the 0.075 mm sieve; and \(F\) = 0–2% (this factor is based on the absorption of bitumen and, in the absence of other data, a value of 0.7 is suggested).

Once the quantity of graded aggregate is fixed, the batched aggregates are heated to constant weight and the binder is heated to the mixing viscosity. MS-2 requires that temperatures at which the bitumen attains kinematic viscosities of 170 ± 20 cSt kinematic and 280 ± 30 cSt kinematic shall be established as the mixing and compaction temperatures, respectively. The Marshall method does not require any ageing/curing of the asphalt prior to compaction. The Marshall specimens are prepared using the standard Marshall compactor that consists of a flat faced hammer that is 98.3 mm in diameter, weighs 4.5 kg and falls freely by a distance of 457 mm onto the asphalt placed within the specimen mould. The Marshall compactor is required to impart a fixed number of blows per face on the specimen, usually 75 blows (for details refer to Table 12.5). After compaction, the specimens are extruded from the mould and then allowed to cool to room temperature (preferably overnight).

4. **Mechanical testing of the asphalt.** The Marshall method requires that the specimens are tested for stability and flow. Testing for stability and flow utilises the *Marshall stabilometer*, which applies a load at a constant rate of vertical strain rate of 51 mm per minute through cylindrical testing heads, as shown in Figure 12.9. The maximum load taken by the specimen at 60°C is noted as the Marshall stability value, and the amount of deformation at the maximum load is designated as
the flow. The stability and flow are measured using calibrated proving rings, and the values are reported in newtons and millimetres, respectively. Prior to placing the extruded Marshall specimen in the Marshall stabilometer breaking head, the specimens are placed in a water bath at $60 \pm 1^\circ C$ for 30–40 min prior to the test, and the stability and flow test should be completed within 30 s of extracting the specimen from the water bath.

5. Density and voids analysis. The Marshall asphalt specimens and loose asphalt specimens are then tested to determine the following parameters
   a. The average bulk density of the Marshall asphalt specimens ($G_{mb}$) (AASHTO, 2013a) and the average specific gravity of the binder ($G_b$) (ASTM, 2009).
   b. The theoretical maximum specific gravity ($G_{mm}$) as per ASTM D2041 (ASTM, 2011) at two binder contents close to the optimum binder content. The $G_{mm}$ value will be used to determine the effective specific gravity of the aggregate ($G_{se}$) (Equation 12.9).
   c. Using the three aggregate specific gravities, the specific gravity of the binder, the $G_{mm}$ value and the three key volumetric parameters (i.e. air voids ($V_a$) (Equation 12.1), voids filled with bitumen (VFB) (Equation 12.5) and voids in the mineral aggregate (VMA) (Equation 12.4)). The formulae for calculating the above can be found earlier in this chapter in section 12.2 on mixture volumetrics – corresponding equation numbers and references have been appended.

6. Data interpretation. The test data are interpreted as follows
   a. The stability values are standardised based on the specimen height.
   b. The following plots are developed
      i. $V_a$ versus binder content
      ii. VFB versus binder content
      iii. VMA versus binder content

![Schematic diagram of load application in the Marshall stability test](image-url)
iv. stability versus binder content
v. flow versus binder content
vi. $G_{mb}$ of the mix versus the binder content.

Typical results from the above tests are given in Figure 12.10, which shows typical trends of the plots as noted above. These graphs are used to determine the optimum binder content, by plotting them as

![Graphs showing typical trends](image)

**Figure 12.10** Typical plots from a Marshall volumetric analysis
shown in Figure 12.11, which helps the mixture designer to observe
the influence of binder content on the various mixture volumetric
parameters at a glance and to determine the binder content suitable
for the project.

7. *Selection of the optimum binder content.* The test data and plots are
further analysed, and the initial binder content is selected as a binder
content that satisfies all strength and volumetric requirements. The
Asphalt Institute (1997) recommends a starting point as the median of
the percentage of air voids, say the binder content at 4% air voids.
Specimens are prepared at this binder content and evaluated for the
strength and volumetric requirements, and further adjusted.

8. *Selection of the final mixture design.* The final mixture should be that
which is the most economical and which meets all the criteria.
However, the mixture designer must use his expertise to ensure that all
mixture properties are balanced. Low binder contents are financially
attractive. However, the designer must ensure that unduly low binder
contents are avoided, as this can have an adverse effect on the
properties of the asphalt. The Asphalt Institute (1997) suggests that a
narrow range of binder content satisfying all binder requirements
should be selected. This is shown in Figure 12.11. However, it must
be borne in mind that the requirements of most projects are unique,
and the mixture designer must make the necessary adjustments to
ensure that these requirements are satisfied prior to deciding on the
value of the optimum binder content. The plot shown in Figure 12.11
enables the mixture designer to view all the responses of the change in
binder content versus all volumetric parameters. This unique plot helps the mixture designer not only to validate the design but also to decide on the values of tolerances that should apply to a particular project.

The Marshall method has enjoyed widespread usage the world over for the design of dense graded asphalts. Indeed, a number of authorities continue to use it today in conjunction with their local specifications.

12.4.4 Smith or Asphalt Institute method

The Hubbard Field, Hveem and Marshall mixture design methods are all empirical, with criteria developed from the correlation of laboratory and field studies. V. R. Smith and the Asphalt Institute applied the principles of the triaxial compression test to the testing of asphalts (Goetz and Wood, 1960). Smith used a closed system triaxial test wherein the vertical load was applied in static increments and the lateral transmitted pressure was measured when equilibrium had essentially been reached. Specimens of rational size (at least twice their diameter) were formed by spading and double plunger compaction or by the use of the kneading compactor. The test was conducted at room temperature, as the effects of temperature and the rate of loading are virtually non-existent under the conditions of the test. The data obtained were used to generate a plot of vertical applied load versus lateral transmitted pressure, from which the parameters angle of internal friction ($\phi$) and cohesion ($c$) were calculated. From theoretical considerations, Smith developed the relationship between the angle of internal friction and the cohesion for varying applied stresses, and presented this in the form of supporting power curves.

By testing many mixtures in the laboratory whose performance in the field was known, an evaluation chart was devised that delineated regions of satisfactory and unsatisfactory mixtures in a plot of the angle of internal friction versus the cohesion. Boundary conditions were set on the basis of traffic severity and the minimum angle of internal friction required for satisfactory performance while applying voids criteria.

Goetz (1989) opines on the test while a triaxial test is a rational one that generates the measurement of parameters of a more fundamental nature than those measured previously and does use theoretical concepts in applying these measurements to mixture design, in the final analysis, it depends upon correlation between field and laboratory results for application. The measured parameters can be determined in a rational way for known stress conditions in the laboratory, but the pavement is not loaded in the rational way in service and stress conditions in the pavement are unknown . . . . In spite of such shortcomings from a design point of view, a design based on fundamental
measurements was considered progressive in anticipation of the day when stress conditions in the pavement could be more accurately defined and rational strength test data applied.

Schaub and Goetz (1961) were of the view that

the triaxial shear strength test has been recognized as the most fundamental in that it uses field-simulated confined loading where stress conditions are known and statically determinant methods of analysis can be applied.

The complete description of the Smith mixture design method can be found in two publications by Smith (1949, 1951), and a complete theoretical consideration of the triaxial state of stress can be found in a paper by McLeod (1950). A comparison of the Marshall and Smith methods can be found in a paper by Goetz (1951).

The Smith or Asphalt Institute method was used more for the purposes of research rather than in the field.

**12.4.5 Superpave**

12.4.5.1 Background to Superpave

The Strategic Highway Research Program (SHRP) Asphalt Research Program was developed in the USA in the early 1980s because of an increasing number of premature asphalt pavement failures. In recognition of this problem, US states initiated the development of a coordinated, well funded, national research effort to develop improved specifications for bitumens and, ultimately, asphalts. The importance of and need for specification development in the SHRP Asphalt Research Program originated in Transportation Research Board Special Report 202, America’s Highways: Accelerating the Search for Innovation (TRB, 1986a), popularly known as the blue book, which presented the conclusions and recommendations of the Strategic Transportation Research Study (STRS). The authors of the blue book stated the objective of the bitumen research programme as follows:

To improve pavement performance through a research program that will provide increased understanding of the chemical and physical properties of asphalt cements and asphalt concretes. The research results would be used to develop specifications, tests . . . needed to achieve and control the pavement performance desired.

This emphasis was reinforced and further defined in the May 1986 Transportation Research Board report Strategic Highway Research Program Research Plans (TRB, 1986b), popularly known as the brown book. This report stated a specific constraint or guideline for the bitumen programme: ‘the final product will be performance-based specifications for asphalt, with or without
modification, and the development of an asphalt–aggregate mixture analysis system (AAMAS) (where ‘asphalt’ means ‘bitumen’).

The brown book further described the programme’s Project 1-4, ‘Preparation of Performance-based Specifications for Asphalt and Asphalt–aggregate Systems’. This project consisted of two tasks: to develop the bitumen specification and to develop the AAMAS.

Finally, the SHRP executive committee in 1987 approved A Contracting Plan for SHRP Asphalt Research (SHRP, 1987). This blueprint became the strategic plan for the SHRP Asphalt Research Program, and took precedence over earlier research plans when issues of proper technical direction arose. The contracting plan combined the many tasks identified in the 1986 brown book into a coordinated, manageable structure of eight main contracts.

The responsibility for the development of the Superpave performance based binder specification, performance based bitumen–aggregate mixture specification and mixture design system, and for the technical direction and coordination of the entire programme, was assigned to the A-001 contractor, The University of Texas at Austin, in October 1987.

Between 1987 and 1993, the Asphalt Research Program was established and carried out through the award of the eight major research contracts and an additional 15 supporting studies. Throughout this period, the goals of the programme remained substantially unchanged from those originally articulated in the blue book, although details were changed as the programme evolved.

Four significant changes to the research and contracting plans evolved through an ongoing dialogue in the highway community that participated in the development of the programme:

- The term asphalt was broadened to asphalt binder in recognition of the fact that the specification encompasses modified as well as unmodified binders.
- The original concept of an SHRP specification for an asphalt–aggregate mix analysis system (AAMAS) evolved to a performance based specification for asphalt–aggregate mixtures supported by a distinct mixture design system. (The term ‘AAMAS’ was introduced by NCHRP Project 9-6 (Von Quintus et al., 1991). The SHRP adopted the term Superpave in order to distinguish the two research efforts and to better describe the specific goal of the SHRP research.)
- The term Superpave was chosen in 1991 to signify the integrated structure of performance based specifications, test methods, equipment and protocols, and a mixture design system. The Superpave system is the principal product of the SHRP Asphalt Research Program.
Also in 1991, the Superpave software was developed to provide a unifying framework for the entire system.

The eight major contracts of the SHRP Asphalt Research Program are listed in Table 12.6.

The major products of the SHRP Asphalt Research Program were

- a performance based specification for asphalt binders and the supporting test methods and equipment
- a performance based mixture design system with supporting test methods and equipment
- a modifier evaluation protocol
- the Superpave specification, design and support software.

These products are an integrated, coordinated system, and are tied together by the Superpave software. The Superpave system provides a means for the rational design of asphalt paving mixtures, and provides a framework and mechanism to tie mixture and structural design together.

Superpave was developed to replace the diverse specifications and mixture design methods used by the 50 US states and other transportation agencies. Superpave provides a single, performance based system that can account for the distinct traffic and environmental conditions found throughout the USA, Canada and other parts of the world. It was developed to address and minimise permanent deformation, fatigue cracking and low temperature

---

Table 12.6 The eight major contracts of the SHRP Asphalt Research Program

<table>
<thead>
<tr>
<th>SHRP contract number</th>
<th>Contract title (contractor)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A-001</td>
<td>SHRP Asphalt Research Program: Technical Direction, Specification and Superpave Development (The University of Texas at Austin)</td>
</tr>
<tr>
<td>A-002A</td>
<td>Binder Characterization and Evaluation (Western Research Institute)</td>
</tr>
<tr>
<td>A-002B</td>
<td>Novel Approaches for Investigation of Asphalt Binders (University of Southern California)</td>
</tr>
<tr>
<td>A-002C</td>
<td>Nuclear Magnetic Resonance Investigation of Asphalt (Montana State University)</td>
</tr>
<tr>
<td>A-003A</td>
<td>Performance-related Testing and Measuring of Asphalt–aggregate Interactions and Mixtures (University of California at Berkeley)</td>
</tr>
<tr>
<td>A-003B</td>
<td>Fundamental Properties of Asphalt–aggregate Interaction Including Adhesion and Adsorption (Auburn University)</td>
</tr>
<tr>
<td>A-004</td>
<td>Asphalt Modification Practices and Modifiers (Southwestern Laboratories)</td>
</tr>
<tr>
<td>A-005</td>
<td>Performance Models and Validation of Test Results (Texas Transportation Institute)</td>
</tr>
</tbody>
</table>

Kennedy et al. (1994)
cracking, and it considers how the effects of ageing and moisture damage contribute to the development of these three distress modes.

12.4.5.2 Overview of Superpave

This section is based on the National Research Council publication SHRP-A-410, *Superior Performing Asphalt Pavements (Superpave): The Product of the SHRP Asphalt Research Program*, which was published in 1994 (Kennedy et al., 1994).

Superpave mixture design encompasses the performance based tests and performance prediction models developed in the SHRP Asphalt Research Program using mixture volumetrics. Superpave has been developed for ready application to all classes of roadway, from rural or urban residential streets to roads carrying substantial numbers of heavy commercial vehicles. The Superpave hierarchical approach is illustrated conceptually in Figure 12.12. It matches the appropriate level of mixture design effort and technology to the pavement being designed.

The Superpave mixture design is an example of a hierarchical mixture design method that utilises defined subsystems to define the requirements with which the mixture design must comply. The three levels of design are defined based on traffic and the use to which the pavement is put. While the suggested boundary values are 1 million and 10 million equivalent single axle loads (ESALs), the actual traffic levels are established by the user. All three design levels include a volumetric mix design phase, while level 2 and level 3 designs require performance based tests to allow the mixture design to be optimised for resistance to deformation, fatigue cracking and low temperature cracking – the three primary distress modes that SHRP binders and mixtures are designed to counter. (Although the criteria for level 2 and level 3 in Figure 12.12 appear to be identical, the detailed requirements that have to be met for level 3 are much more demanding than those for level 2.)
A general flow chart of the Superpave mixture design system is shown in Figure 12.13. For level 1, the procedure involves only volumetric design, which evaluates aggregates and asphalt binders in order to select a grading and binder content that satisfy specified criteria for air voids, voids in the mineral aggregate and voids filled with asphalt. For levels 2 and 3, performance based tests are conducted, and estimates of distress with time are made. This allows the mixture design to be optimised with regard to one or more of the three distress modes: deformation, low temperature cracking and fatigue cracking. Subsequent to completion of the mixture design, modifications may be required because of changes to the material properties once the mixture is produced in a plant. These modifications may or may not require retesting.
During construction, control tests are undertaken at the production plant to verify that the mixture being manufactured accords with the laboratory mixture design. Level 3 also includes an optional proof testing scheme that allows the mixture to be subjected to tests that simulate the actual traffic and environmental conditions to confirm that the mixture actually performs as required.

In level 2 mixture design, a smaller number of tests are performed at a smaller number of temperatures than for level 3 mixture design. Performance based tests for rutting are undertaken at a single effective temperature for deformation. Similarly, tests to predict fatigue cracking are performed at a single effective temperature for fatigue cracking. Effective temperatures for deformation and fatigue cracking are defined as the single test temperature at which the amount of deformation or fatigue cracking that occurs is equivalent to that obtained by considering each season separately throughout a year. These two temperatures will be different from each other, and can be calculated using equations contained in Chapter 4 of SHRP-A-408 (Cominsky et al., 1994). Low temperature tensile strength is measured at a single temperature in level 2 design.

Level 3 mixture design simulates the entire year by breaking it into representative seasons. Performance based tests for deformation and fatigue cracking are performed at a range of temperatures. A greater number of tests are conducted to evaluate more rigorously mixture response across a greater range of stresses. Deformation and fatigue cracking are predicted using mixture properties in each of the representative seasons. A summary comparison of level 2 and level 3 is shown in Table 12.7.

### Table 12.7 Comparison of level 2 and level 3 mixture designs

<table>
<thead>
<tr>
<th>Test types</th>
<th>Permanent deformation/fatigue cracking</th>
<th>Low temperature cracking</th>
</tr>
</thead>
<tbody>
<tr>
<td>Level 3 considers more states of stress, and requires two additional test methods</td>
<td>Level 3 considers a range of temperatures from 4 to 40°C</td>
<td>No difference between level 2 and level 3</td>
</tr>
<tr>
<td>Level 3 uses one effective temperature for fatigue cracking and one for permanent deformation</td>
<td>Level 3 considers three temperatures</td>
<td>Level 2 considers tensile strength at one temperature only</td>
</tr>
<tr>
<td>Level 3 breaks the year into seasons</td>
<td>Level 2 considers the entire year as a single season</td>
<td>No difference between level 2 and level 3</td>
</tr>
</tbody>
</table>

Kennedy et al. (1994)

12.4.5.3 Laboratory compaction

Superpave marked the departure from impact compaction systems to a gyratory mechanism for the preparation of laboratory specimens. The entire mixture design system, including field control, is based on the use of the
Superpave gyratory compactor (SGC). The adoption of the gyratory mechanism for the laboratory compaction of asphalt specimens followed consideration of the topic including use by the French of this method of compaction. The SHRP committee defined the following as being the key advantages of the SGC (Kennedy et al., 1994):

- It is relatively inexpensive, portable and capable of quickly moulding specimens with minimal variation between individual samples.
- The performance properties of the compacted specimen simulate the performance properties of cores from pavements constructed with the same bitumen–aggregate combination.
- It allows the compactibility of the mixture to be evaluated including an estimate of the final air voids content under traffic (the probability of the mixture becoming plastic under traffic) and a measure of the grading of the aggregate in the mixture.

The SGC (Figure 12.14) has the following characteristics:

- an angle of gyration of $1.25 \pm 0.02^\circ$
- a rate of 30 gyrations per minute
- a vertical pressure during gyration of 600 kPa
- the capability of producing specimens that are 150 mm in diameter having a height of 150 mm.

The trend line that is obtained by plotting the change in relative density with the increasing number of gyrations, known as the densification curve, also defines the three different values of compactive effort specified in the Superpave mixture design procedure:

- $N_{\text{ini}}$, the initial compactive effort
- $N_{\text{design}}$, the design compactive effort
- $N_{\text{max}}$, the maximum compactive effort.
where $N$ is the number of gyrations to which the asphalt specimen is subjected (Figure 12.15).

The compactive efforts, $N_{\text{ini}}$ and $N_{\text{max}}$, are used to evaluate the compactability of the mixture, while $N_{\text{design}}$ is used to select the bitumen content. Densities that correspond to each value of compactive effort are designated $C_{\text{ini}}$, $C_{\text{design}}$, $C_{\text{max}}$.

These densities are expressed as a percentage of the maximum theoretical specific gravity (AASHTO T 209 (AASHTO, 2011a)).

In the original Superpave manual (Kennedy et al., 1994) the values of $N_{\text{ini}}$, $N_{\text{design}}$ and $N_{\text{max}}$ were defined as indicated below.

The number of gyrations for $N_{\text{ini}}$ varies from 7 to 10, and is given by the following equation

$$\log N_{\text{ini}} = 0.45 \log N_{\text{design}} \quad \{12.20\}$$

The $N_{\text{design}}$ values are selected from a matrix that defines $N_{\text{design}}$ values corresponding to traffic (in millions of standard axles (msa)) and the average design air temperature (in °C). It will be recalled that Superpave applies to all classes of road, from residential streets to heavily trafficked expressways, thus, the traffic values varied from 0.3 to >100 msa, with the temperature varying in the following ranges: <39°C and 39–45°C with increments of 2°C.

The number of gyrations for $N_{\text{max}}$ varies from 104 to 287, and is given by the following equation

$$\log N_{\text{max}} = 1.10 \log N_{\text{design}} \quad \{12.21\}$$

![Figure 12.15 Typical densification curves using the SGC](image-url)
The importance of aggregate structure (at the same binder content) is shown in Figure 12.16, and is indicated by the slope of the densification curve. Mixtures exhibiting relatively steep slopes and low $C_{ini}$ values are indicative of mixtures that have developed a good aggregate structure or internal resistance to densification. To ensure adequate structure, it is suggested that the specifications require that

$$C_{ini} \leq 89\%$$

where the number of gyrations $N_{max}$ varies from about 7 to 10 according to Equation 12.21.

A maximum density requirement at $N_{max}$ ensures that the mixture will not compact excessively under the anticipated traffic, become plastic and result in excessive deformation. Thus, the specification requires that

$$C_{max} \leq 98\% \text{ or air voids } \geq 2\%$$

As $N_{max}$ represents a compactive effort that would be equivalent to traffic that greatly exceeds the design traffic (ESALs), excessive compaction under traffic will not occur. Thus, the air voids will not drop below 2%, and the mixture will not become susceptible to plastic deformation.

12.4.5.4 Selection of aggregate

Consensus properties

In order to finalise the aggregate properties, the SHRP committee adopted the Delphi method for developing consensus. It was the view of the SHRP pavement researchers that certain aggregate characteristics were critical and needed to be achieved in all cases to arrive at asphalts that performed well. These characteristics were described as consensus properties because there was wide agreement in their use and specified values. Those properties are
**coarse aggregate angularity (CAA)**

This property ensures a high degree of aggregate internal friction and rutting resistance. It is defined as the percentage by weight of aggregates larger than 4.75 mm with one or more fractured faces. The test procedure for measuring the CAA is described in ASTM D5821 (ASTM, 2013). The procedure involves manually counting particles to determine the number of fractured faces for a defined size of sample. A fractured face is defined as any fractured surface that occupies more than 25% of the area of the outline of the aggregate particle visible in that orientation.

**Fine aggregate angularity**

This property ensures a high degree of fine aggregate internal friction and rutting resistance. It is defined as the percentage of air voids present in loosely compacted aggregate smaller than 2.36 mm. Higher void contents mean more fractured faces. The test procedure used to measure this property is described in AASHTO T 304 (AASHTO, 2011b). Both the CAA and FAA limits are defined by the location of the layer in the pavement and the traffic intensity to which it will be subjected.

### 12.4.5.5 Grading

Superpave uses the 0.45 power chart to define a permissible grading. Superpave uses a standard set of ASTM sieves. The definitions below are related to the size of the aggregate:

- **maximum size**: one sieve size larger than the nominal maximum size (see Figure 12.17)
- **nominal maximum aggregate size (NMAS)**: one sieve size larger than the first sieve to retain more than 10% (see Table 12.6).

Superpave specifies that two additional features are added to the 0.45 power chart: control points and a restricted zone (Figure 12.17). Control points function as master ranges through which gradings must pass. They are placed on the nominal maximum size, an intermediate size (2.36 mm) and the dust size (0.075 mm).

The restricted zone resides along the maximum density grading between the intermediate size (either 4.75 or 2.36 mm, depending on the maximum size) and the 0.3 mm size. It forms a band through which gradings should not
Gradings that pass through the restricted zone have often been called humped gradings because of the characteristic hump in the grading curve that passes through the restricted zone. In most cases, a humped grading indicates a mixture that possesses too much fine sand in relation to total sand. Such a grading practically always results in tender mixture behaviour, which manifests itself as a mixture that is difficult to compact during construction and offers reduced resistance to deformation during its service life. Gradings that violate the restricted zone may possess weak aggregate skeletons that depend too much on binder stiffness to achieve mixture shear strength. These mixtures are also very sensitive to bitumen content and can easily become plastic. Superpave recommends, but does not require, mixtures to be graded below the restricted zone. However, in 2002 the Transportation Research Board (TRB) issued its Technical Circular E-C043, which recommended the discontinuation of the restricted zone (TRB, 2002).

The term used to describe the cumulative distribution of aggregate particle sizes is the design aggregate structure. A design aggregate structure (Figure 12.18) that lies between the control points and avoids the restricted zone meets the Superpave grading requirements. Superpave defines five mixture types by their nominal maximum aggregate size (Table 12.8).

12.4.5.6 Summary of Superpave mixture design
This section describes, in broad terms, the Superpave mixture design procedure. For a step by step procedure, reference should be made to AASHTO M 323 (AASHTO, 2013b).

The Superpave asphalt mixture design procedure consists of the following four key steps
select the materials
select the design aggregate structure
select the design binder content
evaluate the moisture resistance.

Step 1: select the materials

1. Binder requirements
   a. The binder shall be a performance-graded (PG) binder, meeting the requirements of AASHTO M 320 [AASHTO, 2010].

2. Aggregate requirements
   a. Superpave requires that the combined aggregate shall have an NMAS of 4.75–19.0 mm for asphalt surface courses, and not larger than 37.5 mm for asphalt subsurface courses. The NMAS definitions in section 12.3.2.1 of this chapter shall apply.
   b. The control points for the various grading requirements are defined in Table 3 of AASHTO M 323 [AASHTO, 2013b], reproduced here as Table 12.9.
   c. The aggregates are then checked for compliance with the consensus properties as per Table 12.10. Some consensus properties are defined by the location of the layer within the pavement (e.g. the base).

<table>
<thead>
<tr>
<th>Superpave mixture designation</th>
<th>Nominal maximum size: mm</th>
<th>Maximum size: mm</th>
</tr>
</thead>
<tbody>
<tr>
<td>37.5 mm</td>
<td>37.5</td>
<td>50</td>
</tr>
<tr>
<td>25 mm</td>
<td>25</td>
<td>37.5</td>
</tr>
<tr>
<td>19 mm</td>
<td>19</td>
<td>25</td>
</tr>
<tr>
<td>12.5 mm</td>
<td>12.5</td>
<td>19</td>
</tr>
<tr>
<td>9.5 mm</td>
<td>9.5</td>
<td>12.5</td>
</tr>
</tbody>
</table>
Step 2: select the design aggregate structure

1. The grading is classified (Table 12.11) by using the primary control sieve (PCS) control point as defined in Table 12.9. The combined aggregate grading shall be classified as coarse graded when it passes below the PCS control point, and fine graded for all other combinations.

2. Superpave requires that a minimum of three trial blend gradings are prepared. Each of the blends is plotted on the 0.45 power chart to check whether they conform to the AASHTO M 323 requirements.

3. The compliant trial blends are tested against the requirements for aggregates in section 6 of AASHTO M 323.

4. Replicate asphalt specimens are prepared using the initial trial binder content and the chosen trial aggregate blend grading. The asphalt

Table 12.9 Aggregate grading control points per AASHTO M 323 (© 2013, the American Association of State Highway and Transportation Officials, Washington, DC. Used by permission)

<table>
<thead>
<tr>
<th>Sieve size: mm</th>
<th>Nominal maximum aggregate size: mm</th>
</tr>
</thead>
<tbody>
<tr>
<td>37.5</td>
<td>100</td>
</tr>
<tr>
<td>25.0</td>
<td>90</td>
</tr>
<tr>
<td>19.0</td>
<td>–</td>
</tr>
<tr>
<td>12.5</td>
<td>–</td>
</tr>
<tr>
<td>9.5</td>
<td>–</td>
</tr>
<tr>
<td>4.75</td>
<td>–</td>
</tr>
<tr>
<td>2.36</td>
<td>15</td>
</tr>
<tr>
<td>0.075</td>
<td>0</td>
</tr>
</tbody>
</table>

AASHTO (2013b)

Table 12.10 Superpave aggregate consensus property requirements (From AASHTO M 323. © 2013, the American Association of State Highway and Transportation Officials, Washington, DC. Used by permission)

<table>
<thead>
<tr>
<th>Design traffic: (10^6) ESALs</th>
<th>Coarse aggregate angularity, minimum: % with one fractured face/% with two fractured faces</th>
<th>Uncompacted void content of fine aggregate, minimum: %</th>
<th>Sand equivalent, minimum: %</th>
<th>Flat and elongated, maximum: %</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Thickness (\leq 100) mm</td>
<td>Thickness &gt;100 mm</td>
<td>Thickness (\leq 100) mm</td>
<td>Thickness &gt;100 mm</td>
</tr>
<tr>
<td>&lt;0.3</td>
<td>55/-</td>
<td>–/–</td>
<td>–/–</td>
<td>40/–</td>
</tr>
<tr>
<td>0.3 to &lt;3</td>
<td>75/-</td>
<td>50/-</td>
<td>40/–</td>
<td>40/–</td>
</tr>
<tr>
<td>3 to &lt;10</td>
<td>85/80</td>
<td>60/-</td>
<td>45</td>
<td>40/45</td>
</tr>
<tr>
<td>10 to &lt;30</td>
<td>95/90</td>
<td>80/75</td>
<td>45/40</td>
<td>45/45</td>
</tr>
<tr>
<td>(\geq30)</td>
<td>100/100</td>
<td>100/100</td>
<td>45/45</td>
<td>45/50</td>
</tr>
</tbody>
</table>
specimens are prepared using the SGC following the guidelines per AASHTO T 312 (AASHTO, 2014b). The number of gyrations is given in Table 12.12.

5. The mixture volumetrics are then determined for the compacted specimens.

Step 3: select the design binder content

The replicate specimens are prepared at four binder contents using the number of gyrations from Table 12.12

- the estimated design binder content, \( P_b \) (design)
- 0.5% below \( P_b \) (design)

<table>
<thead>
<tr>
<th>Nominal maximum aggregate size: mm</th>
<th>Primary control sieve: mm</th>
<th>PCS control point: % passing</th>
</tr>
</thead>
<tbody>
<tr>
<td>37.5</td>
<td>9.5</td>
<td>47</td>
</tr>
<tr>
<td>25.0</td>
<td>4.75</td>
<td>40</td>
</tr>
<tr>
<td>19.0</td>
<td>4.75</td>
<td>47</td>
</tr>
<tr>
<td>12.5</td>
<td>2.36</td>
<td>39</td>
</tr>
<tr>
<td>9.5</td>
<td>2.36</td>
<td>47</td>
</tr>
</tbody>
</table>

An aggregate gradation that passes below the PCS control point is classified as 'coarse graded'. A gradation that passes above the PCS control is classified as 'fine graded'.

Table 12.11 Grading classification defining the PCS concept per AASHTO M 323 (© 2013, the American Association of State Highway and Transportation Officials, Washington, DC. Used by permission)

<table>
<thead>
<tr>
<th>Nominal maximum aggregate size: mm</th>
<th>Primary control sieve: mm</th>
<th>PCS control point: % passing</th>
</tr>
</thead>
<tbody>
<tr>
<td>37.5</td>
<td>9.5</td>
<td>47</td>
</tr>
<tr>
<td>25.0</td>
<td>4.75</td>
<td>40</td>
</tr>
<tr>
<td>19.0</td>
<td>4.75</td>
<td>47</td>
</tr>
<tr>
<td>12.5</td>
<td>2.36</td>
<td>39</td>
</tr>
<tr>
<td>9.5</td>
<td>2.36</td>
<td>47</td>
</tr>
</tbody>
</table>

Applications include roadways with very light traffic volumes such as local roads, country roads and city streets where truck traffic is prohibited or at a very minimal level. Traffic on these roadways would be considered local in nature, not regional, intrastate or interstate. Special purpose roadways serving recreational sites or areas may also be applicable to this level.

Applications include many collector roads or access streets. Medium trafficked city streets and the majority of country roadways may be applicable to this level.

Applications include many two-lane, multilane, divided and partially or completely controlled access roadways. Among these are medium to highly trafficked city streets, many state routes, US highways and some rural Interstates.

Applications include the vast majority of the US interstate system, both rural and urban in nature. Special applications such as truck weighing stations or truck climbing lanes on two-lane roadways may also be applicable to this level.

Table 12.12 Number of gyrations for varying levels of traffic (From AASHTO M R35. © 2014, the American Association of State Highway and Transportation Officials, Washington, DC. Used by permission)
The objective is to determine the binder content that produces a target air void content ($V_a$) of 4% at $N_{\text{design}}$. For this, the following steps are followed:

1. The mixture volumetric parameters (i.e. $V_a$, VMA and VFB at $N_{\text{design}}$) are calculated using the standard equations as given in AASHTO R 35 (AASHTO, 2014a).

2. For the duplicate specimens, the average $V_a$, VMA, VFB and relative density at $N_{\text{design}}$ against binder content are plotted (Figure 12.19).

3. The binder content to the nearest 0.1% at which the target $V_a$ is equal to 4.0% is determined by graphical or mathematical interpolation.

4. A check is carried out using interpolation where necessary to ensure that volumetric parameters are satisfied at the binder content determined in the previous step.

5. A densification curve is prepared to check its slope and so on, followed by a determination of the corrected specimen relative densities at $N_{\text{initial}}$ (percentage of $G_{\text{mm initial}}$) to confirm that its value satisfies the design requirements in AASHTO M 323 (AASHTO, 2013b) (Table 12.13).

**Figure 12.19** Results of plotting average $V_a$, VMA, VFB and relative density at $N_{\text{design}}$ against binder content
6. Eight duplicate specimens are prepared having the design aggregate structure at the design binder content, to confirm that the percentage of $G_{mm\, max}$ satisfies the design requirements in AASHTO M 323.

Step 4: evaluate the moisture resistance

The moisture resistance of the asphalt is evaluated in accordance with AASHTO T 283 (AASHTO, 2007b). The mixtures are then adjusted after which a report is issued.

12.4.5.7 Use of Superpave

The Superpave system of mixture design uses gradings that pass below the maximum density grading. Research at the National Center for Asphalt Technology and, especially, results of the WesTrack study have shown that some coarse graded Superpave mixtures can exhibit very poor rut resistance (Brown et al., 1998; Kandhal et al., 1998). At the same time, durability problems have been observed in a significant number of pavements constructed using Superpave surface course mixtures. A study in Florida has documented relatively high permeability of Superpave surface course mixtures (Choubane et al., 1998). NCHRP Project 1-42 has been initiated to evaluate the increasing occurrence of top down cracking in asphalt pavements since the implementation of Superpave.

The TRB published a report in 2005 on the status of implementation of Superpave, entitled Superpave Performance by Design (TRB, 2005). The report notes that the first ever Superpave pavement was constructed on

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**Table 12.13 Superpave mixture design requirements (© 2013, the American Association of State Highway and Transportation Officials, Washington, DC. Used by permission)**

<table>
<thead>
<tr>
<th>Design ESALs $\times 10^6$</th>
<th>Required relative density: % of the theoretical maximum specific gravity</th>
<th>VMA, minimum: %</th>
<th>Nominal maximum aggregate size: mm</th>
<th>VFB range: %</th>
<th>Dust-to-binder ratio range</th>
</tr>
</thead>
<tbody>
<tr>
<td>$N_{initial}$</td>
<td>$N_{design}$</td>
<td>$N_{max}$</td>
<td>37.5</td>
<td>25.0</td>
<td>19.0</td>
</tr>
<tr>
<td>&lt;0.3</td>
<td>$\leq 91.5$</td>
<td>96.0</td>
<td>$\leq 98.0$</td>
<td>11.0</td>
<td>12.0</td>
</tr>
<tr>
<td>0.3 to &lt;3</td>
<td>$\leq 90.5$</td>
<td>96.0</td>
<td>$\leq 98.0$</td>
<td>11.0</td>
<td>12.0</td>
</tr>
<tr>
<td>3 to &lt;10</td>
<td>$\leq 89.0$</td>
<td>96.0</td>
<td>$\leq 98.0$</td>
<td>11.0</td>
<td>12.0</td>
</tr>
<tr>
<td>10 to &lt;30</td>
<td>$\leq 89.0$</td>
<td>96.0</td>
<td>$\leq 98.0$</td>
<td>11.0</td>
<td>12.0</td>
</tr>
<tr>
<td>$\geq 30$</td>
<td>$\leq 89.0$</td>
<td>96.0</td>
<td>$\leq 98.0$</td>
<td>11.0</td>
<td>12.0</td>
</tr>
</tbody>
</table>

$^a$ Design ESALs are the anticipated project traffic level expected on the design lane over a 20-year period. Regardless of the actual design life of the roadway, determine the design ESALs for 20 years.

$^b$ For 37.5 mm nominal maximum size mixtures, the specified lower limit of the VFB range shall be 64% for all design traffic levels.

$^c$ For 4.75 mm nominal maximum size mixtures, the dust-to-binder ratio shall be 0.9–2.0.

$^d$ For 25.0 mm nominal maximum size mixtures, the specified lower limit of the VFB range shall be 67% for design traffic levels.

$^e$ For design traffic levels $\geq 3$ million ESALs, the specified VFB range for 9.5 mm nominal maximum size mixtures shall be 73–76% and for 4.75 mm nominal maximum size mixtures shall be 75–78%.
8 July 1992, when the Mathy Construction Company of Onalaska, Wisconsin, and the Wisconsin Department of Transportation placed the first 150 m of asphalt conforming to the then prototype Superpave bitumen and mixture specifications. The report summarises that, in 2005, the Superpave binder specifications were adopted by 52 US state transportation organisations, with the mixture design method being adopted by 36 state transportation organisations.

12.4.5.8 Asphalt mixture design quotations
The following are quotes from famous asphalt designers on various aspects of asphalt mixture design and performance.

Leahy and McGennis (1999) provide a rare quote of Marshall’s own mixture design philosophy

The ultimate result in the improvement of aggregate gradation is the reduction of the VMA. VMA should be reduced to the lowest practical degree. This reduction results in a superior pavement structure as well as to reduce the quantity of asphalt required in the mixture. No limits can be established for VMA, for universal application, because of the versatile application of bituminous materials to many types and gradations of aggregates.

White (1985) concludes his paper on the history of Marshall mixture design by stating that

the two variables that stand out in the design and performance of pavement mixtures are asphalt content and density. In the field it is the highest satisfactory asphalt content at a density achieved under traffic that is significant. In the laboratory an important feature is selecting a compaction procedure that represents traffic induced density and then selection of responsive properties that can be averaged to give an asphalt content that will produce satisfactory performance even if further densification takes place under traffic … producing a mix density comparable to traffic induced densities appears to be the key to Marshall or any other mix design procedure.

Cominsky et al. (1994) note that

compaction is considered the single most important factor affecting the performance of asphalt pavements.

Hughes (1989) states that

It is important that the density of laboratory-compacted specimens approximate that obtained in the field in terms of (a) the structure of the mix and (b) the quantity, size, and distribution of the air voids.
Monismith et al. (1989) opine that

Proper selection of the mix components and their relative proportions, that is, asphalt or ‘binder’ content, requires a knowledge of the significant properties and performance characteristics of asphalt paving mixtures and how they are influenced by the mix components.

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Road pavements are often placed into three main categories:

- flexible pavements
- rigid pavements
- composite pavements.

Flexible pavements are constructed with several layers of asphalt. The description arises because, when a heavy commercial vehicle traverses the pavement, it flexes vertically by an imperceptible amount. In the UK, the maximum gross vehicle weight of a commercial vehicle is 44 tonnes, with a maximum axle load of approximately 8 tonnes. However, these values vary widely throughout the world. Rigid pavements are constructed entirely with a cementitious material. Composite pavements consist of a combination of asphalt and cementitious layers.

Flexible pavements are, by far, the most common pavement type in the UK. There is a wide range of asphalts available, and appropriate materials can be chosen based on a range of factors, including the nature of the traffic to be carried and the availability of materials locally.

Although this chapter discusses design practices for flexible pavements as adopted in the UK, with a short explanation of how pavement design is undertaken in the USA, the principles have universal application.

### 13.1 Introduction

In the UK, there are three generic types of pavement:

- flexible
- rigid (continuous)
- rigid (jointed).
There are two types within each of these categories, and these are illustrated in Figure 13.1. As can be seen, there are two types of flexible pavement. ‘Flexible with HBM base’ is constructed entirely in asphalt, except for the lower part of the base (called the ‘lower base’), which is a hydraulically bound mixture (HBM). The upper base, the binder course and surface course are all asphalts. A wide range of materials is acceptable as HBMs with many being cementitious mixtures. Notwithstanding, a cross section through a road having the most common type of pavement is that shown in Figure 13.2.

The functions of each of the layers shown in Figure 13.2 are as follows.

The foundation is the platform on which the more expensive layers are placed. On a unitary basis, the cost of each layer is usually higher than its predecessor. The foundation carries the load bearing layers of the pavement.

The base is the main structural layer in the pavement, and its main function is to distribute the stresses generated within it efficiently to the foundation. It must be able to sustain the stresses and strains generated within itself without excessive or rapid deterioration of any kind.

The binder course is necessary to allow asphalt contractors to achieve the very high standards of surface regularity required on modern road pavements. (Surface regularity is a measure of rideability. In other words, an assessment of the smoothness of the journey experienced by persons travelling in a vehicle across a particular pavement. It is a very important feature of a finished pavement.) Asphalts are laid by specialist machines called pavers, and one of the results of each pass of a paver is the reduction of any irregularities in the upper surface of the material being covered. The binder course may well also provide a waterproof layer to protect the base from the ingress of water. Water and pavements simply do not mix. Pavements must be designed such that water is prevented from entering the pavement layers.
or indeed the foundation (unless it is a pavement specifically designed to allow water to pass through without being trapped, such as is the case with a porous asphalt surface course and specialist SuDS (sustainable drainage systems) or reservoir pavements). Many thin surface course systems (TSCS) (defined in Clause 942 of Specification for Highway Works (Highways Agency et al., 2008)), often incorrectly described as ‘stone mastic asphalts’, are permeable, and water simply flows through them. If the water reaches the base, it may strip the bitumen from the aggregate in the base asphalt and lead to premature failure of the weakened asphalt. The current UK pavement design standard HD 26 allows, per Note 8 to Figure 2.1 (Highways Agency et al., 2006a), omission of the binder course if the surface course is a chipped hot rolled asphalt surface course. Many engineers wisely insist on the inclusion of a binder course to give asphalt contractors the opportunity to achieve the very tight surface regularity tolerance. Although binder courses are often more expensive than their base versions because they have some 0.6% more bitumen than the base material (e.g. an HDM50 binder course

Figure 13.2 A road constructed with a ‘flexible with asphalt base’ pavement: SC, surface course; TSCS, thin surface course system; PA, porous asphalt
has around 0.6% more than an HDM50 base), many engineers feel that a binder course is essential in an asphalt pavement. (Indeed, most binder course materials (DBM125, HDM50 and DBM50) have an additional 0.6% or so more bitumen than the base equivalent, the exceptions being EME2 and HRA50, which have the same bitumen content regardless of whether they are used in a base or a binder course.)

The surface course has to provide a smooth skid resistant running plane having good surface regularity and adequate resistance to the onset of defects.

The most common structural defects (a structural defect is one that requires replacement of the surface course, binder course and base to correct the defect) are deformation and cracking. Thicker pavements, according to Transport Research Laboratory (TRL) Report 250, are those in which the depth of asphalt is at least 180 mm – see Figure 13.3, which is based on Figure C2 in the TRL report (Nunn et al., 1997). This is a very important figure. It explains the basis of long-life pavements (i.e. pavements that have an indeterminate life). Examination of Figure 13.3 indicates that if the thickness is above about 180 mm, then the effect of loading in producing deformation is reduced by a factor of about 200, given that the y axis is logarithmic.

How long is indeterminate? Who knows? It is expected to be perhaps 80, 100 or 120 years, maybe longer. It is simply not known. However, what TRL Report 250 established and states, in paragraph 1 of section 13 (Nunn et al., 1997), was that

A well constructed, flexible pavement that is built above a defined threshold strength will have a very long structural service life provided that

![Figure 13.3 Rate of rutting versus thickness](image-url)
distress, in the form of cracks and ruts appearing at the surface, is detected and remedied before it begins to affect the structural integrity of the road.

Note that prompt action to deal with cracks and ruts (the two most common reasons for structural failure) is vital if long life is to be achieved.

From Figure 13.3 it can be seen that the minimum asphalt thickness for long life is 180 mm. Different authorities in the UK, and indeed all over the world, specify different thicknesses for a road to be considered a long-life pavement (‘perpetual pavement’ in the USA).

13.2 Background to the design of pavements

The aim of pavement design is to produce a structure that will distribute traffic loads efficiently while minimising the whole life cost of the pavement. The term ‘whole life’ when applied to a pavement refers to all the costs incurred in connection with a pavement and throughout its useful life (i.e. the period during which it will be ‘in service’). Thus, it includes the works costs (construction, maintenance and residual value) and user costs (traffic delays, accidents at roadworks, skidding accidents, fuel consumption/tyre wear and residual allowance).

Designing a pavement is essentially a structural evaluation process, and is needed to ensure that traffic loads are distributed such that the stresses and strains developed at all levels in the pavement and the subgrade are within the capabilities of the materials at those levels. It involves the selection of materials for the different layers and the calculation of the required thicknesses. The load-carrying capacity of a pavement is a function of both the thickness of the material and its stiffness. Consequently, the mechanical properties of the materials that constitute each of the layers in a pavement are important in designing the structure. As moisture may affect the subgrade and the subbase (and also the base if it is unbound), and temperature affects the bitumen bound layers, it is essential that the design process takes account of the prevailing climatic conditions. As has been suggested in other chapters, pavements and water simply do not mix. The drainage is a vital element of the system, and must be designed to ensure that the pavement and the foundation remain dry.

Figure 13.4 illustrates the two classic modes of failure caused by trafficking of a pavement. It assumes two failure modes

- fatigue cracking at the underside of the base
- cumulative deformation.

It is the modes illustrated in Figure 13.4 that pavement engineers have traditionally sought to hold to acceptable limits within the design life. The concept of a design life was regarded as being particularly important for pavements,
because it is known that they do not fail suddenly but gradually deteriorate over a period of time. However, TRL Report 250 (Nunn et al., 1997) demonstrated that these failure modes did not apply to thicker pavements. It was explained above that thicker pavements are those of at least 180 mm in thickness. TRL Report 250 states that

No evidence of conventional roadbase [now ‘base’] failure or structural deformation was found in well constructed pavements. The observed rutting and cracking were found to originate in the surface layers and, as long as it is treated in a timely manner, it is unlikely to lead to structural deterioration.

The findings of TRL Report 250 were incorporated in the UK’s pavement design standard HD 26/01 (Highways Agency et al., 2001) and its current successor, HD 26/06 (Highways Agency et al., 2006a). The latter states in paragraph 2.15 that

Monitoring the performance of all types of flexible pavements that are heavily trafficked has indicated that deterioration, in the form of cracking or deformation, is far more likely to be found in the surfacing, rather than deeper in the structure. Generally for ‘long life’ it is not necessary to increase the pavement thickness beyond that required for 80 msa (millions of standard axles), provided that surface deterioration is treated before it begins to affect the structural integrity of the road.

In this statement, ‘surfacing’ means the combination of the surface course and the binder course. The number of commercial vehicles carried by a pavement is usually measured in terms of millions of standard axles (msa). In the UK, a standard axle is one imposing, exerting or applying a force of 80 kN
13.3 The importance of stiffness

Stiffness can be defined as a measure of the load spreading ability of a material, and applies whether the material is granular, asphaltic or cementitious. It is a fundamental and important parameter that must be fully appreciated by the pavement designer. It can be readily understood by considering the situations depicted in Figure 13.5.

In Figure 13.5(a), material 1 has a lower stiffness value than material 2. If both are laid at the same thickness, then the pressure resulting from a particular load will be higher at the underside of the layer of material 1 compared with the pressure resulting from the same load at the underside of material 2. This, in turn, subjects the layer underneath to a higher value of stress. If this value exceeds that which can be tolerated by the underlying layer, then it will fail. In order to get the same pressure on the underside, it will be necessary to lay a greater thickness of material 2, which has a lower stiffness. Compare that situation with that shown in Figure 13.5(b), where a thicker layer of the material having a lower stiffness is required to produce an acceptable value of the resultant pressure at the underside.

This explains why different pavement materials require different thicknesses for a particular design. An example of such a design is shown in Figure 13.6.

Figure 13.6 illustrates what happens when contract tenders for pavements are issued. Usually, the client will offer tendering contractors the opportunity to price any of a number of options. There may be four or more options, and not all of them will necessarily be flexible pavements. All the options are deemed to be structurally equivalent. Figure 13.6 shows two options. One is based on the binder course and base being EME2 while the other is based on the binder course and base being HDM50. The materials used in these example pavements are as follows.

- TSCS is a thin surface course system, which is a proprietary surface course conforming to Clause 942 of the UK Specification for Highway Works (Highways Agency et al., 2008).
- EME2 is a relatively high bitumen content (usually about 6% by weight of the total mixture) asphalt based on a French specification. EME is an abbreviation of ‘enrobé à module élevé’, which translates into English as ‘high modulus base’. It uses an unmodified bitumen with a penetration of 15–20.
- HDM50 is ‘heavy duty macadam’ asphalt concrete, and is probably the most common UK binder course and base asphalt. It has a mid-
Figure 13.5 Stiffness of pavement layers: (a) equal thicknesses, different material stiffnesses, different resultant pressures; (b) different thicknesses, different material stiffnesses, equal resultant pressures. (Courtesy of Dr Robert N Hunter)
range bitumen content of typically 4% in HDM50 base and 4.5% in HDM50 binder courses, and is usually made with crushed rock. It uses an unmodified bitumen with a penetration of 40–60, and following modern convention should be called HDM40/60, but tradition remains, and it is still called HDM50.

Type 1 unbound mixture is a graded crushed rock (it can also be crushed slag, crushed concrete, recycled aggregates or well burnt non-plastic shale) previously known as type 1 subbase (subbase was previously called ‘sub-base’) complying with Clause 803 of the Specification for Highway Works (Highways Agency et al., 2008).

The base is 60 mm thicker in the HDM50 variant compared with its value in the EME2 pavement. EME2 has a higher stiffness than HDM50. A pressure is applied over a relatively small area by heavy commercial vehicles running over the surface courses, and Clause 2.3.2 of TRL Report 615 (Nunn, 2004) states that the UK design method uses a multi-layer, linear elastic response model to calculate the critical stresses or strains induced under a single standard wheel load (40 kN) that is represented by a circular patch (0.151 m radius) with a uniform vertical stress.

The additional thickness in the HDM50 pavement is necessary because this pressure reduces as it travels down through the pavement. Because EME2 has a higher stiffness than HDM50, the pressure reduces more quickly as it passes through the EME2 pavement. Accordingly, a thinner pavement is

![Figure 13.6 Comparison of two fully flexible pavements, one constructed using EME2 and the other constructed with DBM50](image-url)

Design of flexible pavements
needed with an asphalt having a higher stiffness to give structural equivalence.

It will be recalled from other chapters that the value of stiffness is affected by temperature and the period of loading. UK design methods assume a standard frequency in service (i.e. a standard vehicle speed). Thus, UK roads carrying slow-moving traffic will be under-designed because the stiffness decreases as the speed decreases. The publication *The London Bus Lane and Bus Stop Construction Guidance* states in section 2.2 that ‘where traffic is slow moving and/or channelised this traffic load has to be increased by a factor of 3’ (London Technical Advisors Group, 2013). Australian specifications take account of vehicle speed, and Table 13.1 is an extract from Table Q6.5 of *Pavement Design Supplement* issued by the State of Queensland (2013).

### 13.4 The elements of a flexible pavement

As was shown in Figure 13.2, a road consists of three elements

- the subgrade
- the foundation
- the pavement.

#### 13.4.1 The subgrade

The subgrade is the material below the foundation, and will either be existing soil or imported fill. In areas of cut, it will be the existing soil unless the existing soil is very weak and has to be improved by some means: see Clauses 5.16–5.21 in *Interim Advice Note (IAN) 73* (Highways Agency et al., 2009) for advice on the options available to do this. In areas of fill, the subgrade will be imported fill made of material that has characteristics suitable for its intended role, laid and compacted in layers. It may be very tempting to think that the characteristics of the subgrade will have little effect on the useful lifespan of a pavement. However, nothing could be further from

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**Table 13.1** Presumptive values for elastic characterisation of asphalt mixtures

<table>
<thead>
<tr>
<th>Asphalt mixture type</th>
<th>Binder type</th>
<th>Volume of binder: %</th>
<th>Asphalt modulus at heavy vehicle operating speed: MPa</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>10 km/h</td>
</tr>
<tr>
<td>OG10</td>
<td>A5S</td>
<td>N/A</td>
<td>800</td>
</tr>
<tr>
<td>OG14</td>
<td>A5S</td>
<td>N/A</td>
<td>800</td>
</tr>
<tr>
<td>SM14</td>
<td>A5S</td>
<td>13.0</td>
<td>1000&lt;sup&gt;a&lt;/sup&gt; (600)</td>
</tr>
<tr>
<td>DG10</td>
<td>C320</td>
<td>11.5</td>
<td>1000&lt;sup&gt;a&lt;/sup&gt; (900)</td>
</tr>
<tr>
<td>DG10</td>
<td>A5S</td>
<td>11.5</td>
<td>1000&lt;sup&gt;a&lt;/sup&gt; (600)</td>
</tr>
</tbody>
</table>

From State of Queensland (2013), Table Q6.5

<sup>a</sup> Indicated values have been limited to a value of 1000 MPa
the truth, as is demonstrated by Table 13.2. The key role of the subgrade material can be demonstrated by considering Table 13.2, which has been taken from Table A1 in TRL Report 250 (Nunn et al., 1997).

It has long been the case that maintenance engineers will consider that a carriageway begins to warrant consideration for replacement or renewal when deformation (‘deformation’ and ‘rutting’ are interchangeable terms) reaches 10 mm (what was sometimes described as a ‘critical condition’). This was established following a study published in 1972 (see Table 1 in TRL Report LR 375) (Lister, 1972).

Table 13.2 relates to pavements where the base is either hot rolled asphalt or asphalt concrete (DBM; dense bitumen macadam), which will have been the vast majority of pavements at the time when the investigation associated with TRL Report 250 was undertaken. What these data suggest is that if the California bearing ratio (CBR; a measure of the strength of a soil, a test devised by US engineers and used all over the world for roads and airfield construction) is less than 5%, then the pavement will carry some 17 msa before a 10 mm depth of rut is formed (BSI, 1990a, 1990b). However, if the CBR exceeds 5%, then the pavement will carry some 28 msa before a 10 mm depth of rut is formed. Thus, the increase in the CBR results in the pavement carrying an additional 65% or so of traffic before a 10 mm rut is generated. This is stark proof that the subgrade will have a significant effect on the service life of the pavement.

This book does not address subgrades to any extent. Suffice to say that great care needs to be taken in constructing the subgrade and/or preparing it to receive the foundation.

### 13.4.2 The foundation

The foundation will also play a very important role in determining the performance of the pavement, including the length of its service life.

The foundation has to perform in two different sets of circumstances.

- During construction it has to protect the subgrade and withstand the relatively high stresses generated within it by construction traffic (i.e. excavators, lorries, pavers, etc.). Although the number of stress repetitions will be lower than when it is in service and the line travelled...
by plant is not as channelised as that taken by traffic when the pavement is in service, the intensity of individual loads will be much higher because site traffic travels directly on the foundation. In addition, it may well be exposed to adverse weather conditions without the protection of the overlying pavement layers.

In service, traffic, in terms of the line travelled, is much more disciplined. The load intensity will be reduced because of the effect of the intervening pavement. However, the number of stress repetitions will be very much higher than was the case during construction.

Like the subgrade, constructing the foundation requires great care. The material chosen must be appropriate for the role it has to fulfil. In addition, it has to be placed in strict conformance with the specification if it is to perform adequately (layer thickness, compaction passes, etc.).

As can be seen from Figure 13.2, the foundation can now be formed of a subbase on capping, entirely capping or entirely subbase.

Materials used in UK foundations range from low grade fill to a number of hydraulically bound mixtures that may contain cement. However, the material most commonly used in the UK is type 1 unbound mixture (formerly type 1 subbase), a graded crushed rock.

This book does not consider foundations in any detail, but their importance cannot be overstated. Advice on foundations can be found in IAN 73 (Highways Agency et al., 2009) and Chapter 32 of the ICE Manual of Highway Design and Management (Payne, 2011).

13.4.3 The pavement

Fully flexible pavements invariably consist of the base, on top of which is the binder course, and above which is the surface course. Each of these is discussed below.

13.4.3.1 Bases

The base is the main structural element in the pavement, although the binder course and surface course contribute to the strength of the pavement. The function of the base is to distribute the imposed loading so that the underlying materials are not overstressed. It must resist deformation and cracking caused by fatigue through repeated loading. It must also be capable of withstanding stresses induced by temperature gradients through the structure. The base in a flexible pavement is usually a dense asphalt.

In the UK, the options for base asphalts are

- EME2
- DBM50 or HDM50
In pavements carrying heavy traffic, the base thickness will typically be 200 mm or more. Layers of this thickness would normally be laid in two layers, with the first layer being described as the ‘lower base’ and the second layer being described as the ‘upper base’.

13.4.3.2 Binder courses
The combination of the binder course and the surface course is described as the surfacing (see Figure 2.1 in HD 23 (Highways Agency et al., 1999)). The purpose of the binder course is to allow surfacing contractors to achieve the very high standards of level control (often described in contracts as the ‘surface regularity’) required by current highway maintenance and construction contracts. It may also act as an impermeable layer protecting the base from the ingress of water.

Pavement layers are usually placed by a specialised piece of equipment called a ‘paver’. The nature of these machines is such that, as each layer is placed, they tend to reduce any irregularities on the top of the layer being laid. A major factor in determining the quality of the ride experienced by persons travelling across a pavement is its ‘surface regularity’. The smoother the ride, the better the surface regularity. It is for this reason that surface regularity requirements are often very tight: for example, in the UK the surface course more often than not must be within $\pm 6\text{ mm}$ of the designed finished level at the top of the pavement (Highways Agency et al., 2009, Table 7.1). Indeed, some contracts require compliance to $+6\backslash -0\text{ mm}$.

The thickness of the binder course is often around 60 mm. The thickness adopted should be the minimum because, unless it is EME2 or HRA50, the binder course variant will have around 0.6% more bitumen than the equivalent base material.

In the UK, the options for binder course asphalts are

- EME2
- DBM50 or HDM50
- HRA50
- DBM125
- SMA.

The difference between the options available for bases and binder courses is that SMA (stone mastic asphalt – a material originally developed in Germany) can be used. The reason for this is that SMA can be laid as thin as 30 mm, while the other options cannot be laid much thinner than 50 mm. (The minimum thickness of an asphalt layer is governed by a rule of thumb that it should be at least 2.5–3 times the nominal stone size, with the nominal...
13.4.3.3 Surface courses

The layer on which traffic runs is termed the surface course. It is this layer, and this layer only, that is visible to the road user. The surface course has to meet a formidable list of requirements. It must

- resist deformation by traffic
- be durable, resisting the effects of weather, abrasion by traffic and fatigue
- provide an acceptable level of skid resistance
- not result in unacceptable noise levels being generated
- provide a surface of acceptable riding quality
- except in the case of porous asphalt, preferably be impermeable to the passage of water through the layer, thus protecting the lower layers of the pavement
- contribute to the strength of the pavement structure
- not cause unacceptable levels of spray to be produced.

Surface courses on heavily trafficked UK roads (i.e. those carrying 80 msa or more in their design lives) are usually either a TSCS, as described above, or a chipped hot rolled asphalt surface course (complying with Clause 911 or 943 of the UK’s Specification for Highway Works (Highways Agency et al., 2008). (TSCSs should be described as such. In the UK, they are often described as SMAs, but that is incorrect because they are not stone mastic asphalts.)

The pavement design standard mostly commonly used in the UK (HD 26) makes a binder course optional where the surface course is a chipped hot rolled asphalt surface course. However, this standard states that a binder course is always required when the surface course is a TSCS, many of which are permeable. The reason for this approach is to ensure that pavement bases are protected from the ingress of water through the upper layers of the pavement. Notwithstanding, in order to give surfacing contractors the best chance of achieving surface regularity compliance, many engineers advocate the use of a binder course in all asphalt pavements. This is a wise approach. However, it is a more expensive option, and the client’s approval for their inclusion should be canvassed when necessary.

13.5 Stages involved in pavement design

The stages involved in designing a UK pavement are shown in Figure 13.7, and are based on the use of HD 24 (Highways Agency et al., 2006b),
IAN 73 (Highways Agency et al., 2009), HD 26 (Highways Agency et al., 2006a) and HD 36 (Highways Agency et al., 2006c), which are all standards published by the UK government. However, the approach depicted in Figure 13.7 could well be applied in any country by application of the appropriate applicable standard. The steps in carrying out this process are discussed below.

### 13.5.1 Design traffic

The first stage in most structural designs is to estimate the loading that the structure is expected to carry, and that approach is no different in the case of the design of pavements. The load carried by a UK pavement is described as the ‘design traffic’.

![Figure 13.7 Pavement design procedure. (Courtesy of Dr Robert N Hunter)](image-url)
The design traffic is the total loading that the structure (i.e. the pavement) is designed to carry throughout its service life. In the UK, calculation of the design traffic is usually undertaken on the basis of the method outlined in HD 24 (Highways Agency et al., 2006b). The service life of the pavement is 40 years on the basis that this period has generally proved to be the most economic, particularly where traffic flow is high, all as per paragraph 2.14 of HD 24.

Loads are applied to the pavement through contact between vehicle tyres and the surface course. Obviously, the degree and nature of the traffic loading is one of the major factors affecting the design and performance of pavements. Both the magnitude and the number of loadings contribute to the overall damage to the pavement. The tyre pressure primarily affects the stresses (and strains) developed at the surface and within the upper layers of the pavement. (In road pavement design, a typical commercial vehicle tyre pressure would be between 5 and 9 bar (0.5 and 0.9 MPa), whereas aircraft tyres inflict pressures up to 30 bar (3.0 MPa). These values have a significant influence on the design of the materials used in road and airfield pavements.)

Although a great deal of publicity has been given to the physical size and gross weight of commercial vehicles, it is the axle load that is critical in pavement design and performance. Heavier vehicles are usually carried on a larger number of axles, thereby maintaining or reducing the axle load.

Most countries limit the axle loadings that can be carried on their carriageways. However, these limits are difficult to enforce, and there is the possibility that some lorries carry a higher payload than their design or the applicable regulatory framework permits.

Some countries base their pavement design on the total number of vehicles of any size carried by the pavement. However, that approach is incorrect. The loading exerted by cars and lighter goods vehicles does not contribute to any structural deterioration in a pavement. Only commercial vehicles, which in the UK for pavement design purposes are defined as vehicles having a gross vehicle weight in excess of 3.5 tonnes, cause any structural deterioration.

Normal traffic on conventional roads is mixed in composition and the magnitudes of axle loads, and it is therefore necessary for design purposes to simplify the real situation by converting actual axle loads to an ‘equivalent’ loading system. This idea originated with the American Association of State Highway Officials (AASHO) road test in the USA, which was undertaken in 1958 (Highway Research Board, 1962; Liddle, 1962). The mixed axle load spectrum is converted to an equivalent number of ‘standard axles’, an example being the value of 80 kN adopted in the UK (Road Research
Laboratory, 1970). This is based on a concept of equivalent damage to the pavement. Thus, on an axle of a heavy lorry having a single tyre at each end, the very common super single tyre, the load on each tyre will be 40 kN (i.e. approximately 4 tonnes). The loading exerted (and thus the damage) for each vehicle can then be computed in terms of a number of equivalent standard axles by using the ‘fourth power law’ given below

\[
\text{wear/axle} \propto \text{axle load}^4
\]

It has been shown (Curren and Connor, 1979) that on weak pavements the exponential power can be as high as 6 if trafficked by heavy axle loads at or above the current legal limit. However, if the UK is used as an example again, the value of 4 is accepted in the national traffic assessment standard HD 24 (Highways Agency et al., 2006b), and has been so for many years. This approach has proved to lead to the construction of highways that have successfully withstood the enormous growth in traffic in the UK.

Thus, using the above relationship, under normal conditions it takes the application of 16 40 kN axle loads to cause the same damage as a single application of an 80 kN axle load. Furthermore, one application of a 160 kN axle load induces 16 times the damage of a standard 80 kN axle load.

13.5.2 Foundation
This is an extremely important element of pavement design, and must be undertaken by experienced, qualified engineers. However, it is not discussed in any detail in this book (see section 13.4.2).

13.5.3 Pavement design
Once the design traffic has been ascertained, the pavement thickness can be designed. In the UK this is done by reference to Figure 2.1 of HD 26 (Highways Agency et al., 2006a). Flexible pavement design in HD 26 is based on TRL Report 615 (Nunn, 2004).

Figure 13.8 is an excerpt from Figure 2.1 of HD 26. The procedure for designing the pavement is as follows:

1. on the upper x axis, select the calculated design traffic
2. drop vertically to the point that cuts the applicable Foundation Class
3. turn right until the EME2 line is intercepted
4. drop vertically down to the lower x axis
5. read off the applicable ‘total asphalt thickness’ for EME2
6. repeat step 3 above, but turn right until the DBM50/HDM50 line is intercepted
7. drop vertically down to the lower x axis
8. read off the applicable ‘total asphalt thickness’ for DBM50/HDM50
Figure 13.8 Design thickness for flexible pavements
9 repeat step 6 above, but turn right until the HRA50 line is intercepted
10 drop vertically down to the lower x axis
11 read off the applicable ‘total asphalt thickness’ for HRA50
12 repeat step 9 above, but turn right until the DBM125 line is intercepted
13 drop vertically down to the lower x axis
14 read off the applicable ‘total asphalt thickness’ for DBM125
15 repeat for any other applicable foundation classes.

An example is a road with a design traffic of 40 msa on a class 3 foundation

1 go to the upper x axis at the point where the ‘traffic’ (it should actually be labelled ‘design traffic’) is 40 msa
2 drop down to where the Foundation Class 2 line is encountered
3 turn right to the point where the EME2 line is intercepted
4 drop down to the lower x axis, and read off the ‘total asphalt thickness’ which is around 240 mm
5 return to the point where the EME2 line was crossed, and move horizontally right to the point where the DBM50/HDM50 line is intercepted
6 drop down to the lower x axis, and read off the ‘total asphalt thickness’, which is just below 300 mm and would be rounded up to 300 mm
7 repeat the process for HRA50 and DBM125.

The reading for EME2 is 240 mm which is the ‘total asphalt thickness’. The EME2 pavement option will then be 40 mm of surface course, 70 mm of AC 14 EME2 binder course (this minimum thickness value of EME2 is stated in Table 1A of BS 594987 (BSI, 2010)) and 130 mm of AC 20 EME2 base, making a ‘total asphalt thickness’ of 240 mm. Table 13.3 shows the full set of results for the entire range of asphalt options.

Figure 2.1 in HD 26 (Highways Agency et al., 2006a) has, attached to it, a series of notes relating to the materials featuring in Figure 2.1. These notes must always be considered as part of the design process.

Inclusion of the above thicknesses in a tender allows the pavement alternatives to be put out to tender as part of a submission for highway maintenance.

<table>
<thead>
<tr>
<th>Table 13.3 Example of pavement design results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Asphalt</td>
</tr>
<tr>
<td>Total asphalt thickness: mm</td>
</tr>
<tr>
<td>Surface course thickness: mm</td>
</tr>
<tr>
<td>Binder course thickness: mm</td>
</tr>
<tr>
<td>Base thickness: mm</td>
</tr>
</tbody>
</table>
or highway construction or as an element of contract for other work types.

It is worth considering some aspects of Figure 2.1 and the notes attached thereto:

- the thickness is to be rounded up to the next 10 mm
- where the asphalt design thickness is 300 mm or less, the material is to be laid with no negative tolerance
- the figure assumes that the binder course and base will be constructed of the same asphalt (e.g. HDM50)
- once the design traffic reaches a value of 80 msa, then the same design applies up to a design traffic of 400 msa
- for low traffic values up to 5 msa, the thickness for different asphalts is the same
- the minimum design traffic is 1 msa.

Finally, it is worth noting the following about the asphalt options available.

- Although permitted, it may not be wise to use a DBM125 asphalt, as the penetration, being somewhere between 100 and 150, is significantly softer than the 40/60 bitumen used in DBM50/HDM50 and HRA50 and the 15/20 penetration bitumen in EME2.
- In roads carrying heavy traffic, it may be wise to be very wary of specifying HRA50 as a binder course because it is prone to deforming far more rapidly than the other options.

In summary, the UK pavement design method is a mechanistic–empirical method, and, therefore, that fact alone means that the method can only ever be an approximation to the thicknesses necessary to carry the applied loading. This notwithstanding, it has served the UK well, having provided roads that have carried very high traffic levels for many years.

13.6 Pavement designs in the USA

The most widely used empirical pavement design methods for roadway pavements were developed from data collected in the AASHO road test, which took place over a 25 month period starting in October 1958 near Ottawa, Illinois. Among the objectives of the road test were the determination of relationships between axle load applications and the performance of different types and thicknesses of pavement layer materials; and the organisation and storage of data that could be further analysed were essential for the development of the AASHTO Interim Guide for Design of Pavement Structures (AASHTO, 1972) and the revisions and versions that followed. The most recent version was published in 1993, and AASHTO has developed software for using the design procedure (AASHTO, 1993).
The basic pavement performance models developed from road test data are used in the 1993 guide. To design a flexible pavement structure using the AASHTO method, the designer must characterise:

- the pavement performance (serviceability change over time)
- the traffic (characterised as equivalent single axle loads)
- the roadbed soil (subgrade) resilient modulus, $M_R$
- the pavement layer materials (layer coefficients, $a_i$)
- the drainage
- reliability.

The AASHTO flexible pavement design procedure ultimately results in a structural number for the pavement that provides a pavement design adequate for the assumed conditions. The structural number for a specific pavement cross-section is determined using the following equation:

$$SN = a_1D_1 + a_2D_2m_2 + a_3D_3m_3$$

where $a_i$ is the layer coefficient for layer 1, 2 or 3 (dimensionless); $D_i$ is the thickness for layer 1, 2 or 3 (inches); and $m_2$ and $m_3$ are the drainage coefficients for the base and subbase layers, respectively (dimensionless).

Commonly used values for layer coefficients are shown in Table 13.4.

Many combinations of different pavement layer materials and thicknesses can result in the same structural number for a pavement section that would probably perform very differently. Consequently, pavement designers using the 1993 AASHTO guide must apply engineering judgement to develop appropriate pavement cross sections to consider for a particular project. One way to evaluate and compare different pavement sections is to use an analytical or mechanistic–empirical procedure.

The AASHTO Mechanistic–Empirical Pavement Design Guide (MEPDG) was developed to overcome many of the shortcomings of the empirical method (TRB, 2014). This procedure requires the use of the computer program AASHTOWare Pavement ME to analyse a pavement structure and predict its performance from the input variables.

<table>
<thead>
<tr>
<th>Layer material</th>
<th>Layer coefficient range</th>
<th>Typical value assumed</th>
</tr>
</thead>
<tbody>
<tr>
<td>Asphalt surface courses, less than 4 in. (100 mm) thick</td>
<td>0.40–0.55</td>
<td>0.44</td>
</tr>
<tr>
<td>Asphalt base</td>
<td>0.20–0.34</td>
<td>0.30</td>
</tr>
<tr>
<td>Unbound aggregate base</td>
<td>0.12–0.18</td>
<td>0.14</td>
</tr>
<tr>
<td>Unbound aggregate subbase</td>
<td>0.08–0.12</td>
<td>0.11</td>
</tr>
<tr>
<td>Portland cement treated base</td>
<td>0.10–0.28</td>
<td>0.20</td>
</tr>
</tbody>
</table>
Inputs to this design method include the pavement layer thickness, the layer material properties (e.g. stiffness, strength), project climate/environmental data and traffic axle load spectra. Accumulated damage is predicted, with the output from the analysis including predicted levels of roughness (as the international roughness index, IRI), rutting, load-associated cracking and low temperature cracking.

The MEPDG has been implemented for use by several state departments of transportation, and is currently being evaluated for implementation by most highway agencies in the USA and Canada. The evaluation requires calibration of the various performance models for conditions specific to the state or province.

There are other pavement design methods used in the USA, such as the MnPave (Mechanistic–Empirical Flexible Pavement Design) program or the California Department of Transportation empirical method among others.

13.7 Empirical and analytical pavement design

There are two approaches to pavement design: empirical and analytical. Most design methods in current practice around the world are empirical, being based on experience accumulated in practice and from specially constructed test sections. There is an assumption within this approach that the performance of the pavement will be the same irrespective of the properties of the pavement or the constituent materials. This approach to pavement design is somewhat removed from engineering principles, and cannot cope with circumstances beyond those included in the trial section on which the method is based. In contrast, the analytical approach uses theoretical analysis of the mechanical properties of materials, and is capable, in principle, of dealing with any design situation. Efficient pavement design can only be achieved through the use of accurate analytical methods. Hopefully, in the foreseeable future, pavements will be designed analytically based on an accurate assessment of their actual physical properties.

13.8 Analytical pavement design

The philosophy of an analytical approach to pavement design is that the structure should be treated in the same way as other civil engineering structures. The basic procedure is as follows

1. assume a form for the structure, usually involving rational simplification of the actual structure to facilitate analysis
2. specify the loading
3. estimate the size of the components
4. carry out a structural analysis to determine the stresses, strains and deflections at critical points in the structure
5 compare these values with the maximum allowable values to assess whether the design is satisfactory
6 adjust the materials or geometry repeating steps 3, 4 and 5 until a satisfactory design is achieved
7 consider the economic feasibility of the result.

13.9 The contribution of Shell Bitumen to the development of pavement design

Shell Bitumen has a long association with the subject of pavement design that stretches back many decades.

The nomographs for asphalt characteristics (stiffness modulus and fatigue properties) are based on those originally developed by Shell in the 1950s. The bitumen stiffness nomograph was developed by Van der Poel in 1954 (Van der Poel, 1954), and is shown in Figure 13.9. This has proved to be an excellent tool for predicting bitumen stiffness over a wide range of loading conditions. The stiffness of the bitumen is calculated at a specific temperature and loading time (or frequency of loading) from the penetration and softening point of the bitumen. It has been found that there is good agreement between the measured and predicted stiffnesses of asphalts containing conventional penetration grade bitumens. Accordingly, the nomograph is still considered accurate enough for engineering purposes.

The Van der Poel nomograph was later superseded by PONOS, a computer program that was able to calculate the bitumen properties based on the nomographs, which enabled a wider range of bitumen characterisation to be used as input. PONOS has been adapted for use in the BANDS (bitumen and asphalt nomographs developed by Shell) software package.

BANDS was devised to predict the bitumen properties as described above, using the Van der Poel nomograph and PONOS. However, the improvement with BANDS is that it has been developed to be used on modern computer systems such as Windows. Thus, BANDS is far easier to use than its predecessor.

Over time, there have been ongoing efforts to develop methods for the prediction of the mechanical properties of asphalts from the properties and the relative volumes of the constituent materials, hence the interest in the Van der Poel nomograph.

In the late 1950s, there was a great deal of interest in analytically-based pavement design procedures. This interest grew, and, in 1963, Shell published a set of design charts that allowed the calculation of stress and strain in a structure based on a multi-layer linear elastic analysis. (A multi-layered elastic analysis calculates stresses, strains and deflections at any point in a pavement structure resulting from the application of a surface load.)
To determine the stiffness modulus of a 50 pen bitumen at the test conditions, connect 0.02 s on the loading time-scale with (53.5 + 1.5 – 5 = 50ºC) on the temperature scale. Stiffness modulus is 1.5 × 10^8 Pa at a PI of 0.

**Figure 13.9** Van der Poel nomograph for determining the stiffness modulus of bitumens
Layered elastic models assume that each pavement structural layer is homogeneous and linearly elastic throughout the layer. A multi-layered elastic analysis examines a structure made up of a number of structural layers. In 1978, this system was extended to incorporate all relevant major design parameters that were derived from empirical design methods and the AASHO road test and laboratory data, and published as the Shell Pavement Design Manual (SPDM) (Shell International Petroleum Company, 1978, 1985). A photograph of the first delivery of these design charts is shown in Figure 13.10. SPDM allowed for the effects on the pavement of temperature, traffic density, and the physical properties of the bitumen and aggregates while standardising the asphalts with respect to stiffness and fatigue properties. SPDM was presented in the form of graphs, charts and tables, and, to keep the number of graphs, charts and tables manageable, the number of variable parameters used was limited, with the pavement engineer being expected to interpolate using these data.

In the early 1970s, Shell developed BISAR (bitumen stress analysis in roads) mainframe computer program. This program was designed to calculate and analyse the stresses and strains within the layers of a pavement. BISAR, whose forerunner was the 1963 design charts, was used in drawing up the SPDM design charts that were issued in 1978.

![Figure 13.10 First delivery of Shell’s pavement design charts in 1978](image-url)
In 1992, the first release of SPDM-PC (Shell pavement design method on a
cersonal computer) was issued. This software followed the same design
method as the 1978 manual. However, the computer program allowed
the use of a wide variety of data without the need for cumbersome
interpolations.

Since then, with the huge increase in computer power available to virtually
every individual within an organisation and at home, the limitations of the
original Shell analytical software and the manual procedures introduced
by Shell over the last 40 years have now been removed and replaced with
user-friendly computer packages that can be integrated with other computer-
based applications.

As already mentioned, one of the benefits of the analytical approach is the
ability to design for individual sets of conditions. Therefore, it is very impor-
tant that the characteristics and properties of the materials that are used in the
design are appropriate for the loading and climatic conditions.

13.10 Example pavement designs using SPDM-PC

Figure 13.11 shows a pavement designed for 20 msa (Strickland, 2000)
using three different methods.

- The first pavement was designed in accordance with the current UK
design method (DMRB) using this assumed value of design traffic

<table>
<thead>
<tr>
<th>DMRB, Volume 7 Design</th>
<th>Shell pavement design method - 'conventional design'</th>
<th>Shell pavement design method - 'alternative design'</th>
</tr>
</thead>
<tbody>
<tr>
<td>50 mm 35/14 HRA with 50 pen</td>
<td>50 mm HRA-40/60 pen 7.7% binder content 4% air voids stiffness modulus 382 MPa</td>
<td>30 mm SMA with Carphalte TS 5.3% binder content 4% air voids stiffness modulus 898 MPa</td>
</tr>
<tr>
<td>270 mm DBM with 100/150 pen</td>
<td>270 mm DBM with 100/150 pen 4.8% binder content, 6% air voids stiffness modulus 296 MPa</td>
<td>185 mm 28 mm HDM with 30/45 pen 3.7% binder content, 5% air voids stiffness modulus 10 100 MPa</td>
</tr>
<tr>
<td>225 mm subbase</td>
<td>225 mm subbase stiffness modulus 56 MPa</td>
<td>235 mm subbase stiffness modulus 57 MPa</td>
</tr>
</tbody>
</table>

Figure 13.11 Example pavement designs
followed by the procedures set out in IAN 73 (Highways Agency et al., 2009) and HD 26 (Highways Agency et al., 2006a).

- The second pavement was designed with the SPDM-PC computer software with ‘conventional materials’.
- The third pavement was designed with SPDM-PC using ‘alternative materials’. SPDM-PC requires criteria for subgrade strain and fatigue failure to be input as part of the design process, and the values of these parameters were taken from RRL Report 1132 (usually described as ‘LR 1132’) (Powell et al., 1984). The subgrade strain and fatigue failure criteria given in LR 1132 provide details for the conditions that prevail in the UK. However, this can easily be changed within SPDM-PC for the criteria that apply in different countries. Thus, by adopting the applicable criteria, SPDM-PC can be used in any country.

The subgrade is taken to have a CBR of 5%, a value that is typical for a silty clay. The CBR has been converted to an equivalent stiffness modulus, \( E \), using the relationship specified in paragraph 5.11 of IAN 73 (Highways Agency et al., 2009). The relationship is

\[
E = 17.6(CBR)^{0.64} \text{ MPa}
\]

This relationship was used for all three designs, which calculated the stiffness modulus of the subgrade as 49 MPa.

The designs of foundations for the three pavements are shown in Table 13.5.

For the three pavement constructions, the designed asphalt thicknesses are shown in Table 13.6.

The above example demonstrates that pavements designed in accordance with the current UK pavement design method (DMRB) and pavements designed with SPDM-PC are very similar. However, as SPDM-PC is an

<table>
<thead>
<tr>
<th>Table 13.5 Foundation design</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Current UK pavement design (DMRB)</strong></td>
</tr>
<tr>
<td>By following IAN 73, two subbase designs were possible:</td>
</tr>
<tr>
<td>150 mm subbase on 250 mm capping</td>
</tr>
<tr>
<td>225 mm subbase, no capping</td>
</tr>
</tbody>
</table>

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analytical procedure, it has the advantage of being able to take into account variations in material properties. The example also shows that, with the SPDM-PC computer package, thinner pavements constructed using alternative materials can be designed to carry the same number of standard axles (i.e. the same design traffic). This can significantly reduce the cost of the pavement or, alternatively, provide a pavement that, having the same thickness, will have a longer service life.

13.11 A glimpse of the future for pavement design?

It can be seen from the section above that true analytical design offers a much more accurate method of designing pavements. This fact coupled with the flexibility that such a method offers means it has many advantages over the current empirical methods in use around the world. Figure 13.12 shows a section through what is probably the most common pavement construction in the UK.

Figure 13.12 is based on a flexible pavement (i.e. the surface course, binder course and base are all asphalts). The load is provided by a heavy goods vehicle equipped with super single tyres pressurised to 8.5 bar. The values of pressure down through the pavement and foundation layers were calculated by BISAR. Figure 13.12 shows how the pressure is reduced as it passes through each of the layers. As can be seen, 2.5% of the applied stress is lost as the pressure passes through the surface course, and 17.4% as it passes through the binder course. As would be expected, the majority of the pressure is lost as the loading passes through the base. Finally, 2.8%,
1.9% and 1.1% are lost through the subbase, capping and subgrade respectively. Note that 94.1% of the pressure is lost by the time the pressure reaches the foundation.

Thus, this powerful program enables engineers to begin to design pavements in the way they should be designed as set out in section 13.10 above.

Other analytical pavement design software packages are available, including Amadeus, CIRCLY, ALYSEE and CAPA 3D.

**References**


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<table>
<thead>
<tr>
<th>Material in layer</th>
<th>Elastic modulus: MPa</th>
<th>Poisson’s ratio</th>
<th>Thickness: mm</th>
<th>Pressure drop in layer</th>
<th>% Drop in layer</th>
</tr>
</thead>
<tbody>
<tr>
<td>HRA SC</td>
<td>3100</td>
<td>0.35</td>
<td>40</td>
<td>0.215 bar</td>
<td>2.5%</td>
</tr>
<tr>
<td>HDM50</td>
<td>4700</td>
<td>0.35</td>
<td>60</td>
<td>1.479 bar</td>
<td>17.4%</td>
</tr>
<tr>
<td>HDM50</td>
<td>4700</td>
<td>0.35</td>
<td>200</td>
<td>6.309 bar</td>
<td>74.2%</td>
</tr>
<tr>
<td>Type 1 unbound mixture</td>
<td>150</td>
<td>0.35</td>
<td>200</td>
<td>0.239 bar</td>
<td>2.8%</td>
</tr>
<tr>
<td>Capping</td>
<td>75</td>
<td>0.35</td>
<td>600</td>
<td>0.162 bar</td>
<td>1.9%</td>
</tr>
<tr>
<td>Soil</td>
<td>40</td>
<td>0.35</td>
<td>N/A</td>
<td>N/A</td>
<td>1.1%</td>
</tr>
</tbody>
</table>

**Figure 13.12** Pressure–stress dissipation down through a typical UK flexible pavement. (Courtesy of Dr Robert N Hunter)


Chapter 14

Asphalt production plants

John Moore
Consultant (UK)

14.1 Introduction

The principles of making asphalt have not changed since the middle of the nineteenth century (Jones, 1986). Different size aggregates are dried, heated and then mixed with bitumen to make a saleable product. The aggregates require drying and heating to enable the bitumen to adhere to the surface of the aggregate.

However, although the principle has not changed, the methods of manufacture, and therefore the production plants, have changed significantly in their design. Capacities have increased from just a few tonnes per hour up to 800 tonne/h of asphalt. Worldwide environmental regulations have resulted in particulate emissions from current asphalt production plants being undetectable with the naked eye, whereas early asphalt plants had a bad reputation due to clouds of dust surrounding the works. The main recent developments have been in the controls: early plants were controlled manually but modern asphalt plants are completely computer controlled. The quality of the asphalt produced has also improved beyond recognition from the early days, when ‘hot, black and sticky’ were the criteria. Now, the quality of a mixed asphalt can be guaranteed, with an accurate bitumen content and a consistent asphalt product.

In the first production of asphalt, coal was used as the fuel to dry and heat the aggregates and the bitumen. Later, oil was used, and now there are numerous fuels available as alternatives to oil, including natural gas, liquid propane gas, butane and coal. The choice of fuel is influenced by the fluctuating cost of oil and the availability of the alternatives.

Each of the different types of asphalt plant is available in fixed, free standing and mobile configurations. The fixed plants are bolted to reinforced concrete foundations. The free standing plants are mounted on a flat, levelled surface that can withstand a predetermined pressure, and no concrete foundations
are required. Mobile plants are also mounted on a flat, levelled surface, and the tyres are usually positioned on steel base plates; again concrete foundations are not required. In some parts of the world the term ‘portable’ is used instead of ‘mobile’, as the plants are not mobile during the production of asphalt. The choice of using a fixed, free standing or a mobile asphalt plant will depend mainly on the size and location of the contract.

14.2 Types of mixing plants
There are two basic categories of asphalt plant: batch and continuous. The terminology can be confusing, as both categories are continuous in operation. The categorisation refers to the method of mixing the dried and heated aggregates with the bitumen. In a batch plant the ingredients are mixed in discrete batches, whereas in a continuous plant the ingredients are mixed in an uninterrupted continuous process.

All of the batch and continuous asphalt plants operating in Europe are designed to meet the European safety standard BS EN 536:1999 (BSI, 1999).

The selection of which type of asphalt plant to use is influenced by many factors.

- **Location.** Is the plant to be located in a quarry, where the aggregate source is consistent and can be controlled, or is the plant to be located on a depot site, where the aggregate supply could change?
- **Market requirements.** How many different mixtures are to be produced? What type of mixtures and how many different mixtures will be required in a day’s production?
- **Plant capacity.** What is the estimated output required from the plant, and what are the operating conditions, including the moisture content of the feed materials and the final mixture temperatures?
- **Competition.** What other types of asphalt plant are in the surrounding area? Can the plant offer a niche product mixture?
- **Budget.** How much funding is available for the purchase of the plant?
- **Operational costs.** Does the plant selected manufacture an economical product based on the operational costs, including maintenance, fuel, labour etc.?
- **Site restrictions.** Does the type of plant meet all the site restrictions, such as height limits, space available, noise parameters and emission regulations?
- **Customer preference.** Is there a preference in the country or area in which the plant is to be located for a specific type of asphalt plant? Different customers and different continents have different requirements and different preferences.
Asphalt batch plants can be divided into two further categories: conventional asphalt batch plants and batch heaters. Batch heater plants are found only in the UK. Continuous asphalt plants consist of the drum mix, the counter flow drum mix, the double drum and the continuous mix plant. Each of the different types of asphalt plant is described in detail below, highlighting the influences on selection, the benefits of each type of plant, and where appropriate, the limitations.

14.2.1 Asphalt batch plants

14.2.1.1 Conventional asphalt batch plant

The conventional asphalt batch plant is the most common type of asphalt plant in Europe, Central Africa, China, the Middle East and Australasia. Although the process is continuous in operation, each batch of mixed material is made separately and the production then repeated. The batch size relates to the quantity of material being mixed in the paddle mixer. Capacities of from 50 up to 450 tonne/h are achievable, depending on the size of the equipment, the moisture content of the feed aggregates and the final mixture temperature. The rotary dryers range from 1 up to 3.1 m in diameter, and paddle mixer batch sizes range from 500 kg up to 5 tonnes and even larger. A conventional asphalt batch plant is shown in Figure 14.1.

Figure 14.1 A 320 tonne/h conventional asphalt batch plant. A TBA-4000 with a 4 tonne mixer and a 2.8 m dryer. (Courtesy of Benninghoven UK Ltd.)
The mixture recipe being produced provides the percentages and the tolerance limits of each aggregate size, the filler content and the bitumen content in the final mixture.

There are two methods of feeding the aggregates into an asphalt batch plant. The first, and most common, is to store the different aggregate sizes in separate stockpiles, or loading bays, and then transfer the aggregates using a loading shovel into the plant feed hoppers. The feed hoppers are usually fitted with safety grids to prevent people accidentally falling into the hoppers. In Europe, the loading bays are generally covered to help reduce unwanted water reaching the aggregates, especially the fines, and therefore improve plant efficiency. The feed hoppers are also covered on three sides in Europe to help prevent the aggregates and fines becoming airborne when transferred from the loading shovel to the hopper, this covering also adds protection from the elements.

The second method of feeding the aggregates and fines into the asphalt plant is direct from the crushing plant storage bins. The asphalt plant must obviously be located in a quarry for this method to be adopted. This method is the ideal solution, as the crushed stones and fines have the minimum possible moisture content, usually less than 2%, and the efficiency of the drying and heating process is therefore considerably increased.

A flow diagram of a conventional asphalt batch plant is shown in Figure 14.2. Each different aggregate size is fed into a dedicated feed hopper. The number of feed hoppers typically ranges from eight to 20, or more, depending on the different mixture recipes to be produced. Variable speed belt feeders are predominantly used to control the flow of the feed materials from the hoppers, or the crushed stone bins, onto the collecting conveyor before they progress to the dryer. The individual belt feeder speed is set in the controls to match the mixture recipe percentage of the particular aggregate being handled. The aggregates and fines are fed through the plant in the proportions required in the final mixture, thus ensuring economic use of the feed materials.

The dryer consists of a rotating steel cylinder inclined at 3–5°, with a burner mounted at the discharge end. The inside of the dryer cylinder is fitted with lifters, which aid the passage of the feed materials along the dryer, while also exposing the aggregates to the heat from the burner flame. The lifters at the feed end create a curtain of the aggregates, and the lifters at the discharge end form an annulus of the aggregates inside the dryer cylinder around the burner flame, which enables the flame to develop fully. The rotating dryer incorporates a two stage drying and heating process: first the aggregates are dried to remove the water in the feed materials, and then the aggregates are heated to the required temperature, usually between 120°C.
and 200°C, depending on the mixture recipe. The dryer cylinders can be insulated to prevent heat loss, with the added advantage of also reducing noise levels.

In order to support the burner flame, to remove the products of combustion and to remove the water vapour from the feed aggregates, an exhaust system is connected to the dryer. The exhaust is extracted from the feed end of the dryer. As the feed aggregates are passing along the dryer in the opposite direction to the exhaust air, this is termed a ‘counter flow air system’. The counter flow air system enables the hot exhaust gases to preheat the cold incoming feed materials. The exhaust system creates a flow of air inside the dryer, which consequently also picks up dust particles from the feed materials. These dust particles must be removed before releasing the exhaust air to the atmosphere.

The exhaust system generally consists of a two stage process. The first stage removes the coarse particles of dust, usually plus 75 micron, and returns these to join the heated product from the dryer discharge. In the UK the first stage or primary dust collector is usually a skimmer, which has the added benefit of being able to alter the coarse/fine dust split. Other primary collectors include knock out boxes, drop out boxes and cyclones. The second stage is the collection of fine dust using a bag filter. The bag filter consists of a large number of filter bags that collect the fine dust. The cleaned air then
passes through the exhaust fan and is discharged to the atmosphere through a high level stack. Particulate emission levels of less than 20 mg/m³ are easily achieved using a bag filter, and these dust emissions are not visible with the naked eye. The fine dust is removed from the filter bags by reverse air cleaning, a blower or pneumatic jets, and is transferred to a reclaimed filler silo for later addition to the mixture, or for disposal to waste, again depending on the final mixture recipe.

The heated aggregates, together with the coarse dust, are elevated from the dryer discharge by a hot stone elevator to a vibrating screen. The vibrating screen separates the feed materials into the sizes specified in the mixture recipe, and the sized aggregates are discharged into individual hot stone bin compartments. There are usually six individual compartments in the hot stone bins, although in some plants there are four or five compartments. There is a trend towards separating the dust and sand in the hot stone bins by way of a diverter door under the screen fines deck, resulting in seven hot stone compartments. A separate compartment is also usually incorporated in the hot stone bins for the screen bypass material – material that is a blended product and does not require screening. The hot stone bins incorporate integral overflow chutes that enable the plant to continue production if the individual compartments are full. Continuous level indicators are fitted to each of the hot stone bin compartments to provide information on the trend in the filling and discharging criteria of the sized aggregates. The operator can then adjust the feed rates accordingly, to control the overflow of aggregates and unwanted wastage of the hot materials. The overflow is collected in a separate storage hopper together with the screen rejects, or discharged to the ground. The hot stone bins can be insulated to prevent heat loss; again this also reduces the noise levels.

The aggregates are discharged separately from each hot stone bin compartment into the aggregate weigh hopper in accordance with the percentages specified in the mixture recipe. The aggregate weigh hopper is mounted on load cells, and positioned under the hot stone bins and directly above the paddle mixer. At the same time as the aggregate is being weighed off, the bitumen and filler are also weighed in dedicated weigh hoppers. In some US plant designs, the filler is weighed together with the aggregates in the aggregate weigh hopper.

The bitumen is stored in horizontal or vertical heated tanks, with each grade of bitumen being stored in a different tank. The heating of the tanks can be indirect, by using a hot oil heater with transfer oil, or electric heating elements inside the tanks. The tanks are insulated to reduce heat losses, and the piping through which the bitumen passes is also heated and insulated. The bitumen is circulated in a ring main at tank level; alternatively, each grade of bitumen
can be circulated up to the bitumen weigh hopper on the plant. A three way valve is used to discharge the circulating bitumen into the bitumen weigh hopper. The bitumen weigh hopper is mounted, or suspended on load cells, and the quantity of bitumen is delivered according to the amount specified in the final mixture. Two stage weighing of the bitumen can sometimes be used to provide a bitumen top-up relative to the actual amount of aggregates and filler being held in the aggregate and filler weigh hoppers. The weighed bitumen is either gravity fed to the paddle mixer, usually through a spray bar, or pumped under pressure through a spray bar across the mixer.

Reclaimed filler from the bag filter and imported filler to the plant are stored in separate filler silos. These silos discharge by means of screw conveyors to the load-cell-mounted filler weigh hopper, the amount discharged depending on the quantity specified in the mixture recipe. In Europe, if the reclaimed filler is to be discharged to waste, it is handled through a dust conditioner, where water is mixed with the reclaimed dust to prevent airborne particles.

The aggregate, bitumen and filler are weighed simultaneously, and then discharged into the paddle mixer in a pre-set order. The paddle mixer consists of two rotating shafts with arms and tips attached. The ingredients are mixed until a homogeneous mixture is produced, which is usually achieved in 30–40 s. The mixing and weighing cycles overlap, so that while one batch is being mixed the next batch is being weighed. Most asphalt batch plants are rated using a 45 s mixing cycle, which is the time from when the mixer discharge door opens to the next time it opens. The 45 s cycle results in 80 batches of mixture produced in one hour. A 3 tonne paddle mixer can therefore produce 240 tonne/h. The mixing cycle time is variable, and can be adjusted within the control system. However, the mixing time should be sufficient to enable a thorough coating of the bitumen, but not too long in order to avoid oxidation of the bitumen in the mixture. Recovered bitumen contents of ±0.1% of the target bitumen value can be achieved in an asphalt batch plant. The mixer discharges the finished product either directly to trucks, or to a mixed material storage system.

The mixed material storage can be positioned directly underneath the paddle mixer, using flap doors to direct the mixed material into different storage hoppers, or the paddle mixer can discharge into a horizontal skip that traverses and discharges into individual storage hoppers. This design can be found in many European asphalt batch plant sites where space is limited. Alternatively, the mixed material storage hoppers can be positioned adjacent to the mixing tower, where the paddle mixer discharges into an inclined skip that then feeds the storage hoppers. The mixed material silos are insulated and heated with electric elements or hot oil, to help prevent heat loss. European asphalt plants usually store mixed materials for a maximum of

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12–24 h, the heating and insulation ensuring minimum heat loss. Extended storage times of up to 4 days are possible in the mixed material silos without loss of mixture quality if the top and bottom doors are sealed to prevent oxidation of the mixture. This extended storage is a requirement in the USA, but not all mixtures can be stored for extended periods: polymer modified and open graded mixtures are excluded from plant manufacturers’ long term storage guarantee (for literature on storage silos, see Astec Inc., 2014). Mixed material storage silos can also incorporate automatic load out systems into the trucks. The silos are mounted on load cells, or a weighbridge is positioned underneath the silos, and the operator can pre-select the amount to be discharged into a truck.

In the majority of developed countries, the asphalt batch plant tower is enclosed, or sheeted. The sheeting contains any airborne dust inside the mixing tower, to comply with environmental regulations, and also results in a more aesthetically acceptable design. However, if employed in countries that have very high ambient temperatures, sheeted mixing towers must be vented, as the sheeting also contains the heat generated in the process, which can lead to premature bearing failures.

A variant of the asphalt batch plant is the multiple hot stone bin plant. The concept is to store two different types of hot material for weighing; for example, six hard stone aggregates and six soft stone aggregates. Instead of the standard six or seven hot stone bin compartments located underneath the vibrating screen, a central division plate enables the storage of 12, 13 or more hot aggregates. The vibrating screen has a large diverter chute underneath, which directs the aggregates to the hard or soft stone compartments. An alternative is to use two vibrating screens instead of one, and the hot stone elevator then incorporates a two way diverter chute. The asphalt batch plant would be fed first with soft stone to fill the relevant hot stone compartments, and then with hard stone. Usually, the hot stone bin capacity is larger in a multiple hot stone bin plant, and the increased surge capacity results in a more economical drying process. This type of plant is ideal for complete flexibility of production, allowing quick recipe changes without having to empty the hot stone bins if a hard stone mixture is required directly after a soft stone mixture. In Europe, this type of plant is used instead of the batch heater type, which is found only in the UK. However, multiple hot stone bin plants are gradually replacing the batch heater type in the UK. A multiple hot stone bin plant is shown in Figure 14.3.

Up to 25% reclaimed asphalt pavement (RAP) can be added to the batch plant directly into the paddle mixer, the percentage depending on the moisture content of the RAP and the mixture temperature. A flow diagram for a cold RAP feed is shown in Figure 14.4. The RAP must be in manageable
pieces of less than 30 mm size, and may need to be processed through a breaker to reduce the size of the sections being handled. The RAP is fed into a steep sided feed hopper, through a belt feeder to a conveyor that is fitted with a belt weigher. The conveyor discharges directly into the paddle mixer. The conveyor operates on a start–stop basis, depending on the percentage of RAP required. The weighed virgin aggregates are superheated and mixed with the RAP in the paddle mixer until the specified mixture temperature has been reached and a homogeneous mixture is produced. The paddle mixer requires an evacuation system to remove the water vapour produced from the heated RAP. This evacuation system is connected to the dryer exhaust system.

A small percentage of RAP (5–10%) can be added directly to the aggregate dryer by way of a recycling collar or a slinger conveyor positioned at the discharge end. The heated RAP is elevated to the vibrating screen, separated into the hot stone bins, weighed off in the aggregate weigh hopper and then mixed in the paddle mixer with the bitumen and filler. It is common to find a dual RAP feed to an asphalt batch plant, utilising the cold feed method direct to the paddle mixer and the RAP feed to the aggregate dryer. Percentages as high as 35% RAP can be added to the batch plant using the dual feed method.

Figure 14.3 A multiple hot stone bin plant with 360 tonne/h capacity. A Universal S plant with a single vibrating screen and a 14-compartment hot stone bin, located in Rubigen, Switzerland. (Courtesy of Ammann)
Higher percentages of RAP, up to 60%, can be added to a batch plant using a RAP dryer. Figure 14.5 shows a flow diagram for a typical hot RAP feed. The RAP dryer consists of a parallel flow drum, with the burner mounted at the feed end. The flow in the dryer is termed ‘parallel flow’, as the RAP and the exhaust air travel in the same direction through the drum. The RAP, which must be less than 30 mm in size, is fed into a steep sided feed hopper by means of a belt feeder onto a conveyor, or a vertical elevator, and discharged into the RAP dryer. The dryer heats the RAP to between 80°C and 120°C to remove the moisture, and it is then stored in a surge hopper before being discharged into a dedicated RAP weigh hopper. The RAP is then discharged directly into the paddle mixer, where it is mixed with the virgin aggregates, bitumen and filler. The RAP dryer has a large drop-out box mounted at the discharge end to allow the fine particles to be returned to the RAP. A scavenger fan removes the exhaust air, complete with the fumes, from the RAP dryer, and feeds the air and fumes into the virgin dryer burner flame for incineration. A high level RAP dryer is shown in Figure 14.6.

The conventional asphalt batch plant is the asphalt plant of choice in Europe, Central Africa, China, the Middle East and Australasia, and therefore the majority of development for this type of plant has emanated from Europe. As the plant incorporates a vibrating screen, one of its major advantages is that high quality mixed material can still be produced even if the sizing
of the feed aggregates cannot be guaranteed. Although the capacity suffers, this makes the plant universally accepted in any working environment. There are many benefits of an asphalt batch plant:

- the ability to manufacture all European (BS EN 13108:2006 (BSI, 2006a)) and US individual state Department of Transportation’s

![Figure 14.5 Flow diagram for a hot RAP feed (Hunter, 2000)](image)

![Figure 14.6 A high percentage RAP dryer on a 320 tonne/h TBA-4000-U asphalt plant, with a 4 tonne mixer, a 2.8 m dryer and a 2.4 m high level RAP dryer. (Courtesy of Benninghoven UK Ltd.)](image)
(DOT) mixture specifications, and therefore worldwide mixture specifications

- the emissions to the atmosphere are within European and US emission regulations, and therefore worldwide emission regulations
- inconsistent feed aggregate sizes can be accommodated while maintaining mixture quality, although plant capacity is reduced
- small and large tonnages are possible
- mixed material storage is not essential
- a high percentage of RAP can be added to the mixture.

Asphalt batch plants have few shortcomings, and these are only relative to the other types of production plant available

- capacity is restricted to the mixer size and the mixing cycle
- heat is wasted from the overflow and rejected materials
- high production costs relative to a drum mix plant.

14.2.1.2 Batch heater plant

Batch heater plants are only found in the UK. Each batch of material is produced individually throughout the process. Capacities range from 50 tonne/h up to about 200 tonne/h, with paddle mixer capacities typically ranging from 1 tonne up to 5 tonnes. The mixing cycle times can be up to 2 min, depending on the feed moisture content and the mixture temperature. A 3 tonne paddle mixer would, therefore, perform 30 cycles/h and achieve 90 tonne/h. Figure 14.7 shows a batch heater plant during construction, and completed and ready for production.

Pre-graded aggregates must be used in a batch heater plant as no further screening will take place during the process. The batch heater plant is fed from feed hoppers, the number of which is determined by the number of different aggregate sizes and different types of aggregate to be used. The number of feed hoppers can range from eight up to 20, or more, and each is fitted with a safety grid and a weather cover. Alternatively, the batch heater plant can be fed directly from the crushing plant storage bins. The latter is the ideal feed arrangement for this type of plant, as the moisture content in the feed aggregates is low, usually less than 2% for crushed stone. Batch heater plants have limited drying and heating capabilities compared with conventional batch plants.

The pre-graded aggregates are fed into the plant in a single batch using time-controlled belt feeders. Each feeder is calibrated for a specific aggregate type and size, the time set in the control system for each feeder depending directly on the final mixture recipe. Variable speed belt feeders can be used, but this is considered to be uneconomical and unnecessary. The batch is
layered onto the collecting conveyor and fed to a check weigh hopper. A flow diagram of a batch heater is shown in Figure 14.8.

The check weigh hopper is mounted on load cells and is used to measure the wet weight of the batch. An allowance is made in the control system for
the moisture content in the feed material to determine the dry weight, the bitumen and filler contents are calculated from the dry weight of the batch. The weighed batch is then discharged into the batch heater drum, which is a large diameter rotating cylinder, short in length, with a burner mounted at the opposite end. Inside the rotating drum are lifters that expose the aggregates to the heat from the burner flame. The aggregates are dried and heated inside the drum. The retention time inside the drum determines the discharge temperature.

An exhaust system is attached to the batch heater drum to remove the water vapour and to support the combustion process. The coarse dust is collected by a skimmer and fed back into the paddle mixer in discrete batches. The fine dust is collected in a bag filter. The different types of bag filter have been described in section 14.2.1.1 on conventional asphalt batch plants.

At the same time as the aggregates are being dried and heated, the bitumen and filler are weighed off in separate weigh hoppers using the calculated dry batch weight. The bitumen and filler systems are similar to those in conventional asphalt batch plants.

The heated aggregates are discharged from the rotating batch heater from the same end as the material feed, directly into the paddle mixer. In a pre-set order, the bitumen and filler are also discharged into the paddle mixer until a
homogeneous mixture is produced. The mixed material can discharge from the paddle mixer either directly into trucks or to a mixed material system. The mixed material systems on a batch heater plant can be high level or low level, and these systems feature the same characteristics as described for conventional asphalt batch plants.

Some later batch heater plants incorporate an aggregate weigh hopper after the batch heater drum, and a surge hopper replaces the wet batch weigh hopper. The bitumen and filler contents are then calculated from the true dry weight of the batch, reducing the possibility of errors.

If the fines or sands being used in a batch heater plant have a high moisture content, greater than 2%, then a separate dryer is used. The wet fines are dried in a counter flow dryer and stored in silos. The dried fines are then considered as an additional ingredient, and join the batch being made at the conveyor stage.

Between 10% and 15% RAP can be added to a batch heater plant using a weigh conveyor and fed directly into the paddle mixer. The process is described in more detail in section 14.2.1.1 on conventional batch plants. The percentage of RAP used is less than in a conventional batch plant because of the difficulty of superheating the aggregates in the batch heater drum.

The batch heater plant is ideal for complete production flexibility, where small batches of many different material specifications are required in a day’s work. The batch heater is unique to the UK and initially emerged because of the many different material specifications that existed prior to the advent of the European material specifications. The multiple hot stone bin plant has been developed as the European alternative to the batch heater. The benefits of the batch heater are

- an ability to manufacture all European (BS EN 13108:2006 (BSI, 2006a)) mixture specifications
- the emissions to atmosphere are within European emission regulations
- small batches can be economically produced
- quick recipe change possible
- there is no material wastage
- small and medium tonnages are possible
- mixed material storage is not essential
- a low percentage of RAP can be added to the process.

The limitations of the batch heater plant are relative only to the conventional asphalt batch plant, as the batch heater does not compete with the continuous plants.
pre-graded single-sized feed materials are essential, otherwise the mixture quality cannot be guaranteed
- a maximum 2% moisture content of the feed materials is necessary, or a pre-dryer is required for the fines
- the capital cost is high, especially if a pre-dryer is required.

### 14.2.2 Continuous asphalt plants

#### 14.2.2.1 Drum mix plant

Drum mix plants were used in the USA as early as the 1920s, although they did not come into common use until the 1970s. A US patent was applied for as early as 1929 by Donald McKnight Hepburn, and was granted in 1931 for a process that mixed the aggregates and bitumen inside a rotating drum (US Patent, 1931). The early drum mix plants were used for the production of ‘warm mixtures’, and further developments enabled drum mix plants to be used for the majority of mixture recipes.

Drum mix plants are ideal for high capacity production ranging up to 800 tonne/h. The equipment on a drum mix plant is minimal by comparison to a batch type asphalt plant, there is no grading screen, no hot stone bins, no weigh hoppers and no paddle mixer. The maintenance and running costs on a drum mix plant are therefore less than for a batch plant. However, it is essential on all continuous plants that pre-graded, single-sized feed aggregates are used, otherwise the final mix will not meet the necessary specification requirements and rejection of the finished product will result.

Variable speed volumetric belt feeders are used to control the flow of aggregates to the plant. Each feeder is used for a specific aggregate size and type of aggregate. The number of feed hoppers and feeders can be from five up to 20 or more. A flow diagram for a drum mix is shown in Figure 14.9. As for batch plants, in Europe the feed hoppers are fitted with safety grids and weather covers. The feeders are calibrated and set during production to achieve the percentage of the individual ingredients in the final mixture. As the aggregate feed to a drum mix plant is so important to ensuring a good mixture quality, the feeders are fitted with ‘no flow’ indicators to confirm to the operator that the feed is being discharged. A ‘no flow’ would result in the plant being stopped. Wet sands and fines can be difficult to control on a continuous plant, and on these occasions weigh feeders can be used.

A scalping screen is usually fitted between the feed section and the drum mix feed conveyor, which removes any unwanted oversized material or possible debris from the feed. This is especially useful if the aggregates have been stored as ground stockpiles prior to being fed to the plant.

The drum mix feed conveyor is fitted with a belt weigher that accurately determines the wet weight of the feed aggregates. The operator manually inputs
the overall moisture percentage of the feed material into the control system, and the controls then calculate the dry weight of the feed aggregates being delivered to the plant. This dry weight is used to calculate the bitumen and filler contents that are to be added later to the mixture.

The aggregates are fed into a rotating steel cylinder with a burner mounted at the feed end. The drum is angled at between $3^\circ$ and $5^\circ$, and is fitted internally with a series of lifters to expose the aggregates to the heat from the burner flame, to aid the aggregate flow through the drum and finally to mix the ingredients inside the drum. There are two distinct zones within the drum: the drying and heating zone; and the mixing zone, where the hot aggregates are mixed with the bitumen and filler. The two zones can be physically separated with a flame/target plate, or the design of the heating lifters separates the two zones by creating a dense material curtain out of the cascading aggregates. The length of the rotating cylinder is longer on a drum mix plant than on a batch plant dryer to cater for the mixing zone.

The exhaust system is connected to the discharge end of the rotating cylinder, and the opposite end to the burner. The exhaust system removes the water vapour, the products of combustion and the collected dust from the process. This is a parallel flow drum mix, as the aggregates and the exhaust air flow in the same direction through the drum. As the exhaust air passes through the mixing chamber on a drum mix plant the percentage of dust
collected is considerably less than in a batch type plant, as a large quantity of the airborne dust particles are absorbed by the bitumen in the mixing zone. Dust collection usually consists of a large drop-out box at the discharge end of the drum, the coarse dust particles dropping back into the process. The fine dust is collected in a bag filter, the different types of bag filter are as described in section 14.2.1.1 on conventional batch plants.

The proportion of bitumen and filler to be added to the mixture is calculated from the dry weight of the aggregates passing over the dryer weigh conveyor. The bitumen flow rate is controlled by a variable speed pump, or variable orifice control valves. The bitumen is added into the drum mixing chamber from the discharge end. It is important not to expose the bitumen to the naked burner flame, hence the reason for fitting the burner at the feed end of the drum. Degradation of the bitumen can occur if direct contact is made with the burner flame, and blue smoke results.

The imported filler and/or the reclaimed collected fine dust can be either blown or screwed back into the drum mixing chamber from the discharge end. These fillers can be accurately fed back into the drum using separate ‘loss-in-weight’ systems. The filler is conditioned to ensure a constant density, and is continuously reverse weighed against a known calibration rate. The bitumen and filler are added at the beginning of the mixing zone inside the drum, ensuring that the fillers mix with the bitumen and do not become airborne.

The mixed material discharges from the end of the drum mix, usually to a mixed material system. Small capacity drum mix plants can discharge into a surge hopper and then to trucks but, being continuous plants, the majority discharge into mixed material systems as described for conventional asphalt batch plants. The US manufacturers’ preference is to use slat conveyors instead of a skip to transport the mixed material; alternatively, vertical elevators can be used if site restrictions apply. The slat conveyor is ideal for handling high capacities of mixed material.

Between 15% and 20% of RAP can be added in a drum mix plant by means of a collar mounted around the rotating cylinder at the start of the mixing zone. The RAP must be less than 30 mm in size. It is fed into a steep sided feed hopper through a variable speed feeder, and onto a RAP weigh conveyor. The signal from the drum mix feed conveyor controls the amount of RAP added to the process. The virgin aggregates are superheated in the drying and heating section, and the RAP is mixed with the virgin aggregates, the filler and the bitumen in the mixing section. The percentage of RAP is limited because of its tendency to produce fumes or blue smoke when the RAP is mixed with the superheated virgin aggregates.
Drum mix plants have been the plant of choice in the USA, India, Malaysia, New Zealand, Norway and France, where long production runs of one material specification are required. In addition, drum mix plants have been, and still are, used for smaller capacities where funds are limited and where mixture specifications are not of paramount importance. As stated above, the success of drum mix plants, and all continuous plants, relies totally on using pre-graded single-sized aggregates as feed materials. If this cannot be guaranteed, the finished mixed material product is likely to be out of specification and rejected. The need to produce high temperature mixtures and to add high percentages of RAP has led to the development of counter flow drum mix, double drum and continuous mix plants, which are all variants of the drum mix plant. The benefits of drum mix plants are

- they are economical for long production runs of one material specification
- high capacities are readily achieved
- they have relatively low maintenance costs
- the amount of dust removed from the process is minimal
- they are easily adapted to a mobile design
- a low percentage of reclaimed material can be added to the mixture.

Drum mix plants are still in use currently in many different countries, and are performing a useful role. However, the production of high temperature mixtures and the requirement to add high percentages of RAP can result in degradation of the bitumen on a drum mix plant, and blue smoke results. These emissions adversely affect the performance of the bag filter, production rates suffer considerably over a period of time, and the fumes, or blue smoke, emitted into the atmosphere are not acceptable. The limitations of drum mix plants are

- pre-graded, single-sized feed materials are essential, otherwise mixture quality cannot be guaranteed
- they are unable to produce high temperature mixtures
- they are unable to produce a high percentage of RAP
- there is wastage of heated material at the beginning and end of production
- there is a possibility of blue smoke emissions
- small quantities are uneconomical
- a mixed material system is essential on higher capacity plants.

14.2.2.2 Counter flow drum mix plants
Counter flow drum mix plants are a direct development of the drum mix plant, and overcome some of the shortcomings so that high temperature mixtures can be produced and high percentages of RAP can be added to the mixture.
The counter flow drum mix consists of three basic categories: the counter flow drum mix, the double drum and the continuous mixture plants. All the ancillary equipment, such as the feed section, the weigh conveyor, the dust collector, the bitumen supply, the filler feed and the mixed material system, is the same as in a drum mix plant. The only difference is the drum mix rotating cylinder. The internal structure of a counter flow drum mix cylinder is shown in Figure 14.10.

The counter flow drum mix consists of a long steel rotating cylinder with an extended burner mounted at the discharge end. The burner extends inside the rotating cylinder so that the flame develops towards the centre of the drum. The rotating cylinder consists of two distinct and separate zones: the drying and heating zone, and the mixing zone, which is mounted behind the burner flame. The drying and heating section is based on the same principle as in a conventional batch type dryer, the exhaust air travels in the opposite direction to the aggregate flow, hence the term ‘counter flow drum mix’. The heating and drying is therefore more efficient than in a standard drum mix, as the feed aggregates are preheated by the hot exhaust air. The bitumen and fillers are added in the mixing zone and discharged when a homogeneous mix is produced. The mixing time depends on the length of the mixing chamber, the design of the mixing lifter pattern and the angle of the drum, which is usually between 3° and 5°. Typical mixing retention times, which is the time the constituents are mixed, can range from 45 s up to 60 s or more. The mixing zone incorporates a scavenger extraction system for any fumes or blue smoke that may be created, and feeds this into the extended burner. The fumes are then incinerated in the burner flame.

Figure 14.10 Flow diagram of a counter flow drum mix cylinder. (Hunter, 2000)
Up to 60% RAP that has been reduced to less than 30 mm in size can be added to the mixing zone when called for in the mixture recipe. The reclaimed material is fed into the rotating cylinder through a collar wrapped around the drum. The aggregates are superheated to enable heat to be transferred to the RAP, first removing any moisture present in the RAP, and then heating the combined mixture. The water vapour, together with any fumes from the RAP, is extracted by the scavenger fan and fed into the extended burner for incineration.

Another counter flow drum mix solution is the double drum (Astec Inc., 2014). Figure 14.11 shows a double barrel drum mix plant. The double drum consists of a rotating dryer with the burner mounted at the discharge end. After passing through the dryer, the dried and heated aggregates are fed into an outer stationary shell of the drum. The rotating dryer has mixing paddles attached to the outside of the drum. The bitumen, filler and RAP are added to the outer stationary shell. Up to 60% RAP can be added to the double drum. Water vapour, together with any fumes created by the RAP addition, are drawn into the burner and incinerated. The internal structure of the double drum is illustrated in Figure 14.12.

The counter flow drum mix and the double drum are both US design solutions for the new breed of drum mix plant. They can both also be used as conventional dryers, with the dried and heated aggregates being discharged.
Virgin material enters the drum by means of an air-sealed flop gate and gravity chute.

1. Recycle entry
2. Bitumen entry
3. Fines entry (not shown)

Dried aggregate leaves the drum through aggregate exit openings.

Final mix exits out by means of a discharge chute to the silo drag conveyor.

Mixing paddles on the outer shell ensure bitumen thoroughly coats all of the aggregate.

**Figure 14.12** The internal structure of a double drum, showing (a) the inner drum drying zone and (b) the outer drum mixing zone. (Courtesy of Astec Inc.)
before the mixing zone. The discharged hot aggregates can then be fed to a conventional asphalt batch plant. This type of arrangement is termed a ‘convertible’ or ‘combination’ plant, and provides versatility to the customer where a batch and a continuous plant are required.

The European development of the drum mix plant is the continuous mix plant. The drum mix rotating cylinder is replaced by a standard conventional asphalt batch plant dryer and a continuous mixer. The continuous mixer is an extended paddle mixer with a weir plate at one end. The weir plate can be adjustable to control the mixing time, and the mixer shaft speeds can also be varied, again to control the mixing time. The bitumen, imported filler, reclaimed filler and RAP are added to the continuous paddle mixer. The continuous mixer is usually vented to the dryer to incinerate the fumes. The maintenance and operational costs of a continuous mix plant are higher than for a counter flow drum mix or double drum plant, as a paddle mixer is incorporated in the design. A continuous mix plant is shown in Figure 14.13.

The counter flow drum mix, double drum and continuous mix plants enable high temperature mixtures and the utilisation of high percentages of RAP for asphalt production in a continuous plant. The benefits are

- they are economical for long production runs of one material specification
- they can be used to manufacture all European (BS EN 13108:2006 (BSI, 2006a)) and US individual state DOT mix specifications, and therefore worldwide mixture specifications
- high capacities are readily achieved
- high temperature mixtures can be produced
- maintenance costs are relatively low
they are easily adapted to a mobile design
- a high percentage of RAP can be added to the mixture
- fumes or blue smoke are incinerated by the burner.

Although the counter flow drum mix, double drum and continuous mix plants overcome two of the main shortcomings of the drum mix plant, it is still necessary that the feed is composed of pre-graded, single-sized aggregates, otherwise the final mixed material product may not meet relevant specifications. The limitations are

- pre-graded, single-sized feed materials are essential, otherwise mixture quality cannot be guaranteed
- there is wastage of heated material at the beginning and end of production
- small quantities are uneconomical
- a mixed material system is essential on higher capacity plants.

14.3 Plant capacities

There are two main constraints on the capacity of an asphalt plant: the drying capacity and the mixing capacity (Hunter, 2000). Both these process constraints are complex. Factors affecting the drying capacity are

- altitude
- ambient air temperature
- type and grading of feed material
- feed material temperature
- moisture content of the feed material
- moisture content of the discharge material (usually less than 0.5%)
- dryer discharge temperature
- exhaust gas temperature from the dryer
- angle of inclination of the dryer
- calorific value of the fuel
- available exhaust air volume
- lifter pattern inside the dryer
- burner efficiency.

Factors affecting the mixing capacity to ensure a homogeneous mixture are

- mixture recipe
- mixer live zone (if applicable)
- paddle tip speed (if applicable)
- size of paddle tips and arms (if applicable)
- mixer arm configuration (if applicable)
- mixing zone lifter configuration (if applicable)
- method of feeding ingredients
Either the drying process or the mixing process determines the capacity of an asphalt plant. Both must be considered separately, and the capacities compared. The lower restricting capacity is then selected as the overall plant capacity.

The capacity of a batch plant is easy to determine, as the drying and mixing processes are separate. In drum mix, counter flow drum mix and double drum plants, the drying and mixing processes are combined, and therefore more detailed consideration is required. The constraints due to drying in a drum mix plants and its variants are similar to those for a conventional asphalt plant dryer. The mixing time in a drum mix type plant can be controlled by adjusting the throughput, altering the drum angle, adjusting the position of the bitumen injection pipe within the mixing zone, altering the design of the mixer lifters, or altering the speed of the rotating cylinder. However, in reality, once these variables have been set during commissioning, or at the design stage, they are not usually altered. The constraint due to the mixing process in a drum mix type plant is, therefore, usually determined from firsthand operational experience with the equipment used.

The following information is required to determine the size of an asphalt batch or continuous plant:

- maximum plant capacity
- type of feed material
- average moisture content of the feed material
- types of mixtures to be produced
- temperature of mixtures
- special constraints (e.g. minimum mixing time).

### 14.4 Additive systems

There are many additive systems that can be incorporated in an asphalt production plant, either batch or continuous. However, an additive ingredient is usually easier to control in the batch process.

Asphalts can be produced in many different colours. The most common coloured additive is red, although blue, green, gold and clear mixtures are also available. The different coloured mixtures can be achieved by adding a pigment, in bag form, direct into the paddle mixer, or by adding the pigment into the bitumen supply.

Rubber can be added to the mixture in several formats. Latex can be added by means of a pump into the paddle mixer, or into the mixing zone in a drum...
mix type plant. Rubber granulate can be added, by weight, directly into the mixer in a batch plant, or through the RAP collar in a continuous plant.

Fibres are also required in some stone mastic asphalts and thin surface course systems. The fibres can be in loose form, which are small fibre ‘strings’ or pellets. The pellets can also be impregnated with bitumen. The fibres can be introduced in bulk form or as bags. The use of bulk fibres requires the inclusion of silos with associated handling equipment, and the bagged fibres require the use of bag splitters. The fibres are added to the mixture by volume or weight, directly into the paddle mixer.

14.5 Cold mix plants
Cold mix plants can be of batch or continuous design, although the majority in use now are continuous plants. The term ‘cold mix plant’ refers to the aggregates being fed into the plant, which are not heated. The bitumen and the bitumen emulsions added to the mixture are heated. The equipment is therefore very basic, as there are no dryers, burners, dust collectors, grading screens, hot stone bins or weigh hoppers, and mixed material storage is optional. The simplicity of the cold mix plants results in low level arrangements, which lend themselves to being mobile or skid mounted. A continuous cold mix plant is shown in Figure 14.14.

Cold mix plants can be used to produce cold mix asphalt with foamed bitumen, cold mix asphalt with emulsion, and hydraulically bound mixtures.

Figure 14.14 A continuous cold mix asphalt plant with capacity of up to 400 tonne/h. (Courtesy of Ammann)
for use as a base. Foamed bitumen is created by spraying water under pressure through nozzles into the hot bitumen. The water expands when the pressure is released, resulting in a fine bitumen foam that is used to coat the cold aggregate feed (for an overview of the literature on asphalt mixing plant, see Ammann Schweiz AG, 2014).

In a continuous cold mix plant, pre-graded single size aggregates are fed into the plant using pre-calibrated, dedicated variable speed belt feeders. The feeders are set at the percentage required in the finished mixture specification, and the aggregates are then fed onto a belt weigh conveyor. Prior to the weigh section on the conveyor, the cement, hydrated lime or imported filler is added by way of a silo and a loss-in-weight system onto the aggregate feed. The belt weigher is therefore weighing the total of the wet weight of the aggregates and the dry filler weight. The aggregates and filler are discharged into a continuous paddle mixer. This is an extended paddle mixer with a weir plate fitted at the discharge end. The signal from the belt weigher controls the metering of the bitumen and/or water or bitumen emulsion feed into the continuous mixer through a spray bar. The cold mixture discharges directly into trucks, by means of a conveyor into trucks, or into an optional cold mix surge hopper. Direct truck loading involves the control system being set to produce the specific quantity required in the truck.

A cold mix batch plant has a paddle mixer with a bottom discharge instead of a continuous mixer, and a bitumen weigh hopper instead of a bitumen metering system. Each batch is made separately, whereas in a cold mix continuous plant each truck load or surge bin capacity is specifically set for production. Continual discharge into trucks from the surge hopper would equate to continuous plant production. Capacities of up to 400 tonne/h can easily be achieved in cold mix plants.

RAP can be directly added in the cold mix plants, either into the continuous mixer or the paddle mixer.

Cold mix plants are becoming increasingly popular in some parts of the world for the production of bitumen bonded bases and hydraulically bound mixtures, where only aggregates, cement and water are used. The initial cost of the plant is low, as are the operational and maintenance costs, because no heating or dust collection is required, and therefore the cost of the cold mix is economical. However, pre-graded, single size aggregates must be used to feed the plant, and the performance of the finished cold mix product is limited. Further development of these cold mixes is currently being undertaken. The benefits of cold mix plants are

- low initial cost of equipment
- low maintenance and operational costs
low cost of the finished mixture
reduced environmental impact, as no heating and drying of aggregates is necessary
simple low level design
easy to adapt to a skid mounted or mobile design
high capacities are readily achieved
small batches are possible
no material wastage
mixed material storage is not essential
RAP can be added.

The limitations of cold mix plants are

- pre-graded, single size feed materials are essential, otherwise mixture quality cannot be guaranteed
- restricted performance of the finished cold mixture product.

### 14.6 Warm mix plants

The need to reduce the ‘carbon footprint’ when producing asphalt has led to the development of warm mix asphalt. Warm mix asphalt is categorised as being produced and mixed at temperatures roughly between 100°C and 140°C (EAPA, 2010; Nicholls et al., 2013). Warm asphalt mixtures are produced in several ways

- the incorporation of chemical additives, such as amines, or the addition of waxes to reduce the bitumen viscosity (Hauguel, 2013; Walsh, 2013)
- the introduction of foamed bitumen, or the addition of moisture releasing additives, such as zeolites to the mixture (Nicholls et al., 2013).

Whichever method is adopted the asphalt production plant can be adapted to accommodate the changes. Existing production plants can also easily be retrofitted with the necessary equipment.

The most common method used to produce warm mix asphalt is by the introduction of foamed bitumen. Foamed bitumen is created by injecting cold water into the flow of bitumen, which is usually at a temperature of 150°C (for more information on foamed bitumen, see section 22.1). The water is heated above boiling point in the hot bitumen and some of it evaporates, and the result is a mixture of steam, water and bitumen, which is then mixed with the low temperature aggregates. The foaming of the bitumen enables the aggregates to be coated by the bitumen at a lower temperature.

The quantity of water added is small when making foamed bitumen; for example, 1.25–2% of water by weight of the total bitumen content. The
water is added to the process using a positive displacement pump and the quantity is controlled by a flow meter.

The benefits of producing warm mix asphalts compared with traditional hot asphalts are considerable:

- lower manufacturing costs
- lower fuel consumption
- reduced carbon dioxide emissions to the atmosphere
- reduced fumes and odours at the production plant
- reduced fumes and odours at the laying site, and therefore a safer working environment for paving crews.

The major advantage with warm mix asphalts is the significant reduction in emissions. Research has shown that for every 11°C reduction in mixing temperature the emissions to the atmosphere are halved (Hauguel, 2013; Walsh, 2013). This is a substantial reduction in the carbon footprint for the asphalt production plant considering that the existing equipment can still be used.

### 14.7 Asphalt plant control

The controls on all asphalt plants are now fully automatic, usually with a PLC (programmable logic controller) to control the process and a PC (personal computer) to act as the interface with the operator. Plants are equipped with manual control mainly for maintenance purposes. The direction now being taken is to use a PC for manual control, instead of individual push buttons to start/stop each motor. The PC is then augmented by a backup PC, which is feasible due to the relatively low cost of a PC compared with the push buttons and associated wiring.

A PC can provide a dynamic image of the plant layout, with the separate components of the plant changing colour depending on whether or not they are active, and the flow of material through the plant is simulated using line graphics and interactive animated symbols. The image will also show the weights, tonnages and temperatures throughout the plant, so that the operator can see at a glance the overall plant performance. A dynamic image for a conventional asphalt batch plant is shown in Figure 14.15.

The control system has several functions:

- plant operation
- fault diagnostics
- recipe storage
- production data
- daily production schedule
- operator interface
printing, storing and transferring production data

quality assurance.

The control system detects and confirms whether the mixture has been produced in automatic or manual mode. This is important for quality assurance purposes. In fact, if the control system is in automatic mode, the correct data have been input, the correct aggregate sizes and bitumen grades have been fed into the plant, and no faults have been recorded, then the control system will ensure that the asphalt mixture is within specification. The plant operator just needs to observe the process, and intervention is only required if there is a problem. The quality of the mixture is therefore not dependent on the operator.

Process faults such as starved feeders, stopped motors, broken chains, tare weight faults, sticking doors, sticking valves etc. can be monitored by the control system and alarms raised when there is a problem. This reduces downtime and improves the overall plant efficiency. The control system can also be set to stop the plant if temperatures or weights are not within pre-set parameters, thus providing quality assurance.

The information from the production schedule can be printed, stored and/or transferred to another PC, which may be in the laboratory, at the weighbridge or in the accounts department.
Each aspect of the asphalt plant production process is or can be controlled and monitored automatically using the flow rates, the weights being processed, the levels in the hoppers, bins and silos, and the temperatures throughout the plant.

The variable speed volumetric belt feeders have a turndown ratio of 20 : 1 to control the flow of aggregates to the plant. The feeders are calibrated during plant commissioning to achieve the required capacities. The feeders can therefore be set to operate at between 5% and 100% capacity to ensure optimum aggregate feed to the plant with minimum material and heat wastage, providing the hoppers are filled with the correct size aggregates. The individual belt feeder speed is set in the controls to match the mixture recipe percentage of the particular aggregate being handled.

Low level indicators can be fitted to the feed hoppers, with elevated warning lights above the hoppers to alert the shovel driver of the need to charge that particular aggregate hopper. No flow indicators can also be fitted to the feeders to warn the operator that the hopper is either empty or a blockage has occurred. A no flow condition can either activate an alert, or after a time delay the plant can be automatically stopped in a controlled sequence.

The burner on a batch or continuous plant can have a turndown ratio of up to 10 : 1. This means that for a plant with a maximum burner flow rate of 3000 l/h the minimum flow rate can be reduced to 300 l/h. This enables an economic use of fuel for a whole range of different flow rates and various discharge temperatures. The turndown ratios are achieved by variable speed fuel pumps and variable speed blowers fitted to the burner, the motor speeds and settings are individually controlled by the PLC. The discharge temperature of the material leaving the dryer, or the drum mix is measured by a non-contact infrared pyrometer, which is used to alter the burner fuel and air settings in a closed loop control.

The volume of exhaust air from the dryer, or drum mix is controlled by a pressure sensor mounted in the burner end box. The volume of exhaust air is increased or decreased to maintain a set pressure at the burner in order to sustain combustion and remove the products of combustion. The exhaust air is varied by employing a variable speed exhaust fan.

The levels of the materials within the hot stone bins and filler silos are monitored by the use of continuous level indicators within each hot stone bin compartment and in each filler silo. The hot stone bin level indicators provide the operator with trending information on the filling and discharge rate of each hot stone bin compartment. The operator can then adjust the feeders according to this information if the levels are not within the mixture specification requirements. This enables the plant to perform more economically with
reduced overflow from the hot bins. The feeders are not automatically controlled from the hot bin level indicators because the levels are a trend and only accurate to within 10% of the actual level. In addition, and more importantly, the feed aggregates from each feed hopper will be distributed into several hot bins and the specification of these aggregates can be inconsistent, so automatic control is not feasible. The levels in the filler silos are for information only; however, they provide useful data instead of just ‘full’ or ‘empty’.

The accuracy of the batch plant weigh gear for the aggregate, bitumen and filler ingredients is consistently repeatable to within ±0.3%. Weighing the bitumen relative to the actual weight of aggregate and filler recorded in the weigh hoppers enables the bitumen, which is the most expensive ingredient in the mixture, to have a recovered accuracy of ±0.1% of the specification target value. Self-learning in-flights from the hot stone bins discharge to provide accurate aggregate weights, and alternative hot bin selection are features incorporated in batch plant control systems to help sustain plant capacity while still maintaining accuracy. The load cells on the belt weigher in continuous plants have guaranteed accuracies of ±1%.

Temperatures are monitored throughout the plant prior to the asphalt being mixed. Thermocouple probes are located in the hot stone bins and the bitumen ring main to ensure that the temperatures are within the specification limits before mixing takes place. Non-contact infrared pyrometers are also positioned under the paddle mixer on batch plants and at the drum mix discharge to record the actual mixture temperature. Pyrometers can also be fitted at each mixed material silo discharge point to record the temperatures of the asphalt in the trucks.

The mixed material storage silos can be fitted with load cells or strain gauges to determine the weight in each silo, and to offer an accurate automatic load out facility to trucks, thus preventing wastage and saving time from excessive visits to the weighbridge. Alternatively, weighbridges can also be mounted underneath the mixed material silos to provide an automatic load out system; however, this does not record the weight in each silo.

14.8 Sampling

Testing of samples taken from an asphalt plant should only be carried out by skilled and experienced personnel using well maintained and calibrated test equipment (Forde, 2009). It is impractical to test every tonne produced, but measures must be taken to ensure that the aggregate grading and the bitumen content are in accordance with the mixture specification. Therefore, samples are taken at predetermined intervals, making sure that the samples are representative of the whole production. Recommendations
about minimum sampling frequencies are provided in BS EN 13108-21:2006 [BSI, 2006b], and care must be taken to ensure that the sampling frequency provides evidence that the plant is under control.

In the UK, methods of sampling and achieving representative samples are given in PD 6682-2:2009 + A1:2013 [BSI, 2009/2013] for the incoming aggregates, sands and fillers to the asphalt plant, and are described in PD 6692:2006 [BSI, 2006c] for the asphalts being produced. These are useful UK guidelines on the use of BS EN 13043:2002 [BSI, 2002] and the various BS EN 12697 testing standards [BSI, 2005–2012], respectively.

Materials being delivered to the asphalt plant should also be sampled, including the aggregates, bitumens and fillers. If the asphalt plant is connected directly to the crushing plant storage bins then part of this control will already be in place.

Samples can also be taken from within the production process. On batch plants, sampling facilities are usually located beneath each hot stone bin compartment. Alternatively, these samples can be taken directly from the paddle mixer discharge. These samples confirm the aggregate grading within each hot stone bin compartment, helping to warn of material contamination or screen mesh breakages. Sampling points are also incorporated in the reclaimed and imported filler discharge points, and usually for each bitumen tank. These individual samples should be taken at regular intervals to help control the aggregate grading and to confirm that the correct bitumens and fillers are in use.

During the calibration and commissioning of asphalt plants, or if troubleshooting, samples are taken from the paddle mixer, drum mix discharge and the mixed material discharge points. The samples from the paddle mixer or drum mix confirm whether a homogeneous mixture has been produced. The samples from the mixed material discharge indicate whether the material has been stored successfully, or if segregation, excessive loss of temperature or some other deterioration has occurred. During full production, the samples are usually taken from the truck.

Sampling and testing in the laboratory takes time, and asphalt plants producing at high capacities can result in a large quantity of asphalt being laid on a highway or airport runway before the results are available. For this reason, small production runs and tests are usually carried out prior to large production batches being undertaken. Sampling of asphalt production plants is essential to show compliance with the European standard for asphalts, and is usually carried out in accordance with components of BS EN 12697 [BSI, 2005–2012]. The positive results of sampling ensure confidence in the performance of an asphalt plant, and provide quality
control of the production process. Sampling on an asphalt plant is essential, and must be performed regularly.

References
Chapter 15

Transport, laying and compaction of asphalts

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15.1 Transport
It is important that heat loss during transportation is minimised. Accordingly, delivery vehicles must have a body constructed with an acceptable insulating material on all surfaces in contact with the asphalt, together with a means of covering the top of the asphalt in a manner that minimises heat loss. This can be achieved using either a mechanical sheeting system, or a sheet positioned under the asphalt coupled with a waterproof sheet over the top of the hot asphalt. A typical modern delivery vehicle used for transporting asphalts is shown in Figure 15.1. Journey times and standing time on site should be as short as possible, and this is particularly important during inclement weather.

Care should be exercised during the loading of delivery vehicles at the asphalt production plant to ensure that opportunities for cold spots are minimised. This, in turn, will reduce the risk of inadequate compaction and early failure of the asphalt.

Any releasing agent used in the wagon body to assist discharge should be such that it will not contaminate or adversely affect the performance of the asphalt. The amount of releasing agent used should be kept to a minimum.

15.2 Preparatory works
Asphalts should only be laid on surfaces free from detritus and deleterious material. Prior to installation, the underlying course should be prepared to produce a stable surface of appropriate profile on which the asphalt can be placed. Any weak areas of the substrate should be strengthened, and major variations in profile corrected. Crazing and narrow cracks in the substrate are acceptable; however, wide cracks should be sealed using an appropriate treatment. Badly cracked areas should be removed completely.
Where improvements are required to the receiving layer, the addition of a regulating course or cold milling may be necessary to achieve the designed profile. Cold milling (Figure 15.2), also described as ‘planing’, should be undertaken using a suitable milling machine that can complete the work to the required depth both effectively and efficiently. Depending on their size, modern cold milling machines can remove asphalt at depths of between 0 and 350 mm, and at operating widths of between 0.35 and 3.8 m. The cut edges should be left neat, vertical and in straight lines. The existing ironwork should not be disturbed by the milling process. Where necessary, surfacing in the vicinity of ironwork and in small or irregular areas should
be cut out using pneumatic tools or other suitable methods and removed. Immediately after milling, surplus materials should be removed using a small loading shovel or back hoe loader, and the milled surface mechanically swept to remove all dust and loose debris. The finished surface should have a uniform texture.

Any water ponding on the receiving substrate must be removed.

Where asphalt is to be installed on existing concrete pavements with defective joints, the joints should be made good by cleaning out and refilling with a suitable joint treatment. This treatment should finish flush with the surface, and should be of a type that will not adversely affect the overlying asphalt.

Prior to the installation of the asphalt, the vertical faces of any access chamber covers, gullies, kerbs, channels and similar projections against which the asphalt is to abut should be cleaned and painted with a uniform coating of hot applied paving grade bitumen, cold applied thixotropic bitumen emulsion of similar grade, or polymer modified bitumen emulsion bond coat. The choice of material to be used will be dictated by the job specification. Many clients understandably believe that there is no substitute for paving grade bitumen.

15.3 Application of bond coats

It was explained in Chapter 9 that a tack coat is a bituminous emulsion used to promote adhesion of contiguous layers. It was further explained that bond coats perform the same function but are proprietary products manufactured with polymer modified bitumen. Compared to a tack coat, a suitable bond coat is much more likely to enhance adhesion of contiguous layers. It is for this reason that, in the UK, bond coats are invariably specified rather than the traditional tack coats as per the Note to Clause 3.2 of BS 594987:2010, which states: ‘While tack coats have traditionally been used, they are no longer regarded as best practice’ (BSI, 2010b).

The bond/tack coat should be hot applied either by using equipment separate from the paver (Figure 15.3), in which case it should be permitted to break (i.e. to turn from brown to black in colour) prior to paving, or by using a paver with an integrated spray bar (sometimes described as an ‘integral paver’), in which case breaking occurs on contact with the overlying hot asphalt. Breaking is essential because lorries reversing into the paver hopper and the paver itself will lift the bond coat on their tyres.

Before spraying is started, the surface should be free of all loose material and standing water, which would affect the adhesion between the layers.

Rates of spread for bond coats or tack coats are typically found within the applicable specification and/or proprietary product guidelines. In the
UK, it is currently specified in Clause 5.5.2 of BS 5949-87:2010 (BSI, 2010b) as 0.35 kg/m² of residual binder. This is currently being reviewed with a recommendation that it should be reduced to 0.20 kg/m². Selection of an appropriate tack coat or bond coat will be dictated by the nature of the substrate. The aim is to leave an even coating of bitumen on the receiving layer. Permeable (dry and/or open) surfaces and hydraulically bound substrates may require a higher rate of spread than that required for asphaltic substrates.

Care should be taken to avoid over application or ponding, as this could result in a degree of slippage or instability in the overlying layer and, in some circumstances when overlaying with an open material, bleeding. Consequently, any emulsion accumulating in hollows should be dispersed by brushing.

In the UK, until 2006, the inclusion of a tack or a bond coat was normally only required below surface courses on areas of overlay or at tie-ins to existing pavements. However, the 2006 edition of the UK pavement design standard HD 26, which is still current, states that ‘a tack or bond coat is required between all layers’ (Highways Agency et al., 2006). In recent years, there has been some discussion in the UK about the role played by tack coats and bond coats in aiding efficient pressure dissipation across asphalt layer interfaces. Although there have been no studies that have provided proof that a lack of bond may, of itself, cause a pavement to fail, there is no doubt that the application of a tack coat or a bond coat at all asphalt to asphalt interfaces is sound practice (Nicholls et al., 2008).
15.4 Delivery and discharge
Asphalt deliveries should be coordinated primarily to avoid interruption to the laying operation but also to avoid any unnecessary waiting on site. Any such delay may result in the delivered asphalt cooling to a point at which workability becomes an issue and, in turn, have an adverse effect on the long term performance of the pavement.

Where the paver cannot be operated in a continuous mode, consideration should be given to the possibility of material at the extremes of the hopper and at the end of loads, where cooler material is often found, being discarded. Cool material can result in inadequate compaction and subsequent premature failures, such as ravelling. This type of failure can also occur when material is dropped from delivery vehicles in front of the paver and is not removed, as it will cool rapidly on the cold substrate before being overlaid with asphalt from the paver. Delivery vehicles should be completely empty before withdrawing from the hopper of the paver to avoid the tailgate dragging asphalt off the hopper onto the area in front of the paver.

15.4.1 Material transfer vehicle
In many countries, material transfer vehicles, often colloquially known as ‘shuttle buggies’, are used in tandem with a paver or pavers to remix the asphalt immediately prior to installation (Figure 15.4). This reduces the incidence of material segregation and temperature variations caused by subjecting the material to long haul distances or excessive waiting times.

Material transfer vehicles, in conjunction with a paver hopper insert, can provide additional receiving capacity of the delivered asphalt, with the result

Figure 15.4 Material transfer vehicle. (Courtesy of Eurovia SA)
that the waiting times of delivery vehicles on site are reduced and continuous paving operations are achievable. Additionally, during material discharge, the material transfer vehicle serves as a buffer between the truck and paver, eliminating any bumping and resulting surface regularity issues.

15.5 Pavers
Pavers can install the full range of asphalts as well as granular, hydraulically bound and cement bound materials. They can lay at depths up to 400 mm at widths up to 16 m. Pavers are fitted with powerful engines that provide traction through crawler tracks (Figure 15.5) or pneumatic tyres (Figure 15.6).

Crawler tracked pavers provide a large contact area with the substrate, allowing them to achieve a high tractive effort, and making them ideal for installing materials on difficult terrain or when paving at large widths.

Wheeled pavers are much easier to transport than tracked pavers, and, with a travel speed of around 20 km/h, moving them between sites in close proximity is also an option. Compared to a tracked paver, a wheeled paver has excellent manoeuvrability, a typical turning radius being 6.5 m.

Conventional pavers are capable of laying asphalt on the inclines typically found on the UK road network up to a gradient of $20^\circ$ (36%). For specialist applications, such as dam walls or the steep banks of a racing circuit, a conventional paver will require modifications to lay asphalt either vertically

Figure 15.5 Tracked paver with integral spray bar. (Courtesy of Eurovia UK Limited)
or horizontally on the incline. Alternatively, specialist ‘slope’ pavers can be used to lay asphalt vertically on an incline of up to 32° (62.5%).

After deposition into the paver hopper at the front of the machine, slat conveyors transfer the material through a tunnel in the paver to the augers, located in front of the screed. The augers distribute the material evenly in front of the screed. The width of the augers can be adjusted to ensure uniform compaction of the material by the screed at all times.

15.5.1 Screed
The screed is the main difference distinguishing a paver from other construction machinery capable of laying asphalt (e.g. a grader). Pivoting from tow points on each side of the paver, the screed generally ‘floats’ on the uncompacted material during the paving process. The screed alters with every change in the balance of forces (e.g. as a result of greater resistance from the paving material caused by variations in its temperature and/or inconsistent material properties/composition). The layer thickness should only change as a result of an alteration in the ‘angle of attack’ of the screed or an adjustment of the height of the screed tow points, which are located on each side of the paver. By operating in this manner, irregularities in the layer being overlaid are reduced in each subsequent layer.

All screed components that are in contact with the asphalt should be heated to approximately 90°C before the operation starts. Failure to do so may result in the asphalt sticking to the underside of the screed, resulting in the
formation of strips and irregular surface texture in the laid material. In addition, the floating behaviour of the screed may be affected, resulting in irregular layer thickness.

The screed is capable of laying a wide range of transverse profile shapes in the paved layer. Adjustments allow for a positive crown profile up to +5% and a negative crown down to −2.5%. ‘M’ and ‘W’ profiles are also possible (so called because these are profiles that loosely take an ‘M’ or ‘W’ shape in cross section).

The screed also houses the compaction system, which pre-compacts the asphalt to the greatest possible extent and consequently reduces the influence of layer thickness on the amount of subsequent compaction by the roller(s). Different pre-compaction systems are available.

15.5.2 Tamping screed

Driven by an eccentric shaft, the tamper bar employs a vertical high amplitude movement at comparatively low frequencies, typically between 300 and 1800 revs/min, to assist the flow of material under the screed as well as providing optimum pre-compaction. Correct set up of the tamper to suit the layer thickness and paving speed is crucial in ensuring optimum installation. Two factors determine the rate of compaction: tamper speed and the length of the tamper stroke. Both can be set up precisely, and adjusted to match perfectly the flow of mixture, type of mixture and layer thickness.

The tamper speed can be regulated: the higher the tamper speed, the greater the compaction achieved. In addition, the length of the tamper stroke should be adjusted depending on the thickness of the layer to be paved. For thin layers or ultra-thin overlays, a short tamper stroke of 2 mm is recommended, along with a very low tamper speed. For medium layer thickness, typically binder courses and surface courses, best results are achieved with a stroke length of 4 mm. When laying thick layers (i.e. those in excess of 120 mm), a maximum tamper stroke of 7 mm will provide optimum compaction.

15.5.3 Vibrating screed

Vibrators acting on screed plates and the screed frame cause them to vibrate. This movement reduces the frictional resistance between the asphalt particles, and thus promotes particle realignment and consequently aids compaction. Like the tampers, the speed of the vibrators can be regulated, the vibrating intensity rises with an increase in vibrator speed.

Typically, for thin layers (i.e. up to 30 mm) or for asphalts having a small nominal aggregate size (up to 6 mm), a low vibrator speed between 1000 and 1200 revs/min is required. For medium layer thickness in the
range 30–100 mm, best results are achieved using a medium vibrator speed of between 1200 and 2100 revs/min. Thick layers (i.e. those in excess of 100 mm thick) require a high vibrator speed. Depending on the type of screed, the maximum vibrator speed can be up to 3000 revs/min.

15.5.4 Combination screed
Combination screeds provide the versatility of both tamping and vibratory pre-compaction, either independently or in combined mode. The additional screed weight from housing both systems also contributes to higher levels of pre-compaction being achieved. Most modern pavers are fitted with a combination screed.

15.5.5 High compaction screed
When laying thick asphalt layers, a high compaction screed may be considered. Compared to a conventional screed, these types of screed contain double tamper and vibration systems, which add to the weight of the unit. This results in higher levels of pre-compaction being achieved.

Some manufacturers also incorporate pressure bars into the rear of these types of screed. These operate at between 40 and 130 bar depending on the depth of the asphalt layer, and force the mixture down until it cannot be compressed further and the maximum density is achieved (Figure 15.7).
15.5.6 Fixed or variable paving width

Extending screeds and fixed width screeds are capable of paving widths from 0.5 to 16 m. Besides asphalt, they can handle a variety of other materials, such as concrete, crushed stone or special mixtures.

Fixed width screeds (Figure 15.8) are ideal where large paving widths are required. Their modular design permits the use of bolt-on extensions up to a maximum width of 16 m.

Due to its flexibility, the most common type of screed is the telescopic extending screed (Figure 15.9). Depending on the size of the paver, the screed width can be adjusted to suit the site conditions. The narrowest screed can typically operate at widths of between 0.5 and 2.0 m, while the widest screed can be from 2.0 up to 6.0 m. Similar to fixed width screeds, the operating widths can be extended further by the addition of bolt-on extensions (e.g. the widest screed can be extended up to 9.5 m).

Figure 15.8 Fixed screed. (Courtesy of Joseph Vögele AG)

Figure 15.9 Variable width screed. (Courtesy of Joseph Vögele AG)
15.5.7 Angle of attack/planing angle
The angle between the bottom plate of the screed and the paved layer surface is called the ‘angle of attack’ or ‘planing angle’. The level of the screed above the surface on which material is being laid determines the laid thickness. As explained previously, the screed moves until it is in a state of equilibrium. This is dictated by the angle of attack and the height of the tow points of the screed. By adjusting the height of the tow points, the state of equilibrium is affected, resulting in the depth of material being increased or decreased (Figure 15.10).

15.6 Laying in adverse weather conditions
Laying is not recommended if standing water is present on the receiving layer, as the water may form a barrier and prevent bonding between adjacent layers. If water vapour results then, as the vapour attempts to escape, it produces cavities, which impair the stiffness of the layer. It may also cause the asphalt to cool rapidly, reducing the level of pre-compaction that is effected in the asphalt. Additionally, it may have a negative effect on the behaviour of the screed.

Installation should be carried out with due regard to ambient weather conditions so that materials can be properly compacted. Wind speed has a greater effect on the rate of cooling than does ambient temperature, particularly when laying thin layers.

Furthermore, the ground temperature should also be taken into consideration, as a cold receiving layer will similarly cause the asphalt to cool more rapidly. Asphalt should not be laid on any surface that is frozen or covered with ice or snow.

Figure 15.10 Angle of attack/planing angle. (Courtesy of Joseph Vögele AG)
15.7 Machine laying

During paving, when the position of the screed tow points are stationary, an equilibrium of forces resulting from a combination of paving speed, screed weight and tamper speed is established. If any of these parameters alter, there will be an immediate effect on the floating behaviour of the screed. Accordingly, machine laying should be planned so that the asphalt layer can be installed continuously, in order to produce an even and compact surface to the required width, thickness, profile and camber or crossfall without causing defects or irregularities such as segregation, dragging and/or burning. The topography of the site should be taken into consideration when selecting the plant required to undertake the work.

It is essential that the paver is set up correctly to lay the particular asphalt being laid. The flow gates and/or the material handling conveyor should be adjusted to deliver an adequate supply of material and to provide and maintain a constant head of material in front of the screed.

The auger height and speed should be adjusted to suit the laying depths and the required rate of delivery. Typically, the lower edge of the auger flight should be set at a height above the receiving layer that is equivalent to approximately five times the nominal aggregate size of the asphalt being placed. If the auger height is excessive, the head of material in front of the screed may not extend to the outer extremities of the screed. If the auger height is inadequate, the pre-compacted material may have an open appearance, which may cause compaction deficiencies and subsequent durability issues. Extension augers should be fitted if the width of the screed is increased. This is necessary to minimise material segregation.

The screed should be checked for true alignment across its width prior to installation. Particular attention should be paid to the adjustment of the screed compaction unit, to ensure it pre-.compacts the material sufficiently and does not cause any damage to the material.

The finished surface should be continuously inspected as it is laid, and any defects rectified immediately and before any rolling takes place. A finished asphalt surface that looks good (i.e. homogeneous and properly compacted both longitudinally and laterally) is much more likely to perform properly than one of variable appearance. There should be no scattering back by hand of additional asphalt on machine-laid work where it has an open appearance. The reason for the laid asphalt appearing open should be ascertained and addressed.

It is particularly important that asphalt in the vicinity of longitudinal joints (such joints are sometimes called ‘rip joints’) is laid properly and compacted adequately. Longitudinal joint failure is probably the most common defect in asphalt highways.
15.7.1 Head of material

The height of the asphalt in front of the screed is called the head of material. The head of material should be constant across the full paving width, and it should remain so as the asphalt is laid. A constant head of material is a precondition for perfect floating behaviour of the screed.

Figure 15.11 illustrates the importance of ensuring that the head of material is correct.

(a) If the head of material is too high, then the result is a force tending to lift the screed. If the screed lifts, then the laid thickness will be greater than that required.

(b) If the head of material is too low, then the result is a force tending to push the screed down. If the screed moves down, then the laid thickness will be less than that required.

(c) This shows the screed in equilibrium, with the result that the thickness of mat laid is the required value.

Figure 15.11 The importance of the head of material
The possibility of the head of material being incorrect can be reduced by the use of strike off and limiting plates for the auger tunnel, and sensors to regulate the auger feed.

15.7.2 Paving speed

Ideally, the paver should be operated at a constant speed, as any variations may result in surface irregularities. Stoppages can also cause profile irregularities and localised imperfections within the finished asphalt layer. After a stoppage, a paver screed tends to rise when work restarts. This creates a bump in the finished surface. To counter this tendency, modern pavers are fitted with a device called a screed stop or screed freeze.

The paving speed also determines the impact of the screed’s pre-compaction system on the asphalt layer, so the speed should be set at a value that ensures optimum pre-compaction. To minimise the possibility of irregularities occurring, a small angle of attack should be adopted.

Material supply logistics and the capability of the compaction plant should also be considered when determining the paving speed.

15.7.3 Material segregation

Segregation of asphalts is the separation of the larger particles of coated aggregate in the mixture (i.e. the coarser fractions in the material). A compacted asphalt has to be homogeneous if it is to perform as expected. Segregation tends to be prevalent in asphalts of larger nominal sizes, which is why 40 mm nominal size asphalts were dropped from the UK asphalt production standard BS 4987-1:2003 (BSI, 2003). With 40 mm mixtures, it was always a concern that segregation would occur during loading and transportation. In the case of the latter, this was due to vibration of the load as it was transported. If segregation occurs in the hopper of the paver, the solution is to ensure the conveyors are covered with homogeneous material when the hopper sides are folded in (hopper sides are folded in from time to time during laying by the paver operator to move asphalt at the sides of the hopper into the area where the slats will convey the material through the tunnel to the augers).

If segregation occurs in front of the screed, it may be possible to rectify the issue by adjusting the auger height. According to the Vögele Booklet on Paving (Joseph Vögele AG, 2012), if, after adjustment, there is no improvement, additional smaller or different auger blades may be required. In these circumstances, the auger needs to rotate more quickly or more continuously, resulting in the asphalt being mixed more effectively.

If segregation occurs around the centre of the auger box (the auger box is the volume around the vicinity of the auger), the screed should be moved further
back in order to increase the head of the material in front of the screed. Additionally, the height of the auger should be increased, and one or two auger blades in the central area of the auger box should be turned around to cause a reverse flow, which should ensure that all the asphalt fractions are conveyed behind the auger in the vicinity of its centre.

15.8 Hot-on-hot paving

Two layer construction of asphalt pavements by hot-on-hot paving is an efficient method of constructing durable roads. This method consists of simultaneously installing the binder course and surface course layers hot-on-hot (i.e. in a single pass), with both layers being compacted together. This approach results in perfectly bonded layers with a strong interlock between the courses, without the use of a bond coat or tack coat (Figures 15.12 and 15.13).

The process requires simultaneous supply of both materials, usually from two asphalt plants. On site, the materials are alternately fed into separate hoppers by a material transfer vehicle before being individually transferred and installed by two floating screeds running one behind the other. A high compaction screed is used to install the binder course, resulting in densities of 98% being achieved. Due to this high level of pre-compaction on the binder course, only a medium weight tandem roller is required to achieve the final density of the installed layers.

15.9 Hand laying

Hand laying is normally only permissible where machine laying is impractical or where small quantities of asphalt are being laid. Hand laying will

Figure 15.12 InLine Paving® system. (Courtesy of Joseph Vögele AG)
never achieve the quality of finished pavement produced using a paver. Thus hand laying should be adopted only where there is no other option. When areas of handwork are unavoidable, installation should be undertaken quickly and efficiently, with the compaction plant being deployed as promptly as is safely possible.

The asphalt should be spread in a loose layer of uniform thickness and even texture, and thoroughly compacted immediately. Every precaution should be taken to minimise segregation and to avoid contamination. An inspection should be carried out after the initial rolling, with any rugous areas made good by carefully applying small quantities of hot material, commonly described as ‘spotting up’.

15.9.1 Patching and repairs
As is always the case, the asphalt should continue to be protected from heat loss and the effects of adverse weather conditions. The material should be taken directly from the delivery vehicle, which should be kept sheeted for as long as possible. It should, preferably, be protected in a portable hot box.

As a means of ensuring reasonable riding quality (specifications often describe this aspect as ‘surface regularity’) after patching operations, the surface of any compacted patch should be flush with or slightly higher than the surrounding material. The finished surface should not be left below the surrounding surface level, as ponding may occur, and such depressions are more noticeable to vehicle drivers. (A slight bump upwards in the direction of travel is always less apparent to vehicle occupants than a drop down.)
15.10 Layer thickness and surface regularity

The nominal thickness of an asphalt layer can usually be found within the relevant transport, laying and compaction specifications. Typically, this is equivalent to multiplying the nominal aggregate size of the asphalt by around 2.5 or 3. However, there are exceptions to this rule. In the UK, the nominal and minimum compacted layer thickness for asphalts is specified in Clause 6.6 of BS 594987:2010 (BSI, 2010b).

Excessive thickness can lead to instability, causing surface regularity and level control problems. As a result, poor deformation resistance can be a potential long term issue in binder courses and surface courses of thicker pavements (total asphalt ≥180 mm).

Insufficient thickness can lead to aggregate damage (fracturing) within the asphalt, making the layer difficult to compact. As a result, the laid material may have poor durability. Such difficulties are often encountered when tying into an existing surface at the ends of an area of new construction or maintenance where the depth of asphalt laid falls below what is an acceptable thickness if the material is to be properly compacted. The solution is to ensure that the existing surface is milled out to an adequate depth to avoid this possibility.

Surface regularity (often described as ‘rideability’) can affect the durability of an asphalt pavement. The finished road profile is extremely important, because it is the main factor controlling ride quality, and hence user perception of the road condition. Indeed, many clients will judge the quality of a finished pavement solely on the quality of the ride. A sudden change in profile over a short length will significantly reduce the ride quality of a pavement. The profile of a road can also influence vehicle interaction. When a vehicle is moving along a road, unevenness will cause the vehicle to move up and down. In turn, this will cause a dynamic variation in the loading on the pavement, above and below their static values. This dynamic effect of road unevenness may accelerate the deterioration of the pavement, and its effect should not be underestimated.

The accuracy and quality of the surface regularity can be improved by using mechanical and electronic levelling devices, which are available for many modern pavers. These vary in design, and include devices described as contactable sensors, single cell sensors, sonic grade non-contacting sensors, variable length multiplex skis, averaging beams and laser receivers. All these systems detect level changes in the receiving layer and, depending on the readings, adjust the depth of asphalt laid by the paver accordingly. Each type of device can be used alone, on either side of the paver, or on both sides of the paver simultaneously (Figure 15.14). Another option for larger projects is the use of three dimensional positioning systems. These use a global
positioning system and specific digital design data for the site and automatically control the gradient, slope, paving width and direction of the paver.

### 15.11 Joints

Pavement durability can be extended by minimising the number of longitudinal and transverse joints. Thus, careful planning of the paving works should always be undertaken. Where possible, longitudinal joints should not be located within wheel tracks: wheel tracks are typically centred around 0.9 and 3 m from the nearside edge of the carriageway. In the UK, the wheel track zones are defined in Clause 903.21 of the *Specification for Highway Works* (Highways Agency *et al.*, 2009), which states that they ‘shall be taken to be between 0.5 m and 1.1 m and between 2.55 m and 3.15 m from the centre of the nearside lane markings for each traffic lane (or, in the absence of lane markings, lane edges)’.

Transverse joints typically occur as a consequence of interrupted paving work. Such joints create a localised weakness within the pavement, resulting in a reduced service life as well as having a detrimental effect on ride quality. Accordingly, everything should be done to keep transverse joints to a minimum. On multi-layer pavements, the joints between the individual layers should be offset generally by 300 mm, or whatever is practicable if the applicable specification allows, from the joints in the course below. This minimises the possibility of water ingress throughout the pavement depth. It should always be remembered that water and pavements simply do not mix, and every possible step must be taken to prevent water getting into the pavement.
This is why the faces of kerbs should be painted, preferably with 40/60 penetration grade bitumen, although cold-applied joint paint or polymer modified emulsion may be permissible. This is essential to prevent rainwater, shed as a result of the camber, crossfall or superelevation of the carriageway into the channel, running down the face of the kerb and into the pavement.

Where joints have to be formed, it is imperative that care and consideration is given to their formation and compaction.

All transverse joints that occur when tying into adjacent surfacing or the day joint in new surfacing should be cut using a floor saw to provide a good vertical edge. The exposed edge should be thoroughly cleaned, and all loose material removed and discarded, before being sealed with an application of hot bitumen, cold applied joint paint or polymer modified emulsion, which should coat the entirety of the exposed vertical face. As above, many specifications understandably stipulate hot bitumen for joints. This bitumen seal is intended to ensure that the interface between adjacent layers is not permeable. For the same reason, transverse joints in base and binder course layers should be sealed, preferably with hot bitumen.

It is deemed best practice to form all exposed longitudinal joints. Indeed, many specifications will require this in future. This is most readily undertaken
by either using a chamfered cutting wheel, attached to the roller to form an edge at an angle of 45–60° (Figure 15.15). Alternatively, the longitudinal joint can be formed by cutting back the edge of the layer using a cutting wheel attached to the roller (Figure 15.16). This forms a near vertical face that exposes the full thickness of the layer.

Once the longitudinal joint has been formed, as with transverse joints the exposed edge should be thoroughly cleaned, and all loose material removed and discarded, before being sealed with an application of hot paving grade bitumen, cold-applied joint paint or polymer modified emulsion. It is important that the coat is applied to the entire exposed face. On base and binder course layers, the sealing of the joint at the surface with an additional application of a bitumen seal is also essential. As before, the use of properly applied penetration grade bitumen, either 40/60 or 100/150, is the most effective way of ensuring a waterproof seal.

It is not uncommon in car parks, including those at shopping centres, for joints to be left devoid of any bitumen seal. This is poor practice. All asphalt joints must be properly constructed if the asphalt is to last as long as possible, and this applies as much to a car park as it does to a highway.

When installing new asphalt layers adjacent to existing cold asphalt, the new layer must be thicker (surcharge) than the adjacent surface by the reduction in
thickness that will occur as a result of the hot asphalt being compacted. This is essential to avoid the barrel of the roller being supported by the cold asphalt, resulting in the hot asphalt being compacted to a lesser degree than is required. In addition, the end plate of the screed should overlap the cold asphalt layer by 20–30 mm. Subsequently, the overlapped material should be manually pushed back into the area of the joint on the new asphalt layer prior to compaction by the roller. This procedure should not require any additional material, and keeps the required level of handwork to a minimum. This reduces the risk of material segregation and consequently improves the level of compaction at the joint (Figure 15.17). If the overlap is too narrow, then poor compaction at the joint will result. Alternatively, if the overlap is too wide, the paver will ride on the existing asphalt, causing damage to the cold material and resulting in poor compaction of the adjacent hot asphalt at the joint. This is a very common occurrence in asphalt surfacing.

Laying with two or more pavers working in echelon produces a hot-to-hot integral bond between the paved asphalt materials without the need to form a longitudinal joint (Figure 15.18). Elimination of the longitudinal joint is a very attractive proposition. However, as can be seen in Figure 15.18, the pavers cannot operate side by side. In these circumstances, the distance between the individual pavers should be minimised so that the joint face is still sufficiently hot to form a perfect bond with the adjacent asphalt.

Another way of hot jointing surface courses is to reheat the edge of the cold lane and bring it up to a plastic state using a joint heater prior to the new,
adjacent hot asphalt being laid. The heater must raise the temperature of the full depth of the surface course to within the specified range of minimum and maximum rolling temperatures for a width of not less than 75 mm. Care must be taken to ensure that the heated asphalt is not damaged in the process, as this would result in premature ageing.

15.12 Chipping of hot rolled asphalt surface course

In order for the finished surface to have an acceptable level of skid resistance, hot rolled asphalt surface course mixtures containing 35% or less nominal coarse aggregate have coated chippings rolled into them. This is achieved by the addition of 14/20 or 8/14 mm aggregate coated with not less than 1.5% by mass of 30/45 or 40/60 penetration grade bitumen. These are placed at a specified rate of spread on top of the newly laid asphalt, and then rolled into the asphalt using the compaction plant. Chippings of size 14/20 mm always have a superior appearance and result in less plucking than 8/14 mm chippings, and thus 14/20 mm chippings should be used wherever circumstances permit.

After initial compaction by the paver or, in the case of hand application, prior to the first pass of the roller, the installed asphalt surface should be covered with a uniform layer of coated chippings. Where possible, this operation should be undertaken using a mechanical chipping spreader (Figure 15.19). The chipped surface is then rolled to embed the chippings and compact the material. When spreading chippings on roads, an
unchipped channel not less than 150 mm wide should be left adjacent to the kerb to facilitate the rapid flow of surface water into the gullies.

The degree of embedment and adhesion of the coated chippings is adversely affected by surface chilling of the asphalt layer. This occurs rapidly in windy conditions, even more so when combined with a low ambient temperature. Accordingly, all steps should be taken to ensure that embedment of the chippings and compaction of the layer is completed before the material reaches the relevant minimum rolling temperature.

15.13 Installing gussasphalt

Gussasphalt is a type of mastic asphalt, and consists of aggregate and high proportions of bitumen and limestone filler. The aggregate takes the form of sand or crushed rock fines. Typically, it contains between 6.8% and 7.5% binder. Once placed, the finished layer is impervious.

Gussasphalt is delivered in mobile cookers that maintain the material temperature between 200 and 260 °C, while also agitating the mixture in order to retain its workability and stability. Differences in the temperature and thickness of installed layers significantly affect the quality of the finished layer, so uniformity in both aspects is essential.

The paver, or as it is more typically known the ‘finisher’ (Figure 15.20), is fitted with a powerful engine that provides traction through crawler tracks,
rail wheels or pneumatic tyres. Pavers are available in varying operational widths, typically between 1.0 and 14.5 m, and consist of a heated non-vibrating screed bar and a material distributor. Typical laying speeds range between 0.25 and 3.5 m/min.

Gussasphalt is typically laid at a depth between 20 and 40 mm. Due to its low viscosity, it must be laid between two restrained edges. These can be in the form of an adjacent existing layer of gussasphalt or other asphalt, a pre-constructed strip of gussasphalt or rails on which some finishers run. As a result, the restraining edges determine the laying depth.

The gussasphalt is deposited onto the substrate directly in front of the finisher. Subsequently, the finisher distributes the material across the width of a heated blade, which levels the surface to the pre-determined depth by travelling over it. The low viscosity of gussasphalt makes it self-levelling, so no vibration is required.

When installed as a surface course, gussasphalt is immediately gritted, either manually or mechanically, with aggregate coated with bitumen. The grit is subsequently embedded in the surface using a roller.

In common with other asphalts, gussasphalt should not be installed on a wet substrate, as the high installation temperatures and the impervious properties of the material trap the moisture beneath the layer, causing bubbles to appear on the surface.
15.14 Compaction

In a surfacing operation there is a myriad of activities that are undertaken before, during and after the asphalt has been laid. These range from cutting joints to the final rolling. Each of these elements is important in producing a finished road surface that will give many years of satisfactory service. However, compaction is perhaps the single most important activity in dictating the lifespan of a pavement. Accordingly, it is very important that all personnel involved ensure that compaction is carried out effectively and efficiently.

Compaction is the action of removing air from an asphalt, and is effected by rollers traversing the material, usually several times (called passes). Asphalt must be compacted in such a way that the particle displacement reduces the void content of the asphalt layer. Asphalts that are properly compacted will dissipate loads from traffic more quickly because compaction increases the stiffness of the mixture. High levels of compaction also reduce the rate of ageing of the binder, and, in doing so, prolong the life of the asphalt.

Compaction can be undertaken using two methods, or a combination of both

- static compaction
- dynamic compaction.

Static compaction uses the weight of the roller to exert downward pressure on the uncompacted material. Weight only operates vertically. This pressure causes the internal friction in the asphalt to be overcome, resulting in the mineral particles displacing air in the mixture and, as a consequence, reducing the void content.

The depth of the layer affected by static compaction is comparatively shallow. The higher the static linear load, the greater the depth to which static compaction has an effect. However, caution is warranted, as excessive linear loading may subsequently result in cracking and distortion in the layer being compacted.

Dynamic compaction penetrates to a much greater depth than static compaction. It is a more efficient compaction mode than static compaction. Eccentric weights cause the roller drum to vibrate, resulting in vibrations being transferred to the individual particles in the asphalt. This overcomes the frictional resistance between the particles, resulting in particle displacement. The effect of the vibrations together with the static load of the roller means that very high compaction can be achieved.

Some roller manufacturers now offer rollers that are capable of a particular variant of vibratory mode described as ‘oscillation’. When a roller operates
in oscillatory mode, the drum is in contact with the asphalt at all times, unlike normal vibratory rollers. Manufacturers claim that oscillation rolling only produces about 10% of the vibration-related stresses on the surrounding environment compared to those typically generated by a standard vibrating roller. This attribute makes their use more appropriate in situations where close control is required (e.g. in the vicinity of structures).

The amplitude and frequency at which rollers operate is important when compacting materials, including asphalt, in vibratory mode. The frequency is the rate at which the drum impacts the asphalt and is measured in hertz (Hz). The amplitude is the distance the drum moves from its axis, and is usually measured in millimetres. In the case of asphalt, the frequency would typically range from 50 to 70 Hz, while the amplitude will often be between 0.25 and 0.8 mm (Hamm AG, 2008). These parameters may be varied during compaction as the mixture densifies. It is important that the values of these parameters are correct in order to avoid problems that range from under-compaction to over-compaction, with the latter resulting in aggregate particles being crushed. Both under-compaction and over-compaction may well result in the asphalt having a shorter service life.

15.14.1 Types of rollers

The following roller types are typically used in compacting asphalts:

- three wheeled static
- pneumatic tyred
- tandem.

Three wheeled rollers (Figure 15.21) compact asphalt layers through high static loading. The basis of this is a combination of their heavy operating weight, typically between 10.6 and 13.2 tonnes and the narrow width of the steel drums. These rollers are particularly suitable for levelling the asphalt surface and operating on sites where dynamic compaction may cause damage to the surrounding environment.

Pneumatic tyred rollers (Figure 15.22) achieve high levels of compaction through a combination of vertical loading from their heavy operating weight, typically between 9.5 and 30 tonnes, and a kneading effect caused by the wheels and, more specifically, their tyre pressure. This can be adjusted, within a range of 2 to 8 bar, to provide an even tyre-to-surface contact area. Curved tyre profiles should be avoided. These rollers are typically used in combination with a steel drum roller on asphalt possessing low stability properties, the latter being used purely to obtain an acceptable surface evenness on the compacted asphalt.

Tandem rollers (Figure 15.23) are the most common type of roller used for the compaction of asphalt. They comprise two steel drums, and are available
with operating weights of between 1.5 and 18 tonnes and corresponding working widths of between 0.8 and 2.14 m. They are fitted with either vibratory or oscillatory compaction modes, as well as having the capacity to operate as a deadweight roller. Additional options of articulated or pivot steering and all-wheel drives make these roller types very versatile.
Combination rollers (Figure 15.24) have a row of rubber wheels mounted on one axle and a smooth drum mounted on the other. As a consequence, these rollers combine the advantages of both a pneumatic tyred and tandem roller in one machine.
15.14.2 Intelligent rollers
Compaction quality can be monitored and controlled more effectively by using rollers equipped with sensors. These sensors count each roller pass, plot the position of each pass and measure the surface temperature of the asphalt. All these data are available to the operator during compaction, and can be downloaded on completion of the work.

The advantages of intelligent rollers are
- uniform compaction
- detection of soft/weak spots
- optimising the number of passes
- efficient compaction
- potential cost saving.

15.14.3 Compaction of machine laid asphalt
The compaction of each layer should be carried out to a predetermined rolling pattern until the required density is achieved, and, in the case of surface courses, the layer has a homogeneous appearance.

As well as the size and weight of the roller(s) being sufficient to undertake the work effectively, the topography of the site should also be taken into consideration, so that the work is completed efficiently and safely. Smaller rollers should be considered for sites with restricted access, limited working area, restricted width of surfacing or where the underlying construction will not support a heavier roller.

The number of rollers available for use on a particular site should be such that, while travelling at a low but steady speed, sufficient passes can be made in order to compact the asphalt adequately in the time available. It is important to ensure that the frequency and amplitude of vibration or oscillation, along with the speed of travel of the rollers, are correctly matched to layer thickness and mixture composition.

Compaction should start without delay, and the rollers should operate as close as is safely possible to the paver. When using a tandem roller, compaction typically starts with an initial static compaction pass before the main compaction process is undertaken with vibration or oscillation. Care should be taken, as excessive vibration on a cooled material may fracture the aggregate. The final pass is typically undertaken using static compaction. To ensure that there are no uncompacted areas, the rolled tracks should overlap by at least 150 mm. Unsupported edges cannot be compacted. Accordingly, the edge should be either cut-back, thus removing the inadequately compacted material, or the edge should be supported during compaction. This can be achieved using a temporary restraint or an edge compactor fitted to the side.
of the roller, which compacts the asphalt leaving the edge sloping at 45–60°
(see Figure 15.15).

There are many variables that affect the time available for compaction. One
important parameter is the thickness of the layer being compacted. Figure 15.25 shows, in effect, the time available for compaction of two
layers (20 and 150 mm thick). If the minimum temperature at which compac-
tion can be effected is 80°C, then compaction has to be completed within
3 minutes of the 20 mm layer being placed, while compaction can still be
effective after 30 minutes in the thicker layer. Thus, layer thickness is a very
important factor in determining the time available for compaction (Hunter,
1986).

**Figure 15.25** Cooling time in asphalt of (a) 20 mm thickness and (b) 150 mm
thickness. (Courtesy of Dr Robert N Hunter)
Another crucial parameter is the speed of wind acting on the surface, as wind takes heat by convection from the hot asphalt. Convection losses are much more significant than the air temperature. The safest option is always to compact as quickly as possible. If the material moves unduly because it is too hot to be compacted, then the rollers can wait until the material has cooled sufficiently to permit compaction without damaging the surface. However, once a material has cooled to a point where further compactive effort is ineffective, nothing can be done to correct that shortcoming.

Due to the number of variables involved (up to 22), it is impossible to generalise about the best combination of rolling and roller pattern to use in all scenarios. There is no substitute for proper training and experience, combined with advice and support from roller manufacturers.

15.14.4 Compaction of hand laid and patching work
In areas where it is impractical to operate a large roller, such as footpaths, restricted access sites and limited working areas, either a 2.5 tonne static deadweight roller or vibrating roller having a minimum static weight of 750 kg would typically be used. The procedure adopted should closely replicate that used for larger scale work. However, more attention may be required with regard to material and rolling temperatures.

15.15 Application of grit to surface courses
Some contracts stipulate that the finished surface course should be gritted to improve initial skid resistance. This involves accurately applying a grit, which may be uncoated or coated with bitumen, at a specified rate of spread to the surface by way of a rear-mounted spreader fitted to a tandem roller. Grit application is undertaken after an initial compaction pass by the roller. The rolling pattern should, as far as is practicable, provide a single application of grit to the full width, with no overlap.

The rate of spread for the grit is typically no less than 600 g/m². The rate of spread should be automatically adjusted to take account of the roller speed, and should stop automatically when the roller halts or reverses.

Once the asphalt surface has reached ambient temperature, any surplus grit should be carefully removed using a mechanical sweeper prior to the application of road markings and/or being opened to traffic.

15.16 Opening to traffic
In order to avoid damage to newly laid sections of asphalt, it should not be overlaid or opened to traffic until the surface temperature has fallen to 40°C.
**15.17 Specification and quality control**

Compaction should be continuously assessed using an indirect density gauge, with readings taken at regular intervals in alternate wheel tracks. The gauge should be initially calibrated, and subsequently rechecked and updated based on correlations between gauge readings and core densities at the same locations. For each location, the in situ void content can be determined using the bulk density from the gauge reading compared with the maximum theoretical value ascertained in a laboratory. The average in situ void content should be calculated from any six consecutive indirect gauge readings.

Tests on pairs of cores can also be used to determine the achieved density. Cores have the additional advantage that they can be used to assess the void content at points below which the density gauge is effective. Cores are typically taken from each wheel track. In addition, a pair can be taken at a location centred 100 mm from the final joint position at any unsupported edge.

Other tests include longitudinal regularity and transverse regularity. On roads with crossfalls, this can be determined by measuring the depth of the depressions under a 3 m straight edge placed parallel to or at right angles to the centre line of the carriageway.

The most common method of determining the in situ texture depth of surface course materials is by means of the volumetric patch test, which involves taking ten individual measurements on a diagonal line across the lane width at a spacing of 5 m (BSI, 2010a). Although many UK authorities rely heavily on this parameter, published work suggests that such faith may well be misplaced (Hunter, 1996).

**References**


The properties of asphalts need to be known for a variety of reasons, including performance evaluation, mixture or pavement designs, and production and/or construction specification compliance. In situ testing of material properties in full scale trial sections or in-service pavements is impractical or uneconomical in most cases, so engineers generally have to rely on laboratory testing to characterise or predict material properties. In addition, testing may also be necessary to ensure that specified requirements are met.

Laboratory tests should reproduce the anticipated in situ conditions as closely as possible (i.e. in terms of temperature, loading time, stress conditions, degree of compaction etc.). However, in situ conditions may be the subject of change, and selection of appropriate testing conditions may be difficult. Figure 16.1 (Pell, 1988) shows a simplified model depicting a representation of a pavement element, and shows the stresses to which it is subjected when a wheel load approaches. In practice, the stresses are applied three dimensionally: horizontal and shear stresses occur in planes that are perpendicular to those shown in Figure 16.1. As the wheel passes over the element, these stresses change with time, and this is shown in Figure 16.2 (Pell, 1988). Given the difficulty of reproducing such complex stress regimes accurately in the laboratory, simplified tests have been introduced that can reproduce certain aspects of the in situ behaviour. Such tests are also used to correlate the laboratory mixture design with in situ performance in relation...
**Figure 16.1** Stresses induced on a typical pavement element under an approaching wheel load

**Figure 16.2** Stress state changes induced by a moving wheel load
to aspects such as moisture susceptibility, resistance to deformation, stiffness, fatigue and tensile strength.

Many laboratory tests have been proposed over the years. Efforts to standardise particular tests for routine usage are ongoing, and over time these have been improved by advances in equipment technology that measures material behaviour more accurately. In general, laboratory tests can be divided into the following three groups:

- **Fundamental**
  - static creep test
  - repeated load test
  - dynamic stiffness and fatigue tests
  - indirect tensile tests.

- **Simulative**
  - wheel tracking tests
  - gyratory compaction
  - durability
  - cracking.

- **Empirical**
  - Marshall
  - indirect tensile strength.

### 16.1 Fundamental tests

#### 16.1.1 Static creep test

In the 1970s, Shell developed a simple creep test whereby a static uniaxial compressive load was applied to unconfined cylindrical samples of asphalt to assess the resistance to deformation of the material (Hills, 1973). This test gained wide acceptance due to the ease of specimen preparation, the simplicity of the test procedure and the low cost of test equipment. The only requirements for the test specimen are that it should be prismatic, with flat and parallel ends normal to the axis of the specimen, while the test procedure involves the application of a constant axial stress of up to 500 kPa to a 100 mm dia test specimen for up to 10 000 s at a constant temperature, and measurement of the resultant deformation, as described in a now withdrawn British Standard (BSI, 1995).

The obvious limitation of the simple creep test is that the static loading mechanism does not simulate the dynamic traffic loading conditions to which asphalts are subjected in service, and it was found that rut prediction based on this test underestimated rut depths measured in trial pavements (Hills et al., 1974). Furthermore, static loading could not capture the improved performance of binder modifiers that enhanced the elastic recovery properties of an
asphalt, whereas, in contrast, this could be demonstrated under repeated loading (Valkering et al., 1990).

### 16.1.2 Repeated load tests

Repeated loading more closely simulates actual traffic loading. In the repeated load axial test, an unconfined test specimen is subjected to repeated block pulse loadings of 1 s duration separated by 1 s duration rest periods for up to 10,000 pulses, as described in a now withdrawn British Standard (BSI, 1996). A European Standard, EN 12697-25:2005 (BSI, 2005b), improved the position with the application of a confining pressure to the test specimen. This allows test conditions to reproduce more closely the stress conditions in an in-service pavement, and has the advantage that the applied loading is closer to the levels predicted in the pavement.

EN 12697-25:2005 (BSI, 2005b) describes two methods to achieve the confinement. In the first method, the cylindrical test specimen of 150 mm diameter is larger than the loading platens of 100 mm diameter, so the confinement is provided by the material along the outer cylindrical surface of the test specimen. In the second method, the confining pressure is exerted by a gaseous or liquid medium held in a triaxial cell, which makes the test equipment more expensive and test procedures more complex. A cyclic block pulse load can be used for both methods, and an alternative option to apply a sinusoidal compression (i.e. haversine, which is always positive) is also included for the second method with the triaxial cell. The results are typically reported as a cumulative strain curve, as shown in Figure 16.3. The load repetition number at which there is a turning point or an inflection point between the secondary flow and tertiary flow stages is identified as the flow number, which can be correlated with rutting potential.

In the USA, the National Cooperative Highway Research Program (NCHRP) developed a simple performance procedure using compact servo-hydraulic test equipment with a built-in test chamber capable of controlling temperature and confining pressure (Bonaquist, 2008a, 2008b, 2011; Bonaquist et al., 2003; Witczak et al., 2002). Renamed the asphalt mixture performance tester (AMPT), this equipment (Figure 16.4) can be used to evaluate the rut resistance of an asphalt by subjecting a cylindrical test specimen of 100 mm diameter and 150 mm height to a repeated haversine axial compressive load pulse of 0.1 s every 1.0 s at a specified test temperature to find the flow number that can serve as the performance parameter to complement the Superpave mix design method (AASHTO, 2013b; FHWA, 2013).
Figure 16.3 Typical cumulative creep curve

Figure 16.4 Asphalt mixture performance tester set up
16.1.3 Dynamic tests

Dynamic tests are more complex than repeated load tests in terms of the loading cycles and frequencies – dynamic tests apply repeated cyclic loads (usually a haversine wave) over a range of frequencies. This necessitates more accurate load application and deformation measuring systems.

Several dynamic test methodologies have been developed to determine stiffness and fatigue resistance. Bending tests using beams or cantilevers subjected to repeated applications of load have been used since the 1970s (Bonnaure et al., 1977; Brown, 1983; Cooper and Pell, 1974; Monismith et al., 1985). In such tests, the maximum stress occurs at a point on the surface of the specimen, and its calculation, using the standard beam bending formula, depends on the assumption of linear elasticity. The European Standards (BSI, 2012a, 2012b) have brought together various dynamic fatigue resistance and stiffness test methodologies, including the flexural bending tests as shown in Table 16.1.

In the two-point cantilever bending test, trapezoidal and square prismatic sample types are both included in the European standards (BSI, 2012a, 2012b), and different maximum aggregate sizes of the asphalts are also considered in the test sample dimensions. The base end of the sample is glued, and thus held rigidly while a sinusoidal load is applied at the head. For stiffness modulus measurements, the applied force or deflection at defined temperatures and frequencies should only cause up to 50 microstrain in the most heavily stressed part of the test sample so as to remain within the linear range. The load is applied for a minimum duration of 30 s and a maximum duration of 2 min, while the force, deflection and phase angles are measured and recorded over the last 10 s of the test. The test can be repeated for at least four temperatures at 10°C intervals and three

<table>
<thead>
<tr>
<th>Test mode</th>
<th>Fatigue resistance</th>
<th>Stiffness</th>
</tr>
</thead>
<tbody>
<tr>
<td>Two-point bending</td>
<td>✔ EN 12697-24</td>
<td>✔ EN 12697-26</td>
</tr>
<tr>
<td>Three-point bending</td>
<td>✔ EN 12697-24</td>
<td>✔ EN 12697-26</td>
</tr>
<tr>
<td>Four-point bending</td>
<td>✔ EN 12697-24</td>
<td>✔ EN 12697-26</td>
</tr>
</tbody>
</table>

* EN 12697-24 (BSI, 2012a); EN 12697-26 (BSI, 2012b); ASTM D7460-10 (ASTM, 2010)
frequencies at each temperature, in order to determine the master curve. For fatigue measurements, the European standards specify loading under constant displacement or strain for the trapezoidal samples, with at least one-third of the test set of 18 specimens able to reach one million test cycles, while the prismatic samples are tested under constant stress to a displacement of 280 μm. The French mixture design method has five levels that are chosen according to the type of asphalt, loadings and intended use, with additional tests required as the level increases. The two highest levels require fundamental tests for stiffness and fatigue using the two-point cantilever bending test (Delorme et al., 2007). Figure 16.5 illustrates the two-point cantilever bending test set up with a trapezoidal sample.

For the three-point and four-point bending tests, a beam is subjected to periodic bending through vertical movements in the central load point(s), while the vertical positions of the two end points are kept fixed. Free rotation and horizontal movement are allowed at all load and reaction points in order to prevent the development of horizontal and torque stresses that can affect the behaviour of the material during the test. The sample width and height should be at least three times the maximum aggregate size of the asphalt, while the effective sample length should be at least six times the width or height. The European standard, EN 12697-24:2012 (BSI,
The various fatigue tests are carried out until ‘failure’ occurs in the test specimen. ‘Failure’ can be an arbitrary end point, not where the test specimen literally fails. In a constant-strain test, the sample is usually deemed to have ‘failed’ when the load required to maintain that level of strain has fallen to 50% of its initial value. Because of the scatter of test results associated with fatigue testing, it is normal to test several specimens at each stress or strain level, and to plot the results plotted as stress or strain against cycles.
to failure on a log-log graph, as shown in Figure 16.8 (Read, 1996). In EN 12697-24:2012, a minimum of 18 samples should be tested, generally over at least three selected levels of loading, while ASTM D7460-10 requires nine replicate samples for a complete fatigue curve.

The AMPT described in section 16.1.2 can also be used to apply controlled sinusoidal compressive stresses to determine the dynamic modulus. The test is undertaken at multiple temperatures and loading frequencies because these factors have a direct impact on material stiffness – the dynamic modulus values decrease with higher temperatures and lower loading frequencies. The multiple test results are then used to develop master curves to be input into mechanical–empirical pavement design methods. The American Association of State Highway and Transportation Officials (AASHTO) has
a provisional standard test method, TP 79-13 (AASHTO, 2013b), and a standard practice, PP 61-13 (AASHTO, 2013a), for determining the dynamic modulus and developing the master curve, respectively, while work is in progress to develop and standardise a direct tension test using the AMPT to assess fatigue and top down cracking resistance (Christensen and Bonaquist, 2009; Hou et al., 2010).

### 16.1.4 Indirect tensile tests

In addition to the uniaxial and triaxial compression tests and flexural bending tests described in the previous sections, indirect tensile tests have also been developed to evaluate the fundamental stiffness and fatigue properties. In such tests, a repeated loading is applied in the vertical diametrical plane of a cylindrical specimen using a loading strip, as shown in Figure 16.9. This vertical loading produces both a vertical compressive stress and a horizontal tensile stress on cylinders of the specimen. The magnitudes of the stresses vary across the cylinder, as shown in Figure 16.10, but are at a maximum in the centre of the specimen.

The situation, as depicted in Figure 16.10, enables the calculation of strain based on the following assumptions

- the specimen is subjected to plane stress conditions ($\sigma_z = 0$)
- the material behaves in a linear elastic manner
- the material is homogeneous
Figure 16.9 Indirect tensile stiffness modulus test set up

Figure 16.10 Stress distributions in the indirect tensile test mode
the material behaves in an isotropic manner
Poisson’s ratio for the material is known
the force is applied in the vertical diametrical plane.

16.1.4.1 Indirect tensile stiffness modulus
The indirect tensile stiffness modulus (ITSM) test, as defined in the European standard EN 12697-26:2012 (BSI, 2012b), is a simple test that can be completed quickly. The operator selects the target horizontal deformation and a target load pulse rise time (the time from the start of load application to the peak load). The force applied to the specimen is then automatically calculated by a computer, and a number of conditioning pulses are applied to the specimen. These conditioning pulses are used to make any minor adjustments to the magnitude of the force needed to generate the specified horizontal deformation and to seat the loading strips correctly on the specimen. Once the conditioning pulses have been completed, the system applies five load pulses. This generates an indirect movement on the horizontal diameter and, as the diameter of the specimen is known beforehand, the strain can be calculated. As the cross-sectional area of the specimen is also known and the force applied is measured, the applied stress can be calculated. Thus, as the stress and the strain are now known, the stiffness modulus of the material can be calculated.

Standard test conditions and requirements for the ITSM test are

- peak horizontal deformation of at least 5 μm
- rise time 124 ms – equivalent to a frequency of 1.33 Hz
- specimen diameter 80, 100, 120, 150 or 200 mm
- specimen thickness between 30 and 75 mm.

As material stiffness depends on the test frequency and temperature, EN 12697-26:2012 (BSI, 2012b) also defines a cyclic indirect tension test to be done at multiple temperatures and loading frequencies. A sinusoidal compressive loading is applied, without rest periods, under the following test conditions

- initial horizontal strain of up to 0.1%
- load frequency range 0.1–10 Hz
- specimen diameter 100 or 150 mm
- specimen thickness 40–90 mm
- minimum of four test temperatures: −10, 0, 10 and 20°C.

16.1.4.2 Indirect tensile fatigue test
The indirect tensile fatigue test (ITFT) uses a repeated controlled stress pulse to damage the specimen, and the accumulation of horizontal deformation
against the number of load pulses is continually measured and recorded at preselected intervals.

Standard test conditions and requirements for the ITFT according to the European standard, EN 12697-24:2012 (BSI, 2012a), are

- initial strain range 100–400 μm/m
- loading amplitude: 250 kPa
- failure criterion – obvious vertical cracks or when the dynamic tensile strain increases to twice its initial value.

16.2 Simulative tests

As the stress conditions in a pavement loaded by a rolling wheel are extremely complex and cannot be replicated in a laboratory test on a sample of asphalt with any precision, simulative tests have been used to compare the performance of different materials.

16.2.1 Wheel tracking tests

Wheel tracking tests are based on the general operating procedure of tracking a load repeatedly over an asphalt sample to simulate in-service behaviour. The equipment ranges from large full scale pavement test facilities such as those at the UK’s Transportation Research Laboratory (TRL), France’s Laboratoire Central des Ponts et Chaussées (LCPC) and the USA’s National Centre for Asphalt Technology (NCAT), to mid-scale pavement test facilities to the smaller laboratory scale wheel tracking machines (Figure 16.11). Full scale and mid-scale facilities can be used to test entire pavement structures to provide inputs for mechanistic pavement analyses, whereas laboratory tests are not mechanistic but do seem to simulate field behaviour (Brown et al., 2001) so they can be employed to evaluate the relative resistance of particular asphalts to deformation.

The laboratory wheel tracking devices shown in Figure 16.11 have different sets of testing parameters and conditions. The European standard, EN 12697-22:2007 (BSI, 2007a), has brought together several wheel tracking test methodologies covering the Hamburg wheel tracking device (HWTD), the French rutting tester (FRT) and the UK wheel tracking machine WTM, while AASHTO also has test standards for the HWTD, T 324-11 (AASHTO, 2011b), and the asphalt pavement analyzer (APA), T 340-10 (AASHTO, 2010), another wheel tracking device more commonly used in the USA. Table 16.2 shows the test parameters commonly used for these four devices.

The test temperatures and initial air void contents have the greatest effects on test results (Cooley et al., 2000) – rutting increases with higher test temperatures and air void contents. In relation to the former, all four test
devices are capable of testing at a range of higher temperatures (around 60°C), at which rut resistance is the critical performance requirement for asphalts in pavements. For the latter, the initial voids contents are commonly set at 7%, which is taken as the typical as-constructed condition, or 4% to assess actual rutting due to shear failure. The specimen density and aggregate orientation are influenced by the specimen types, sizes and preparation methods. The two common specimen types (beam/slab and cylinder) provide different rut depths but generally similar mixture ranking. With beam/slab specimens, roller compaction better simulates field practice compared to vibratory or kneading plate compaction. With cylindrical
<table>
<thead>
<tr>
<th>Device</th>
<th>APA</th>
<th>HWTD</th>
<th>FRT</th>
<th>UK WTM</th>
</tr>
</thead>
<tbody>
<tr>
<td>Test temperature: °C</td>
<td>US PG high temperature level, commonly 58 or 64</td>
<td>50 or 60</td>
<td>60</td>
<td>45 or 60</td>
</tr>
<tr>
<td>Specimen size: mm</td>
<td>150 diameter × 75 thickness cylinder, or 300 length × 125 width × 75 thickness slab</td>
<td>150 diameter × 38 to 100 thickness cylinder, or 320 length × 260 width × 38 to 100 thickness slab</td>
<td>500 length × 180 width × 50 or 100 thickness beam</td>
<td>200 dia. cores of various thickness, or 300 length × 300 width × 50 thickness slab</td>
</tr>
<tr>
<td>Compaction method</td>
<td>Gyratory compactor for cylinder specimen, vibratory compactor for beam specimen, 7% voids</td>
<td>Gyratory compactor for cylinder specimen, kneading plate compactor for slab specimen, recommended 7% voids</td>
<td>Roller compactor, compacted, void content depends on mixture type, at low and high ends of field as-compacted range</td>
<td>Roller compactor for slab specimen</td>
</tr>
<tr>
<td>Type of wheel</td>
<td>Stiff rubber hose under concave steel wheel</td>
<td>Solid treadless rubber tyre (EN) or solid steel wheel (AASHTO)</td>
<td>Pneumatic tyre</td>
<td>Solid treadless rubber tyre, rectangular cross-section</td>
</tr>
<tr>
<td>Wheel dimensions: mm</td>
<td>29.5 width (external diameter of rubber hose)</td>
<td>200 diameter × 50 width</td>
<td>400 diameter × 80 width</td>
<td>200 diameter × 50 width</td>
</tr>
<tr>
<td>Applied load: N</td>
<td>445</td>
<td>700 (EN) or 705 (AASHTO)</td>
<td>5000</td>
<td>700</td>
</tr>
<tr>
<td>Number of cycles</td>
<td>8000</td>
<td>10 000</td>
<td>30 000</td>
<td>1000</td>
</tr>
<tr>
<td>Test frequency: cycles/minute</td>
<td>60</td>
<td>26.5</td>
<td>60</td>
<td>26.5</td>
</tr>
<tr>
<td>Test time: minutes</td>
<td>135</td>
<td>380</td>
<td>500</td>
<td>38</td>
</tr>
<tr>
<td>Average test speed: m/s</td>
<td>0.60</td>
<td>0.20</td>
<td>0.82</td>
<td>0.20</td>
</tr>
</tbody>
</table>
specimens, gyratory compaction is usually used to achieve lower voids contents more easily.

The loading methods, magnitudes, effective stress applied and tracking speeds will also significantly affect the results. The FRT wheel is the largest (400 mm diameter), and is fitted with an 80 mm wide pneumatic tyre that contributes to higher surface shear and which simulates actual field conditions. The APA attempts to simulate the tyre pressure effect with a metal wheel running over a pressurised hose to transmit the load, but its representation of the pneumatic rubber tyres used in real life is limited, as are the solid rubber tyres of the UK machine and the HWTD’s steel wheel. The steel wheel on the HWTD does not deform, resulting in a relatively high 0.73 MPa contact stress, and the HWTD test is also usually conducted with test samples submerged under water to assess moisture sensitivity, further increasing its severity.

The common test load applied for the FRT (5000 N) is significantly higher than that used with the other devices, but this is offset by its wider wheels and relatively high average tracking speed of 0.82 m/s. The APA has the same test frequency of 1 Hz but its average tracking speed is a lower 0.60 m/s. The average tracking speeds of the HWTD and UK machine are notably lower, and hence more severe than those of the other two devices.

Due to the variation in testing parameters and conditions, the different devices produce different results. However, comparative tests with full scale accelerated test facilities have shown that they are generally capable of assessing and ranking relative mixture performances similar to those in the field, with no clear superiority of any particular equipment. The various devices are also able to differentiate between binders of different grades (Cooley et al., 2000). As such, these devices are useful tools to highlight the contributions to improved mixture rut resistance when using modified binders.

Various transportation agencies have applied particular devices to set rut performance specifications. This requires tests to be conducted with local materials and mixtures to develop the reference results to compare against, as well as the performance criteria to be met. It should also be noted that, beyond the most basic level of the French mixture design method, a minimum rut resistance performance using the FRT is required for most mixtures (Delorme et al., 2007).

In China, the wheel tracking test equipment, sample sizes and test conditions are similar to the UK wheel tracking machine, with some modifications, such as test duration (60 versus 45 min) and analysis method – the results are
analysed as the dynamic stability (DS) value calculated using the following formula, with higher DS values representing better rut resistance at 60°C.

\[
    DS = \frac{N_{15}}{D_{60} - D_{45}}
\]

where \(N_{15}\) is the number of loading passes after 15 min (i.e. \(15 \text{ min} \times 42 \text{ passes/min}\)), and \(D_{60} - D_{45}\) is the change in rut depth in the last 15 min of the test.

**16.2.2 Gyratory compaction**

The gyratory compactor (Figure 16.12) simulates, to a reasonable degree, the compaction that actually takes place in service. The test procedure consists of placing a sample of hot asphalt in a cylindrical mould and applying a static pressure of a controlled magnitude over a round metallic insert covering the entire upper surface of the sample. The mould is then gyrated through a small angle in order to allow the aggregate particles to reorientate.

![Gyratory compactor](image)

*Figure 16.12 Gyratory compactor*
themselves under the loading. A schematic diagram of the gyratory motion is shown in Figure 16.13.

After compaction, the volumetric properties of the asphalt can be assessed against the mixture design specifications. This is a basic mixture design requirement in France (Delorme et al., 2007) and in the Superpave mixture design method used in the USA (Asphalt Institute, 2001). The test conditions commonly used are a vertical stress level of 600 kPa and a gyration speed of 30 revs/min, but the angle of gyration used in France is 1.0° (Delorme et al., 2007), whereas the Superpave gyratory compactor adopts a standard of 1.25° (ASTM, 2009) in order to achieve the higher level of densification of 4% design air voids (Harman et al., 2002).

The gyratory compactor also allows a measure of the compactability of a mixture to be assessed by monitoring the vertical movement of the loading ram and the number of gyrations of the mould. Hence, if one mixture takes fewer gyrations in comparison to another mixture for the same vertical movement of the loading ram, then the former would be said to be more easily compacted.

Additionally, slab compaction using rollers and/or kneading plates is becoming more widespread, and is specified for the preparation of test specimens for certain tests such as wheel tracking (see Table 16.2). The European standard, EN 12697-33:2007, has brought together several methodologies for test specimen preparation by roller compactor (BSI, 2007b). A photograph of this equipment is shown in Figure 16.14.
16.2.3 Durability testing
The durability of asphalts can be defined as the ability of the mixtures to resist the effects of water, ageing and temperature variations, for a given traffic loading without significant deterioration for an extended period of time. Accordingly, for durability evaluation, there is a need to be able to age artificially and/or to simulate the effect of water damage on compacted asphalt samples, as opposed to merely the components of the mixtures. A European standard, EN 12697-45:2012 (BSI, 2012d), specifies a saturation ageing tensile stiffness conditioning regime to age asphalts in the presence of water, while the comparison of the indirect tensile strength (AASHTO, 2011a) before and after a water-conditioning regime is one of the methods commonly used to assess moisture susceptibility. This test and other durability related moisture resistance tests are covered in section 19.4.4.

16.2.4 Low temperature cracking
The cracking of asphalt pavements in winter, or thermal cracking, is a common mode of distress in asphalt pavements in cold climates. While the asphalt binder is the main determinant in such failures, a European standard, EN 12697-46:2012 (BSI, 2012e), specifies several uniaxial tension tests
with different stress–strain regimes to assess the resistance of asphalts to such cracking.

- In the uniaxial tension stress test, a specimen is pulled at a constant strain rate until failure while the temperature is kept constant. This yields the maximum stress (tensile strength) and the corresponding tensile failure strain at the test temperature.

- In the thermal stress restrained specimen test, as the test temperature is decreased at a constant rate, a test specimen is restrained from shrinking (as the name suggests), inducing an increase in the level of cryogenic stress in the test specimen. This yields the results of the progression of the cryogenic stress over the temperature and, finally, the failure stress at the failure temperature.

- In the relaxation test, a spontaneous strain is applied to the test specimen and held constant. Over time, relaxation causes the tension stress to decrease. The remaining tension stress and time of relaxation are monitored.

- In the tensile creep test, the test temperature is kept constant while subjecting the test specimen to a constant tension stress. The progression of the resultant strain is measured, and the stress is later withdrawn at a given time. From the measured strain data, the elastic and viscous properties of the asphalt are interpreted.

- In the uniaxial cyclic tension stress test, a cyclic tensile stress is applied as a sinusoidal stress to simulate the dynamic loading condition by traffic in combination with a constant stress, which acts as a surrogate for the cryogenic stress. The strain response is monitored as the test progresses, while the stiffness is also recorded until fatigue failure occurs. The resultant number of load cycles to failure is reported.

In China, for the cracking resistance performance of asphalts at low temperature, a bending beam test is carried out as shown schematically in Figure 16.15. A beam specimen of 250 mm length \(l\) by 30 mm breadth \(b\) and 35 mm height \(h\) is subjected to a vertical load applied at its centre at a rate of 50 mm/min at \(-10^\circ C\) until it fails. The load and vertical displacement \(d\) are recorded, and their values at failure are used to calculate the maximum tensile strain using the following equation

\[
\varepsilon_b = \frac{6hd}{l^2}
\]

### 16.2.5 Resistance to crack propagation

The fundamental fatigue tests described in sections 16.1.3 and 16.1.4 cover the initiation of cracking as the first phase of the failure mechanism during dynamic loading. The second part involves the propagation of the
cracks, and the resistance of asphalts to such crack propagation can be measured using the semi-circular bending test method described in a European standard, EN 12697-44:2010 (BSI, 2010). As the name suggests, a half cylinder test piece, with a 0.35 mm wide by 10 mm deep notch specially cut into the middle of the specimen, is loaded in three-point bending in such a way that its base is subjected to a tensile stress (Figure 16.16). A load is then applied at a rate of 5 mm/min until the test sample fails. The force and vertical displacement are recorded to calculate the maximum strain, stress at failure and fracture toughness.

Figure 16.15 Low temperature bending beam test

Temperature = -10ºC
Load speed = 50 mm/mn

Figure 16.16 Semi circular bending test set up

16.3 Empirical

Empirical tests do not provide results such as fundamental material stresses and strains, but many have been widely used historically. One example, the Marshall test, is still used in many countries now, even though the complex stress system set up in the material is quite unrelated to the actual pavement condition in situ under traffic loading, and the tests do not provide basic information on the stress–strain characteristics of the material being tested.
16.3.1 The Marshall test

The concepts of the Marshall test were developed by Bruce Marshall, formerly bituminous engineer with the Mississippi State Highway Department. In 1948, the US Corps of Engineers improved and added certain features to Marshall’s test procedure, and ultimately developed mixture design criteria (Asphalt Institute, 1997). Since 1948, the test has been adopted by organisations and government departments in many countries, sometimes with modifications either to the procedure or to the interpretation of the results.

In the ASTM and European standards (ASTM, 2006; BSI, 2012c), the Marshall test entails the manufacture of cylindrical specimens 102 mm in diameter by 64 mm high using a standard compaction hammer (Figure 16.17) and a cylindrical mould. The specimens are tested for their resistance to deformation at 60°C under a loading jig moving vertically at a constant rate of 50 mm/min. The jaws of the loading rig confine the majority, but not all, of the circumference of the specimen, the top and bottom of the cylinder being unconfined (see Figure 16.17). Thus, the stress distribution in the specimen during testing is extremely complex. Two properties are determined: the maximum load carried by the specimen before failure (‘Marshall stability’), and the amount of deformation of the specimen before failure occurred (‘Marshall flow’). The ratio of stability to flow is known as the ‘Marshall quotient’.

Figure 16.17 Marshall impact compactor and loading jig
Although the Marshall test is very widely used, it is important to recognise its limitations. Research at the University of Nottingham (Brown et al., 1982) comparing the mechanical properties of various mixtures, using repeated load triaxial tests, triaxial creep tests, uniaxial unconfined creep tests and Marshall tests, suggested that the Marshall test was a poor measure of resistance to deformation and did not rank mixtures in order of their deformation resistance. In that study, the fundamental repeated-load triaxial tests gave more realistic results.

16.3.2 Indirect tensile strength test

In this test, the asphalt specimen is loaded diametrically, as shown in Figure 16.18, under a constant compression rate of 50 mm/min until it breaks (BSI, 2003; ASTM, 2012b). The indirect tensile strength, taken as the highest stress derived from the peak load applied at the break point, can be used as an indicator of cracking or rutting potential. The comparison of the indirect tensile strength before and after a water conditioning regime is
also commonly used to assess moisture susceptibility (see section 19.4.4 for more information on this subject).

16.4 Determination of recovered bitumen properties

In addition to routine production control tests, there are a number of procedures available to assist highway engineers in the investigation of carriageway defects. One useful test is to recover the bitumen from the mixture and determine its properties, usually the penetration and softening point.

In this test, a sample of the asphalt is soaked in a solvent, such as trichloroethylene or dichloromethane (methylen chloride), to remove the bitumen from the aggregate into solution. The bitumen/solvent solution is separated from the fine mineral matter by filtration and centrifuging. The solvent is then evaporated under controlled conditions, using the apparatus shown in Figure 16.19, ensuring that all of the solvent is removed without

![Figure 16.19 Distillation apparatus used for recovery of bitumen (BSI, 2005a)](image-url)

**Key**
1. Fit stirrer/CO₂ tube here
2. Thermometer with bulb opposite bottom of flask
3. Enclosed electrical heater
4. Condenser
5. Fractioning column
6. Vacuum gauge
7. Water trap
8. Pump
9. 500 ml bottom flask
10. Oil bath
11. Jack
taking away any of the lighter components of the bitumen. When the recovery procedure has been completed, the penetration and softening point of the recovered bitumen can be determined using the standard tests. The method is described in detail in the European standard, EN 12697-4:2005 (BSI, 2005a). Alternatively, the solvent can be removed using a rotary evaporator (Figure 16.20) (ASTM, 2012a; BSI, 2013), which has the advantage of removing the solvent very quickly from the solution.

The test is very operator sensitive, and deviation from the standard method will almost certainly result in misleading data (Wadelin, 1982). It should also be noted that the extraction at high temperatures may cause the bitumen to harden, whereas residual solvent that remains after recovery may lead to some degree of softening, and these two opposite effects are usually taken to cancel each other out approximately (BSI, 2005a, 2013). For these reasons, caution should be exercised when using the test for contractual purposes. Notwithstanding, it is a very useful tool for investigating defects in service. With sensible interpretation of the test results, the bitumen recovery method can be used to determine if the bitumen in the mixture is abnormally hard or soft.
Figure 16.21 Chromatographic plots of bitumen (a) with diesel and (b) without diesel
In addition to the determination of the penetration and the softening point, chromatographic examination of the recovered bitumen can be useful to detect the presence of any contaminants. Figure 16.21(a) and (b) shows chromatographic plots, with and without the presence of diesel, respectively, generated using a combination of gas chromatography and mass spectrometry. The presence of the contaminant is very clear. Identification, however, is less precise, because diesel can weather, losing lighter fractions, and the composition can change between summer and winter. Although the technique is only qualitative and not quantitative, it is a useful tool for examining recovered bitumens that are abnormally soft.

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Asphalt Institute (1997) Mix Design Methods for Asphalt Concrete and Other Hot-Mix Types (MS-2). Asphalt Institute, Lexington, KY, USA.


Valkering CP, Lancon DJL, de Hilster E and Stoker DA (1990) Rutting resistance of


Chapter 17

Properties of asphalts

The analytical design of asphalt pavements involves consideration of two aspects of material properties. These are

- the load/deformation or stress/strain characteristics used to analyse critical stresses and strains in the structure
- the performance characteristics of the materials that show the mode, or modes, of failure.

The two principal structural distress modes are cracking and deformation. The former applies to bound materials only in the pavement, while the latter is valid for all materials in the pavement, both bound and unbound.

17.1 Stiffness of asphalts

As discussed previously, asphalts behave visco-elastically (i.e. they respond to loading in both an elastic and a viscous manner). The proportion of each depends on the time of loading and on the temperature at which the load was applied. The complexity of this behaviour is increased by the proportion of the components of the mixture, the bitumen being responsible for the visco-elastic properties, while the mineral skeleton influences elastic and plastic properties. Mixture components and compositions can be extremely diverse, which makes prediction of the properties of a particular mixture difficult.

Asphalt stiffness can be divided into elastic stiffness, which dominates under conditions of low temperatures or short loading times, and viscous stiffness, which dominates at high temperatures or long loading times. The former is used to calculate critical strains in the structure in analytical design. The latter
is used to assess the resistance of the material to deformation. It has also been shown that, at intermediate temperatures, when the stiffness has both an elastic and a viscous component, the stiffness is stress dependent (Read, 1996). High stresses result in lower stiffness, and low stresses result in higher stiffness, making the assessment of performance even more difficult. However, this stress dependency is less important than the effects of both the time of loading and temperature.

Stiffness at a particular temperature and time of loading can be measured by a variety of methods

- bending or vibration tests on a beam specimen
- direct uniaxial or triaxial tests on cylindrical specimens
- indirect tensile tests on cylindrical specimens
- dynamic loading on the asphalt mixture performance tester.

Identical samples from the same mixture will return different modulus values, as the results depend on the measuring criteria as described in Chapter 16.

Different types of loading can be used in the tests but for elastic stiffness of materials that carry traffic, sinusoidal or pseudo-sinusoidal repeated loading at high frequency is most appropriate (Raithby and Sterling, 1972).

### 17.1.1 The prediction of asphalt stiffness

Asphalt stiffness can be measured quickly and easily using tests such as the indirect tensile stiffness modulus test (BSI, 2003). However, when testing is not feasible, such as in the design office, then the stiffness of a particular mixture at any temperature and time of loading can be estimated using empirical methods, to an accuracy that is acceptable for most purposes.

#### 17.1.1.1 Shell method

In 1977, Shell produced a nomograph (Figure 17.1) for predicting the stiffness of asphalt (Bonnaure et al., 1977). The data required for this nomograph are

- the stiffness modulus of the bitumen (in Pa)
- the percentage volume of bitumen
- the percentage volume of the mineral aggregate.

The University of Nottingham has also developed a method for calculating mixture stiffness (Brown and Brunton, 1986) and the data required are

- the stiffness modulus of the bitumen (in Pa x \(10^6\))
- the voids in the mineral aggregate (VMA in %).

These two procedures can only be applied when the stiffness modulus of the bitumen exceeds 5 MPa (i.e. under high stiffness conditions appropriate to
traffic when the response is predominantly elastic and, for the Nottingham method, values of VMA between 12% and 30%). These two methods assume that the grading, type and characteristics of the aggregate affect only the elastic stiffness of the mixture, as they influence the packing

Example
Stiffness modulus of the recovered bitumen, $2.2 \times 10^8$ Pa
$V_b$: Volume of bitumen, 13.1%
$V_g$: Volume of mineral aggregate, 80.5%

Stiffness modulus of the mixture $1.1 \times 10^{10}$ Pa

Figure 17.1 Nomograph for predicting the stiffness modulus of asphalt (Bonnaure et al., 1977)
characteristics of the aggregate and, thus, the state of compaction of the material.

17.1.1.2 Asphalt Institute method
Hwang and Witczak (1979) developed the DAMA computer program for the Asphalt Institute. They applied regression formulae to determine the dynamic modulus of asphalts (what is described in the USA as 'hot mix asphalt')

\[ E^* = 100000 \times 10^{\beta_1} \]
\[ \beta_1 = \beta_3 + 0.000005\beta_2 - 0.00189\beta_2 f^{-1.1} \]
\[ \beta_2 = (\beta_4)^{0.5} + T^{\beta_5} \]
\[ \beta_3 = 0.553833 + 0.028829(P_{200}^{f-0.1703}) - 0.03476V_a 
+ 0.070377\lambda + 0.931757f^{-0.02774} \]
\[ \beta_4 = 0.483V_b \]
\[ \beta_5 = 1.3 + 0.49825 \log f \]

where \( \beta_1 \) to \( \beta_5 \) are temporary constants, \( f \) is the load frequency (in Hz), \( T \) is the temperature (in °F), \( P_{200} \) is the percentage by weight of aggregate passing through a no. 200 sieve, \( V_a \) is the percentage volume of air voids, and \( \lambda \) is the bitumen viscosity at 70°F (21.1°C) in 10^6 poise.

Note: if insufficient viscosity data are available, the following equation may be used

\[ \lambda = 29508.2(P_{77°F}) - 2.1939 \]

where \( P_{77°F} \) is the penetration at 77°F (25°C) and \( V_b \) is the percentage volume of bitumen.

It can be seen that the factors considered by Hwang and Witczak (1979) for the Asphalt Institute model are very similar to those considered by Shell, with the following differences.

- The percentage of fines passing the no. 200 sieve is used in the Asphalt Institute method but not in the Shell method.
- The viscosity or penetration of bitumen utilised in the Shell method is derived from recovered binder from the mixture, while the Asphalt Institute method uses virgin bitumen.
- The temperature and viscosity of bitumen feature in the Asphalt Institute method, while the Shell method employs the penetration index.
17.1.1.3 Witczak equation for MEPDG

Witczak’s equation is based on a non-linear regression analysis using the generalised reduced gradient optimisation approach. This model incorporates mixture volumetric and aggregate gradation, and is currently one of two options for level 3 analysis using the NCHRP 1-37A Mechanistic–Empirical Pavement Design Guide (MEPDG) program (Advanced Research Associates, 2004). The new Mechanistic–Empirical Pavement Design Guide adopted by the American Association of State Highway and Transportation Officials (AASHTO) represents a fundamental advance over the current 50 year old empirical pavement design procedures derived from the AASHTO road test. A hierarchical input data scheme has been implemented in the MEPDG to permit varying levels of sophistication for specifying material properties, ranging from laboratory measured values (level 1) to empirical correlations (level 2) to default values based on the prediction model (level 3).

In a study by Andrei et al. (1999), the original Witczak predictive equation was revised as follows

\[
\log_{10}|E^*| = -1.249937 + 0.02923P_{200} - 0.001767(P_{200})^2 - 0.002841P_4 - 0.05809V_a - 0.082208\left(\frac{V_{\text{beff}}}{V_{\text{beff}} + V_a}\right) - 3.871977 - 0.0021P_4 + 0.0003958P_{3/8} + 0.000017(P_{3/8})^2 + 0.00547P_{3/4} \frac{1}{1 + \exp(-0.603313 - 0.313351 \log f - 0.3935321 \log \eta)}
\]

where \(P_{200}\) is the percentage of aggregate passing a no. 200 sieve, \(P_4\) is the percentage of aggregate retained on a no. 4 sieve, \(P_{3/8}\) is the percentage of aggregate retained on a \(\frac{3}{8}\) in. (9.56 mm) sieve, \(P_{3/4}\) is the percentage of aggregate retained on a \(\frac{3}{4}\) in. (19.01 mm) sieve, \(V_a\) is the percentage of air voids (by volume of mixture), \(V_{\text{beff}}\) is the percentage of effective bitumen content (by volume of mixture), \(f\) is the loading frequency (Hz) and \(\eta\) is the binder viscosity at the temperature of interest (10⁶ poise or 10⁵ pa·s).

The limitations of Witczak’s equation, acknowledged by Bari (2005), include relying on other models to translate the dynamic shear modulus into binder viscosity. Because the original Witczak equation is based on regression analysis, extrapolation beyond the calibration database should be restricted. Bari also mentions that there is limited volumetric influence (precision) when the model is compared to the Shell method. Dongre et al.
2005) have also noted the need for improved sensitivity to volumetric values, such as the percentage of voids in mineral aggregate (VMA), the percentage of voids filled with bitumen (VFB), the bitumen percentage ($V_b$) and the percentage of air voids ($V_a$).

To include the dynamic shear modulus of the binder $|G^*|_b$ in the predictive model, Witczak reformulated the model to include the binder variable directly. The updated model is as follows

$$
\log_{10}|E^*| = -0.349 + 0.754(|G^*|_b)^{0.0052} \\
\times \left[ 6.65 - 0.032P_{200} + 0.0027P_{200}^2 + 0.011P_4 \\
- 0.0001(P_4)^2 + 0.006P_{3/8} - 0.00014(P_{3/8})^2 \\
- 0.08V_a - 1.06 \left( \frac{V_{\text{beff}}}{V_{\text{beff}} + V_a} \right) \right] \\
2.558 + 0.032V_a + 0.713 \left( \frac{V_{\text{beff}}}{V_{\text{beff}} + V_a} \right) + 0.124P_{3/8} \\
+ \frac{-0.0001(P_{3/8})^2 - 0.0098P_{3/4}}{1 + \exp(-0.7814 - 0.5785 \log |G^*|_b + 0.8834 \log \delta_b)}
$$

where $|G^*|_b$ is the dynamic shear modulus of asphalt binder (lb/in.$^2$) and $\delta_b$ is the binder phase angle associated with $|G^*|_b$ (degrees). This equation is one of two options for level 3 analysis in the most current NCHRP 1-40D MEPDG program.

Because some of the mixtures in this database do not contain $|G^*|_b$ data, the Cox–Mertz rule, using correction factors for the non-Newtonian behaviour (the relationship between the shear stress and the shear rate is different, and can even be time dependent) is used to calculate the dynamic shear modulus of the binder $|G^*|_b$ from $A$:VTS values, where $A$ is the intercept of the temperature susceptibility relationship and VTS is the slope of temperature susceptibility relationship as follows

$$
|G^*|_b = 0.0051f_s\eta_{\text{f}}(\sin \delta_b)^{7.1542 - 0.4929f_s + 0.0211f_s^2} \\
\delta_b = 90 + (-7.3146 - 2.6162VTS)\log (f_s\eta_{\text{f}}) \\
+ (0.1124 + 0.2029VTS)\log (f_s\eta_{\text{f}})^2 \\
\log[\log(\eta_{\text{f}})] = 0.9699f_s^{-0.0527}A + 0.9668f_s^{-0.0527} \times \text{VTS} \log T_R
$$

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where $f_s$ is the dynamic shear frequency (Hz), $\delta_b$ is the predicted binder phase angle (°), $\eta_{b,T}$ is the viscosity of asphalt binder at a particular loading frequency ($f_s$) and temperature ($T$) (centipoise (cP)) and $T_R$ is the temperature using the Rankine scale (i.e. absolute temperature) (°R).

17.1.1.4 Hirsch model
Christensen et al. (2003) examined four different models based on the law of mixtures parallel model, and chose the model that incorporates the binder modulus, VMA, and voids filled with asphalt (VFA) because it provides accurate results in the simplest form. The other more complicated forms attempt to incorporate the modulus of the mastic or the film thickness, which are difficult parameters to measure. The suggested model for estimating the dynamic modulus of asphalt (again what would be described in the USA as ‘hot mix asphalt’) ($|E^*|$) is as follows

$$|E^*|_m = P_c \left[ 4 200 000 \left( 1 - \frac{\text{VMA}}{100} \right) + 3 |G^*|_b \left( \frac{\text{VFA} \times \text{VMA}}{10 000} \right) \right]$$

$$+ \frac{(1 - P_c)}{\frac{\text{VMA}}{4 200 000} + \frac{\text{VMA}}{4 200 000} + 3 |G^*|_b \text{VFA}}$$

$$\phi = -21(\log P_c)^2 - 55 \log P_c$$

$$P_c = \frac{20 + 3 |G^*|_b \text{VFA}/\text{VMA}^{0.58}}{650 + 3 |G^*|_b \text{VFA}/\text{VMA}^{0.58}}$$

where $|E^*|_m$ is the dynamic modulus of the asphalt (lb/in.²), $P_c$ is the aggregate contact volume (in.³) and $\phi$ is the phase angle of the asphalt (°).

A strength of this model is the empirical phase angle equation, which is important for the inter-convension of the dynamic modulus to the relaxation modulus or creep compliance. Weaknesses of the model include a lack of a strong dependence on volumetric parameters, particularly under conditions of low air void and voids filled with bitumen. Also, questions arise as to whether the dynamic shear modulus is capable of being altered to take account of the possible beneficial effects of modifiers (Al-Khateeb et al., 2006).

17.1.1.5 Al-Khateeb model
Based on their findings from the Hirsch model, Al-Khateeb et al. (2006) suggest the law of mixture parallel model as follows

$$|E^*|_m = 3 \left( \frac{100 - \text{VMA}}{100} \right) \left[ \frac{90 + 10 000 |G^*|_b / \text{VMA}}{1100 + (900 |G^*|_b / \text{VMA})^{0.66}} \right] |G^*|_g$$
where \( G^* \) is the dynamic shear modulus of the asphalt binder at the glassy state (assumed to be 145 000 psi (999 050 kPa)).

Like the Hirsch model, this formulation is based on the law of mixtures for composite materials. In this model, the different material phases (aggregate, asphalt binder and air) are considered to exist in parallel. Therefore, this model is a simpler interpretation of the Hirsch model. The researchers note that their model addresses one of the primary shortcomings of the Hirsch model (namely, the inability of the Hirsch model to predict accurately the dynamic modulus of the mixture at low frequencies and high temperatures).

The strengths of this model include the improved prediction of high temperature and low frequency dynamic modulus. Weaknesses include a lack of model verification and the fact that the researchers who developed this model did so based on dynamic modulus values obtained from tests at higher than recommended strain amplitudes (200 microstrain versus the recommended maximum of 75–150 microstrain).

### 17.2 Deformation of asphalts

In order to determine the deformation characteristics of asphalt, the low stiffness response of the material (i.e. its response at high temperature or long loading times) must be analysed. When the stiffness of the bitumen is <0.5 MPa, mixture behaviour is much more complex than it is in the elastic zone (the area where the elastic behaviour of the mixture is more prominent than its viscous behaviour). Under these conditions, the stiffness of the mixture not only depends on the bitumen stiffness and the voids in the mixed aggregate and bitumen, but also on a variety of other factors. These include the aggregate grading, its shape, texture and degree of interlock, and the degree of compaction. This is illustrated in Figure 17.2.

The simplest test used to study the deformation behaviour of asphalts is the creep test. Figure 17.3(a) shows the deformation behaviour of a mixture tested at different temperatures. If these same results are plotted on a graph of mixture stiffness \( S_{\text{mix}} \) against bitumen stiffness \( S_{\text{bit}} \), the test results form a single continuous ‘master curve’ for the mixture, as shown in Figure 17.3(b). Thus, the effect of testing at different temperatures is combined in a single curve. Similarly, the effect of using different grades of bitumen or the application of different stress levels can also be combined.

Thus, the \( S_{\text{mix}} \) versus \( S_{\text{bit}} \) curve is a means of assessing the resistance to deformation that is independent of arbitrarily chosen test variables for a particular asphalt. For example, Figure 17.4 shows the deformation characteristics of two asphalts having identical aggregate grading but containing 5% and
11% bitumen, respectively. It can be seen that the stiffness of the leaner mixture levels out with decreasing bitumen stiffness. In contrast, the stiffness of the richer mixture continues to decrease with decreasing bitumen stiffness. Clearly, therefore, maintaining a high value of $S_{\text{mix}}$ when $S_{\text{bit}}$ is decreasing is a desirable characteristic for long term resistance to deformation. Similarly, Figure 17.5 demonstrates the effect of aggregate shape on asphalt having the same aggregate grading and bitumen content (Hills et al., 1974). As would be expected, crushed aggregate increases the degree of aggregate interlock, and this results in an increase in the value of $S_{\text{mix}}$ and higher resistance to deformation.

The resistance of mixtures to deformation can be determined by tests such as the unconfined creep test and the repeated load uniaxial or triaxial test, their results being plotted as $S_{\text{mix}}$ versus $S_{\text{bit}}$. These tests reproduce stress conditions on the road more accurately and, as a result, are gaining widespread popularity.

### 17.2.1 The prediction of deformation

When attempting to calculate the amount of deformation on a highway, account has to be taken of a range of wheel load spectra, contact areas and pressures, lateral distribution of the wheel loads and ambient temperature gradients within the asphalt layers. Clearly, the situation is very complex.
Figure 17.3 Results from a creep test (Hills et al., 1974): (a) effect of temperature on deformation; (b) master curve for a particular mixture
17.2.1.1 Shell method

The final step in the Shell Pavement Design Manual (Claessen et al., 1977; Shell, 1978) when layer thicknesses have been determined is to predict the rut depth from

\[ \text{Rut depth} = C_m \times h \times \frac{\sigma_{\text{ave}}}{S_{\text{mix}}} \]

where \( C_m \) is the empirical correction factor, \( h \) is the thickness of the layer, \( \sigma_{\text{ave}} \) is the average stress, and \( S_{\text{mix}} \) is the stiffness of the mixture.

**Figure 17.4** The effect of bitumen content on creep properties (Shell, 1978)

**Figure 17.5** The effect of aggregate shape on creep properties (Hills, 1973)
However, as temperature gradients occur within the asphalt layers and different mixture types are often used in the surface course, binder course and base, the total asphalt thickness must be subdivided and each layer considered individually. The rut depth is the sum of the deformation in each of these layers.

17.2.1.2 VESYS method
VESYS is a linear computer program that can be used to determine the deformation based on resilient modulus (FHWA, 1978). The VESYS method of prediction of rut depth is based on an assumption that permanent strain is proportional to resilient strain.

\[ \varepsilon_p(N) = \mu \varepsilon N^{-\alpha} \]

where \( \varepsilon_p(N) \) is the permanent or plastic strain due to a single load application at the \( N \)th application, \( \varepsilon \) is the elastic or resilient strain at the 200th repetition, \( N \) is the load application number, \( \mu \) is the constant between permanent and elastic strain, and \( \alpha \) is the rate of decrease in deformation.

The total deformation can be obtained by integrating the earlier equation

\[ \varepsilon_p = \int_0^N \varepsilon_p \mu \varepsilon(N) \, dN = \varepsilon \mu \left( \frac{N^{1-\alpha}}{1-\alpha} \right) \]

Converting the above equation into a log–log relationship gives the following relationship

\[ \log \varepsilon_p = \log \left( \frac{\varepsilon \mu}{1-\alpha} \right) + (1-\alpha) \log N \]

where the slope of the plot is \( S = 1 - \alpha \) (i.e. the rate of decrease of deformation \( \alpha = 1 - S \).

The intercept \( I = \varepsilon \mu / (1 - \alpha) \) or constant \( \mu = IS/\varepsilon \).

VESYS also has an option to determine the deformation parameter of a layered pavement.

17.2.1.3 MEPDG rutting model
Similar to the stiffness prediction model, the new MEPDG also includes a rutting prediction model. The analysis using the rutting model was simplified, and separate rutting at the asphalt concrete surface course was used, in contrast to the earlier version that assessed deformation of the full depth of the asphalt layers. The current version of the MEPDG model (National
Cooperative Highway Research Program, 2004) for rutting of asphalt concrete is based on the following relationship

\[ \frac{\varepsilon_p}{\varepsilon_r} = k_1 10^{-3.35412 T^{1.5606}} N^{0.4791} \]

where \( \varepsilon_p \) is the plastic strain, \( \varepsilon_r \) is the elastic strain, \( T \) is the temperature (°F), \( N \) is the number of load applications, and \( k_1 \) is a constant that depends on the depth at which the elastic strain is calculated based on the equation

\[ k_1 = (C_1 + C_2 \times \text{depth}) \times 0.328196^{\text{depth}} \]

\[ C_1 = -0.1039 h_{ac}^2 + 2.4868 h_{ac} - 17.342 \]

\[ C_2 = -0.0172 h_{ac}^2 - 1.7331 h_{ac} - 27.428 \]

in which ‘depth’ is the depth to the point of strain calculation and \( h_{ac} \) is the thickness of the asphalt layer. The total rutting in the asphalt layer is calculated by integrating the calculated plastic strain over the thickness of the layer.

17.3 Fatigue characteristics of asphalts

Fatigue can be defined as: ‘The phenomenon of fracture under repeated or fluctuating stress having a maximum value generally less than the tensile strength of the material’ (Pell, 1988). However, this has been generally accepted as referring to tensile strains induced by traffic loading, and, because other means of generating tensile strains in a pavement exist, a better definition may be: ‘Fatigue in bituminous pavements is the phenomenon of cracking. It consists of two main phases, crack initiation and crack propagation, and is caused by tensile strains generated in the pavement by not only traffic loading but also temperature variations and construction practices’ (Read, 1996).

Under traffic loading, the layers in a flexible pavement are subjected to flexing that is virtually continuous. The size of the strains is dependent on the overall stiffness and nature of the pavement construction, but analysis confirmed by in situ measurements has suggested that tensile strains of the order of \( 30 \times 10^{-6} \) to \( 200 \times 10^{-6} \) for a standard wheel load occur. Under these conditions, the possibility of fatigue cracking exists (Pell, 1988).

As described in section 16.1.3, dynamic bending tests are normally used to measure the fatigue characteristics of asphalts. An example of the constant-stress fatigue life characteristics for the same mixture at different temperatures is shown in Figure 17.6. The lines are essentially parallel, and show longer fatigue lives at lower temperatures. If the tests are carried out at a different frequency, the result is similar. Thus, fatigue lives are longer at higher frequencies. The fatigue results could also be presented in terms
of initial strain, as shown in Figure 17.7. When the criterion of failure is strain, the temperature and time of loading will affect the mixture stiffness. This failure mechanism or effect is known as the ‘strain criterion’ (Pell and Taylor, 1969).

The general relationship defining the fatigue life, in terms of initial tensile strain, is

\[ N_f = c(1/\varepsilon_i)^m \]

**Figure 17.6** Typical fatigue lines – stress criterion (Pell, 1988)

**Figure 17.7** Fatigue lines from Figure 17.6 – strain criterion (Pell, 1988)
where $N_f$ is the number of load applications to initiate a fatigue crack, $\varepsilon_t$ is the maximum value of applied tensile strain, $c$ and $m$ are factors depending on the composition and properties of the mixture, and $m$ is the slope of the strain/fatigue life line.

However, the strain criterion does not account for the differences in fatigue characteristics predicted using controlled stress and controlled strain experiments. Accordingly, researchers have tried to find alternative failure criteria that take account of the differences in fatigue characteristics.

It has been suggested by Van Dijk and others (Van Dijk, 1975; Van Dijk and Visser, 1977; Van Dijk et al., 1972) that the differences in the fatigue life of asphalt determined under conditions of stress and strain control can be explained by the dissipated-energy concept. This is the amount of energy that is lost from the system, due to fatigue damage, per cycle summed for the entire life of the specimen. Van Dijk and others stated that, for a given mixture, the relationship between dissipated energy and the number of load repetitions to failure is valid, independent of testing method and temperature. This work has been progressed by Himeno et al. (1987), who developed the dissipated-energy concept for three dimensional stress conditions and applied it to the failure of an asphaltic layer in a pavement, and by Rowe (1993), who has shown that dissipated energy can be used to predict the life to crack initiation accurately. The dissipated-energy concept shows considerable promise as a failure criterion that encompasses all the variables of fatigue.

The fatigue characteristics of a mixture can be influenced significantly by its composition. Mixture stiffness is also influenced by the composition of the asphalt, and distinguishing between the two is important. For example, adding filler or reducing the void content will increase mixture stiffness, resulting in increased fatigue life at a given level of stress, as the resulting strain is smaller. In other words, a point has been reached lower down on the strain–fatigue life line. At a particular strain level, however, it has been found that the entire line moves to the right (Pell and Cooper, 1975) (i.e. the fatigue life is improved if, for example, the volume of bitumen is increased). Basic fatigue performance is influenced by other mixture variables only insofar as these affect the volume of bitumen in the mixture. If rounded aggregate is used, for instance, a denser mixture with a lower void content will result, and this will increase the relative volume of bitumen, and thus improve the fatigue performance.

**17.3.1 The prediction of fatigue deformation**

Traditionally, the establishment of the fatigue line for a particular mixture involved specialised testing equipment. A simpler procedure to predict fatigue performance with sufficient accuracy for pavement design purposes was clearly needed.
17.3.1.1 Shell method
A method for predicting the fatigue life of asphalts was developed by Shell (Bonnaure et al., 1980) using the nomograph shown in Figure 17.8. The required data are

- the percentage volume of bitumen
- the penetration index of the bitumen
- the stiffness modulus of the mixture (in Pa)
- the initial strain level.

The equations used to develop the nomographs are as follows.

For constant stress tests

\[ N_f = [0.0252P - 0.00126P(V_b) + 0.00673V_b - 0.0167]^5 \varepsilon_t^{-5} S_m^{-1.4} \]

For constant strain tests

\[ N_f = [0.17P - 0.0085P(V_b) + 0.0454V_b - 0.112]^5 \varepsilon_t^{-5} S_m^{-1.8} \]

Other prediction techniques based on some or all of the above input data (Asphalt Institute, 1981; Cooper and Pell, 1974) have also been developed, but these empirical techniques are unable to deal with material developments such as polymer modified mixtures, as they are based on historical data. This has led to the usage of other types of fatigue measurement as described in detail in Chapter 16.

The application of laboratory determined fatigue lives to predict actual pavement performance in practice is a complex problem that is likely to yield conservative results. This happens because simple continuous cycles of loading neglect the beneficial effects of rest periods that occur in practice between axle loads. Longer lives are also likely in practice because of the lateral distribution of wheel loads in the wheel track, and the fact that a degree of crack propagation will occur before the performance of the pavement is adversely affected. As a result of these problems, laboratory fatigue lives have to be calibrated to correlate with actual pavement performance, and the calibration factor is likely to depend on environmental and loading conditions. However, these factors have been incorporated in a prediction technique (Read, 1996) for use in analytical pavement design, based around work carried out at the University of Nottingham.

17.3.1.2 Asphalt Institute method
The Asphalt Institute fatigue equation (Asphalt Institute, 1969) is based on constant stress criteria only.

\[ N_f = 0.00432C \varepsilon_t^{-3.291} |E^*|^{-0.854} \]
Stiffness modulus of the mixture $S_{mix}$: Pa

Fatigue life: cycles

Initial strain

Bitumen volume, $V_b$

Penetration index

Example
Stiffness modulus of the mixture, $3.3 \times 10^9$ Pa
Bitumen volume, 13%
Penetration index, $-0.7^*$

*Recovered bitumen

Initial strain for a lifetime of:
$10^6$ cycles – constant stress test, $1.1 \times 10^{-4}$
$10^8$ cycles – constant strain test, $4.5 \times 10^{-4}$

Figure 17.8 Nomograph for predicting the laboratory fatigue performance of asphalt (Bonnaure et al., 1980)
where \( C \) is a correction factor, where \( C = 10M \)

\[
M = 4.84 \left( \frac{V_b}{V_a + V_b} \right) - 0.69
\]

where \( V_b \) is the percentage binder content and \( V_a \) is the percentage of air voids.

### 17.3.1.3 MEPDG method

Besides providing a stiffness and rutting prediction model, the new *Mechanistic–Empirical Pavement Design Guide* includes a fatigue prediction model (Advanced Research Associates, 2004). The MEPDG fatigue model uses the modified Asphalt Institute method with an additional correction factor for the layers. This fatigue equation predicts the number of load repetitions to fatigue cracking failure.

\[
N_f = 0.00432k_1C \left( \frac{1}{E_1} \right)^{3.9492} \left( \frac{1}{E} \right)^{1.287}
\]

where \( \varepsilon_1 \) is the tensile strain at the bottom of the asphalt surface layer, and \( E \) is the elastic modulus of the asphalt layer.

\( k_1 \) and \( C \) are parameters calculated using the following equations

\[
k_1 = \frac{1}{0.000398 + \frac{0.003602}{1 + e^{11.02 - 3.49h_a}}}
\]

\[
C = 10^M
\]

\[
M = 4.84 \left( \frac{V_b}{V_a + V_b} - 0.69 \right)
\]

where \( V_a \) is the percentage air voids and \( V_b \) is the percentage binder content.

### 17.4 Failure theories for cracking in asphalt layers

In the process of designing a structural member, the designer has to ensure that the member under consideration does not fail under service conditions. As empirical failure methods are adopted in the design of flexible pavements, there is little emphasis on research related to the applicability of failure theories to asphaltic layers. Failure theories represent a fundamental approach for evaluating the susceptibility to failure of a material under given loading conditions. Under uniaxial loading, failure occurs when the applied stress reaches the tensile strength of the material. However, under complex,
three dimensional stress fields, the situation is more complex, as both normal and shear stresses of varying magnitude may be present. There are five main failure theories, and these are discussed briefly in the following paragraphs. These theories are named after the principal researcher involved in the study.

17.4.1 Maximum principal stress theory (Rankine)
According to this theory, the maximum principal stress in the material determines failure, regardless of the value of the other two principal stresses, as long as they are algebraically similar. This theory is not supported by experimental results (Raju, 2008). It is considered to be reasonably satisfactory for brittle materials that do not fail by yielding. In uniaxial tension or compression modes, failure occurs when the maximum principal stress at any point reaches a value equal to the tensile or compressive elastic limit or yield strength of the material obtained from the uniaxial test. Thus, if \( \sigma_1 \), \( \sigma_2 \) and \( \sigma_3 \) are the principal stresses at a point and \( \sigma_1 > \sigma_2 > \sigma_3 \) and \( \sigma_y \) is the yield stress for the material under a uniaxial test, then failure occurs when \( \sigma_1 \geq \sigma_y \).

17.4.2 Maximum shearing stress theory (Tresca)
Observations made in the course of extrusion tests on the flow of soft metals through orifices lent support to the assumption that the plastic state in such metals is created when the maximum shear stress just reaches the value of the resistance of the metal against shear. Assuming \( \sigma_1 > \sigma_2 > \sigma_3 \), yielding, according to this theory, occurs when the maximum shearing stress reaches a critical value. The maximum shearing stress theory is accepted as being fairly well justified for ductile materials.

If \( \sigma_1 > \sigma_2 > \sigma_3 \) are the three principal stresses at a point, failure occurs when

\[
\tau_{\text{max}} = \frac{\sigma_1 - \sigma_3}{2} \geq \frac{\sigma_y}{2}
\]

where \( \sigma_y/2 \) is the shear stress at yield point in the uniaxial test (Raju, 2008).

17.4.3 Maximum principal strain theory (Saint Venant)
Maximum principal strain theory is analogous to maximum principal stress theory, but in this case failure is expected when the maximum principal strain exceeds the tensile yield strain, or the minimum principal strain exceeds the compressive yield strain (Raju, 2008).

\[
|\sigma_1 - \mu(\sigma_2 + \sigma_3)| = \sigma_{yt} \text{ or } |\sigma_3 - \mu(\sigma_1 + \sigma_2)| = \sigma_{yc}
\]

where \( \sigma_2 \) is the intermediate principal stress. This failure theory has found limited use in the design of thick walled cylinders (Raju, 2008).
17.4.4 Maximum distortion energy failure theory (von Mises)
The maximum distortion energy failure theory states that a material will fail when, at any point in the material, the distortion energy per unit volume reaches the distortion energy occurring in a uniaxial tension test at failure (Raju, 2008). The failure criteria in terms of normal as well as principal stresses are

\[
(s_x - s_y)^2 + (s_y - s_z)^2 + (s_z - s_x)^2 + 6(t_{xy}^2 + t_{yz}^2 + t_{zx}^2) = s_{yp}^2
\]

where \(s_x, s_y\) and \(s_z\) are the normal stresses in the \(x, y\) and \(z\) directions, and \(t_{xy}, t_{yz}\) and \(t_{zx}\) are the shearing stresses on the \(xy, yz\) and \(zx\) planes.

This can also be stated in terms of principal stresses.

\[
(s_1 - s_2)^2 + (s_2 - s_3)^2 + (s_3 - s_1)^2 = 2s_{yp}^2
\]

where \(s_1, s_2\) and \(s_3\) are the principal stresses, and \(s_{yp}\) is the yield stress.

The maximum distortion energy theory predicts a failure envelope (as defined when plotted using the principal stresses as coordinates) that is very similar to that predicted by maximum shear stress theory, and it is widely applied to failure in ductile materials (Raju, 2008).

17.4.5 Octahedral shearing stress theory

According to this theory, the critical quantity is the shearing stress on the octahedral plane. The plane that is equally inclined to all the three principal axes is called the ‘octahedral plane’. The octahedral shear stress theory is, in essence, a different way of stating the von Mises theory. Octahedral shear stress is given by the following equation (Raju, 2008)

\[
\tau_{oct} = \frac{1}{6}[(s_x - s_y)^2 + (s_y - s_z)^2 + (s_z - s_x)^2 + 6(t_{xy}^2 + t_{yz}^2 + t_{zx}^2)]^{1/2}
\]

where \(\tau_{oct}\) is the octahedral shear stress and the other variables are as the von Mises theory. Combining this relationship with the equation gives the failure criterion for the maximum octahedral shear stress theory (Raju, 2008)

\[
\tau_{oct} = 0.47s_{yt}
\]

Because the maximum distortion energy theory and the octahedral shear stress theory are equivalent, they are applicable to similar situations – triaxial loading conditions in ductile materials.

Figure 17.9 shows failure envelopes obtained by maximum shear stress and von Mises (octahedral shear stress) theories. Ameri-Gaznon and Little (Raju, 2008) applied octahedral shear stress theory to the design of asphalt concrete overlays.
Properties of asphalts

![Image of failure envelope](image-url)

**Figure 17.9** Failure envelope

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Chapter 18

Influence of binder properties on the performance of asphalts

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This chapter considers the relationship between bitumen properties and pavement performance, and a number of failure mechanisms, with particular emphasis on the role that is played by the binder.

18.1 Introduction

The performance of asphalts, in both the short and the long term, is influenced by the binder properties, aggregate and grading characteristics, as well as the composition, with the binder playing a critical role.

Despite the wide range of applications to which asphalts are put and the substantial variations in weather and loading to which they are subjected, the vast majority of asphalt pavements perform well for many years. However, failures such as rutting, cracking, fretting, fatting up etc. do occur. An understanding of the mechanisms that cause asphalt pavements to fail is important if designers, contractors and producers are to employ specifications, manufacturing techniques, equipment and methods that will minimise the possibility of defects occurring.

The performance of asphalts in service is influenced significantly by the rheological (or mechanical) properties and, to a lesser extent, the chemical constitution of the binder. These factors are, in turn, influenced by changes due to the effects of air, temperature and water on the binder.
The chemical constitution of the binder is particularly important at the road surface because it influences the rate of oxidation and, as a result, how rapidly the binder is eroded by traffic.

There are, of course, many other factors influencing behaviour, including the nature of the aggregate, mixture composition, binder content (i.e. binder film thickness), degree of compaction, etc. – all of which influence long term durability.

Binders are visco-elastic materials, and their behaviour varies from purely viscous to wholly elastic depending on loading time and temperature. During the mixing and compaction of asphalts and at high service temperatures, the properties can be considered in terms of viscosity, but for most service conditions, binders behave visco-elastically, and their properties can be considered in relation to their stiffness modulus.

The rheological requirements for binders during mixing, compaction and in service are illustrated in Figure 18.1, and the critical requirements are summarised in Table 18.1.
As the traditional methods of characterising bitumens (i.e. penetration, ductility and softening points) are empirical, they cannot be used to predict how an asphalt or an asphalt pavement will perform in service.

Some researchers have, in the past, used penetration, softening point and viscosity data to assess the complex properties of bitumen. Values of parameters such as penetration indexes (PIs) and penetration-viscosity numbers (Anderson et al., 1983; McLeod, 1972) were believed capable of reflecting the effect of temperature on rheological behaviour. However, it is now known that these parameters cannot accurately describe the effect of time of loading and temperature on the stress–strain response of bitumen.

In the USA, the Strategic Highway Research Program (SHRP) was initiated in the 1990s because specifications based on penetration value and viscosity could not be used to describe asphalt pavement performance fully. One of the major objectives of the SHRP project was to develop test methods that could be used to characterise the performance related physical properties of bitumen binders, so that a performance related binder specification could be developed based on rational relationships between bitumen properties and pavement performance.

The SHRP asphalt research program addressed the major distress modes encountered in asphalt concrete pavements:

- rutting
- low temperature cracking
- fatigue cracking.

<table>
<thead>
<tr>
<th>Condition</th>
<th>Significant property of the binder in the mix</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temperature: °C</td>
<td>Time of loading: s</td>
</tr>
</tbody>
</table>

### Table 18.1 Engineering requirements for binders during application and in service (Dormon, 1969)

<table>
<thead>
<tr>
<th>Behaviour during application</th>
<th>Temperature: °C</th>
<th>Time of loading: s</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mixing</td>
<td>(&gt;100°C)</td>
<td>–</td>
</tr>
<tr>
<td>Laying</td>
<td>High</td>
<td>–</td>
</tr>
<tr>
<td>Compaction</td>
<td>High</td>
<td>–</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>In-service deformation</th>
<th>Temperature: °C</th>
<th>Time of loading: s</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fatting up</td>
<td>(&gt;30°C)</td>
<td>Long (&gt;10²)</td>
</tr>
<tr>
<td>Cracking – traffic stresses</td>
<td>Low road temperature</td>
<td>Short</td>
</tr>
<tr>
<td>Cracking – thermal stresses</td>
<td>Low road temperature</td>
<td>Long</td>
</tr>
<tr>
<td>Fretting</td>
<td>Low road temperature</td>
<td>Short</td>
</tr>
</tbody>
</table>

Maximum viscosity determined by penetration and the softening point of the binder.
The parameters considered against these distress modes in the SHRP programme (Petersen et al., 1994; Kennedy et al., 1990) are

- for rutting – the ratio of the complex modulus ($G^*$) and the sine of the phase angle ($\sin \delta$) measured on the original binder at the maximum pavement temperature

- for low temperature cracking – creep stiffness and the $m$ value, which is the slope of the master stiffness curve at 60 s, measured on the residue from the pressure ageing vessel (PAV) test at the minimum pavement temperature plus 10°C

- for fatigue cracking – the product of the complex modulus ($G^*$) and the sine of the phase angle ($\sin \delta$) measured on the residue aged in the PAV at an intermediate pavement temperature.

More detailed information about the SHRP programme can be found in sections 5.7, 7.2.3.2, 12.1 and 12.4.5.

In Europe, highway engineers faced similar challenges in trying to ascertain which binder properties dictated the performance of asphalts and pavement performance. In 1995, Eurobitume hosted a workshop on rheology, and subsequently held an international workshop on performance-related properties of bituminous binders in 1999. The workshop identified key properties of binders associated with pavement performance, as listed in Table 18.2 (Eurobitume, 2002).

In 2000, CEN Technical Committee 336, Bituminous Binders, was assigned the task of drafting the second generation of European standards for bituminous binders. The aim of these standards was to define the relevant binder properties affecting the performance of asphalts, surfacing dressings and other road or industrial applications.

In 2012, Eurobitume published an updated position report to provide the views of the bitumen industry on the development of performance-related bitumen specifications. It proposed bitumen test methods appropriate for

<table>
<thead>
<tr>
<th>Table 18.2</th>
<th>Binder properties linked to asphalt performance (Eurobitume, 2002)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Binder properties</td>
<td>Performance requirements for pavement/mix</td>
</tr>
<tr>
<td>Rheological property at elevated service temperature</td>
<td>Resistance to deformation</td>
</tr>
<tr>
<td>Ageing behaviour: short term and long term</td>
<td>Resistance to surface cracking due to binder ageing</td>
</tr>
<tr>
<td>Rheological property: complex modulus</td>
<td>Structural strength</td>
</tr>
<tr>
<td>Combination of rheological and failure properties</td>
<td>Resistance to low temperature cracking</td>
</tr>
<tr>
<td>Failure property</td>
<td>Resistance to fatigue cracking</td>
</tr>
<tr>
<td>Viscosity vs temperature, storage stability</td>
<td>Manufacturing and laying</td>
</tr>
</tbody>
</table>
Influence of binder properties on the performance of asphalts

performance-based binder specification at some time in the future (Eurobitume, 2012). This initiative focused on the properties that Eurobitume considered to be directly related to pavement performance, namely

- mechanical properties of the binder
- bearing capacity
- fatigue cracking
- resistance to fretting
- resistance to low temperature thermal cracking.

The background to the relevance of these parameters is as follows.

- **Mechanical properties of binder.** The most commonly used properties are rheological parameters such as viscosity, stiffness, phase angle and deformation energy, because these parameters change dramatically depending on test conditions such as temperature, frequency/loading time, stress/strain level, number of loading cycles etc. By selecting sets of test conditions, it is possible to simulate the loading actually applied on the binder in the pavement and, in such circumstances, the response of the binder will provide an indication of its performance in the pavement.

- **Bearing capacity.** Bearing capacity is related to the stiffness of the pavement structure. The stiffness of the pavement is determined by the stiffness of the individual constituents of the asphalt. Therefore, bearing capacity is related to binder stiffness.

- **Fatigue cracking.** Fatigue cracking is the result of the accumulation of damage due to a large number of loading cycles. Even though there have been many attempts to relate the fatigue behaviour of bituminous binders to a simple rheological parameter, no clear relationship has yet been established. There are currently no harmonised test methods for determining the fatigue behaviour of bituminous binders. However, some laboratories are developing different methods. RILEM (the International Union of Laboratories and Experts in Construction Materials, Systems and Structures [Réunion Internationale des Laboratoires et Experts des Matériaux, systèmes de construction et ouvrages]) is evaluating the use of a dynamic shear rheometer (DSR) to determine the fatigue behaviour of binders. IFSTTAR (the French Institute of Science and Technology for Transport, Spatial Planning, Development and Networks [Institut français des sciences et technologies des transports, de l’aménagement et des réseaux]). IFSTTAR is evaluating a tension–compression fatigue test based on diabolo shaped specimens. The University of Catalunya in Barcelona is also evaluating a tension–compression fatigue test with cylindrical specimens.
Resistance to fretting. Resistance to fretting is related to the cohesive properties of the binder, which can be determined by measuring the deformation energy using the tensile test or force ductility test.

Resistance to low temperature thermal cracking. The resistance of binders to low temperature cracking can be evaluated using different methods: stiffness at low temperatures, bending beam rheometer test, and fracture toughness and tensile test.

Eurobitume (2012) concluded that ‘A new specification is required only for rheologically ‘complex’ bitumens such as polymer modified and hard paving grade bitumen. Rheologically ‘simple’ bitumen that meets the EN 12591 specification does not need new specifications as EN 12591 is considered adequately related to performance.’ EN 12591 is, of course, the European standard for paving grade bitumens (BSI, 2009).

18.2 The influence of binder properties during construction
18.2.1 Mixing and transport

During mixing, the dried hot aggregate has to be coated by the hot binder in a relatively short mixing time (typically 30–60 s). While the mixing temperature must be sufficiently high to allow rapid distribution of the binder on the aggregate, the use of the minimum mixing time at the lowest temperature possible is advocated. The higher the mixing temperature, the greater the tendency of the binder exposed in thin films on the surface of the aggregate to oxidise. This is illustrated in Figure 18.2, where an increase of 5.5°C in the mixing temperature, for a mixing time of 30 s, results in an increase of 1°C in the softening point of the binder (Whiteoak and Fordyce, 1989). There are, therefore, upper and lower limits of mixing temperatures. If asphalt concretes are mixed too hot, drainage of binder from the aggregate may occur during hot storage or transport to site, leading to variations in binder content. In such circumstances, the filler serves an important function. As it is generally added to the mixture after coating of the aggregate has been completed, the filler ‘stabilises’ or increases the apparent viscosity of the binder, reducing binder drainage. In thin surface course systems, stone mastic asphalts and porous asphalts, binder drainage can be prevented by the addition of fibres to the mixture, or by using a polymer modified binder.

These different considerations combine to give an optimum binder viscosity of 0.2 Pa·s at the mixing temperature. It has been shown (Jacobs, 1981) that the temperature required to achieve a viscosity of 0.2 Pa·s can be crudely estimated by simply adding 110°C to the softening point of the binder (Figure 18.3). The disadvantage of this method is that it does not take account of the PI of the binder (Figure 18.4). A more precise estimate can be determined using both the penetration and softening point of the binder on
the binder test data chart. The various conditions described above are summarised in Figure 18.4.

When materials are being laid at low ambient temperatures, or if haulage over long distances is necessary, mixing temperatures are often increased to offset these factors. However, increasing the mixing temperature will tend to accelerate the rate at which the binder oxidises, and this increases the viscosity of the binder. Thus, a significant proportion of the reduction in viscosity achieved by increasing the mixing temperature may be lost because of additional oxidation of the binder. If asphalts are transported in properly sheeted, well-insulated vehicles, the loss in temperature is very low, about 2°C/h.

18.2.2 Laying and compaction
Discharging the asphalt into a paver and spreading the material on to a substrate will reduce the temperature by about 20–30°C. The temperature loss will be dependent on a number of factors, including the thickness of the layer, ambient temperature, wind speed and the temperature of the substrate on which the new material is being placed. The two most crucial factors

Figure 18.2 Relationship between the temperature of the mixture and the change in softening point (Whiteoak and Fordyce, 1989)
influencing the cooling of the layer are wind speed and layer thickness (Daines, 1985).

Once the asphalt has been laid, it must be sufficiently workable to enable the material to be satisfactorily compacted with the available equipment. For effective compaction to take place, the viscosity of the binder should be between 5 and 30 Pa·s. At viscosities lower than 5 Pa·s, the material will probably be too mobile to compact, and at viscosities greater than 30 Pa·s the material will be too stiff to allow further compaction. The minimum rolling temperature can be estimated by adding 50°C to the softening point of the binder (Jacobs, 1981) (see Figure 18.3).

The slope of the viscosity–temperature relationship, or PI, in this temperature region is particularly important because it determines the temperature range.

Figure 18.3 Relationship between the softening point of the bitumen and equiviscous temperatures for mixing and rolling of a hot rolled asphalt surface course
Influence of binder properties on the performance of asphalts

over which the viscosity of the binder remains at a suitable level for compaction. Figure 18.4 clearly shows that binders with a high temperature susceptibility (i.e. those with a low PI) have a much narrower temperature ‘window’ within which satisfactory compaction of the material can be achieved.

It should be noted that the principles and observations discussed above apply to traditional hot asphalts but may not apply to warm asphalts due to these materials containing a range of additives not found in traditional asphalts.

18.3 The influence of binder properties on the performance of asphalts in service

Once an asphalt has been successfully manufactured, laid and compacted, its behaviour in service (i.e. its low and high temperature stability, moisture sensitivity, durability etc.) can be forensically analysed by considering the circumstances that prevailed while the material was in service and the relevance of those circumstances to specific categories of pavement defect.

18.3.1 Cracking

Cracking of pavements is a complex phenomenon that can be caused by several factors. It is associated with stresses induced in the asphalt layers by wheel loads, temperature changes or a combination of the two. Furthermore, the volume of binder in the mixture and its rheological behaviour have a major bearing on the susceptibility of the asphalt to cracking. An asphalt, by virtue of the binder it contains, displays visco-elastic behaviour. If an asphalt test specimen is strained to a predetermined point and the amount of strain held constant, a stress will be induced. Depending on temperature, this stress will dissipate more or less quickly. This process is called ‘relaxation’. At high temperatures, the viscous component dominates, and total stress relaxation may take a few minutes. At very low temperatures, relaxation can take many hours or even days.

Cracking occurs when the tensile stress and related strain induced by traffic and/or temperature changes exceed the breaking strength or breaking strain of the mixture. At elevated temperatures, stress relaxation will prevent these stresses reaching a level that can cause cracking. On the other hand, at low temperatures, the tensile condition will persist and, therefore, pavement cracking is more likely.

It is also recognised that binder in an asphalt ages during its service life, and this is called ‘curing’. This results in a progressive increase in the stiffness modulus of the asphalt, together with a reduction in its stress relaxation capability. This will further increase the likelihood of the pavement cracking.
Figure 18.4 The influence of the penetration index (PI) on behaviour during construction and subsequent performance in service
During mixing and laying, the penetration of binder generally reduces to about 70% of the pre-mixed value. This is known as ‘short term ageing’. This ageing continues during its service life, albeit at a lower rate, and is known as ‘long term ageing’. The most dominant of these mechanisms is oxidative hardening, and this is influenced by the thickness of the binder film, the air voids in the mixture and the temperatures experienced.

Pavement cracking can take several forms. The most frequent are

- longitudinal cracking, which occurs generally in the wheel track
- transverse cracking, which can be in any area and is not necessarily associated with the wheel path
- alligator or crocodile cracking, where longitudinal and transverse cracks link up to form a network of cracks
- reflective cracking, which results from an underlying defect (typically joints in concrete layers or cracks in hydraulically stabilised bases).

Traditionally, pavement design has only considered load-associated cracking, in which cracks are initiated at the underside of the asphalt base caused by repeated pavement flexure under traffic. This has led to the design of pavements of greater thickness to withstand the higher flow rates predicted for future commercial traffic. A programme of research carried out at the Transport Research Laboratory (TRL) (Nunn et al., 1997) to investigate how these thicker asphalt pavements (thicker pavements are those having at least 180 mm of total asphalt in the base, binder course and surface course – see Figure A3 in TRL 250 (Nunn et al., 1997)) were performing revealed that cracks in thick asphalt pavements invariably initiate at the surface and propagate downwards. It should be noted that such cracking in thicker pavements was confined to the surfacing (i.e. in the surface course and binder course only). Similar observations have been made by others (Schorak and Van Dommelen, 1995; Uhlmeyer et al., 2000).

18.3.1.1 Thermal cracking

Cracking that results from extreme cold is generally referred to as low temperature cracking, whereas cracking that develops from thermal cycling is normally referred to as thermal fatigue cracking. Thermal cracking will occur when the binder becomes too stiff to withstand the thermally induced stress, and it is related to the coefficient of thermal expansion and the relaxation characteristics of the mixture. Both these properties are related to the nature of the binder, and the risk of thermal cracking increases with the age of the pavement because the binder hardens as a result of oxidation or time-dependent physical hardening.

Two different thermal cracking mechanisms can occur. At low pavement temperatures, transverse cracks that run the full depth of the pavement can
suddenly appear. Pavement temperatures generally have to fall below about 
\(-30^\circ\text{C}\) to induce this form of cracking. Accordingly, documented cases of
low temperature cracking in the UK are extremely rare.

Under milder conditions, cracks may develop at a slower rate, taking several
seasons to propagate through the asphalt layers. This form of cracking
initiates at the surface and propagates relatively slowly with each thermal
cycle. This is generally described as ‘thermal fatigue cracking’.

The general mechanism responsible for these two forms of cracking is
considered to be similar. The main differences are as follows.

- Low temperature cracking is a single event phenomenon that is the
  result of the full depth of asphalt being put into thermal tension under
  conditions where stress relaxation cannot occur.
- Thermal fatigue cracking is more dependent on the properties of the
  surface course, and cracks first have to initiate at the surface and
  propagate through the surface course before they affect the lower
  asphalt layers.

### 18.3.1.2 Low temperature cracking

The mechanism of low temperature cracking is illustrated in Figure 18.5. The
asphalt layer is subjected to a tensile stress distribution with depth. These
stresses are caused by the contraction of the asphalt as it cools, and they are
a function of the temperature change and the relaxation characteristics,
stiffness modulus and coefficient of expansion of the asphalt. They are not
necessarily uniform, because pavement temperature can vary with depth.
These stresses can, potentially, cause a crack to propagate down from the
surface.

The coefficient of thermal expansion of the binder is an order of magnitude
higher (i.e. ten times higher) than that of the aggregate in the mixture. A
rough estimate for the thermal expansion coefficient of the mixture can be
determined using the equation (European Commission, 1999)

\[
\alpha_m = \frac{\alpha_b v_b + \alpha_a v_a}{v_b + v_a}
\]

![Figure 18.5 Thermal cracking mechanism](image-url)
where $a_m$ is the coefficient of expansion of the mixture, $v_b$ is the proportion of the binder by volume, $v_a$ is the proportion of the aggregate by volume, $a_b$ is the coefficient of thermal expansion of the binder, and $a_a$ is the coefficient of thermal expansion of the aggregate.

The coefficient of volumetric expansion of bitumen is approximately $6 \times 10^{-4}/\degree C$. Typically, the linear coefficient of thermal expansion of an asphalt is between $2$ and $3 \times 10^{-5}/\degree C$.

As an asphalt becomes colder, its tensile strength ($\beta_z$) initially increases and then begins to decrease as a result of micro-cracking in the binder matrix. This is caused by the differential contraction that results from the large difference between the coefficients of thermal expansion of the aggregate and binder. These fractures can be detected as acoustic events, using a sensitive microphone to record their increasing occurrence with falling temperature (Valkering and Jongeneel, 1991).

At the same time, thermal stress ($\sigma$) builds up as the material loses its relaxation ability and, at some point, the thermal stress will exceed the strength of the material. This defines the probable low temperature fracture temperature (spontaneous cracking). The difference between the tensile strength and the low temperature stress is known as the tensile strength reserve ($\Delta \beta_z$), and it is this reserve that is available to accommodate additional superimposed stresses (e.g., traffic-induced stresses). This is illustrated in Figure 18.6.

**Figure 18.6** Low temperature stress ($\sigma$) and tensile strength of asphalt ($\beta_z$) as a function of temperature
The curves shown in Figure 18.6 can be derived using the thermal stress restrained specimen test (TSRST) and the isothermal direct tensile test over a range of low temperatures (Jung and Vinson, 1992). Although expensive and time consuming, these tests are considered (King et al., 1993) to offer a reliable means of predicting the temperature at which a pavement will crack due to excessive thermal stresses. The results are reasonably reproducible, and the relative ranking of materials is consistent with field performance. For example, for asphalts manufactured using 60/80 pen bitumen, the TSRST tested cracking temperature is about −30°C (Fan et al., 2012).

Various studies of low temperature cracking have generally concluded that, in order to reduce cracking, the binder stiffness must not exceed some defined limit at the coldest pavement temperature (Bahia, 1991).

The bending beam rheometer (BBR) can be used to produce a master curve for the binder stiffness as a function of loading time and temperature. This curve is used to predict the temperature at which the binder has a stiffness of 300 MPa at a loading time of 60 s (Anderson et al., 1994).

In 1996, a research project was undertaken (Kandhal et al., 1996) to verify whether the BBR test results could have predicted the low temperature cracking of six AC-20 bitumens (ASTM, 2014). One bitumen sample was found to have had a stiffness exceeding 300 MPa at the minimum design temperature but it had not cracked during its 7 year service life.

Marasteanu (2004) investigated the relationship between the stiffness and the $m$ value calculated from the BBR experimental data and the development of thermal stresses in asphalt pavements. The analysis showed that the benefit of a high $m$ value for low temperature performance is not obvious. But it was also pointed out that

- in climates where the temperature stays at reasonably low values for extended periods of time, higher $m$ value binders may perform better as they allow more relaxation to occur
- in climates characterised by extremely cold temperatures, higher $m$ values result in worse performance, as thermal stresses develop faster and can result in the occurrence of cracking before relaxation takes place.

Jung and Vinson (1992) concluded that the TSRST provided an excellent indicator of low temperature cracking. Furthermore, King et al. (1993) demonstrated that results obtained using the TSRST correlated very well with the temperature prediction using the BBR ($R^2 = 0.96$). However, later work (Anderson et al., 2001) suggests that fracture toughness should be included in the applicable criteria when a polymer modified binder is used.
Low temperature cracking is of particular concern in Canada, North America, northern and eastern Europe and northern Asia, where temperatures can fall as low as $-40^\circ C$. To counter this problem, softer binders are often used. It is recognised that the risk of low temperature cracking is related primarily to the properties of the binder, with viscosity and temperature susceptibility of the binder being the most important properties. The risk increases as the hardness of the binder rises. Variations in the grading and type of aggregate have very little effect, and increasing the binder content reduces only slightly the susceptibility of the mixture to thermal cracking.

18.3.1.3 Thermal fatigue and load-associated surface cracking

Cracking in the surface of asphalt pavements is relatively common. These cracks can be transverse or longitudinal, and are usually located in the wheel tracks. Longitudinal wheel track cracking has often been regarded as evidence of conventional fatigue, in which cracks are assumed to have initiated at the bottom of the base and then propagated to the surface. However, investigation by coring thick asphalt pavements has invariably found that either the cracks partially penetrate the thickness of asphalt or, if the crack is full depth, the propagation is downwards rather than upwards. A typical example of this type of cracking in a major UK motorway is shown in Figure 18.7. This crack extended for several hundred metres. Coring showed that only at the most seriously cracked locations were there cracks that had progressed further than the top 100 mm of the asphalt (i.e. cracking was restricted to the surface course and binder course only). None of the cracks had penetrated the full thickness of the asphalt layers.

Investigations of a number of relatively thick UK motorways showed that this behaviour was typical. However, crack investigations by coring in the USA (Uhlmeyer et al., 2000) and Holland (Schorak and van Dommelen, 1995) have shown that, in fully flexible pavements with less than 160 mm of asphalt, the cracking is likely to be full depth. Above this thickness, an increasing proportion of the pavements have cracks that are confined to the asphalt surfacing (i.e. the binder course and the surface course). In the Dutch study, all pavements less than 160 mm thick had full depth cracking, and all pavements with 300 mm or more of asphalt had cracks that were confined to the top 100 mm of asphalt.

Penetration tests on the binder recovered from hot rolled asphalt surface courses in the UK study showed somewhat lower values in cracked areas compared with uncracked lengths. However, binder penetration is not always a good indicator of the susceptibility of the surface course to surface cracking, and it is evident that more detailed studies will be required to understand the loading, material and environmental factors that determine the initiation and propagation of cracks.
Surface cracking is not always longitudinal: transverse or block cracking can occur. This is not confined to wheel tracks, and it can occur at any location across the carriageway. Figure 18.8 shows an example of transverse cracks in a UK motorway that had carried traffic totalling some 30 msa (million standard axles) since it was laid 24 years previously. During this time, it had not been resurfaced or overlaid. The penetration of recovered binder from the surfacing was 15 dmm, and it was concluded that the cracking was not traffic related but due to brittle surfacing resulting from aged binder. As with longitudinal cracking, transverse cracks generally only penetrated to a depth of about 100 mm.

The phenomenon of surface cracking has received relatively little attention. However, there are a considerable number of observations of surface cracking of flexible pavements that span all climatic regions, from cold to tropical regions. This suggests that higher temperatures associated with the warmer
season cause binder to harden with age, and this reduces its capacity to withstand the thermal stresses generated during the cooler nights, particularly during the coldest season.

With all types of surface cracking, age hardening of the binder in the surface course will play a part. As stated above, this hardening over time progressively reduces the ability of the surface course to withstand the thermal and traffic generated stresses at the surface. Binder rheological studies (Nunn et al., 1997) have shown that the binder in the top few millimetres of the surface course becomes particularly hard with age.
Figure 18.9 shows an example of the results obtained using a DSR on binder extracted from the top 10 mm of the surface course and from between 20 and 30 mm below the surface.

Figure 18.9 shows that the binder close to the surface is substantially harder than binder that is deeper in the layer. Investigations of four sites that were 18 to 24 years old indicated that the penetration of the binder recovered from the top 10 mm of surface course was approximately 50% of that obtained from the lower layer. This aged, and hence hard, upper skin of the surface course is considered to be a major factor in the initiation of surface cracks.

18.3.1.4 Modelling surface cracking
The mechanism of surface cracking is complex and, to date, there is no completely satisfactory explanation of this phenomenon. It is now being recognised that tyre stresses can induce a tensile condition at the pavement surface and this can initiate a crack. De Beer et al. (1997) have shown that the tyre can transmit non-uniform horizontal and vertical stresses to the pavement, and pavement stresses in the locality of the tyre need to be taken into account in modelling crack behaviour. Finite element modelling predicts that a tensile condition can be created in the surface course close to the edge of
the tyre as shown in Figure 18.10. Similar conditions are also predicted near the edge of tyre treads (Myers and Roque, 2001).

These near-tyre tensile stresses, shown in red in Figure 18.10, extend only approximately 10 mm into the asphalt. Consequently, they may initiate a longitudinal crack in the wheel track, but another mechanism is required to account for crack propagation to any depth. There is increasing conviction that these surface cracks are propagated by thermal stresses.

Groenendijk (1998) concluded that the ageing of asphalt concrete at the surface combined with the non-uniform contact stress could result in critical tensile stress on the surface.

Traditionally, it was considered that there were two categories of cracking: top down and bottom up. As the names suggest, top down cracking starts at the surface, with bottom up cracking beginning at the underside of the base. Nunn et al. (1997) discovered that in thicker pavements cracking was likely to start at the surface and was found only in the surfacing (i.e. the surface course and the binder course) – there was no cracking in the base.

There is also a view that top down cracking is caused by the shear stress/strain in the surface course rather than by tensile stress. Bensalem et al. (2000) concluded that the shear strains on the vertical plane play a dominant role in top down cracking. In a research report from Michigan State University, Svasdisant (2003) pointed out that the cracks caused by shear failure along a vertical plane should incline at an angle from the...
vertical plane. Wang et al. (2003) suggested that both tensile type and shear type cracking could initiate top down cracking. Using strength theory and a finite element method to analyse stress/strain in asphalt pavements under non-uniformly distributed tyre loads, Jia et al. (2008) stated that top down cracking could be caused by the combination of tensile stress and shear stress.

Thermal fatigue is the mechanism responsible for the development of transverse surface cracks, with crack propagation resulting from the cyclic, daytime thermal stresses. However, transverse rolling cracks, sometimes seen at the time of compaction, are likely to play a major part in the initiation of transverse cracks.

The models predict (Nesnas and Merrill, 2002) that, for a thick asphalt pavement, a thermally propagated crack will stabilise at some point and will not propagate through the full thickness of the asphalt. On the other hand, the prediction for a thin pavement is that the rate of crack propagation will accelerate as the remaining thickness of asphalt is subjected to greater thermal and traffic loading.

18.3.1.5 Fatigue cracking
Fatigue cracking has received more attention from the research community than any other deterioration mechanism. Fatigue is the phenomenon of cracking under the repeated application of a stress that is less than the tensile strength of the material.

When a wheel load passes over a point in an asphalt pavement, the pavement flexes and a tensile strain is induced at the underside of the base layer. Continuous flexure and relaxation over many years carries with it the possibility of fatigue cracks initiating at the underside of the asphalt base and propagating upwards.

Fatigue is a major component of all modern analytical pavement design methods. These methods use a simple shift factor, or calibration factor, to relate the laboratory-determined fatigue life to the performance of the material in the road. The conditions and the behaviour of material in service are much more complex (Thrower, 1979) than those used in laboratory studies. For example, the nature of the stress system and long term physico-chemical changes in the binder are not considered.

The results of studies on the fatigue performance of the asphalt indicated that asphalt fatigue life was a function of the initial loss of stiffness in sinusoidal loading. Thus, it seemed logical to use the binder loss stiffness as the parameter to control mixture fatigue, as fatigue is attributable to cracking in the binder phase of the mixture. Therefore, SHRP uses the product of the complex modulus ($G^*$) and the sine of the phase angle ($\sin \delta$) to measure the fatigue
cracking (Deacon et al., 1997). This product is known as the SHRP fatigue parameter or fatigue parameter.

The fatigue parameter \( G^* \sin \delta \) is measured after a relatively small number of load cycles, and therefore cannot reflect the non-linear behaviour of a binder. Many experts agree that fatigue damage resistance cannot be predicted from linear visco-elastic properties alone, as \( G^* \sin \delta \) (Bahia et al., 2010). A number of studies have shown that there is a poor correlation between the linear visco-elastic \( G^* \sin \delta \) and mixture fatigue performance (Stuart and Mogawer, 2002; Tsai et al., 2005). In NCHRP Report 459 (Bahia et al., 2001), an analysis method was outlined that uses the ratio of dissipated energy of the binder to replace the SHRP fatigue parameter to quantify the amount of fatigue damage in an asphalt.

The development of the long-life pavement concept (Nunn et al., 1997) has demonstrated that long term changes in the binder properties can result in an increased fatigue life of the pavement. Also, in the USA, where this concept was subsequently developed, it has been suggested that, at low strain amplitudes, asphalt has a fatigue endurance limit (Newcombe et al., 2002).

The fatigue resistance of an asphalt is especially sensitive to binder volume. The simplest means of increasing the predicted fatigue life of the pavement is to construct the pavement using a binder-rich lower asphalt layer (Harm and Lippert, 2002).

18.3.1.6 Reflective cracking
A composite pavement consists of a continuously laid cementitious base under asphalt surfacing, or an existing cementitious pavement overlaid with asphalt. A regular pattern of thermally induced transverse cracks can appear in the cementitious base soon after laying, and these cracks begin to appear as reflection cracks in the asphalt surfacing several years later, as shown in Figure 18.11.

For some composite pavements, particularly those in which an existing cementitious pavement has been overlaid with asphalt and carries heavy traffic, as observed in China on the Hanyi expressway (Liao and Chen, 2010), these cracks were caused either by the crack in the cementitious layer opening and closing as a result of thermal expansion and contraction, or by a flexing and shearing action caused by wheel loads passing over the crack. These mechanisms will induce a stress concentration in the asphalt immediately above the crack, and this causes a crack in the asphalt to initiate and propagate towards the surface.

However, for some other composite pavements, particularly those consisting of a continuously laid cementitious base under asphalt surfacing, as
observed in the UK, extensive coring of in-service roads has shown that reflection cracks often start at the surface of the road and propagate downwards to meet an existing crack or joint in the underlying concrete layer (Figure 18.12). Furthermore, this study (Nunn, 1989) has shown that environmental effects, rather than traffic loading, cause reflection cracks in as-laid composite pavements. When the pavement is new, the surface course is ductile enough to withstand thermally induced stresses but, as the pavement ages, it will progressively lose this capability. The study showed that the occurrence of reflection cracking in as-laid pavements correlated with the amount of strain that the surface course could accommodate before cracking. This reduced with age, and it is related to the type and volume of binder used.

In their early stages of development, the cracks are not considered to present a structural problem. However, once they propagate through the asphalt layer, water infiltration and the pumping action of the traffic will weaken the foundation layers. At the same time, the load transfer across the slab will deteriorate, and under these conditions the cementitious slabs will move.
under heavy traffic, and further cracking, spalling and general deterioration will result. The reflection cracking of a strengthening overlay over a crack of this nature will be dependent on traffic-induced forces and the severity of the crack.

Modelling reflection cracking in as-laid composite pavements has generally treated the asphalt as a passive layer that has to respond to movements in the concrete layer. These models either assume that the thermal opening and closing of the crack in the cement bound layer or shearing caused by traffic will induce a high stress concentration in the asphalt immediately above the crack, causing it to initiate and propagate upwards. These models have not considered that

- the thermal coefficient of expansion of the asphalt surfacing is several times greater than that of concrete
- greater daytime temperature changes occur close to the surface
- age hardening of the binder is more severe close to the surface.

Figure 18.12 Core of reflection crack initiating from surface
Modelling by Bensalem (2001) takes into consideration all these factors and uses the finite element technique to model the behaviour of the asphalt and cement bound layers as a single system, as illustrated in Figure 18.13(a). Figure 18.13(b) shows the thermally induced stress contours predicted using this model. The model predicts that the highest thermal stresses will occur at the surface immediately above the crack in the cement bound base. Furthermore, it predicts that a compressive stress condition will exist at the underside of the asphalt layer adjacent to the crack in the cement bound layer. This implies that it is not possible for a crack to propagate upwards. This model
provides a qualitative explanation for the observed manner in which cracks initiate at the surface and propagate downwards.

However, for existing cement pavements overlaid with asphalt, the void under the cement concrete slab will cause additional deformation and change the stress/strain status in the pavement structure. Therefore, the modelling needs to take into account the supporting condition beneath the cement concrete slab.

### 18.3.2 Deformation

Deformation (often called ‘rutting’) may be restricted to one or more of the asphalt layers, or it may extend throughout the entire pavement and into the subgrade. It is classified as wearing, post-compaction and shearing rutting. In practice, however, the majority of deformation defects result from plastic deformation of the asphalt layers, mostly in the surface course, or surface course and binder course. Severe rutting can occur under moving or stationary traffic, and particularly under the high shearing stresses imposed by braking, accelerating, turning, overloaded traffic or long and steep sloping sections of road (Figure 18.14). The primary internal factor influencing plastic deformation is the composition of the mixture, while the primary external factors are stress and temperature. Plastic deformation is greatest at high service temperatures, for which 70°C may be taken as a maximum.

**Figure 18.14** Severe rutting in a long and steep sloping section after an extremely hot summer and overloading
in situ temperature. At such temperatures, the cumulative effect of repeated loadings of short duration will be determined by the binder viscosity. It has been estimated that during the long hot summer experienced in the UK during 1976, deformation in the wheel tracks of hot rolled asphalt surface courses was between two and four times the rate of an average UK summer.

Bitumen rheology is one of the most critical factors influencing the propensity of an asphalt to rut. The significance of the PI of the binder was confirmed by an extensive full scale road trial carried out on the Colnbrook bypass. Binders of widely differing rheological characteristics were employed in hot rolled asphalt surface course mixtures used in this project. Figure 18.15 shows the relationship between rut depth on the road after 8 years and the PI of the binder. The advantage of using higher PI binders is evident.

Work in several laboratories has shown that deformation occurring in a wheel tracking test can be predicted using the uniaxial unconfined creep compression test (Van de Loo, 1974, 1976). Creep curves of mixture stiffness \( S_{\text{mix}} \) plotted against binder stiffness \( S_{\text{b}} \) can be used to compare different mixtures and to predict their relative deformations if subjected to the same temperature and loading conditions.

The shape of a creep curve for a typical hot rolled asphalt mixture surface course is such that it can be approximated for most of its length by a straight line having a slope of 0.25. Thus

\[
S_{\text{mix}} = k(S_{\text{b}})^{0.25}
\]

Figure 18.15 Rut depth as a function of in situ penetration index (PI) measured on the Colnbrook bypass after 8 years in service (Dormon, 1969)
For two mixtures having $S_{\text{bit}}$ values in a known ratio,

$$S_{\text{mix}} \text{ ratio} = \left(S_{\text{bit}} \text{ ratio}\right)^{0.25}$$

For asphalts in similar road structures subjected to the same traffic

$$\text{Deformation} = k \left(1 / S_{\text{mix}}\right)$$

For two mixtures having $S_{\text{mix}}$ values in a known ratio,

$$\text{Deformation ratio} = 1 / S_{\text{mix}} \text{ ratio}$$

The above relationships can be used to compare the effect of changing the penetration and/or the PI of the binder.

Calculations made on this basis show that if the deformation of a 50 pen binder is regarded as unity, a mixture manufactured using a 30 pen binder subjected to the same temperature and loading regime will deform by 0.75 units. This assumes that the loading time for determination of viscosity and stiffness are the same. Conversely, if a 100 pen binder is used, 1.5 units of deformation will occur (Table 18.3(a)).

Increasing the PI of the binder significantly improves the resistance to deformation. For example, at $40^\circ C$ a binder with a penetration of 40 and a PI of 0.5 has a viscosity of $4 \times 10^4$ Pa-s, whereas a binder with the same penetration but having a PI of +2.0 has a viscosity of $6 \times 10^5$ Pa-s at $40^\circ C$. This gives a factor of 15 in the viscosity, and hence in the stiffness of the two binders. Thus, applying the 0.25 power formula stated above gives an increase of approximately 2 in mixture stiffness. Accordingly, the deformation using a mixture made with a binder having a PI of +2.0 would be half of one made with a standard binder (Table 18.3(b)). Figure 18.16 shows the above theoretical relationship for relative deformation plotted as a function of PI. This clearly shows that the theoretical relationship is supported by both simulative laboratory wheel tracking tests and full scale road trials.

Correlation of binder properties with both Marshall and wheel tracking tests (Jacobs, 1981) shows that the relationship between binder penetration and Marshall stability is poor, whereas the softening point relates very well to both Marshall stability and deformation in the wheel tracking test (Figure 18.17).

### Table 18.3 (a) Relative deformation of mixtures at $40^\circ C$, effect of binder penetration

<table>
<thead>
<tr>
<th>Binder penetration</th>
<th>PI</th>
<th>Viscosity at $40^\circ C$: Pa-s</th>
<th>Relative $S_{\text{bit}}$</th>
<th>Relative $S_{\text{mix}}$</th>
<th>Relative deformation</th>
</tr>
</thead>
<tbody>
<tr>
<td>100</td>
<td>0.8</td>
<td>$6 \times 10^3$</td>
<td>0.2</td>
<td>0.67</td>
<td>1.5</td>
</tr>
<tr>
<td>60</td>
<td>0.6</td>
<td>$18 \times 10^3$</td>
<td>0.6</td>
<td>0.88</td>
<td>1.1</td>
</tr>
<tr>
<td>50</td>
<td>0.5</td>
<td>$30 \times 10^3$</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>40</td>
<td>0.5</td>
<td>$40 \times 10^3$</td>
<td>1.3</td>
<td>1.07</td>
<td>0.94</td>
</tr>
<tr>
<td>30</td>
<td>0.4</td>
<td>$90 \times 10^3$</td>
<td>3.0</td>
<td>1.32</td>
<td>0.75</td>
</tr>
</tbody>
</table>
For every 5°C increase in softening point, the Marshall stability increased by over 1.3 kN (see Figure 18.17 (b)) and the wheel tracking rate almost halved (see Figure 18.17 (c)).

The SHRP programme has provided a binder parameter, designated the ‘rutting factor’ (the ratio of the complex modulus and the sine of the phase angle, i.e. $G^*/\sin \delta$) representing a measure of the high temperature stiffness of the binder. Laboratory research had found that the binder parameter, $G^*/\sin \delta$, was related to the depth of rutting in wheel tracking results. An earlier indication of the suitability of $G^*/\sin \delta$ as the key specification criterion for rutting was contained in the paper by Petersen et al. (1994).

Mitchell et al. (2004) have used the Federal Highway Administration’s two accelerated loading facility machines to validate the rutting factors of Superpave binder properties. However, no overall relationship between $G^*/\sin \delta$ and pavement rut depth was found.

<table>
<thead>
<tr>
<th>Binder penetration</th>
<th>PI</th>
<th>Viscosity at 40°C, Pa·s</th>
<th>Relative $S_{\text{bit}}$</th>
<th>Relative $S_{\text{mix}}$</th>
<th>Relative deformation</th>
</tr>
</thead>
<tbody>
<tr>
<td>40</td>
<td>−0.5</td>
<td>$4 \times 10^4$</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>40</td>
<td>+0.5</td>
<td>$1.5 \times 10^5$</td>
<td>3.8</td>
<td>1.41</td>
<td>0.71</td>
</tr>
<tr>
<td>40</td>
<td>+2.0</td>
<td>$6 \times 10^5$</td>
<td>15</td>
<td>1.97</td>
<td>0.50</td>
</tr>
</tbody>
</table>

Theoretical line
Field trials – Colnbrook
– Blackwall
Laboratory wheel tracking

**Figure 18.16** Relative deformation as a function of penetration index
Figure 18.17 Relationship between bitumen properties and Marshall stability and wheel tracking rate (Jacobs, 1981)
18.3.2.1 The zero shear viscosity concept

In terms of deformation, it is thought that the binder contribution to this mode of distress is primarily associated with viscosity. However, identifying the relevant viscosity in terms of shear, shear rate and stress is a complex problem, and a single quantifiable regime would have to be established for assessment purposes. Some pavement design methods are based on the assumption that the material response to traffic loading is within the linear regime, and to remain consistent with this premise the corresponding linear viscosity of the binder should be examined. This is the zero shear viscosity, \( \eta_0 \), sometimes termed ‘Newtonian viscosity’ as stress is proportional to shear. In this regime, stress and strain are linearly proportional to each other, so the stiffness modulus is independent of stress or strain level, and the resultant viscosity, \( \eta_0 \), is independent of the shear rate.

By measuring binder properties within the linear regime, conclusions can be made as to the likelihood of a binder contributing to or mitigating the rutting process. Zero shear viscosities can be measured using a dynamic shear rheometer, which also permits the elastic and viscous components to be identified. The higher the value of zero shear viscosity, the lower the influence of the binder in relation to deformation. Research has shown that the useful upper limit on zero shear viscosity of unaged binder is \( 10^5 \) Pa·s, above which further resistance to deformation gives diminishing returns (Phillips and Robertus, 1996). For binders that exhibit zero shear viscosities below this value, the relative value of the zero shear viscosity coupled with the elastic component of the binder can be used to interpret the deformation behaviour of the binder.

18.3.2.2 Multiple stress creep recovery

Another test, called the multiple stress creep recovery test (MSCR), has been suggested, and of particular interest is the resistance to deformation at high temperature under the repetitive action of heavy ‘traffic’ loadings. This test was proposed by D’Angelo et al. (2007) in the USA, and features in ASTM standard D7405 published in 2008 (ASTM, 2008). The MSCR test is a standard rheological test based on the repeated creep recovery test (D’Angelo et al., 2007).

In the MSCR test, a specimen is subjected to a constant load for a fixed time period, \( t_1 \). The resultant deformation is measured as a function of time and applied stress \( \tau \). At a time \( t_1 \) (in the MSCR test \( t_1 = 1 \) s), the specimen is allowed to recover at zero load for a fixed time period, \( t_2 \) (in the MSCR test \( t_2 = 9 \) s). The time dependent and stress dependent recoverable shear deformation or strain is the difference between the maximum shear strain \( \gamma(t_1) \) and the shear strain at \( \gamma(t_2) \) with \( t_2 > t_1 \). The total recoverable deformation \( \gamma_1 \) for \( t_2 - t_1 \) is a measure of the elasticity of the material (i.e. the
Influence of binder properties on the performance of asphalts

mechanical energy stored in the sample during the creep phase). The non-recoverable part relates to the viscosity.

In addition, a property called ‘non-recoverable compliance’, $J_{nr}$, was also derived from the non-recovered strain normalised for the applied stress during the creep portion of the test ($J_{nr} = \gamma_{nr}/\tau$). According to a Federal Highway Administration study (FHWA, 2011), this $J_{nr}$ value at a stress of 3.2 kPa has been shown to be an indicator of the rutting resistance of an asphalt to deformation under repeated load. The study by D’Angelo et al. (2007) showed that reducing $J_{nr}$ by half typically reduced rutting by half. Therefore, the $J_{nr}$ value obtained at a stress of 3.2 kPa, which is based on the response of binders at their performance grade temperature, will be used in the MSCR high temperature specification as an indicator of rutting resistance.

18.3.2.3 The ‘softening point’ of asphalt concept

Considering that the aggregate grading, binder content and design air voids are all important factors that will have an effect on the propensity of a particular asphalt to rut, any attempt to establish the direct relationship between binder parameters and pavement deformation must take account of these factors. Accordingly, any method for predicting rutting must investigate the temperature susceptibility of asphalt.

Xu and Ji (2006) have developed a new method to evaluate the anti-rutting performance of asphalt. A series of wheel tracking tests is carried out at different temperatures. Following this, the relationship of dynamic stability (the number of wheel loads necessary to cause a deformation of 1 mm) versus temperature is plotted. It was found that the whole plot can be divided into three stages as follows.

- **Stage 1**: as the temperature increases, the dynamic stability slowly decreases.
- **Stage 2**: as the temperature increases, the dynamic stability remains constant, with the maximum temperature in this stage being described as the ‘softening point’ of the asphalt.
- **Stage 3**: there is a dramatic increase in dynamic stability.

These stages are illustrated in Figure 18.18.

The explanation for this behaviour is as follows. During stage 1, as the temperature increases, the stiffness and viscosity of the binder drop quickly, which results in deformation of the asphalt. During stage 2, although the stiffness and viscosity of the binder are continually decreasing, the loading is carried by the mechanical interlock of the asphalt, and the dynamic stability remains constant. During stage 3, a further decrease in viscosity causes the binder to
act as a lubricant between the particles of aggregate, with a consequent loss of strength, resulting in greater deformation.

It was found that the ‘softening point’ of a conventional asphalt has good correlation with the softening point of the constituent bitumen, and is about 10°C higher than the softening point temperature of that bitumen. However, the ‘softening point’ of a styrene–butadiene–styrene (SBS) modified asphalt has poor correlation with the softening point of a binder but has a good correlation with the equivalent softening point T800 (the temperature at which the penetration of the binder is 800 dmm) and is also about 10°C higher than the equivalent softening point of binder T800.

The significance of the ‘softening point’ of an asphalt lies in its validity for selection of binder and asphalt to meet the requirement of deformation resistance according to the highest pavement design temperature. The advantage is that it takes into account binder, aggregate and grading characteristics when assessing the deformation resistance of asphalts, rather than only a binder property.

The research on the ‘softening point’ of asphalts has suggested that the grading of the asphalt is a key element in determining anti-rutting performance. An asphalt having a grading in which there is a high degree of stone-to-stone contact always exhibits high rutting resistance, as shown in Figure 18.19. Indeed, this substantiates the view long held by experienced pavement engineers that asphalt concrete mixtures generally have good
resistance to deformation, while hot rolled asphalts perform poorly in relation to rutting. This is why the use of a hot rolled asphalt binder course may cause concern due to its propensity to rut.

18.3.3 Fatting up
Fatting up (sometimes described as ‘flushing up’) is a migration of binder from the asphalt to the top of the pavement, resulting in a smooth, shiny surface that has poor resistance to skidding in wet weather. One reason why fatting up occurs is as a result of secondary compaction of the aggregate in the mixture by traffic. The void content is reduced, eventually squeezing binder from within the structure to the surface. This will be exacerbated if the binder content is too high, resulting in a thicker binder film, or if the void content, after completion of compaction, is too low (Youtcheff, 2002). Fatting up is most likely to occur at high service temperatures. Accordingly, increasing the softening point or viscosity of the binder at 60°C will limit this failure mechanism.

18.3.4 Fretting
Fretting is the progressive loss of interstitial fines from the road surface. It occurs when traffic stresses exceed the breaking strength of the asphalt itself or the asphalt mortar, depending on the nature of the mixture. Fretting is more likely to occur at low temperatures, and at short loading times when the stiffness of the binder is high.
The major factors influencing fretting are the binder content of the mixture and the degree of compaction. Loss of aggregate can be due either to the loss of adhesion between the aggregate and the binder or to brittle fracture of the binder film connecting particles of aggregate. The first condition should not arise if suitable aggregates and/or binders are selected. In the majority of cases, fretting is associated with a low degree of compaction or inadequate binder content. The choice of binder grade may control the degree of compaction that can be achieved. This is especially true if operations are being conducted in adverse weather conditions or if compaction is effected at temperatures below which the asphalt can be properly compacted. The higher the penetration of the binder or the lower the stiffness at low temperatures, the greater will be the resistance to fretting.

Fretting is also often associated with wet weather. It is well known that many types of asphalt defect are caused or increased in severity by the presence of water. Water penetrating the binder–aggregate mixture causes moisture damage: early loss of strength and durability due to loss of adhesion between the binder and the aggregates or the loss of cohesion in the binder.

Over many years, researchers have established that moisture damage, one of the major causes of pavement failure, is affected by many factors. More recent studies are those by Petersen (2002), Thomas et al. (2002), the Pennsylvania Transportation Institute (2002) and Caro et al. (2010). These and other studies have established that the parameters relevant to fretting defects are

- the chemistry of the binder and binder additives
- the rheology of the binder
- the chemistry of the binder–aggregate interaction
- the asphalt film thickness
- the aggregate surface chemistry and shape characteristics
- the surface energy
- the air void distribution and permeability
- the climate.

The most significant of the above factors are the binder chemistry, the binder rheology, the aggregate surface chemistry and the physical properties of the aggregate.

A recent study has addressed the effect of water vapour on asphalt pavement materials and found it to be significant. It was found that water vapour from subgrade soils migrates upwards into the pavement. This water vapour accumulates in the pavement at a rapid rate and reaches near-saturated vapour pressure within a period of 6 months. Moreover, wetting processes
in the pavement layer brought about by sub-surface water vapour diffusion occurs day and night. Therefore, the water presence attributable to vapour diffusion in the asphalt surface layer is one of the major water movement mechanisms in pavement, and it greatly accelerates the deterioration of the asphalt (Tong et al., 2013).

18.3.4.1 The effect of ageing
The ageing of the mortar is attributed to the effects of oxygen, ultraviolet radiation and temperature. The effects of ageing of asphalts can be classified in two major groups:

- short term ageing, which involves the loss of volatiles and oxidation of the bitumen during production, transportation, laying and compaction of the asphalt
- long term ageing, which is predominantly the continuation of the oxidation process during the service life of the asphalt pavement.

The ageing process is influenced by the type of asphalt: for example, a higher rate of ageing is expected in a porous asphalt compared to a dense asphalt (Roberts et al., 1996).

In asphalt surface courses, the bituminous mortar is subjected to shear, tension and/or compression, so the strength and relaxation behaviour of the bitumen are important. However, the effect of ageing increases the stiffness of the asphalt. This ageing also results in a change in creep and relaxation properties of the bitumen, resulting in higher susceptibility to fracture due to brittleness at low temperatures and a decrease in healing potential at intermediate temperatures (Yero, 2012).

Ageing changes the rheological, mechanical and chemical properties of the binder. The changes in the properties of the binder, and therefore the asphalt, can be quantified by means of ageing indicators. Ageing indicators commonly used in practice are either empirical parameters describing changes in binder properties, including changes in the visco-elastic behaviour of the binder, or a measure of the change in the chemical composition of the binder, indicating a change in colloidal structure after ageing (Hagos, 2008).

18.3.4.2 The effect of water
Water infiltration has a negative effect on the material characteristics of the individual components of the asphalt, and damages the bond between the components, which may result in ravelling of the surface. Adhesive failure is largely attributed to damage caused by the effect of water, and cohesive failure is caused when the stress levels in the binder/mortar are exceeded.
The introduction of water into an asphalt increases the chance of cohesive failure due to the ‘softening’ of the binder. Further studies have shown that the water damage phenomenon is an effect of advective transport (a phenomenon causing the mastic films in an asphalt to be washed away when the material is subjected to a moving flow of water) and diffusion, leading to mastic–aggregate interface failure and dispersion of the mastic. The analyses showed that water damage in asphalts is highly dependent on the adsorption characteristics of the aggregate–mastic system, as well as the diffusion and dispersion characteristics of the mastic. It is deduced that, by minimising the susceptibility of the mastic, the damage to the mastic will be solely due to advective transport, and damage due to dispersion phenomena will then be limited (Kringos and Scarpas, 2005).

18.3.4.3 Fretting of asphalt concrete surface courses
The following two different forms of fretting are identifiable in asphalt concrete surface courses.

- **Superficial fretting.** This involves the loss of interstitial fines from the road surface. The material subsequently closes up under traffic without further deterioration or detrimental effect on performance.

- **Severe fretting.** If superficial fretting does not close up under traffic, the loss of interstitial fines results in a reduction of the level of mechanical interlock that is likely to lead to the loss of coarse aggregate on the surface and a substantial reduction in internal cohesion due to the ingress of water. The presence of water breaks down the adhesive bond between the binder and the aggregate resulting in stripping of the binder (i.e. a physical separation of the binder from the aggregate). In addition, the uncoated fines and water form a slurry that is pumped through the material, as a result of pore pressures induced by moving traffic, abrading the coarse aggregate and exacerbating the problem. This will eventually lead to collapse of the material and a lack of internal stability, often visible on the road surface as slight rutting or as potholes that are bowl-shaped holes of various sizes.

Some research results have suggested that adhesive failures and thin film cohesive failures need to be differentiated. Designing binders with improved adhesion properties and/or improved cohesion properties could be an essential step in improving the resistance of the aggregate–binder system to moisture damage (Kanitpong and Bahia, 2003).

A number of procedures and recommendations to eliminate or minimise moisture damage have been formulated. One of those procedures involves treating the asphalt with an anti-stripping agent such as hydrated lime or other commercially available anti-stripping additives (Kennedy and Ping, 1991).
In addition, other studies have identified a range of approaches to this type of defect (Bagampadde et al., 2006; Hefer et al., 2005), namely

- theory of (weak) boundary layers
- mechanical theory
- electrostatic theory
- chemical bonding theory
- thermodynamic theory.

It is hoped that better use of additives, higher testing frequencies and improved testing protocols will result in better in-service performance.

18.3.4.4 Fretting of hot rolled asphalt surface courses

In 1984, a survey was carried out by the British Aggregate Construction Materials Industries (BACMI, now the MPA – Mineral Products Association) (White, 1989) as part of a wider survey into chipping loss by the Institution of Highways and Transportation (Johnson and Salt, 1985a, 1985b). The BACMI survey showed that over half of the locations where chipping loss had occurred were high stress low speed sites such as roundabouts and junctions, where a minimum 1.5 mm texture depth had been needlessly specified.

Fretting of a hot rolled asphalt surface course may well follow a failure to embed the precoated chippings properly. Failure to achieve adequate chipping embedment usually occurs as a result of one or more of the following factors

- the asphalt is unworkable (i.e. too stiff)
- the temperature of the asphalt is too low
- the asphalt cools quickly due to poor weather conditions during laying
- excessive application of precoated chippings
- inadequate compaction of the material (see Figure 18.20).

One method of treating patches where the chippings are inadequately embedded is to reheat the surface using an infrared heater and, when the asphalt is sufficiently hot, embed the chippings further into the asphalt. Used judiciously, this technique should not significantly harden the binder except at the surface. This hardened surface material will be eroded by traffic over time to expose the aggregate in the asphalt, a situation that will not threaten the long term integrity of the surface. However, this technique is often difficult to apply, requires great care and is not always successful. The best way of avoiding such problems is to ensure proper chipping embedment when the surface course is laid.

Some chipping loss may occur as a result of a failure to achieve or maintain bond between the precoated chippings and the asphalt. It has been found that the binder on precoated chippings can be ‘coked’ and made...
non-adhesive by storage of the chippings in large stockpiles immediately after manufacture. To avoid this possibility, chippings should be stored no higher than 1 m. The adhesivity of the binder coating can be checked using the hot sand test (BSI, 2005).

If the chippings have plucked out of the asphalt, the asphalt mortar is exposed to the rigours of the prevailing traffic. It is likely that this material will be poorly compacted and, as a result, it will fret rapidly.

### 18.3.5 Ravelling

Ravelling differs from the two fretting modes described above in that it involves plucking out of the asphalt itself (i.e. not just interstitially). It occurs when individual aggregate particles move under the action of traffic. If the tensile stress (induced in the binder as a result of the movement) exceeds the breaking strength of the binder, cohesive fracture of the binder will occur, and the aggregate particles will be detached from the road surface. Thus, ravelling is most likely to occur at low temperatures and at short loading times, when the stiffness of the binder is high. Obviously, a high binder content (and preferably a bitumen appropriately modified) will increase the ravelling resistance of the asphalt under high frequency traffic stress. A 1977 survey carried out by the Asphalt and Coated Macadam Association (ACMA, subsequently BACMI, then the QPA and currently the MPA) established that the use of higher binder contents and softer grades of binder will limit these fretting occurrences, while binder hardening may also lead to fretting.

Ravelling in asphalt concrete pavements depends very strongly on the bond between bitumen and the aggregate. Some studies found that the chemistry of both the binder and aggregate surface determine the degree of water
sensitivity of the binder–aggregate bond. If the bond is poor, adhesion failure could occur at the binder–aggregate interface, leading to stripping and ravelling in asphalt pavements (Kandhal and Rickards, 2001). A methodology has been developed to establish a relationship between moisture content and the reduction in strength of the mastic–aggregate bond (Copeland et al., 2006).

Recent studies have shown that the adhesive and cohesive characteristics of asphalt surface courses are significant factors in determining whether the asphalt may be susceptible to ravelling (Kanitpong and Bahia, 2003).

The adhesive characteristics are those that are responsible for the adhesion between aggregates and the bituminous mortar, while the cohesive characteristics are those that are responsible for the cohesion in the bituminous mortar. Because of the very nature of asphalt surface courses, the cohesive and adhesive characteristics are strongly influenced by the effects of ageing and water.

18.3.6 Skid resistance

The formal UK definition of skid resistance is ‘the frictional properties of the road surface measured using a specified device under standardised conditions’ (Highways Agency et al., 2004). A less technical definition would be a measure of the ease with which a vehicle stops when the brake is applied. The higher the skid resistance, the shorter the distance a vehicle takes to stop. Improving the skid resistance of a road is often the reason for surface dressing.

Maintaining adequate levels of skid resistance reduces accidents, meaning fewer injuries and deaths in road accidents. In countries where skid resistance standards apply, a loss of skid resistance is likely to be the most common reason for maintenance treatment.

The skid resistance of a road is determined by the microtexture of the aggregate and the macrotexture of the road surface itself. Means of measuring skid resistance are discussed in section 21.1.2.4.

The microtexture, or degree of polish of the exposed aggregate, is primarily dependent on the petrographic characteristics of the aggregate and the intensity of traffic. The macrotexture of the road surface is dependent not only on the type of surface course but also on the characteristics of the binder used in the surface course.

As mentioned above, restoration of an adequate level of skid resistance on an existing road may be achieved by surface dressing. It can also be effected by replacing the surface course. Whatever maintenance measure is chosen, one of the functions of the binder in the surface dressing or surface
course is to provide adequate resistance to the forces between the aggregate particles in the chosen system under conditions of compression, shear and tension. Accordingly, the binder should have high cohesive strength and an excellent ability to adhere to the aggregate, in addition to possessing good visco-elastic properties. Binder durability is also required to ensure retention of adequate rheological, cohesive and adhesive properties during the service life of any replacement system.

An ‘ideal’ asphalt pavement is supposed to be one that possesses exceptional performance, exhibiting minimal surface distress and having a smooth surface over the design period (Von Quintus, 2009). Good materials selection, design procedures and construction practices are essential for ideal asphalt pavements. Eliminating segregation and moisture damage, achieving good compaction (high density and low air voids), constructing good longitudinal joints and maintaining adequate bond between pavement layers improves pavement performance, increasing the service life and reducing maintenance costs.

References
Influence of binder properties on the performance of asphalts


Adhesion of bitumen and moisture damage in asphalts

The primary function of bitumen in an asphalt is to act as an adhesive. Bitumen is required either to bind aggregate particles together or to provide a bond between particles and an existing surface. When used in an asphalt, bitumen binds together the aggregate, sand and filler that comprise the mixture. In a pavement, the bitumen needs sufficient cohesive and adhesive strength to ensure minimum loss of aggregate under the shear stresses induced by traffic. Although the incidence of premature failure attributed to adhesion is relatively rare, failures, when they occur, may involve considerable expense. The need to ensure adhesion between the aggregate and the bitumen is, therefore, very important.

The adhesion of bitumen to most types of dry and clean aggregate presents few problems. However, aggregate is easily wetted by water, the presence of which can result in unexpected problems. These may occur at any time during the life of an asphalt, from the initial coating of the aggregate during the mixing process to its time in service, when it has to maintain an adequate bond between the bitumen and aggregate under traffic conditions. Damage to asphalts in the presence of water is a complex phenomenon and has been the subject of research over many decades. Although it can be described and quantified to an extent using thermodynamic first principles, in practice moisture damage is a complex phenomenon and is commonly assessed using simulative tests in the laboratory.

Measurement of adhesion is notoriously difficult to achieve in a reproducible manner, and many test methods have been employed to measure the combined forces of adhesion and cohesion, often referred to as ‘adhesive strength’. Several laboratory procedures have been developed to assess the dry strength of asphalts and their adhesive strength in the presence of water, typically termed ‘moisture (induced) damage’.
The aim of this chapter is to introduce the fundamentals of adhesion and cohesion, draw attention to aggregate–bitumen adhesion and the ways in which it can be assessed, and how it may be possible to reduce the possibility of premature failures in service resulting from a lack of adhesion.

19.1 Thermodynamic principles of adhesion, cohesion and surface free energy

A relationship exists between Gibbs free energy (the energy associated with a chemical reaction that can be used to do work), work of adhesion and surface energy, which provides a fundamental definition of adhesion and cohesion.

In terms of cohesion, the total work expended per unit of surface area in forming two surfaces is equal to twice the surface energy per unit of surface area

\[ W^c = 2\gamma \]  

(19.1)

where \( W^c \) is the work of cohesion and \( \gamma \) is the surface energy.

When two dissimilar materials form an interface by being in intimate contact this is known as adhesion, and the work of adhesion \( (W^a) \) can be defined by means of the Dupré (1869) equation as

\[ W^a_{12} = \gamma_1 + \gamma_2 - \gamma_{12} \]  

(19.2)

where \( \gamma_i \) is the surface energy of the \( i \)th material and \( \gamma_{12} \) is the interfacial energy between the two materials in contact.

When a drop of liquid is placed on a clean smooth horizontal surface, it either spreads over the solid surface or takes the shape of a drop with a finite contact angle between the solid and liquid phases, as shown in Figure 19.1. Contact angles are commonly used to measure the surface energy of solids based on the relationship between contact angles, surface energy of solids, wetting of solids and thermodynamic considerations.

![Figure 19.1](image-url)
The properties of the solid–liquid, solid–vapour and liquid–vapour interfaces can normally be described as a three phase boundary. Young proposed an equation to obtain surface tension from the contact angle formed when a drop of liquid is placed on a perfectly smooth, rigid solid (Young, 1805). In the contact angle experiment presented in Figure 19.1, the contact angle \( \theta \) is defined as the angle between the solid–liquid interface and the tangent of the liquid–vapour interface. If the liquid under consideration is water, the schematic diagram on the left hand side would be an example of a hydrophilic (water loving) surface, as is evident from the drop that tends to spread over the solid due to a favourable interaction between the interfacial forces. The contact angle on the right hand side, however, is greater than 90°, indicating that water does not spread readily over this surface, and this is thus a hydrophobic (water hating) surface.

The surface energy \( \gamma \) of a solid or liquid can be defined as the reversible work required to create a unit area of new surface. Various theories can be used to describe the interaction or intermolecular forces that contribute to the surface energy of a material (Fowkes, 1962; Good, 1966; Owens and Wendt, 1969; Van Oss et al., 1988). For bitumen and aggregate, the three component acid–base theory, or Good–Van Oss–Chaudhury theory, is most commonly used with the total surface energy being expressed as (Good and Van Oss, 1991)

\[
\gamma = \gamma_{LW} + 2\sqrt{\gamma^+ \gamma^-} \hspace{1cm} (19.3)
\]

where \( \gamma \) is the total surface energy, \( \gamma_{LW} \) is the Lifshitz–van der Waals component of the surface energy, \( \gamma^+ \) is the Lewis acid component of surface interaction, and \( \gamma^- \) is the Lewis base component of surface interaction.

When the system is in equilibrium, the relationship between the contact angle and the surface energies of the solid and liquid is given by Young’s equation as

\[
\gamma_{SV} = \gamma_{SL} + \gamma_{LV} \cos \theta \hspace{1cm} (19.4)
\]

where \( \gamma_{SV} \) is the surface free energy of the solid in equilibrium with the vapour, \( \gamma_{SL} \) is the surface–liquid interfacial free energy, and \( \gamma_{LV} \) is the surface free energy (surface tension) of the liquid in equilibrium with the vapour.

Equations (19.3) and (19.4) can, therefore, be combined to form the Young–Dupré equation, which is the starting point for any method that utilises contact angles to obtain surface energies by relating the contact angle to the work of adhesion.

\[
W_{SL} = \gamma_{LV}(1 + \cos \theta) \hspace{1cm} (19.5)
\]

where \( W_{SL} \) is the work of adhesion between the solid and the liquid.
The work of adhesion between two materials, such as bitumen and aggregate, can be calculated by performing surface energy calculations on both materials and then using Equation (19.2). If the value of adhesion is positive, it means that the two phases of the material tend to bind together, with a lower magnitude dictating the likely mode of failure.

It is also very useful to know the surface energy of adhesion in the presence of water when considering moisture damage. For the general case, the work of adhesion for two different materials in contact within a third medium, \( W_{132} \), can be explained by the following equation

\[
W_{132} = \gamma_{13} + \gamma_{23} - \gamma_{12}
\]  

(19.6)

Using the process proposed by Van Oss and colleagues (Good and Van Oss, 1991; Van Oss et al., 1988), Equation (19.7) can then be used to calculate the adhesion between bitumen and aggregate in the presence of water, where the subscripts 1, 2 and 3 represent bitumen, aggregate and water, respectively.

\[
W_{132} = 2\gamma_3^{1W} + 2\sqrt{\gamma_1^{1W}\gamma_2^{1W}} - 2\sqrt{\gamma_1^{1W}\gamma_3^{1W}} - 2\sqrt{\gamma_2^{1W}\gamma_3^{1W}}
\]

\[
+ 4\sqrt{\gamma_3^+\gamma_3^-} - 2\sqrt{\gamma_3^+(\sqrt{\gamma_1^- + \sqrt{\gamma_2^-}) - 2\sqrt{\gamma_3^-} (\sqrt{\gamma_1^+ + \sqrt{\gamma_2^+})}}
\]

\[
+ 2\sqrt{\gamma_1^+\gamma_2^-} + 2\sqrt{\gamma_1^-\gamma_2^+}
\]  

(19.7)

The two bond energy parameters in the dry condition [Equation (19.2)] and for the bitumen–aggregate system in water [Equation (19.7)] can be used to predict the moisture sensitivity of asphalts.

### 19.2 Factors affecting bitumen–aggregate adhesion

In practice, many factors (in addition to thermodynamics) influence the bitumen–aggregate bond; the importance of aggregate mineralogical composition has been recognised for many years (Saal, 1933; Winterkorn, 1936). Failure of the aggregate–bitumen bond is commonly referred to as ‘stripping’.

One of the main factors is the type of aggregate. This has a considerable influence on bitumen adhesion due to differences in the degree of affinity for bitumen. The vast majority of aggregates are classified as ‘hydrophilic’ (water loving) or ‘oleophobic’ (oil hating). Aggregates with high silicon oxide content (e.g. quartz and granite [i.e. acidic rocks]) are generally more difficult to coat with bitumen than are basic rocks such as basalt and limestone. The majority of adhesive failures have been associated with siliceous aggregates such as granites, rhyolites, quartzites, cherts, etc. The fact that satisfactory performance is achieved with these same aggregates and that
failures occur using aggregates that have good resistance to stripping (e.g. limestone) emphasises the complexity of bitumen–aggregate adhesion, and raises the possibility that other factors may play a role in the failure.

Other factors affecting the initial adhesion and subsequent bond are the surface texture of the aggregate, the presence of dust on the aggregate and, to a lesser extent, the degree of acidity of the water in contact with the interface. It is generally agreed that rougher aggregate surfaces have better adhesion characteristics. However, a balance is required between wetting of the aggregate (smooth surfaces being more easily wetted) and rougher surfaces, which hold the bitumen more tenaciously once wetting has been achieved.

Physio-mechanical adsorption of bitumen into the aggregate depends on several factors, including the total volume of permeable pore space, the size of the pore openings, and the viscosity and surface tension (surface energy) of the bitumen (Thelen, 1958). The presence of a fine microstructure of pores, voids and micro-cracks can bring about an enormous increase in the adsorptive surface available to the bitumen. It has also been shown (Plancher et al., 1977; Scott, 1978) that fractions of the bitumen are strongly adsorbed in the aggregate surface to a depth of approximately 180 Å (18 × 10⁻⁹ m).

It has been suggested that the good mechanical bond achieved on a rough aggregate can be more important than the aggregate mineralogy in maintaining bitumen–aggregate adhesion (Lee and Nicholas, 1957). The properties of the bitumen are also important in the acquisition and subsequent retention of the bitumen–aggregate bond. In particular, the viscosity of the bitumen during coating and in service, polarity and constitution all influence the adhesion characteristics. However, it is the nature of the aggregate that is, by far, the most dominant factor influencing bitumen–aggregate adhesion.

Table 19.1 summarises the main factors that influence bitumen–aggregate adhesion. It is considered that approximately 80% of these factors are controllable during production and construction.

19.3 Disbonding mechanisms in asphalts

Many studies have been carried out to determine the mechanism of bitumen disbonding in asphalts (Asphalt Institute, 1981; Hughes et al., 1960; Taylor and Khosla, 1983). There are two main methods by which the bitumen–aggregate system may fail: adhesive and cohesive mechanisms. If the aggregate is clean and dry and the mixture is effectively impermeable, the mode of failure will be cohesive. However, in the presence of water, the failure mode will almost certainly be due to a loss of adhesion caused by stripping of the
bitumen from the aggregate surface. Several mechanisms of disbonding are possible, and these are discussed below.

The description above is an oversimplification, and the loss of adhesion and the effects of moisture damage are often described as a mixture of adhesive and cohesive damage. The situation is further complicated by scalar effects, in which cohesion of a bitumen–filler mastic is described, whereas such a mastic is a function itself of adhesive strength and physical factors such as bitumen absorption by the filler.

### 19.3.1 Displacement

Displacement theory relates to the thermodynamic equilibrium of the three-phase bitumen–aggregate–water system. If water is introduced at a bitumen–aggregate interface, consideration of the surface energies that are involved shows that the bitumen will retract along the surface of the aggregate.

### 19.3.2 Detachment

Detachment occurs when a thin film of water or dust separates the bitumen and aggregate, with no obvious break in the surface of the bitumen film being apparent. Although the bitumen film completely encapsulates the aggregate particle, no adhesive bond exists, and the bitumen can easily be peeled from the aggregate surface. This process may be reversible (i.e. if the water is removed, the bitumen may re-adhere to the aggregate). A second mechanism of disbonding must occur to allow the ingress of water between the bitumen and the aggregate.

### 19.3.3 Film rupture

Film rupture may occur despite the fact that the bitumen fully coats the aggregate. At sharp edges or asperities on the aggregate surface, where the bitumen film is thinnest, it has been shown (Hughes et al., 1960) that water

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**Table 19.1 Material properties and external factors that can affect the bitumen–aggregate bond**

<table>
<thead>
<tr>
<th>Aggregate properties</th>
<th>Bitumen properties</th>
<th>Mixture properties</th>
<th>External factors</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mineralogy</td>
<td>Rheology</td>
<td>Void content</td>
<td>Rainfall</td>
</tr>
<tr>
<td>Surface texture</td>
<td>Electrical polarity</td>
<td>Permeability</td>
<td>Humidity</td>
</tr>
<tr>
<td>Porosity</td>
<td>Constitution</td>
<td>Bitumen content</td>
<td>Water pH</td>
</tr>
<tr>
<td>Dust</td>
<td>Surface free energy</td>
<td>Bitumen film thickness</td>
<td>Presence of salts</td>
</tr>
<tr>
<td>Durability</td>
<td></td>
<td>Filler type</td>
<td>Temperature</td>
</tr>
<tr>
<td>Surface area</td>
<td></td>
<td>Aggregate grading</td>
<td>Temperature cycling</td>
</tr>
<tr>
<td>Surface free energy</td>
<td></td>
<td>Type of mixture</td>
<td>Traffic</td>
</tr>
<tr>
<td>Absorption</td>
<td></td>
<td></td>
<td>Design</td>
</tr>
<tr>
<td>Moisture content</td>
<td></td>
<td></td>
<td>Workmanship</td>
</tr>
<tr>
<td>Shape</td>
<td></td>
<td></td>
<td>Drainage</td>
</tr>
<tr>
<td>Weathering</td>
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<td></td>
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</tbody>
</table>

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can penetrate through the film to reach the surface of the aggregate. This movement of water to the aggregate surface may occur with the water in either a vapour or liquid form. Once this process has started, it is possible for the water to spread between the bitumen and aggregate surface to produce a detached film of bitumen.

The speed with which the water can penetrate and detach the bitumen film will depend on the viscosity of the bitumen, the nature of the aggregate surface, the thickness of the bitumen film and the presence of filler and other components such as surface active agents. Once significant detachment of the bitumen film from the aggregate has occurred, stresses imposed by traffic will readily rupture the film, and the bitumen will retract, exposing water-covered aggregate. The displacement theory (Blott et al., 1954) relates to the thermodynamic equilibrium of the three phase bitumen–aggregate–water system. If water is introduced at a bitumen–aggregate interface, then consideration of the surface energies that are involved shows that the bitumen will retract along the surface of the aggregate. Figure 19.2 shows an aggregate particle embedded in a bitumen film, with point A representing the equilibrium contact position when the system is dry. When in contact with water, the equilibrium point shifts, and the new interface moves or retracts over the surface to point B. This new equilibrium position has a contact angle that will depend on the type and viscosity of the bitumen used.

19.3.4 Blistering and pitting

If the temperature of a flexible pavement increases, the viscosity of the bitumen within that pavement will reduce. If this is associated with a recent rainfall, the bitumen may creep up the edges of water droplets to form a blister, as shown in Figure 19.3. If the temperature continues to increase, the
A blister will expand, leaving a hollow or a pit, which may allow water to access the surface of the aggregate (Hughes et al., 1960; Thelen, 1958).

### 19.3.5 Hydraulic scouring

Hydraulic scouring or pumping occurs in the surface course, and is caused by the action of vehicle tyres on a saturated pavement surface (i.e. water is forced into surface voids in front of the vehicle tyre). On passing, the action of the tyre sucks up this water, thereby inducing a compression–tension cycle in these surface voids, which may result in disbonding of the bitumen from the aggregate. Suspended dust and silt in the water can act as an abrasive and can accelerate disbonding.

### 19.3.6 Pore pressure

This type of disbonding mechanism is most important in open or poorly compacted asphalts, where it is possible for water to be trapped as the material is compacted by traffic. Once the asphalt becomes effectively impermeable, subsequent trafficking induces a pore water pressure (Lee and Nicholas,
1957). This creates channels around the bitumen–aggregate interface, leading to loss of bond. Higher temperatures acting on the entrapped water result in expansive stresses accelerating water migration and disbonding. Low temperatures may also lead to the formation of ice, which is equally destructive.

**19.4 Methods of measuring and assessing adhesion between bitumen and aggregates and moisture damage**

Given the potential for premature failure due to adhesion related problems and the number of factors at play, the need for predictive laboratory tests is self-evident. A number of different types of test have been developed to compare combinations of aggregate, bitumen and water. However, the problem with many of these methods is a lack of information relating the laboratory prediction to performance in practice. Several techniques have been developed, and these can be categorised broadly as set out below.

- **Surface free energy and work of adhesion calculations**
  - surface energy measurements of bitumen
  - surface energy measurements of aggregate
  - calculation of the work of adhesion with and without the presence of water.
- **Bitumen–substrate tests**
  - peel tests
  - pull off tests.
- **Simulative measurements on bitumen–aggregate combinations**
  - coating and immersion tests
  - adsorption tests.
- **Simulative measurements on asphalts**
  - immersion mechanical tests
  - immersion trafficking tests.

Each of the above is discussed in the following sections.

**19.4.1 Surface free energy and work of adhesion calculations**

The surface energy of solids cannot be measured directly. Indirect methods are therefore used, and the surface energies of solids (including bitumen as well as aggregates) are inferred from the known surface energy values of several liquids or gases. Several techniques can be used, including contact angle measurement methods and vapour sorption techniques.

**19.4.1.1 Surface energy measurements of bitumen**

The Wilhelmy plate can be used to determine the surface energy of bitumen by measuring the contact angle between bitumen and a liquid, based on the kinetic force equilibrium when a very thin plate is immersed or withdrawn.
from a liquid solvent at a very slow and constant speed (Adamson and Gast, 1997). The dynamic contact angle between bitumen and a solvent liquid measured during the immersing process is called the ‘advancing contact angle’; while the dynamic contact angle during the withdrawal process is called the ‘receding contact angle’. The advancing contact angle, which is a wetting process, is associated with the fracture healing process, while the receding angle is associated with the fracture mechanism. The total surface energy and its components of bitumen can be calculated from these advancing and receding contact angles. The surface energy calculated from the advancing contact angles is called the ‘surface energy of wetting’, while the surface energy computed from the receding contact angles is called the ‘surface energy of dewetting’.

Alternatively, bitumen surface free energy can be derived using contact angles measured using a goniometer. In this technique, bitumen is applied evenly to a glass slide and drops of different liquids of known surface free energy are placed onto the surface, and the contact angle measured. Figure 19.4 shows a droplet at rest during a contact angle measurement using a goniometer.

Typical values for surface free energy vary by test method and the choice of solvent but typically fall within the range 25–50 mN/m², some example values (measured using the goniometer method and presented using the Van Oss approach (Van Oss et al., 1988)) are given in Table 19.2.

19.4.1.2 Surface energy measurements of aggregate
A goniometer can be used to measure the surface energy of aggregate but the measurement presents some difficulties. The samples need to be non-porous, and this can be problematic with some aggregates. Specimens are
required to be flat and low in texture, and this requires cutting and often polishing of aggregates in order to carry out the contact angle measurements. This introduces issues around isotropy in the aggregate and also the impact of polishing, and thus deviation from the true nature of the aggregate.

The surface energies of aggregates can be measured in their natural form using various vapour sorption techniques such as dynamic vapour sorption systems and universal sorption methods (Bhasin, 2006; Li, 1997). The basis of surface energy determination in vapour sorption techniques for high energy surfaces such as is found in aggregates is the use of the gas adsorption characteristics of selected solvents, the surface energies of which are known indirectly to measure the surface energies of the aggregate. This method can accommodate the particular shape, size, mineralogy and surface roughness of aggregates.

Again, typical values for surface free energy vary by test method and choice of solvent, and aggregates show a much wider range of values (50–200 mN/m²) than does bitumen, reflecting the wide range of mineralogies found in rocks suitable for aggregate production. Values for some typical aggregate types measured using the vapour sorption technique and adopting the Van Oss approach are presented in Table 19.3.

### Table 19.2 Typical measured values of surface free energy for various bitumen grades

<table>
<thead>
<tr>
<th>Surface free energy, $\gamma$: mN/m²</th>
<th>Lifshitz–van der Waals component of surface energy, $\gamma^{LW}$</th>
<th>Acid component of surface energy, $\gamma^+$</th>
<th>Basic component of surface energy, $\gamma^-$</th>
<th>Acid–base component of surface energy, $\gamma^{AB}$</th>
<th>Total surface energy, $\gamma^T$</th>
</tr>
</thead>
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<tr>
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<td>1.00</td>
<td>3.50</td>
<td>3.74</td>
<td>42.64</td>
</tr>
<tr>
<td>70/100 pen B</td>
<td>26.29</td>
<td>0.00</td>
<td>0.57</td>
<td>0.00</td>
<td>26.29</td>
</tr>
<tr>
<td>40/60 pen A</td>
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<td>0.85</td>
<td>1.90</td>
<td>2.54</td>
<td>35.89</td>
</tr>
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<td>0.00</td>
<td>0.59</td>
<td>0.00</td>
<td>24.62</td>
</tr>
<tr>
<td>10/20 pen A</td>
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<td>0.00</td>
<td>0.59</td>
<td>0.00</td>
<td>34.66</td>
</tr>
<tr>
<td>10/20 pen B</td>
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<td>0.02</td>
<td>0.34</td>
<td>0.16</td>
<td>27.41</td>
</tr>
<tr>
<td>SBS PMB A</td>
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<td>0.14</td>
<td>3.60</td>
<td>1.42</td>
<td>33.32</td>
</tr>
<tr>
<td>SBS PMB B</td>
<td>36.40</td>
<td>3.10</td>
<td>4.90</td>
<td>7.79</td>
<td>44.19</td>
</tr>
</tbody>
</table>

19.4.1.3 Calculation of the work of adhesion with and without the presence of water

Surface free energy measurements for the different types of both bitumen and aggregate and their bond energies provide an improved understanding of the material properties that influence the moisture sensitivity of asphalts but, due to the complex nature of damage in asphalts, cannot provide a full account of the likelihood of moisture damage alone. Similarly, conventional moisture-sensitive asphalt tests are also able to compare mechanical property
results for dry and moisture conditioned specimens, but they are unable to quantify the causes of good or poor performance beyond comparative tests.

Measurement of the surface free energies of bitumens and aggregates allows the calculation of their interfacial work of adhesion and the reduction in free energy of the system (work of disbonding) when water displaces bitumen from the bitumen–aggregate interface. These two parameters can then be used to estimate the moisture sensitivity of asphalts based on the principles of thermodynamics and physical adhesion (see section 19.1).

For an asphalt to be durable and less sensitive to moisture, it is desirable that the work of adhesion $W_{12}$ between the bitumen and the aggregate be as high as possible. In addition to the work of adhesion, the greater the magnitude of work of debonding when water displaces bitumen from the bitumen–aggregate interface, $W_{132}$, the greater will be the thermodynamic potential that drives moisture damage. Therefore, it is desirable that the magnitude of work of disbonding be as small as possible.

### 19.4.2 Bitumen–substrate tests

#### 19.4.2.1 Peel tests

A peel test is a well established and standardised methodology in which a rigid substrate and a flexible substrate are separated and the force required to achieve the separation under set conditions is measured to provide insights into the adhesive strength between the two materials. This approach can be further extended to investigate the influence of conditioning in the presence of water. Various geometries and loading regimes can be adopted, and these have become standardised in many regions (ASTM D1876-08 and ASTM D6862-11 (ASTM, 2008, 2011)). The methods are employed to measure the bonded strength of composites, joints, laminates and related materials.

Recent studies (Blackman et al., 2013) have examined the possibility of adapting peel tests to bitumen–aggregate combinations and a range of

| Material    | Lifshitz–van der Waals component of surface energy, $\gamma^{LW}$ | Acid component of surface energy, $\gamma^+$ | Basic component of surface energy, $\gamma^-$ | Acid–base component of surface energy, $\gamma^{AB}$ | Total surface energy, $\gamma^T$ |
|-------------|---------------------------------------------------------------|-----------------|---------------------------------|-------------------------------|----------------ternalized values of surface free energy for various aggregate types |
| Limestone A | 68.72                                                        | 99.34           | 14.49                           | 75.88                         | 144.60                      |
| Limestone B | 45.00                                                        | 8.00            | 214.00                          | 82.75                         | 127.75                      |
| Basalt      | 51.68                                                        | 14.12           | 59.82                           | 58.13                         | 109.81                      |
| Granite     | 56.55                                                        | 1.58            | 1.90                            | 3.47                          | 60.02                       |
| Gritstone   | 40.00                                                        | 5.00            | 131.00                          | 51.19                         | 91.19                       |
bitumen–aggregate combinations with adhesion promoters. The studies show that adhesion promoters could be successfully applied to bitumen–aggregate combinations.

19.4.2.2 Pull off tests

A pull off test is widely used to measure the mechanical tensile strength of films, paints and other coatings. Generally, a pull off test is conducted by measuring the minimum tensile stress necessary to detach or fracture the coatings of adhesive materials in a direction perpendicular to the substrate. A typical arrangement is shown in Figure 19.5.

The limpet pull off test (Craig, 1991) was developed to measure, quantitatively, the bond strength between the aggregate of a surface dressing and the underlying surface. The test uses a limpet apparatus that was originally developed for measuring the tensile strength of concrete. A 50 mm dia. metal plate is fixed to the road surface and the maximum load to achieve pull off determined. The method may be used both in the laboratory and on the actual road surface.

Bitumen adhesion has been assessed using different types of pull off test. The Instron pull off test (Craig, 1991) uses an Instron tensile apparatus to extract aggregate test specimens from containers of bitumen under controlled laboratory conditions. Test variables such as rock type, dust coatings, test temperature, rate of loading, bitumen type etc., have been shown to alter the results. An example of the data obtained for four different rock types with increasing dust contents is shown in Figure 19.6, and demonstrates how the maximum stress during testing varied for the different aggregates.

19.4.3 Simulative measurements on bitumen–aggregate combinations

19.4.3.1 Coating and immersion tests

This type of testing attempts to assess adhesion between aggregate and bitumen when water is also present. For example, in the immersion tray test

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**Figure 19.5** The pull off test
aggregate chippings are applied to a tray of bitumen covered with a layer of water. By careful examination of the chippings, it may be possible to determine whether surface active agents improve adhesion under wet conditions. Typically, aggregate is coated with bitumen, immersed in water under controlled conditions and the effect of stripping determined after a period of time. The various methods differ in the type of specimen used, the conditions under which the sample is immersed in water and the method that is used to assess the degree of stripping. This is the simplest type of test, and consists of aggregate being coated with bitumen and then being immersed in water. The degree of stripping is estimated by a visual inspection after a period of time. For example, in the total water immersion test, 14 mm single size chippings are coated with a known quantity of bitumen. The coated aggregate is then immersed in distilled water at 25°C for 48 h. The percentage of bitumen stripped off the aggregate is assessed visually.

A static immersion test was standardised [AASHTO T 182-84 (AASHTO, 2002, ASTM D1664 (ASTM, 1985)) and involved coating 100 g of aggregate with bitumen, immersing it for 16–18 h in 400 ml of distilled water with a pH of 6–7, and visually estimating the total visible area of the coated aggregate as above or below 95%. The visual assessment is made while the mixture is still immersed in the water.

The fundamental problem with such methods is their subjective nature and resultant poor reproducibility. An experienced operator may be capable of ranking the aggregates in relation to their performance in situ, but it must be recognised that, in some cases, an aggregate with good laboratory performance may occasionally perform poorly on the road and those with
poor static immersion test results may perform satisfactorily in practice. In summary, although such methods may indicate which combinations of aggregate and bitumen show degrees of water sensitivity, it is doubtful that the long term potential of stripping is adequately addressed, and the static immersion test was discontinued as an ASTM standard in 1993.

A dynamic immersion test is very similar to the static immersion test but the sample is agitated mechanically by shaking, stirring or kneading. Again, the degree of stripping is estimated visually, together with a subjective judgement of whether the mixture remains cohesive or separates into individual particles of aggregate. The reproducibility of this type of test is also very poor. There are several variants of this technique. The rolling bottle (flask) method (BS EN 12697-11 (BSI, 2012a)) is used to determine the adhesion between single size aggregate and bitumen. In the test, 200 spherical aggregate particles (6.3–8 mm) are coated with a 0.1 mm thick binder film. The coated aggregate is then placed in a 250 ml flask containing deionised cold water and a glass rod, and rotated at 40 revs/min for 3 days. The amount of retained bitumen is then visually determined.

The boiling water test (ASTM D3625 (ASTM, 2012)) involves placing a 200–300 g sample of coated aggregate (single size aggregate, or aggregate graded to design specifications) in boiling water (500 ml distilled water) for 1–10 min. For the 10 min test, the mixture is stirred three times with a glass rod while it is being boiled. After boiling, the mixture is dried and the amount of bitumen loss is determined by visual assessment. The test is very subjective and is known to provide inconsistent results in terms of identifying water-sensitive mixtures. In addition, the test only reflects the loss of adhesion, and does not address loss of cohesion.

In the majority of these tests, the coated aggregate is immersed in water. However, salt based or de-icing solutions and those containing fuel oils may also be used. Certain types of test may tend to coat an aggregate in the presence of water. In such cases, the degree of coating obtained is used as an index of adhesivity.

In this type of test, aggregate coated in bitumen is boiled in solutions containing various concentrations of sodium carbonate. The strength of sodium carbonate solution in which stripping is first observed is used as a measure of adhesivity. However, the artificial conditions of the test are unlikely to predict the likely performance on the road.

19.4.3.2 Adsorption tests
The Strategic Highway Research Program (SHRP) in the USA investigated the effect of moisture damage as one of six major distress areas requiring further investigation. The result was the net adsorption test method (Curtis et al.,
1993). This test combined a fundamental measurement of bitumen–aggregate adhesion with a measure of moisture sensitivity. The method was able to match an aggregate and a bitumen to give optimum performance conditions.

In the test, a stock solution of bitumen is first prepared by adding 1 g of bitumen to 1000 ml of toluene. The aggregate is crushed to produce 50 g test samples of 5 mm to dust grading, and then 140 ml of the stock solution is added to a conical flask containing the graded aggregate placed on a rotating table to agitate the mix. After 6 h, a 4 ml sample is removed and diluted to 25 ml using toluene. The absorbance is measured using a spectrophotometer at 410 nm.

The reading is compared with the one obtained for the stock solution, and the percentage initial adsorption calculated. Distilled water (2 ml) is then added to the flask, and the flask is agitated overnight. Another 4 ml is removed and assessed as before. The value obtained at this stage is the percentage net adsorption, which should be lower than the percentage initial adsorption value, as some of the bitumen will have been removed from the aggregate due to the introduction of moisture into the system.

### 19.4.4 Simulative measurements on asphalts
#### 19.4.4.1 Immersion mechanical tests

Immersion mechanical tests involve the measurement of a change in a mechanical property of compacted asphalt after it has been immersed in water. Thus, the ratio of the property after immersion divided by the initial property is an indirect measure of stripping (usually expressed as a percentage or index).

Different types of mechanical properties can be measured, and these include shear strength, flexural strength and compressive strength. The two most common are the AASHTO T 283 (AASHTO, 2011) method sometimes referred to as the ‘modified Lottman procedure’, and the ‘retained Marshall stability test’. Indexes based on retained stiffness are also employed, and these have the advantage of the index measurement being non-destructive, allowing for repeated measurements on the same specimen over time and under different conditions. Common approaches to immersion mechanical testing for asphalts are discussed below.

One of the most globally used procedures is the modified Lottman procedure, which combines features of both the Lottman (1978, 1982) and Tunnicliff and Root (1984) procedures. The Lottman procedure attempts to achieve a 100% saturation level in its specimens, while the Tunnicliff and Root procedure attempts to control the level of saturation between 55% and 80%. Concern that over-saturation induces damage in specimens that is not
associated with moisture damage, but rather with the over-saturation of the specimen, meant that for the modified Lottman procedure the degree of saturation was decreased to between 60% and 80%. As the saturation level achieved by partial vacuum is primarily responsive to the magnitude of the vacuum and relatively independent of the length of time, this reduced saturation was achieved by lowering the partial vacuum from 600 to 508 mmHg.

Moisture damage tests based on Marshall stability are still in widespread use. There are a number of different versions of each type of test. In the version used by Shell, at least eight Marshall specimens are manufactured using a prescribed aggregate type, aggregate grading, bitumen content and void content. Four specimens are then tested using the standard Marshall method to give a standard stability value. The four remaining specimens are vacuum treated under water at a temperature of 0–1°C to saturate the pore volume of the mixture with water. The specimens are then stored in a water bath at 60°C for 48 h, after which their Marshall stability is determined. The ratio of the Marshall stability of the soaked specimens to the standard Marshall stability is termed the ‘retained Marshall stability’. A retained Marshall stability value of >75% is usually regarded as acceptable.

The principle of the retained stiffness test is similar to that of the retained Marshall test. The major difference is that the test specimens are assessed using the indirect tensile stiffness modulus test (BS EN 12697-26:2012 (BSI, 2012b)), which is carried out using a Nottingham asphalt tester (NAT) or similar device. This has the additional benefit that the method is non-destructive, allowing the same test specimens to be used after soaking. A number of versions of the method exist in which the soaking period and water temperature may be varied.

The saturation ageing tensile stiffness (SATS) test (BS EN 12697-45:2012 (BSI, 2012c)) is the first procedure of its kind that combines both ageing and water-damage mechanisms (intended to simulate the conditions to which some asphalt pavements are subjected in service) within a single laboratory test protocol. The conditions used in the SATS procedure were selected in order to reproduce in the laboratory the moisture damage observed in the field, as demonstrated by a decrease in stiffness modulus for particular asphalts as detailed by Airey et al. (2005). The procedure has been found to reproduce successfully the moisture damage observed in asphalts in the field, and to distinguish between poorly performing materials and alternative asphalts incorporating aggregate with good durability track records (Airey and Choi, 2002). The results obtained from the SATS moisture conditioning procedure tend to rank asphalts in terms of moisture sensitivity in the same order as the AASHTO T 283 (AASHTO, 2011) procedure, although the
relative performance of a mixture containing a moisture sensitive aggregate is usually significantly lower in the SATS test (Airey et al., 2005).

The standard SATS procedure involves conditioning five pre-saturated specimens simultaneously in a pressure vessel under 0.5 MPa air pressure at a temperature of 85°C for a period of 24 h. This conditioning is followed by a cooling period of 24 h before the air pressure is released and the vessel opened to remove the specimens for stiffness testing. The pressure vessel used can hold five nominally identical specimens (100 mm diameter and 60 mm thick) in a custom made specimen tray.

19.4.4.2 Immersion trafficking tests
A major problem with most types of adhesion test is that they do not consider the effect of trafficking in causing stripping. A method that considers this factor is the immersion wheel tracking test (Mathews and Colwill, 1962) shown in Figure 19.7. In this test, a specimen is immersed in a water bath and traversed by a loaded reciprocating solid rubber tyre. In the standard method, three specimens that have been compacted in moulds are tracked using a 20 kg load at 25 cycles/min at a water temperature of 40°C until failure occurs.

The development of the rut that forms is measured until stripping starts to occur. This is typically marked by a steep increase in the rut depth and

![Figure 19.7 Immersion wheel tracking device](image-url)
surface ravelling of the test specimen. Good correlation has been shown to exist for stripping failures of heavily trafficked roads. It has been found that factors such as aggregate shape, aggregate interlock, bitumen viscosity and sample preparation affect failure times.

The wet wheel track test is now being reconsidered as a means of assessing asphalts such as porous asphalts and high stone content, thin surface course systems. The method is easily adaptable in which temperature, immersion medium and loading may be varied. The test may be performed for a specific time or until failure occurs.

A locked wheel method was developed to assess the likelihood of ravelling due to the presence of moisture (McKibben, 1987). The only alteration to the equipment involved the provision of a locking ratchet that allowed the wheel to travel freely in one direction but also to be locked in position as it is dragged back across the specimen surface on the remainder of the cycle. Surface ravelling has been related to grading, degree of compaction, bitumen and aggregate characteristics. This method is, however, very aggressive, and may be most suitable for highly stressed trafficking locations.

Similar to the immersion wheel tracker described above, the Hamburg wheel tracking device consists of a pair of samples subjected to wheel loading under water. A sample is typically 260 mm wide, 320 mm long and 40 mm thick. The samples are submerged in water at 50°C, although the temperature can vary from 25°C to 70°C, and subjected to a steel wheel (47 mm wide) with a load of 705 N. The wheel makes 50 passes/min over each sample, with a maximum velocity of 0.34 m/s in the centre of the sample. Testing is undertaken to 20,000 passes or until a 20 mm deep rut develops.

The onset of stripping is termed the ‘stripping inflection point’ and relates to the intersection of the creep slope and the stripping slope where there is a sharp increase in the rate of permanent deformation. Both the stripping slope and the stripping inflection point are related to the moisture damage of the sample. Based on an evaluation of 20 pavements, Aschenbrener (1995) found an excellent correlation between the Hamburg wheel tracking device and pavements of known in-service performance.

19.5 Improving bitumen–aggregate adhesion
Typically, the adhesion of bitumen to aggregate is not a problem. However, in the presence of water, unexpected adhesion related problems may occur. There are a number of traditional methods used to reduce the likelihood of this happening (i.e. using higher viscosity bitumen, hydrated lime or surface active agents that improve the bond between the bitumen and the aggregate). While modifying bitumen viscosity may be easily achieved, this may
result in workability and compaction related problems, particularly for thin surface course systems that use high stone content asphalts.

19.5.1 Hydrated lime
The use of 1–3% of hydrated lime as part of the filler content has traditionally been used as an anti-stripping agent. It has been suggested (Plancher et al., 1976) that the hydrated lime reacts with the carboxyl acids present in the bitumen and allows other carbonyl groups such as ketones to attach themselves to the aggregate surface. These ketones are not as easily removed by water as the acids, and so the mixtures are less susceptible to stripping.

It has also been suggested (Ishai and Craus, 1977) that a hydrated lime solution will result if water is present at the bitumen–aggregate interface. The calcium ions in this solution cause the surface of the aggregate to become basic. The electrochemical balance forces the water away from the aggregate and into an emulsion in the bitumen. The balance will then force attachment to the hydrophobic surface of the aggregate.

19.5.2 Anti-stripping agents
Bitumen–aggregate adhesion may be improved by the addition of chemical additives. These act in two main ways

■ they may change the interfacial conditions between the aggregate and the bitumen so that the bitumen preferentially wets the aggregate, which improves adhesion
■ they may improve the adhesive bond between the aggregate and the bitumen, thus increasing the long term resistance to bitumen detachment due to water.

Typically, 0.1–1.0% of fatty amines is the main type of additive used to improve adhesion. It is believed that the amine groups are attracted to the surface of an aggregate, while the fatty groups remain in the bitumen. The result is an ionically bonded cross-link between aggregate and bitumen. However, these additives may be relatively unstable at bitumen storage temperatures, and can become deactivated. It is also possible that a given additive will not improve the adhesion of all aggregate types (i.e. they may be rock type specific). There is also the issue that, while they may improve initial adhesion, they may have no or limited long term effect.

It is recommended that, prior to their use, laboratory testing should be carried out to optimise the type and amount of additive used for a given bitumen–aggregate combination. This may be undertaken using the wide range of the test methods already mentioned in this chapter. For example, Table 19.4 shows data obtained using the net adsorption test for two different
aggregates and a 70/100 pen bitumen with and without 0.5% adhesion agent. It can be seen that, while the adhesion agent increased initial adsorption by 10.9% for the quartz dolerite, the adsorption only improved by 3% for the greywacke. In terms of improving moisture sensitivity, there was a small improvement for the quartz dolerite, and no improvement for the greywacke. This example shows how aggregate properties influence attempts at improving adhesion. For the greywacke, the addition of adhesion agents essentially did little to improve performance.

### References


### Table 19.4 Net and initial adsorption results with different binders

<table>
<thead>
<tr>
<th>Aggregate type</th>
<th>Initial adsorption: %</th>
<th>Net adsorption: %</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>100 pen bitumen</td>
<td>100 pen bitumen with 0.5% adhesion agent</td>
</tr>
<tr>
<td>Quartz dolerite</td>
<td>52.2</td>
<td>63.1</td>
</tr>
<tr>
<td>Greywacke</td>
<td>64.6</td>
<td>67.6</td>
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Adhesion of bitumen and moisture damage in asphalts

Table 19.4 Net and initial adsorption results with different binders


Adhesion of bitumen and moisture damage in asphalts

Final Report of Cahn Balance Thermogravimetry Gas Adsorption Experiments. Chemical Engineering, Texas A&M University, College Station, TX, USA.
Lottman RP (1978) Predicting Moisture-Induced Damage to Asphaltic Concrete. Transportation Research Board, Washington, DC, USA, NCHRP Report 192.
Bituminous road surfaces are expected to provide serviceability to the road user for many years before a maintenance intervention or rehabilitation is required. Indeed, there are undoubtedly examples of asphalt surfaces in many countries that are 40 or 50 years old and still provide a useable pavement in lightly trafficked environments. The term ‘durability’ is often used to describe the retention of the desirable engineering and serviceability characteristics over the lifetime of an asphalt, and is frequently included in the engineer’s lexicon of performance attributes. Greater clarity of the term is helpful, and one definition of the durability of asphalts is as follows.

The ability of the materials comprising the mixture to resist the effects of water, ageing and temperature variation, in the context of a given amount of traffic loading, without significant deterioration for an extended period. (Scholz, 1995)

While the integrity of the aggregates is an important factor in the long term performance of asphalts, it is primarily aspects associated with the ageing characteristics of bitumen that have the greatest influence on the durability of asphalts, surface dressings, chip seals and sprayed seals.

Quantification of bitumen durability has proved to be a difficult task, and methods of assessing this property tend to be indicative rather than precise. Long term studies have shown that, if an asphalt is to achieve its design life, it is important that the bitumen is not excessively hardened during hot storage, during the asphalt manufacturing process or in service in the road. Bitumen, in common with many organic substances, is affected by the presence of oxygen, ultraviolet radiation and changes in temperature. In bitumen, these external influences cause it to harden, resulting in a decrease in penetration, an increase in softening point and, usually, an increase in penetration index (PI). In recent years, the phenomenon of the hardening
of bitumen and, hence, the hardening of an asphalt has been viewed as being beneficial in pavement layers as it increases the stiffness of the material and, therefore, the load spreading capabilities of the structure. This hardening is known as ‘curing’, and is believed to extend the life of a pavement (Scholz, 1995).

In the surface course, where the material is exposed to the environment, hardening of the bitumen can have a detrimental effect on its performance and can lead to fretting and/or cracking. This effect is still referred to as ‘hardening’, the term implying that the change in bitumen properties is detrimental to the service life of the surface course.

20.1 Bitumen hardening

The tendency for bitumen to harden under the influence of the atmosphere has been known and studied for many years. As many as 15 different factors that influence bitumen ageing have been identified, and these are detailed in Table 20.1.

The most important of these 15 mechanisms of bitumen hardening are

- oxidation
- volatilisation
- steric or physical factors
- exudation of oils.

<table>
<thead>
<tr>
<th>Table 20.1 Mechanisms of bitumen ageing (Traxler, 1963)</th>
</tr>
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<tbody>
<tr>
<td>Factors that influence bitumen ageing</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>Oxidation (in dark)</td>
</tr>
<tr>
<td>Photo-oxidation (direct light)</td>
</tr>
<tr>
<td>Volatilisation</td>
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<tr>
<td>Photo-oxidation (reflected light)</td>
</tr>
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<td>Photochemical (direct light)</td>
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<tr>
<td>Photochemical (reflected light)</td>
</tr>
<tr>
<td>Polymerisation</td>
</tr>
<tr>
<td>Steric or physical</td>
</tr>
<tr>
<td>Exudation of oils</td>
</tr>
<tr>
<td>Changes by nuclear energy</td>
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<tr>
<td>Action by water</td>
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<tr>
<td>Absorption by solid</td>
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<tr>
<td>Absorption of components at a solid surface</td>
</tr>
<tr>
<td>Chemical reactions</td>
</tr>
<tr>
<td>Microbiological deterioration</td>
</tr>
</tbody>
</table>
20.1.1 Oxidation
This is considered to be the most important cause of hardening of bitumen in service.

Like many organic substances, bitumen is slowly oxidised when in contact with atmospheric oxygen. Polar groups containing oxygen are formed, and these tend to associate into micelles (an aggregate of molecules in a colloidal solution) of higher micellar weight, thereby increasing the viscosity of the bitumen. Reaction of oxygen with bitumen molecules causes the formation of carbonyl species, resulting in larger and more complex molecules that make the bitumen harder and less flexible. The degree of oxidation is highly dependent on the temperature, period of exposure and the thickness of the bitumen film. The rate of oxidation doubles for each 10°C increase in temperature above 100°C. Hardening due to oxidation has long been held to be the main cause of ageing, to the extent that other factors have been given scant consideration. However, it has been shown that, although other factors are generally less important than oxidation, they are measurable.

20.1.2 Loss of volatiles by evaporation
The evaporation of volatile components depends mainly on temperature and the exposure conditions. The rate of evaporation is controlled by the diffusion rate and the length or thickness of the diffusion path. Well compacted dense asphalt will have a slow diffusion process, whereas open grade asphalts and surface dressings will present more rapid evaporative conditions. However, paving grade bitumens are relatively involatile, and therefore the amount of hardening resulting from evaporation of volatiles is usually fairly small.

20.1.3 Steric or physical hardening
The phenomenon of steric hardening was first reported in 1944 by Traxler, when it was observed that bitumen samples stored at 25°C showed an increase in viscosity. This physical hardening is usually attributed to a combination of reorientation or restructuring of molecules within the bitumen and the slow crystallisation of waxes. The molecular reorientation that results in the hardening effect is reversible on the application of heat, with Traxler noting that raising the storage temperature to 70°C reversed the changed structure, with the bitumen sample returning to its original viscosity.

20.1.4 Loss of volatiles by exudation
Exudative hardening results from the movement of oily components that exude from the bitumen into the mineral aggregate (Van Gooswilligen et al., 1989). It is a function of both the exudation tendency of the bitumen and the porosity of the aggregate.
20.2 Hardening of bitumen during storage, mixing and in service

The circumstances under which hardening occurs vary considerably. During storage, the bitumen is in bulk at a high temperature for a period of days or weeks. During mixing, hot storage, transport and laying, the bitumen is a thin film at high temperature for a relatively short period. In service, the bitumen is a thin film at a low or moderate temperature for a very long time. The degree of exposure to the air of asphalts in service is important and depends on the void content of the mixture. In dense, well-compacted mixtures, the amount of hardening is relatively small, while asphalts that have a more open constitution, such as porous asphalt, will undergo significant hardening. The degree of exposure to air for surface dressings or sprayed seals is potentially much higher than is the case with dense asphalt layers, and is influenced by the mosaic achieved by the aggregate and the absorptivity of the aggregates (Marais, 1974).

20.2.1 Hardening of bitumen in bulk storage

Very little hardening occurs when bitumen is stored in bulk at high temperature. Data from Shell terminals shows that 3–4 kt of paving grade bitumen stored at 150°C in a 7 kt tank shows virtually no change in viscosity or penetration over a 4 week storage period. This is because the surface area of the bitumen that is exposed to oxygen is very small in relation to the volume. However, if the bitumen is being circulated and is falling from the pipe entry at the top of the tank to the surface of the bitumen, significant hardening may occur. This arises because the surface area of the bitumen will be relatively large as it falls from the entry pipe, exposing it to the action of the oxygen. This effect can be minimised using the storage tank layout described in section 2.4.2 and shown in Figure 2.6.

20.2.2 Hardening of bitumen during mixing with aggregate

This is described as ‘short term ageing’, a term that is also applied to hardening that occurs during laying. During the mixing process, it is estimated that all the aggregate and filler is coated with a thin film of bitumen, usually between 5 and 15 μm thick. If the bitumen from 1 tonne of a dense asphalt concrete was spread at 10 μm thick, it would occupy an area of around 10,000 m², the equivalent of over one and a half average size football pitches. Thus, when bitumen is mixed with hot aggregate and spread into thin films in a paddle mixer, conditions are ideal for the occurrence of oxidation and the loss of volatile fractions within the bitumen. Hardening of bitumen during this process is well known, and is taken into account when selecting the grade of bitumen to be used. As a very rough approximation, during mixing with hot aggregate in a paddle mixer, the penetration of a paving grade bitumen falls by about 30%. However, the amount of
Hardening depends on a number of factors, such as temperature, duration of mixing and bitumen film thickness. Minimising the hardening during mixing depends on careful control of all these factors. Control of the temperature and the bitumen content are particularly critical. Figure 20.1 clearly shows increasing bitumen hardening, measured by higher values of softening point, as mixing temperatures are raised.

Similarly, Figure 20.2 shows that reducing the thickness of the bitumen film significantly increases the viscosity of the bitumen. The latter is measured by the ageing index, which is defined as the ratio of the viscosity of the aged bitumen, $\eta_a$, to the viscosity of the virgin bitumen, $\eta_o$.

Note that the ageing index is not a fundamentally defined parameter – it is usually a ratio of two values (e.g. viscosity, stiffness or penetration) measured at different times.

The type of mixer used also affects the amount of hardening during mixing. It has been recognised for some time that the amount of hardening in a drum mixer is often less than that which occurs in a conventional batch mixer (Haas, 1974). This is due to the presence of steam in the drum, which limits the availability of oxygen. However, the multiplicity of different designs of

![Figure 20.1](image-url)  
**Figure 20.1** Relationship between the temperature of the mixture and change in softening point (Whiteoak and Fordyce, 1989)
drum mixers means that variation in the amount of hardening between different designs of plant is almost inevitable. Modern drum mix plants, notably those with counter flow drum mix and double drum configurations with very high mixing efficiencies, result in minimal hardening of bitumen compared with older drum mix plants. Notwithstanding, a study carried out by Shell Bitumen (1973) on two different drum mixers showed that, for equivalent mixing temperatures, the overall reduction in penetration and increase in softening point can be less than half of that which occurs in a conventional batch mixer.

### 20.2.3 Hardening of bitumen in asphalts during hot storage, transport and laying

Drum mixers, which produce large volumes of asphalts, require silo storage for the mixed material to account for peaks and troughs in demand. In such circumstances, mixtures are stored in hot silos, as well as in the delivery vehicle during transportation.

Some hardening of the bitumen will take place during hot storage, whether it is in a silo or in a truck. It was stated above that the amount of hardening will depend principally on the duration of exposure to oxygen, the thickness of the bitumen film and the temperature of the mixture. When a mixture is discharged into a storage silo, air enters with the mixture and some is trapped in the voids of the material. During the storage period, some of the oxygen in this entrained air will react with the bitumen. If no additional air enters the silo, oxidation of the bitumen will cease.
It is important that the entry and discharge gates are airtight and that there are no other openings where air can enter the silo. If the discharge gate is not airtight, the silo may behave like a chimney, drawing air in at the discharge gate (which exits the loading gate) and resulting in oxidation and cooling of the stored material. In addition, the silo should be as full as is practicable in order to minimise the amount of free air at the top of the silo. Air remaining at the top of the silo will react with the top surface of the material. This reaction forms carbon dioxide which, because it is heavier than air, tends to blanket the surface of the mixture, protecting it from further oxidation. In the USA, some silos have the facility to be pressurised with exhaust gases, containing no oxygen, from a burner. These exhaust gases purge the silo of entrained air and provide a slight positive pressure, preventing more air entering the silo.

Studies carried out in the USA (Brock, 1986) suggest that if oxidation in the silo is limited to that induced by entrained air, then little or no additional oxidation will occur during transportation and laying. It is hypothesised that this is because no significant quantity of fresh air is entrained in the mixture during discharge into the truck. Thus, little or no additional air is available for oxidation. In fact, it was observed that if the mixture was discharged directly from the paddle mixer into the delivery vehicle, the amount of hardening during transportation was very similar to that which occurs during silo storage.

If materials are being laid at low ambient temperatures or if the mixture has to be retained for a period in hot storage, there is a temptation to increase mixing temperatures to offset these two factors. However, increasing the mixing temperature will considerably accelerate the rate of bitumen oxidation, resulting in an increase in bitumen viscosity. Thus, a significant proportion of the reduction in viscosity achieved by increasing the mixing temperature will be lost because of additional oxidation of the bitumen, which may adversely affect the long term performance of the material.

### 20.2.4 Hardening of bitumen in the pavement and on the road

As explained above, a significant amount of bitumen hardening occurs during mixing and, to a lesser extent, during hot storage and transportation. However, hardening of the binder will continue in the pavement until some limiting value is reached. This behaviour is described as ‘long term ageing’ and is illustrated in Figure 20.3, which shows the ageing index of the bitumen after mixing, storage, transport, paving and subsequent service.

#### 20.2.4.1 Bitumen hardening in asphalt surface courses

The main factor that influences bitumen hardening in asphalts, including surface courses, is the void content of the mixture. Table 20.2 shows the properties of bitumens recovered from three asphalt concretes after 15 years’
service. The bitumen recovered from the mixture with the lowest void content had hardened very little. However, where the void content was high, allowing constant ingress of air, substantial hardening had occurred. The PI of the material with the highest void content had increased appreciably. This limits the amount of stress relaxation that can occur, and may result in cracking of the compacted material. Similarly, Figure 20.4 shows the in situ bitumen properties of 5 year old asphalt concrete with void contents ranging from 3% to 12% (Lubbers, 1985). At void contents less than 5%, very little hardening occurred in service. However, at void contents greater than 9%, the in situ penetration fell from 70 dmm to less than 25 dmm.

The bitumen at the surface of the road hardens much more quickly than the bitumen in the bulk of the pavement. There are three reasons for this:

- the existence of a constant supply of fresh oxygen
- the occasional incidence of high temperatures at the road surface
- the occurrence of photo-oxidation of the bitumen by ultraviolet radiation.

Photo-oxidation causes a skin, 4–5 μm thick, to be rapidly formed on the surface of the bitumen film. This is induced by natural ultraviolet radiation.
### Table 20.2 Hardening of bitumen in service (Edwards, 1973)

<table>
<thead>
<tr>
<th>Road</th>
<th>A</th>
<th>B</th>
<th>C</th>
</tr>
</thead>
<tbody>
<tr>
<td>Voids in mix: %</td>
<td>4</td>
<td>5</td>
<td>7</td>
</tr>
</tbody>
</table>

### Properties after mixing and laying

<table>
<thead>
<tr>
<th></th>
<th>Road A</th>
<th>Road B</th>
<th>Road C</th>
</tr>
</thead>
<tbody>
<tr>
<td>Softening point (IP): °C</td>
<td>64</td>
<td>63</td>
<td>66</td>
</tr>
<tr>
<td>Penetration at 25°C: dmm</td>
<td>33</td>
<td>33</td>
<td>30</td>
</tr>
<tr>
<td>Penetration index (Pen, SP)</td>
<td>+0.7</td>
<td>+0.7</td>
<td>+0.9</td>
</tr>
</tbody>
</table>

**Stiffness ($S_0$): Pa (calc.)**

<table>
<thead>
<tr>
<th></th>
<th>Road A ($10^4$ s, 25°C)</th>
<th>Road B ($10^4$ s, 25°C)</th>
<th>Road C ($10^4$ s, 25°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$10^4$ s, 25°C</td>
<td>$1.4 \times 10^3$</td>
<td>$1.4 \times 10^3$</td>
<td>$2.5 \times 10^3$</td>
</tr>
<tr>
<td>$10^4$ s, 0°C</td>
<td>$5.0 \times 10^5$</td>
<td>$5.0 \times 10^5$</td>
<td>$7.0 \times 10^5$</td>
</tr>
<tr>
<td>$10^{-2}$ s, 25°C</td>
<td>$2.5 \times 10^7$</td>
<td>$3.0 \times 10^7$</td>
<td>$3.0 \times 10^7$</td>
</tr>
<tr>
<td>$10^{-2}$ s, 0°C</td>
<td>$3.0 \times 10^8$</td>
<td>$3.0 \times 10^8$</td>
<td>$3.0 \times 10^8$</td>
</tr>
</tbody>
</table>

### Properties after 15 years’ service

<table>
<thead>
<tr>
<th></th>
<th>Road A</th>
<th>Road B</th>
<th>Road C</th>
</tr>
</thead>
<tbody>
<tr>
<td>Softening point (IP): °C</td>
<td>68</td>
<td>76</td>
<td>88</td>
</tr>
<tr>
<td>Penetration at 25°C: dmm</td>
<td>24</td>
<td>15</td>
<td>11</td>
</tr>
<tr>
<td>Penetration index (Pen, SP)</td>
<td>+0.8</td>
<td>+1.1</td>
<td>+2.1</td>
</tr>
</tbody>
</table>

**Stiffness ($S_{15}$): Pa (calc.)**

<table>
<thead>
<tr>
<th></th>
<th>Road A ($10^4$ s, 25°C)</th>
<th>Road B ($10^4$ s, 25°C)</th>
<th>Road C ($10^4$ s, 25°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$10^4$ s, 25°C</td>
<td>$4 \times 10^2$</td>
<td>$20 \times 10^2$</td>
<td>$150 \times 10^2$</td>
</tr>
<tr>
<td>$10^4$ s, 0°C</td>
<td>$13 \times 10^5$</td>
<td>$40 \times 10^5$</td>
<td>$80 \times 10^5$</td>
</tr>
<tr>
<td>$10^{-2}$ s, 25°C</td>
<td>$4 \times 10^7$</td>
<td>$7 \times 10^7$</td>
<td>$8 \times 10^7$</td>
</tr>
<tr>
<td>$10^{-2}$ s, 0°C</td>
<td>$4 \times 10^8$</td>
<td>$6 \times 10^8$</td>
<td>$6 \times 10^8$</td>
</tr>
</tbody>
</table>

**Ageing index ($S_{15}/S_0$)**

<table>
<thead>
<tr>
<th></th>
<th>Road A</th>
<th>Road B</th>
<th>Road C</th>
</tr>
</thead>
<tbody>
<tr>
<td>$10^4$ s, 25°C</td>
<td>2.8</td>
<td>14</td>
<td>60</td>
</tr>
<tr>
<td>$10^4$ s, 0°C</td>
<td>2.6</td>
<td>8</td>
<td>11</td>
</tr>
<tr>
<td>$10^{-2}$ s, 25°C</td>
<td>1.6</td>
<td>2.3</td>
<td>2.7</td>
</tr>
<tr>
<td>$10^{-2}$ s, 0°C</td>
<td>1.3</td>
<td>2.0</td>
<td>2.0</td>
</tr>
</tbody>
</table>

*Note: Ageing index is not a fundamentally defined parameter. It is usually a ratio of two values (e.g. viscosity, stiffness, penetration) measured at different times.*

---

**Figure 20.4** The effect of void content on the hardening of bitumen on the road (Lubbers, 1985)
that, it is believed, is absorbed into the upper 10 μm of a film of bitumen. It has been shown that the skin formation can retard oxygen absorption and loss of volatiles. However, the oxidised material is soluble in rainwater and can be abraded away, resulting in the exposure of fresh bitumen (Dickinson et al., 1958).

Although both continuously graded and gap graded surface course mixtures are regarded as dense, gap graded materials are considered to be more durable. One factor that affects the durability is the permeability of the mixture to air. Gap graded mixtures are generally less permeable to air than continuously graded mixtures for similar air void contents (Marais, 1974). This is because the voids in gap graded mixtures are generally discrete and not interconnecting, as may be the case with continuously graded mixtures.

The speed of surface oxidation is illustrated in Figure 20.5, in which the penetration, softening point and PI of bitumens recovered from the surface and bulk of an asphalt concrete over a number of years are plotted (Lubbers, 1985). After 7 years, the penetration at the surface is 25 dmm, whereas the bitumen in the bulk of the mixture has a penetration of 45 dmm. Similarly, there is a substantial difference in the PI. In the bulk of the mixture, the PI has only marginally changed, whereas the PI of the bitumen at the surface has increased to above 4. Surface oxidation is, in fact, desirable, because it
enables bitumens to be eroded away relatively quickly, exposing new aggregate surfaces and thereby improving the skid resistance of the road surface.

Bitumen content (bitumen film thickness) also plays a very important role in the speed of bitumen hardening on the road. Laboratory work has shown (Blokker and van Hoorn, 1959) that the oxidation of a film of bitumen at high ambient temperatures (i.e. between 40°C and 60°C) is limited to a depth of about 4 μm, as shown in Figure 20.6. The significance of this will depend on the average bitumen film thickness, which, as stated earlier, can range from 5 to 15 μm. It has been suggested (Campen et al., 1959) that a minimum bitumen film thickness of between 6 and 8 μm is required for satisfactory performance of both continuously graded and gap graded mixtures. Thus, in general, the lower the bitumen content, the thinner the bitumen film and the faster hardening will occur.

The bitumen film thickness can be approximated using the methods given in Appendix 4. Typical bitumen film thicknesses are as follows

- dense asphalt concrete  >5 μm
- hot rolled asphalt  >7 μm

![Figure 20.6 Relationship between oxygen absorbed at 50°C and film thickness (Marais, 1974)](image-url)
This method of calculating bitumen film thickness determines surface area factors for the aggregate down to 75 μm. Clearly, some of the material passing this sieve will be substantially finer than 75 μm, particularly for materials in which limestone filler is added to the mixture. Therefore, for high filler content materials such as hot rolled asphalt, the bitumen film thickness will be lower than the value given above. However, if it is assumed that the bitumen and filler coat the coarse and fine aggregate, then the above values underestimate the binder film thickness.

20.2.4.2 Bitumen curing in asphalt bases
Contrary to the effects of bitumen hardening in the surface course, a gradual hardening of the main structural layers of the pavement appears to be beneficial, and is described as ‘curing’. As with all hot mixed asphalts, the penetration of the bitumen in a base will harden by approximately 30% during the mixing and laying process, and, despite asphalt bases being locked inside the pavement construction and shielded from exposure to the environment, the penetration of the bitumen will continue to exhibit varying rates of hardening.

The Transport Research Laboratory (TRL) carried out a wide ranging investigation into the behaviour of asphalt pavements in service (Nunn et al., 1997). This very important study noted in section 2.1 that

One of the findings of this research was that changes occurring in asphalt over the life of the road are crucial in understanding its behaviour. These changes, which are referred to as curing, can help to explain why conventional mechanisms of deterioration do not occur and why, provided the road is constructed above a minimum threshold strength, it should have a very long, but indeterminate, structural life. The increase in the stiffness of the asphalt base causes the traffic-induced strains in the pavement structure, which control fatigue and structural deformation, to decrease with time. Therefore, a road will be more vulnerable to structural damage in its early life, before curing has increased the structural strength of the material. If the road is designed and constructed with sufficient strength to prevent structural damage in its early life, it has been found that curing doubles the stiffness of DBM roadbase [now base, of course] in the first few years in service and this will substantially improve the overall resistance of the pavement to fatigue and structural deformation. The improvement in the bearing capacity of the road, as determined by deflection measurements, provides confirmation of this improvement.
Figure C2 in TRL 250 (Nunn et al., 1997) is reproduced here as Figure 20.7. As can be seen by examining the figure, the stiffness of an asphalt concrete base (the most common type of base in UK roads), which is initially 2 GPa at the time of laying, increases to around 4 GPa 4 years after laying. So, the stiffness has doubled in a period of 4 years. Clearly, the increase in the stiffness of asphalt bases over time will improve the structural competence of a road pavement, and has implications for the structural design of flexible pavements (see section 13.3).

20.2.4.3 Bitumen hardening in surface dressings

Surface dressings probably have the greatest exposure to environmental ageing mechanisms of all bituminous products. Sprayed surface treatments, known as ‘surface dressings’ in the UK, ‘chip seals’ in the USA and New Zealand, and ‘sprayed seals’ in Australia, will present a binder film with a depth of around 1.5 mm on the existing road surface on top of which the chippings will be spread. The mosaic achieved by the chippings will determine the extent of the dressing that is directly exposed to the air but the overall proximity of the surface dressing to the elements means that the binder is aggressively subjected to ageing factors.

Countries such as Australia that rely on sprayed seals for much of the rural road network have paid close attention to the rate of hardening of bitumen in seals. The Australian Road Research Board conducted investigations into
the performance and longevity of sprayed seals, and proposed a relationship between binder ageing and seal life (Dickinson, 1984).

20.3 Bitumen ageing tests
It is clearly desirable that there should be laboratory tests that quantitatively determine the resistance of bitumens to hardening at the various stages during the production process. A number of tests already exist to measure the effect of heat and air on bitumen. The main aim of these tests is to identify bitumens that are too volatile or are too susceptible to oxidation to perform well in service (for more information on this testing see section 5.5.4).

The thin-film oven test (Lewis and Welborn, 1940) simulates practical conditions. In this test, the bitumen is stored at 163°C for 5 h in a layer 3.2 μm thick. It is claimed that in this test the amount of hardening that takes place is about the same as that which occurs in practice. However, diffusion in the bitumen film is also limited, and it is not possible to obtain homogeneous hardening or ageing. Accordingly, the test is far from ideal. This test was adopted initially by the American Society for Testing Materials (ASTM) in 1969 as method ASTM D1754 and has been modified since that time to include improvements (ASTM, 2009).

In 1963, the State of California Department of Public Works, Division of Highways, developed a test that more accurately simulates what happens to a bitumen during mixing. It is called the rolling thin-film oven test (RTFOT) (Hveem et al., 1963). In this test, eight cylindrical glass containers each containing 35 g of bitumen are fixed in a vertically rotating shelf. During the test, the bitumen flows continuously around the inner surface of each container in a relatively thin film, with preheated air blown periodically into each glass jar. The test temperature is normally 163°C for a period of 75 min. The method ensures that all the bitumen is exposed to heat and air, and continuous movement ensures that no skin develops to protect the bitumen. A homogeneously aged material, similar to that which is produced during full scale mixing, is obtained. Clearly, the conditions in the test are not identical to those found in practice but experience has shown that the amount of hardening in the RTFOT correlates reasonably well with that observed in a conventional batch mixer. However, the mixing conditions in efficient modern drum mix plants are less aggressive, and bitumen hardening during asphalt production in such plants is less than predicted by the RTFOT.

The RTFOT was accepted in 1970 by the ASTM as method ASTM D2872-12e1 (ASTM, 2012) and was included as part of the European specification for paving grade bitumens in BS EN 12591:2000 (BSI, 2000) and is part of the Superpave specification used in the USA (AI, 1997).
Australia utilises the RTFOT apparatus artificially to age samples of bitumen to a specified apparent viscosity level (SAVL – taken to be 5.67 log Pa·s at 45°C). The SAVL is considered to equate to the viscosity of a bitumen in a sprayed seal at the end of its service life, and is measured using the Shell sliding plate viscometer (Figure 20.8) (Standards Australia, 1997a).

A correlation between the time taken for a bitumen to reach the SAVL and the in situ service life of a seal was proposed by the Australian Road Research Board (ARRB) from data obtained from bitumen recovered from aged sprayed seals. The assessment is called the ‘durability test’, and is applied by a number of road agencies in Australia as a control point on the suitability of bitumen for use in sprayed seals (Standards Australia, 1997a). Australia is one of the few countries where a specific test is employed to quantify the durability of bitumen (Standards Australia, 1997b).

Over the years there have been a number of attempts to simulate the long term ageing of bitumen in asphalts but this has proved to be extremely difficult because of the number of variables that affect binder ageing – void content, mixture type, aggregate type, etc. The Strategic Highway Research Program (SHRP) investigated accelerated ageing techniques and identified the pressure ageing vessel (PAV) (Anderson et al., 1994) as the preferred apparatus to simulate long term ageing of bitumen. The US Superpave
The Shell Bitumen Handbook

specification [AI, 1996] uses the RTFOT to simulate initial ageing, followed by ageing over 20 h at elevated temperature (90, 100 or 110°C) and pressure 2070 kPa in a PAV. After this ageing procedure, the residue is used for dynamic shear rheometry, bending beam rheometry and direct tension testing. The use of the PAV, using modified conditions, is currently being considered in Europe as a method for ageing bitumens in the laboratory. The artificial ageing of binders in the PAV to simulate ageing in situ has still to be fully validated but the technique is now widely accepted as a satisfactory approach.

References


Surface dressing is a treatment undertaken on roads and other types of pavement. As a maintenance process it has three purposes:

- to provide both texture and skid resistance to the surface
- to seal the road surface against ingress of water
- to arrest disintegration, and hence to extend the life, of the pavement and to assist sustainable development.

The aims listed above are taken from Road Note 39 (Roberts and Nicholls, 2008). Road Note 39 is the default design guide for UK surface dressing activities and would be used for the vast majority of surface dressing designs in the UK.

In addition to the purposes listed above, surface dressing can be used as a treatment to provide:

- a distinctive colour to the road surface
- a more uniform appearance to a patched road.

However, surface dressing cannot restore good surface regularity to a deformed road, nor will it contribute to the structural strength of the pavement.

In a new road, adequate surface texture is designed into the running surface by specifying requirements for both aggregate properties and texture depth. However, during its service life, the surface becomes polished under the action of traffic, and the skid resistance will eventually fall below the acceptable
A major benefit of surface dressing is that the process restores skid resistance to a surface that has become smooth under traffic.

Surface dressing is the most extensively used form of surface treatment on roads. In Europe, the USA, China and Australia, it is widely used for preventive maintenance. In the UK, data suggest that the road lengths of surface dressing are more than 50% of total lengths of maintenance in England and Wales (DfT, 2014). In Australia, about 90% of paved roads are surface dressed (Holtrop, 2008). With care, surface dressing can be used on roads of all types, from a country lane carrying only an occasional vehicle to trunk roads carrying tens of thousands of vehicles a day. However, in some locations, particularly those where vehicles undertake sudden or sharp turning manoeuvres, surface dressing may not be the appropriate surface treatment. This is due to the relatively poor ability of the finished surface to resist tangential forces. In addition, the durability of a surface dressing does not match that of a road resurfaced with traditional asphalt. Some large cities, Hamburg for example, prohibit the use of surface dressings on their carriageways.

Basic surface dressing consists of spraying a film of binder onto the existing road surface, followed by the application of a layer of aggregate chippings as shown in Figure 21.1. The chippings are then rolled to promote contact between the chippings and the binder, and to initiate the formation of an interlocking mosaic. On carriageways having a low value of road hardness, rolling may also cause the chippings to start the process of embedment into...
the existing road surface. (Road hardness is a property that represents the resistance of an existing road surface to the embedment of chippings – see section 7.2.2 of Road Note 39 (Roberts and Nicholls, 2008).)

21.1.1 The design of surface dressings

The design of surface dressings primarily addresses the retention of surface texture. The aim is to design a system that takes account of the hardness of the existing road such that the aggregate is adequately embedded by rolling and, subsequently, traffic.

In the UK, Road Note 39 (Roberts and Nicholls, 2008) and the Code of Practice for Surface Dressing (RSTA, 2014a) are commonly used to provide guidance on the appropriate type of dressing, its design and execution. The principal steps per Road Note 39 are as follows:

- select the size and type of dressing that is appropriate for the particular site and the time of application
- identify the chipping size and basic binder application rate and type of binder
- refine the binder application rate to match the source of chippings selected
- further refine the binder application rate on site immediately before work starts.

Road Note 39 is set out in such a way that the application rate of the binder is only obtainable after completion of the steps outlined above. Tables provide coefficients for application rates of binders for the full range of types of surface dressing (i.e. single, racked-in, double, sandwich and inverted double surface dressings). These coefficients are then adjusted by reference to tables that take into account the type of aggregate and its shape, the condition of the road surface, the gradient, the degree to which the site is shaded by trees and structures, and steps to be taken for sites where traffic volumes are exceedingly light. The coefficients are then translated into actual target application rates of the binder expressed in litres per square metre.

Many countries in Europe employ the CEN (European Committee for Standardization) standards. The CEN specifications for surface dressing are based on the UK’s Road Note 39.

In the USA, individual states have their own specifications. Some states have published documents on the subject (e.g. the Minnesota Seal Coat Handbook 2006 (Wood et al., 2006) and the Texas Seal Coat and Surface Treatment Manual (TxDOT, 2010)). In addition, the Asphalt Institute (a US-based international trade association of bitumen producers, manufacturers and affiliated businesses) and ASTM International (the

Austroads is the association of Australian and New Zealand road transport and traffic authorities. It provides two technical reports for the design of binder rates of application and aggregate spread rates for sprayed seal

- AP-T68/06 (Update of the Austroads Sprayed Seal Design Method) (Austroads, 2006)

Guidance on selection of sprayed-seal-type materials and sprayed-seal construction is provided in the Guide to Pavement Technology

- Part 3: Pavement Surfacing (AGPT03-09) (Austroads, 2009a)
- Part 4F: Bituminous Binders (AGPT04F-08) (Austroads, 2008)
- Part 4K: Seals (AGPT04K-09) (Austroads, 2009b)
- Part 8: Pavement Construction (AGPT08-09) (Austroads, 2009c).

In China, two industry standards dictate the requirements for type selection, materials and construction specifications for surface dressing. These standards are

- JTG D50-2006: Specifications for Design of Highway Asphalt Pavement (RIOH, 2006a)

21.1.2 Factors that influence the design and performance of surface dressings

The principal factors that influence the design, performance and service life of a surface dressing are

- traffic volumes and speeds
- condition and hardness of the existing road surface
- size and other chipping characteristics
- surface texture and skid resistance of the existing road surface
- type of binder
- adhesion
- geometry of site, altitude, latitude and local circumstances
- time of year.

21.1.2.1 Traffic volumes and speeds

The degree of chipping embedment is a function of the quantity of vehicles that use the route being designed. Accordingly, it is necessary to establish
the number of heavier vehicles that regularly use that route and the speed of such traffic. In the UK, the number of medium and heavy vehicles (i.e. those having an unladen weight exceeding 1.5 tonnes) currently travelling in the lane under consideration is normally used for design purposes according to Table 7.2.3 of Road Note 39 (Roberts and Nicholls, 2008).

21.1.2.2 Condition and hardness of the existing road surface
If the existing road surface is rutted, cracked or in need of patching, these defects must be corrected before surface dressing can be undertaken (RSTA, 2014b). Other surface conditions that require special consideration are those in which the binder has flushed up, resulting in a binder-rich surface and, conversely, surfaces that have become very dry, lacking in binder.

The hardness of the road surface affects the extent to which the applied chippings become embedded in the road surface during the life of the dressing. The choice of chipping size is directly related to the hardness of the existing road surface. The use of chippings that are too small will result in early embedment of the chippings in the surface, leading to a rapid loss of texture depth and, in the worst cases, ‘fatting up’ of the binder, which may cover the entire surface of the road. (‘Fatting up’ is where there is an excess of bitumen on the road surface, sometimes but less commonly described as ‘flushing-up’.) The use of chippings that are too large may result in immediate failure of the treatment due to stripping of the binder from the aggregate under the applied stresses of the traffic, and can also lead to excessive surface texture and an increase in the noise generated between the tyre and the road surface. The hardness of the road surface also influences the rate of application of binder required for a given size of chipping. The rate of spread must be reduced where the road surface is soft in order to compensate for the greater embedment of chippings in the road surface under the action of traffic.

In the UK, the categories of hardness of the existing road surfaces for the purposes of surface dressing design are described in Road Note 39. Four graphs take account of the geographical location within the UK and the altitude of the site. The geographical location is not strictly related to latitude but also takes account of climatic conditions. The method of measuring hardness on the site is by use of a road hardness probe, which is described in BS 598-112 (BSI, 2004). Hardness measurements are taken in conjunction with road surface temperature readings. The standard specifies that road surface temperatures must be in the range 15–35°C.

Having identified the applicable surface temperature category and measured the depth of penetration and the surface temperature, the hardness
category of the road can be established by reference to Figure 7.2.2 in Road Note 39 (Roberts and Nicholls, 2008), as shown in Figure 21.2.

In Australia, the ball penetration test is used for testing the pavement hardness. The test method is described in Austroads test method AG:PT/T251 (Austroads, 2010).

In the USA, some guides (TxDOT, 2010; Wood et al., 2006) for surface dressing provide a surface correction factor based on the existing pavement condition.

21.1.2.3 Size and other chipping characteristics
The selection of the size and type of chipping should take into account the following factors.

- The size of chipping has to offset the gradual embedment in road surfaces of different hardness caused by traffic.
- Maintaining both microtexture and macrotexture during the life of the surface dressing by selecting a chipping of appropriate
  - size
  - polished stone value (PSV)
  - aggregate abrasion value (AAV).
- Relating the traffic category for each lane and the road surface hardness category to the size of chipping required.

In the UK, chippings for surface dressing should meet the requirements of BS EN 13043:2002 (BSI, 2002a). Usually, chippings have a nominal size of 2.8/6.3, 6.3/10 or 8/14 mm. Extreme caution should be exercised when using chippings larger than 8/14 mm nominal size due to the risk of premature failure. In China, aggregate requirements are given in JTG F40-2004 (RIOH, 2004), the nominal size is from 3/5 to 15/30 mm, and in Australia it is up to 20 mm. In the USA, the nominal size of the largest grade is normally 19 mm, although in some states it is 12.7 mm.

The presence of dust can delay or even prevent adhesion and this problem is particularly acute at low temperatures and with the smaller sizes of chipping. As well as being of nominal single size and dust free, surface dressing chippings need to satisfy other characteristics. They should

- not crush under the action of traffic or shatter on impact
- resist polishing under the action of traffic
- have an acceptable value of ‘flakiness index’ (the ratio of length to thickness).

Chippings are rarely cubic, and when they fall onto the binder film and are rolled they tend to lie on their longest dimension. If the chippings used for a
Figure 21.2  Hardness categories from depth of penetration and road surface temperature for different surface temperature categories (Roberts and Nicholls, 2008). (Reproduced by courtesy of the Transport Research Laboratory (TRL))
surface dressing have a large proportion of flaky chippings, less binder will be required to hold them in place, and the excess binder may ‘flush-up’ onto the surface of the dressing. Such a dressing will have a reduced texture and life. It is for this reason that the UK sets a maximum limit of flakiness for single size chippings for use in surface dressing. Typical requirements for surface dressing chippings are shown in Table 21.1.

Gritstones, basalts, quartzitic gravels and artificial aggregates such as slag constitute the majority of chippings used for surface dressing. However, not all aggregates with a high PSV are suitable for surface dressing due to them possessing insufficient strength. The PSV of the aggregate defines its skid resistance. In the UK, its determination is dictated by BS EN 1097-8:2009 (BSI, 2009) and the relationship between PSV, traffic and skid resistance is detailed in HD 36 (Highways Agency et al., 2006). Coated chippings have a thin film of bitumen applied at an asphalt plant. This bitumen film eliminates surface dust and promotes rapid adhesion to the bitumen. Coated chippings are used to improve adhesion with cut-back bitumen, particularly in the cooler conditions that occur at the extremes of the season. They should not be used with emulsions, as the shielding effect of the binder will delay ‘breaking’ of the emulsion (emulsions contain water, which evaporates after spraying; the process of ‘breaking’ is discussed in detail in section 21.1.2.5).

21.1.2.4 Surface texture and skid resistance

In the UK, two machines are used for monitoring the skid resistance of roads: the SCRIM (Figure 21.3) and the GripTester (Figure 21.4).

SCRIM is the acronym for the ‘sideways-force coefficient routine investigation machine’. This machine was introduced into the UK in the 1970s. It provides a measure of the wet road skid resistance properties of a bound surface by measurement of a sideways-force coefficient (SFC) at a controlled speed.

| Table 21.1 Typical properties of surface dressing chippings |
|---------------------------------|----------------|
| Property                        | Value          |
| Los Angeles coefficient (LA)    | LA₃₀           |
| Flakiness index (FI)            | FI₂₀           |
| Polished stone value (PSV)      | PSV₅₆ᵃ          |
| Aggregate abrasion value (AAV)  | AAV₁₀          |

ᵃ The required PSV value is usually stated in individual contracts. Because the PSV categories in BS EN 13043:2002 do not always use values that correspond with established UK practice, the standard allows an aggregate supplier to declare an intermediate value.
Surface treatments

Figure 21.3 SCRIM. (Courtesy of PTS Ltd.)

Figure 21.4 GripTester. (Courtesy of PTS Ltd.)
The GripTester was introduced in the 1980s, and is a braked wheel fixed-slip trailer. The resulting horizontal drag and the vertical load on the measuring wheel are continuously measured, and the coefficient of friction, known as the GripNumber, is calculated. The GripNumber can be converted to a SCRM coefficient using the following relationship (Dunford, 2010)

$$SC = 0.89 \times GN$$

where \(SC\) is the SCRM coefficient and \(GN\) is the GripNumber.

In China, besides the SCRM, two other machines are used: one is the Mu-Meter (Figure 21.5) and the other is the dynamic friction tester (Figure 21.6). The Mu-Meter is manufactured in the UK, and the principle is the same as that employed in the SCRM but, like the GripTester, the Mu-Meter is much smaller than the SCRM. The dynamic friction tester is made in Japan and its use is described in ASTM E1911-09ae1 (ASTM, 2009a).

The values of both the SFC and the GripNumber vary with speed. Testing is normally carried out at 50 km/h. Corrections may be made to take account of variations in speed, but for these to be reliable it is necessary for the macrotexture of the surface to be known. Tight radii, such as occur at roundabouts, are normally tested at 20 km/h.

The SFC and GripNumber also vary with traffic flow, temperature and time of year. Network testing is restricted to May to September. Three tests are carried out at fairly similar intervals throughout the season. The average is defined as the ‘mean summer SCRM coefficient’ (MSSC), and skid resistance is normally expressed in terms of the MSSC.

The skid resistance of a road surface is determined by two basic characteristics, the microtexture and macrotexture, as shown in Figure 21.7, which is
Figure 2.1.2 in Road Note 39 (Roberts and Nicholls, 2008). Microtexture is the surface texture of the aggregate. A significant level of microtexture is necessary to enable vehicle tyres to penetrate thin films of water and thus achieve dry contact between the tyre and the aggregate on the carriageway. Macrotexure is the overall texture of the road surface. An adequate value of macrotexure is necessary to provide drainage channels for the removal of bulk water from the road surface.

Figure 21.7 Microtexture and macrotexure (Roberts and Nicholls, 2008). (Reproduced by courtesy of the Transport Research Laboratory (TRL))
TRL Research Report 296 (Roe et al., 1991) considered the relationship between accident frequency and surface texture on roads. It concluded that, on asphalt road surfaces with speed limits greater than 64 km/h, and possibly on roads with lower speed limits

- skidding and non-skidding accidents, in both wet and dry conditions, are fewer if the microtexture is coarse than if it is fine
- the texture level below which accident risk begins to increase is a sensor measured texture depth (SMTD, see below) of around 0.7 mm
- all major types of surfacing provide texture depths across the practical range
- macrotexture has a similar influence on accidents, whether they occur near hazards such as junctions or elsewhere.

Microtexture is the dominant factor in determining the level of skid resistance at speeds up to 50 km/h. Thereafter, macrotexture predominates, particularly in wet conditions. Traditionally, macrotexture has been measured using the volumetric patch technique (BSI, 2010).

Mature surface dressings in good condition typically have texture depths between 1.5 and 2.0 mm; double dressings usually have slightly lower values. Double surface dressings, however, usually stay above intervention levels, particularly on hard surfaces such as those surfaced with a hot rolled asphalt surface course.

Laser texture measurement (technically, SMTD) is a technique that tests areas much more quickly than is the case with volumetric patch testing. Accordingly, it is used as a screening method. The technique is illustrated in Figure 21.8, which is Figure 1 in TRL Research Report 143 (Roe et al., 1988). This method is not directly comparable with measurements resulting from volumetric patch testing. Thus, when quoting texture depth values, the method used to measure the parameter should be stated.

It is the petrographic characteristics of the aggregate that largely influence its resistance to polishing, and this property is measured using the PSV test (BSI, 2009). A high PSV (say 65 or higher) indicates good resistance to polishing. The majority of road aggregates have a good microtexture prior to trafficking and, consequently, most road surfaces have a high skid resistance when new, particularly after the bitumen has worn off the aggregate. The time required for the bitumen to be abraded from the aggregate at the road surface depends on a number of factors, including the type of mixture and, in particular, the thickness of the film of bitumen on the particles of aggregate. Until the latter has been removed by traffic, the microtexture of the aggregate will not be fully exposed to vehicle tyres. Subsequently, the
exposed aggregate surfaces become polished and, within a year, the skid resistance normally falls to an equilibrium level.

The macrotexture is determined by the nominal size of aggregate used and the nature of the asphalt. However, resistance to abrasion is also important. It is expressed as the AAV. Aggregates with poor abrasion resistance, determined using the AAV test (BSI, 2009), will quickly be worn away, with a consequent loss of macrotexture. The lower the AAV, the greater the resistance to abrasion. High speed roads normally require an AAV of 14 or less. Surface dressings generally tend to last longer when the chippings have a lower AAV.

The rate at which skid resistance falls with increasing vehicle speed is influenced by the macrotexture. The TRL has shown that there is a correlation between the macrotexture, expressed in terms of average texture depth, and the skid resistance, measured by the percentage reduction in the braking-force coefficient between 130 and 50 km/h, as shown in Table 21.2, which is Table 4 in Laboratory Report 510 (Salt and Szatkowski, 1973). In the table, ‘flexible’ means fully asphaltic pavements, whereas ‘concrete’ means rigid, cementitious pavements.

The skid resistance necessary at any site will depend on the stresses likely to be induced when the road is in service. Sections of road with minor road junctions and with low traffic volumes are unlikely to require a high value of skid resistance, whereas sharp bends on roads carrying heavy volumes of traffic will require significantly higher skid resistance characteristics. It is

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**Figure 21.8** Principle of the laser-based contactless sensor for measuring texture depth (Roe et al., 1988). (Reproduced by courtesy of the Transport Research Laboratory (TRL))
important to note that roads carrying high volumes of traffic will require aggregates with higher PSVs than will roads where traffic levels are low or moderate.

21.1.2.5 Type of binder
The essential functions of the binder are to seal surface cracks and to provide the initial bond between the chippings and the road surface. The viscosity of the binder must be such that it can wet the chippings adequately at the time of application and prevent dislodgement of chippings when the road is opened to traffic. In addition, the bitumen should not become brittle during periods of prolonged low temperature, and should function effectively for the design life of the dressing.

The subsequent role of the binder is dependent on the traffic stresses applied to the surface dressing, from low stressed, lightly trafficked country roads to highly stressed sites carrying large volumes of traffic. Clearly, a range of binders is necessary if optimum performance is to be achieved under these widely differing circumstances.

The most commonly and widely used binders for surface dressing are

- conventional bitumen emulsion
- polymer modified bitumen
- polymer modified bitumen emulsion.

Other types are also used in some countries

- conventional bitumen
- conventional fluxed or cut-back bitumen
- polymer modified fluxed or cut-back bitumen
- epoxy resin thermoset binder.

**Conventional bitumen emulsion**
Emulsions used for surface dressing are normally rapid setting. In the EU, they should comply with BS EN 13808:2013 (BSI, 2013). In Australia, these
are described in AS 1160-1996 (Standards Australia, 1996), and in JTG F40-2004 (RIOH, 2004) in China.

The relatively low viscosity of bitumen emulsions enables them to be sprayed at temperatures between 75°C and 85°C through either swirl or slot jets (jet types are discussed in section 21.1.4.1). There is the possibility that the binder will flow on road surfaces at the time of spraying. While this is helpful in some respects, migration of the binder from the centre to the side of cambered roads, or downhill on gradients, requires the application of chippings to the binder film as quickly as possible after spraying in order to counter any tendency to flow.

After spraying, the water migrates from the emulsion, leaving a film of bitumen, a process known as ‘breaking’. This can take from as little as 10 min to several hours depending on ambient temperature, humidity, aggregate water absorption, wind speed and the extent to which the dressing is subject to slow moving traffic. Humidity is one of the most important of these factors, and the use of emulsions when humidity is above 80% should only be undertaken on very minor roads and where traffic speeds can be kept within the range 10–20 mph until the emulsion has fully broken. It is simple to check whether the emulsion has broken, as the colour of the emulsion changes from brown to black. In conditions of high humidity, work can continue with emulsions, particularly when the binder is polymer modified, by varying the type of dressing. Good results can be obtained with double and sandwich surface dressings. Precoated chippings should not be used with emulsions, as the coating delays breaking and provides no tangible benefit but carries additional cost.

Binder manufacturers are able to control the rate at which emulsions break within certain limits, but these have to be compatible with the storage, transport and application of the emulsion. On site, breaking can be accelerated by spraying the binder film with a mist of a breaking agent before the application of the chippings. However, this usually requires specially adapted spray bars and the fitting of storage tanks for the breaking agent.

The advantages of emulsion binders are

- low spraying temperature
- low risk of early bleeding
- may be used on higher viscosity base bitumen
- may be used on damp (not wet) surfaces
- do not require precoated chippings.

The disadvantages are

- low initial adhesion
risk of brittle failure in the first winter (employing modified binders will reduce this risk considerably)

- skinning – the aggregates will not be wetted properly resulting in poor adhesion, together with the fact that the aggregate particles tend to sit on top of the emulsion layer rather than penetrating through it. The use of a sandwich or, preferably, a double surface dressing will reduce the film thickness and, thus, the possibility of skinning.

Skinning is the phenomenon that occurs when a thin layer at the surface of the emulsion breaks and forms a ‘skin’, or layer of bitumen, over the rest of the emulsion, which then stops it from breaking.

**Polymer modified bitumen**

The relentless increase in the volume of all types of vehicles on roads throughout Europe has led to the development of modified binders with improved adhesivity and substantially higher cohesive strengths than traditional fluxed/cut-back and emulsion binders. These characteristics are particularly important immediately after chippings have been applied to the binder film, and the use of a modified binder means that the chippings are less likely to be plucked from the road surface by passing traffic than would be the case with conventional unmodified binders. As well as greater adhesive and cohesive strength, modified binders perform satisfactorily over a wider temperature range than is the case with traditional binders. The result is that modified binders do not flush up as rapidly in hot conditions as traditional binders, and are not subject to the same degree of brittleness in winter conditions.

A high proportion of the European road network is lightly or moderately trafficked, and many of the roads can be surface dressed using bitumen emulsion or cut-back bitumen. However, enhanced treatment may be necessary on certain highly stressed sections, such as at sharp bends, road junctions and in shaded areas under trees and bridges and adjacent to trees or buildings. At these locations, it is probable that the same binder can be used, but the specification will change in respect of one or more of the following:

- rate of application of binder
- chipping size and rate of spread
- double chip application
- double binder, double chip application.

The double chip techniques are widely practised in France on relatively heavily trafficked roads, and, by adopting appropriate procedures, the risk of flying chippings and consequential windscreen breakage can be reduced significantly.
As a further aid to consistent performance under variable weather conditions, a number of proprietary adhesion agents are available. These act in two basic ways. Some types (active) modify surface tension at the stone/binder interface to improve the preferential wetting of chippings by the binder in the presence of water. Other types (passive) improve the adhesion characteristics of the binder, resulting in greater resistance to the subsequent detachment of the binder film in the presence of water throughout the early life of the dressing.

Bearing these performance requirements in mind, it is possible to examine the ways in which the various polymers can be used with bitumen to modify its performance. The bitumen properties in which improvement would be sought through the addition of a modifier are as follows:

- Reduced temperature susceptibility, which can be loosely defined as the extent to which a binder softens over a given temperature range. It is clearly desirable for a binder to exhibit a minimum variation in viscosity over a wide range of ambient temperatures.
- Cohesive strength is the property that enables the binder to hold chippings in place when they are subject to stress by traffic. This property can also be coupled with the elastic recovery properties of the binder, to enable a surface dressing to maintain its integrity even when it is subjected to high levels of strain.
- Adhesive power or ‘tack’ is an essential property of binders but it is difficult to define. The adhesion characteristics are particularly important during the early life of a surface dressing before any mechanical interlock or embedment of the chippings has taken place.

In addition, it is desirable for the modified binder to

- maintain its premium properties for long periods in storage and subsequently during its service life
- be physically and chemically stable during storage and spraying and at in-service temperatures
- be capable of being applied using conventional spraying equipment
- be cost effective.

Technically, the choice between an emulsion or cut-back formulation for modified binders is not easy, but health and safety concerns, and applicable regulations, often limit the use of cut-backs. In the UK, the use of cut-backs has dropped enormously because of safety concerns, cut-backs being applied at much higher temperatures than is the case with emulsions.

The development of polymer modified surface dressing binders to complement conventional and epoxy resin systems offers highway engineers a range of binders suitable for all categories of site. Regardless of the improved
properties of a particular binder, it is the performance of the aggregate–
binder system and application mode that will dictate the success or otherwise
of the dressing.

**Polymer modified bitumen emulsion**
This class of materials can be produced using polymer or latex modified
bitumen. Modifying with latex is normally done prior to or after emulsification.
These materials have similar advantages to those of polymer modified bitumen.
In the UK in 2004 about 80% of surface dressings were with polymer modified
dressings (Roberts and Nicholls, 2008). They should meet the requirements of
BS EN 13808:2013 (BSI, 2013). There are three grades based on the mini-
mum levels of peak cohesion of residual bitumen according to Table 5.3.4 in
Road Note 39 (Roberts and Nicholls, 2008). The test method for measuring
peak cohesion is described in BS EN 13588:2008 (BSI, 2008a), and the
grades are shown in Table 21.3. Emulsions having a higher level of cohesion
are appropriate for higher stress situations on the road.

Similar to conventional bitumen emulsion, surface dressing using polymer
modified emulsion is also vulnerable during the early life of the dressing
before the emulsion has fully broken. The rate at which breaking occurs is
influenced by a number of factors, including emulsion formulation, weather
and mechanical agitation during rolling and trafficking. Under certain con-
ditions, the emulsion goes through a state during curing when the bitumen
droplets have agglomerated but not coalesced, and the dressing is extremely
vulnerable. The timing of this condition is variable, and is generally
worsened by the presence of high polymer contents, which can cause skin-
nings and retard breaking of the emulsion. It is essential that the emulsion has
broken fully before uncontrolled traffic is allowed on the dressing. This can be
a serious disadvantage on heavily trafficked roads or where work is
executed under lane rental contracts. Chemical after-treatment sprays have
been shown to be effective in promoting breaking.

**Fluxed or cut-back bitumen**
Fluxed or cut-back bitumen for surface dressing is bitumen that has been diluted
with kerosene, usually 70/100 or 160/220 pen. Cut-back bitumens have
benefits because of their immediate cohesive grip of the road and chippings,
but generally the residual binder properties are more variable due to the differences in the evolution of the diluent and its absorption into the road surface.

Cut-back bitumen is typically sprayed at temperatures in the range 130–170°C (compared with 75–85°C for conventional bitumen emulsion). During spraying, between 10% and 15% of the kerosene evaporates, and a further 50% dissipates from the surface dressing in the first few years after application. At the spraying temperatures, wetting of the road surface and the applied chippings is achieved rapidly. Cut-back bitumen, once applied to the road surface, cools quickly, and it is important to ensure that the chippings are applied as soon as possible after spraying to ensure satisfactory wetting of the chippings.

The trend in most countries is to move away from the use of traditional cut-back/fluxed bitumens where possible.

**Epoxy resin thermoset binder**

This group of binders is bitumen extended epoxy resin binders used with high PSV aggregates such as calcined bauxite. Epoxy resin based binders are used in high performance systems, such as Spraygrip, that are fully capable of resisting the stresses imposed by traffic on the most difficult sites (e.g. roundabouts or approaches to traffic lights and designated accident black spots). It is these systems that are often described as ‘Shellgrip’ but are more correctly generically described as ‘high friction surfaces’. Binders used in such systems are classified as ‘thermosetting’, as the epoxy resin components cause the binder to cure by chemical action and harden, and it is not subsequently softened by high ambient temperatures or by the spillage of fuel. The dressing thus acts as an effective seal against the ingress of oil and fuel, which is particularly important on roundabouts, where spillages regularly occur. Very little embedment of the chippings in the road surface takes place with this type of binder, and the integrity of the surface dressing is largely a function of the cohesive strength of the binder.

The extended life of this binder justifies the use of a durable aggregate with an exceptionally high PSV. The initial cost of these surface dressing systems is high compared with systems based on conventional binders. However, their exceptional wear resistant properties and the ability to maintain the highest levels of skid resistance throughout their service lives make them a cost effective solution for very difficult sites by significantly reducing the number of skidding accidents. Statistics suggest that, in the UK, the average value of prevention per serious accident is far less than a fatal accident (DfT, 2013).

**21.1.2.6 Adhesion**

Some aggregates adhere more rapidly to binder than others, and it is wise to avoid problems by testing the compatibility of the selected binder and
chippings. When using cut-back binder, this can be checked using the total water immersion test. In this test, clean 8/14 mm chippings are totally coated with cut-back bitumen. After 30 min of curing at ambient temperature, the chippings are immersed in demineralised water at 40°C for 3 h. After soaking, the percentage of binder that is retained on the chips is assessed visually. The extent to which the base of the chipping has been coated with the binder after 10 min is then assessed visually. If the coated area is considered to be below 70%, serious consideration should be given to lightly precoating the chippings, as it is much easier to obtain a bond between the binder film and a light coating of binder on the chippings than is the case with uncoated chippings.

The amount of binder required to coat surface dressing chippings lightly varies according to the type of binder and the size of the chipping but it is typically around 1% by weight. A thick coating of binder (e.g. as in the precoated chippings often used with hot rolled asphalt surface course) will make the chippings sticky and will inhibit free flow through chipping machines. When lightly coated chippings are used, this free flow property is critical to a satisfactory surface dressing. Minor pinholes in the coating of the chippings are not detrimental. It is particularly important that the amount of filler is less than the specified level (usually ≤1% passing a 0.063 mm sieve).

Once the chippings have been lightly coated, it is important that they are not allowed to come into contact with dust, as this will impair the bond between the chippings and the applied binder film on the road surface.

21.1.2.7 Geometry of site, altitude and latitude and local circumstances
Sections of road that include sharp bends induce increased traffic stresses on a surface dressing. Three categories of bend are identified in Road Note 39 (Roberts and Nicholls, 2008) – under 100 m radius, 100–250 m radius and over 250 m radius. Gradients that are steeper than 10% will affect the rate at which the binder should be applied due to the fact that on the ascent side of the road traffic, loading will be present for longer than on the descent side. Typically, there is a difference of 0.2 l/m² in the rate of binder application between the ascent and the descent sides.

21.1.2.8 Time of year
This is discussed in section 21.1.6.

21.1.3 Types of surface dressing
The original concept of a normal single layer surface dressing has been developed over the years, and there are now a number of techniques available that vary according to the number of layers of chippings and the number
of applications of binder. Each of these techniques has its own particular advantages and associated cost, and it is quite feasible that along the length of a road a variety of techniques would be used depending on the features of particular sections. Figure 21.9 (a composite of Figures 2.2.2 to 2.2.6 in

**Figure 21.9** Schematic representation of different types of surface dressing prior to embedment (Roberts and Nicholls, 2008). (Reproduced by courtesy of the Transport Research Laboratory (TRL))
Road Note 39 (Roberts and Nicholls, 2008) diagrammatically depicts the different types of surface dressing that are available.

21.1.3.1 Pad coats
A pad coat consists of a single dressing with small chippings, and is applied to a road that has uneven surface hardness, possibly due to extensive patching by utilities. The reason for applying a pad coat is to produce a more uniform surface than would result by simply surface dressing the road. The chipping size for a pad coat is traditionally 2.8/6 mm, with a slight excess of chippings. Pad coats can also be used on very hard road surfaces (such as concrete or heavily chipped asphalts) to reduce the effective hardness of the surface, although using a racked-in system with a polymer modified binder is now the preferred option.

After laying and compaction by traffic, excess chippings should be removed before the road is opened to unrestricted traffic. Pad coats may be left for several months before the application of the main dressing, which may be either a single dressing or a racked-in system. Either system will embed in the pad coat and have immediate significant mechanical strength, reducing the risk of failure. All loose material should be swept from the surface of the road prior to the application of the final dressing.

21.1.3.2 Single surface dressing
The single surface dressing system is still the most widely used type of surface dressing. It consists of one application of binder followed by one application of single size chippings. Its advantages are that it has the least number of operations, uses the least amount of material and is sufficiently robust for minor roads and some main roads where excessive braking and acceleration are unlikely to occur and where speeds are unlikely to exceed 100 km/h.

21.1.3.3 Racked-in surface dressing
This type of dressing uses a single application of binder. A first application of 6.3/10 or 8/14 mm chippings (called the ‘principal chippings’) is spread at approximately 80–90% cover, which leaves a window of binder between the chippings. This window is filled with small chippings of size 2/4 mm (for a 10 mm principal chipping) or 2.8/6.3 mm (for a 14 mm principal chipping) to achieve high mechanical interlock (called a ‘mosaic’). The rate of application of binder on this type of surface dressing is usually slightly higher than that which would be required for a single size dressing.

The initial surface texture and mechanical strength of a racked-in dressing are high. The configuration of the principal chippings is different from that which is achieved with a single dressing. The mosaic, referred to above, cannot be
formed because the principal chippings are locked in place. In time, small chippings that are not in contact with binder are lost to the system without damaging vehicles or windscreens, resulting in increasing surface texture despite some coincident embedment of the larger chipping.

On a racked-in dressing, vehicle–tyre contact is principally on the large aggregate, with little contact on the small aggregate. Thus, the microtexture of the small aggregate is less critical, and consideration should be given to the use of cheaper lower PSV aggregates, such as limestone, for the small chippings.

The advantages of the racked-in system are

- virtual elimination of ‘flying’ 8/14 or 6.3/10 mm chippings, thereby reducing the incidence of broken car windscreens
- early stability of the dressing through good mechanical interlock
- good adhesion of larger size chippings
- a rugous surface texture with an initial texture depth exceeding 3.0 mm.

21.1.3.4 Double surface dressing
In this technique, two surface dressings are laid consecutively, the first consisting of 6.3/10 or 8/14 mm chippings and the second of 2.8/6.3 mm chippings. It is used on main roads as an alternative to the racked-in system to provide a mechanically strong dressing with a texture that is marginally less than a racked-in dressing, and therefore quieter. The method is also suitable for minor roads that have become very dry and lean in binder.

21.1.3.5 Inverted double surface dressing
In this system, two surface dressings are applied consecutively. The first uses 2.8/6.3 mm chippings, followed by a second application of 6.3/10 or 8/14 mm chippings. This technique is appropriate for use on minor concrete roads where chipping embedment does not occur and surface texture is not an important issue, or on sections of road that have been widely patched.

21.1.3.6 Sandwich surface dressing
Sandwich surface dressing is used in situations where the road surface condition is very rich in binder. The first layer of 8/14 or 6.3/10 mm chippings is spread onto the existing road surface before any binder is applied. The binder is then sprayed over these chippings, followed by an application of 2.8/6.3 mm chippings.

Sandwich dressings can be considered as double dressings in which the first binder film has already been applied. The degree of binder richness at the surface has to be sufficient to hold the first layer of chippings in place until the
remainder of the operation has been completed. The sizes of the chippings must be chosen such that they are appropriate for the quantity of excess binder on the surface and the rate of application of the second coat of binder.

### 21.1.4 Surface dressing equipment

Improvements in the design of surface dressings and advances in binder technology and aggregate quality have been major contributory factors in improving the performance of surface dressings. Developments in application machinery in conjunction with these system developments have enabled the whole process to be improved in terms of the speed of operation, compliance with specifications and continued improvements in standards of safety.

#### 21.1.4.1 Sprayers

In the UK, surface dressing sprayers must satisfy the requirements of BS 1707:1989 (BSI, 1989). This sets requirements for evenness of distribution of the spray bar, heat retention, capacity of heaters and other important issues.

In the UK, it has been traditional for spray bars to have swirl jets. In swirl jets, the binder passing through the jets swirls and forms a curtain in the shape of a cone. When using such jets it is advisable to use lower viscosity binders. However, the increased use of polymer modified binders has resulted in some sprayers being equipped with bars having slot jets. In slot jets, the binder travels through the jets in the shape of a flat fan. Slot jets can spray higher viscosity binders. On main roads, the need to undertake work quickly and with the minimum number of longitudinal joints has led to the use of expanding spray bars of widths up to 4.2 m. In the UK, the usual way of checking that the rate of spread across the bar is uniform and within 15% or better of the target rate has been by use of the depot tray test, which is detailed in BS 1707:1989 (BSI, 1989). The rate of spread is normally checked before the start of the surface dressing season.

Figure 21.10 shows a modern surface dressing sprayer. Programmable bitumen distributors are extremely sophisticated spray machines using two bars to achieve differential rates of spread across the width of the machine. Application rates can be changed for areas of patching, wheel tracks, binder rich fatty areas and binder lean areas. The rate of spread is automatically reduced by up to 30% in the wheel tracks, and can be increased between wheel tracks or in shaded areas. Both longitudinal and transverse binder distributions can be preprogrammed. Vehicle speed, binder temperature, spray bar width and application rates are controlled by an on-board computer.
21.1.4.2 Chipping spreaders
Figure 21.11 shows an example of a self-propelled forward driven chipping spreader operating on a site in Estonia. These machines not only allow work to proceed rapidly but also spread the chippings more evenly and accurately at the required rate of spread. The chippings are released much closer to the...
road surface, reducing the likelihood of chippings bouncing either elsewhere on the road surface or off the road surface altogether and exposing the binder.

The latest developments in self-propelled chipping spreaders include

- a four wheel drive for pulling heavy delivery vehicles up steep inclines without juddering
- a mechanism for breaking lumps of lightly coated chippings within the hopper
- methods for spraying additive onto the binder film
- incorporation of a pneumatic tyred roller to ensure that the chippings are pressed into the binder film at the earliest possible moment while it is still hot.

The latest equipment for spraying and chip spreading incorporates two functions within one vehicle. The chippings are spread, followed by the binder. This increases the efficiency of the operation. Figure 21.12 illustrates the latest equipment.

21.1.4.3 Rollers
Rubber covered steel drummed vibratory rollers (see Figure 21.13) are now regarded as the best means of establishing a close mosaic and ensuring
initial bond of the chipping to the binder film without crushing the carefully selected aggregate. Pneumatic tyred rollers are also used, and some rolling is still carried out using steel wheeled rollers. One of the risks of using steel wheeled rollers is that chippings can be crushed as a result of point loading. It should be noted that rolling is largely a preliminary process undertaken before the main stabilisation of the dressing by subsequent trafficking at slow speed.

21.1.4.4 Sweepers
It is necessary to remove surplus chippings before a surface dressing is opened to traffic travelling at uncontrolled speed. This is normally achieved with full width sweepers, which often have a full width sucking capability. Where sections of major roads have been closed to allow surface dressing to take place, sweepers are often used in echelon in order to ensure the rapid removal of surplus chippings. Sweeping that is properly organised and implemented will do much to alleviate the damage caused to surface dressings in their early life through chipping loss.

21.1.4.5 Traffic control and aftercare
Traffic control and aftercare are vital if the dressing is to be properly embedded. Ideally, when traffic is first allowed onto the new dressing, it should be behind a control vehicle that travels at a speed of about
15 km/h. If there are gaps in the traffic, it may be necessary to introduce additional control vehicles. The objectives should be to ensure that vehicles do not travel at more than 15 km/h on the new dressing and to prevent sharp braking and acceleration of these vehicles.

The length of time for which control is necessary will depend largely on the type of binder used and the prevailing weather conditions. It is likely to be longest where emulsions are being used and the weather conditions are humid. The use of wet or dusty chippings or the early onset of rain delays adhesion and necessitates a longer period of traffic control. Thus, ensuring that the chippings are clean, by washing if necessary, and that they comply with grading requirements will minimise the period necessary for traffic control and reduce disruption. Traffic passing slowly over a dressing immediately after completion creates a strong interlocking mosaic of chippings, a result only otherwise obtained by long periods of rolling.

21.1.5 Types of surface dressing failure
The majority of surface dressing failures fall into one of the following five categories

- loss of chippings immediately or soon after laying
- loss of chippings during the first winter
- loss of chippings in later years
- bleeding during the first hot summer
- fatting up in subsequent years.

The type and rate of application of the binder and size of chippings has a large influence on performance. Incorrect application rates are a frequent cause of premature failure. Every care should be taken to ensure that the selected application rates are maintained throughout the surface dressing operation.

Very early loss of chippings may be due to slow breaking in the case of emulsions or poor wetting in the case of cut-back bitumens. The use of cationic rapid setting emulsions with high binder contents (>70% binder) largely overcomes the former problem, and the use of adhesion agents and/or pre-coated chippings should negate the latter difficulty.

Many surface dressing failures only become apparent at the onset of the first prolonged frosts, when the binder is very stiff and brittle. These failures can normally be attributed to a combination of inadequate binder application rate, the use of too large a chipping or inadequate embedment of the chippings in the old road due to the surface being too hard and/or there being insufficient time between applying the dressing and the onset of cold weather. Inadequate binder application and/or the use of excessively large
chippings will exacerbate this problem. Loss of chippings in the long term usually results from a combination of low durability binders and, again, poor embedment of chippings.

Bleeding may occur within a few weeks in dressings laid early in the season, or during the following summer in dressings that were laid late the previous year. It results from the use of binders that have a viscosity that is too low for high ambient temperatures, or from the use of binders that have high temperature susceptibility.

Fatting up is complex in nature and may result from one or more of the following factors

- application of excessive binder
- crushing of chippings
- absorption of dust by the binder.

Each of these factors is considered in more detail below.

21.1.5.1 Application of excessive binder
Gradual embedment of chippings in the road surface causes the relative rise of binder between the chippings to a point where the chippings disappear beneath the surface. This is largely dependent on the intensity of traffic, particularly the proportion of heavy commercial vehicles in the total traffic. The composition of the binder also affects the process. Cut-back bitumens that have a high solvent content can soften the old road surface and accelerate embedment.

21.1.5.2 Crushing of chippings
Most chippings will eventually split or crush under heavy traffic, leading to the loss from the dressing of many small fragments. The binder/chipping ratio therefore tends to increase, adding to the process of fatting up. A dilemma here is that many of the best aggregates for skid resistance often have a poor resistance to crushing.

21.1.5.3 Absorption of dust by the binder
Binders of high durability tend to absorb dust that falls on them. Soft binders can absorb large quantities of dust, thereby increasing the effective volume of the binder. This effect, coupled with chipping embedment, leads to the eventual loss of surface texture.

21.1.6 Surface dressing season
It is clear from the above that the success of a surface dressing operation may well depend on the weather conditions during and following the application of the dressing. Generally speaking, surface dressing is undertaken from
May to September in the UK. This period varies throughout Europe, and there are areas where it is much shorter. In the western isles of Scotland, the period during which ambient temperatures are high is relatively short, with the result that surface dressing using both conventional and polymer modified emulsions can only be carried out from mid-May until the end of June.

At the start of the surface dressing season, environmental factors are critical. The weather conditions during the early life of the dressing have a significant influence on performance. For example, high humidity can significantly delay the break of a surface dressing emulsion, and that, in turn, delays opening the road to traffic. Heavy rainfall immediately after a road has been surface dressed can have a detrimental effect on the bond between the binder and the aggregate, which can result in stripping of the binder from the aggregate.

In the first summer, it is important that some chipping embedment in the existing road surface occurs, as this will aid chipping retention during the first winter, when the binder is at its stiffest and most vulnerable to brittle failure. Therefore, the later in the season the dressing is applied the more likely it is that chipping loss will occur during the first winter, particularly where the traffic volumes are insufficient to promote initial embedment. This explains why, in areas such as the western isles of Scotland, the season finishes at the end of June rather than September.

21.1.7 Safety

Safety in dealing with bitumen is addressed in detail in Chapter 3. The following briefly highlights some of the potential hazards associated with the handling of surface dressing binders. However, this information is no substitute for the health and safety information available from individual suppliers, or the advice contained in the relevant codes of practice (Energy Institute, 2005; RSTA, 2014a).

21.1.7.1 Handling temperatures

Cut-back bitumens are commonly handled at temperatures that are well in excess of their flashpoints. The flashpoint is the temperature at which the vapour given off will burn in the presence of air and an ignition source. Accordingly, it is essential to exclude sources of ignition in the proximity of cut-back handling operations by displaying suitable safety notices.

Every operation should be carried out at a temperature that is as low as possible, to minimise the risks from burns, fumes, flammable atmosphere and fire. Such temperatures must always be less than the maximum values given in Table 21.4, which is part of Table B.1 in Part 11 of the Bitumen Safety Code (Energy Institute, 2005, 2011). Some bitumen emulsions are applied at
ambient temperatures but the majority of high bitumen content cationic emulsions are applied at temperatures up to 85°C.

21.1.7.2 Precautions, personal protective clothing and hygiene for personnel
All operatives should wear protective outer clothing when spraying emulsions or cut-back bitumen binders. This includes eye protection, clean overalls and impervious shoes and gloves to protect against splashes and avoid skin contact. In addition, operatives working near the spray bar should wear oral-nasal fume masks.

Although bitumen emulsions are applied at relatively low temperatures, the protection prescribed above is essential because emulsifiers are complex chemical compounds and prolonged contact may result in allergic reactions or other skin conditions. In addition, most surface dressing emulsions are highly acidic and, therefore, require the use of appropriate personal protective equipment (PPE). Barrier creams, applied to the skin prior to spraying, assist in subsequent cleaning should accidental contact occur but these are no substitute for gloves and other PPE. If any bitumen comes into contact with the skin, operatives should wash thoroughly as soon as is practicable and always before going to the toilet, eating or drinking.

21.2 Slurry surfacing/microsurfacing
Slurry surfacing (which includes slurry sealing) was first developed in the 1950s and was primarily used as a preservative treatment for airfield runways. The introduction of quick setting cationic emulsions and the development of polymer modified emulsions for thick film slurry applications, now termed microsurfacing, has broadened its use in highway maintenance. It was first developed in Germany in the late 1960s. Now, slurry surfacing using conventional bitumen emulsion is usually described as ‘slurry seal’. Figure 21.14 illustrates slurry surfacing.

<table>
<thead>
<tr>
<th>Grade of cut-back binder: s</th>
<th>Temperature: °C</th>
<th>Minimum pumping(^a)</th>
<th>Spraying(^b)</th>
<th>Maximum safe handling(^c)</th>
</tr>
</thead>
<tbody>
<tr>
<td>50</td>
<td>65</td>
<td>150</td>
<td>160</td>
<td></td>
</tr>
<tr>
<td>100</td>
<td>70</td>
<td>160</td>
<td>170</td>
<td></td>
</tr>
<tr>
<td>200</td>
<td>80</td>
<td>170</td>
<td>180</td>
<td></td>
</tr>
</tbody>
</table>

\(^a\) Based on a viscosity of 2 Pa-s.
\(^b\) Based on a viscosity of 0.03 Pa-s.
\(^c\) Based on generally satisfactory experience of the storage and handling of cut-back grades in contact with air. Subject to the avoidance of sources of ignition in the vicinity of tank vents and open air operations.
The purpose of slurry surfacing is to

- seal surface cracks and voids against the ingress of air and water
- arrest disintegration and fretting of an existing surface
- fill minor surface depressions to provide a more regular surface
- give a paved area a more uniform surface appearance
- restore the skid resistance to a polished surface.

In addition, microsurfacing can also be used to fill areas where rutting has occurred.

21.2.1 Applications
21.2.1.1 Footway slurry
Usually, this is mechanically mixed and hand applied using a brush to provide surface texture. This protects the footway and is aesthetically pleasing. Modifiers can be used to give the material greater cohesive strength, and coloured slurries can be used to denote cycle tracks.

21.2.1.2 Thin carriageway slurries
These are usually laid 3 mm thick and are applied by a continuous flow machine. This type of slurry can be laid rapidly with minimal disruption to traffic. The texture depth that is achieved with this material is relatively low, so it is only suitable for sites carrying relatively slow moving traffic.
21.2.1.3 Thick carriageway slurries
These are normally single treatments using 0/6.3 mm aggregates blended with fast breaking polymer modified emulsions. They are more durable than thin slurries and are, therefore, suitable for more heavily trafficked locations.

21.2.1.4 Microsurfacing
Microsurfacing typically uses aggregates up to 6.3/10 mm and provides a new surface course. It can also be placed in two layers. The first layer regulates and re-profiles, including filling ruts, while the second layer provides a dense surface with reasonable texture.

21.2.1.5 Airfield slurries
Slurry surfacing has been used for runways and taxiways for many years. It is an ideal maintenance technique for airfields as the application is relatively rapid and the aggregates used are too small to constitute a significant foreign object damage hazard to jet engines.

Slurry surfacing only provides a thin veneer treatment to an existing paved area. Consequently, it does not add any significant strength to the road structure, nor will it be durable if laid on an inadequate substrate. Slurry seals are usually applied to lightly trafficked situations such as housing estate roads, footways, light airfield runways, car parks, small traffic highways etc. Microsurfacing can be used on heavily trafficked roads because it can provide higher texture depth.

Slurry surfacing can be laid on the surface dressing. It can reduce the noise generated by a surface dressing and inhibit the loss of chippings from a surface dressing. The system of surface dressing followed by slurry surfacing is described as ‘cape seal’.

The use of fibre reinforcement is being developed, and it seems likely that, in the future, use of this technique will help overcome some of the problems of heavily cracked surfaces.

Pigmented slurry surfacings are available as proprietary products in a range of colours. The binder is synthesised, and the pigment and aggregate should be of the same colour. These surfacings are applied in a very thin layer, and are only suitable for pedestrian and lightly trafficked areas.

21.2.2 Materials
The materials used in slurry surfacing are bitumen emulsion, aggregate, mineral filler and water. A chemical additive may be used when the air temperature is much higher or lower than is usual for slurry surfacing application.
21.2.2.1 Aggregate

The aggregate normally used to manufacture asphalts can also be used in slurry surfacing. In China, basalt and diabase are commonly used in micro-surfacing. Other types of aggregate, such as granite, sandstone, limestone etc., can also be used.

The aggregate should meet the same requirements as would be specified for asphalts, but some characteristics warrant special attention.

- **Cleanliness.** The sand equivalent value indicates the aggregate cleanliness and can be obtained by tests described in BS EN 933-8:2012 (BSI, 2012) and ASTM D2419-09 (ASTM, 2009b). Aggregate usually contains some clays, which may require a shorter mix time for the slurry mixture and cause it to be difficult to spread. In addition, the presence of clay can result in long curing times and a material that has poor cohesion. Accordingly, aggregate cleanliness is very important. The higher the sand equivalent value, the cleaner the aggregate.

- **Surface activity.** Aggregates have a surface charge that plays a key role in dictating the nature of the reaction between aggregates and bitumen emulsions. If the surface charge is strong, the reaction is more intense. For freshly crushed aggregates, the amount of surface charge decreases with time, and much more quickly when the aggregates are wet. The type of mineral in the aggregate also has an effect on the surface charge.

The International Slurry Surfacing Association (ISSA) has defined three aggregate grades, type I, type II and type III, in a table in section 4.2.3 of ISSA A105 (ISSA, 2010a). The grading envelopes of these three categories are shown in Table 21.5. Materials having these gradings are widely used in the USA and China.

- **Type I.** The aggregate size is the smallest of the three gradings. It has a higher filler content so it will be capable of a degree of crack

<table>
<thead>
<tr>
<th>Table 21.5 ISSA aggregate gradation envelopes</th>
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<tbody>
<tr>
<td><strong>Sieve size:</strong> mm</td>
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<tr>
<td></td>
</tr>
<tr>
<td>9.5</td>
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<tr>
<td>4.75</td>
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<tr>
<td>2.36</td>
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<tr>
<td>1.18</td>
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<tr>
<td>0.6</td>
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<tr>
<td>0.3</td>
</tr>
<tr>
<td>0.15</td>
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<tr>
<td>0.075</td>
</tr>
</tbody>
</table>
penetration. It can be used to fill surface voids, cracks and correct minor to moderate surface defects. The surface texture with this grading is small because the aggregate is small, so it is normally used on lightly trafficked or low speed roads such as residential streets, rural roads etc. The main function of type I is to prevent pavements weathering.

- **Type II.** This aggregate grading can be used to fill surface voids and correct moderate surface distress, provide sealing and a durable surface. It can provide medium macrotexture and adapt to medium to heavily trafficked roads. It can also be placed on a flexible base or a stabilised base as a sealer prior to placement of the next layer.

- **Type III.** The aggregate size is the largest of the three gradings. As it contains more coarse particles, it can provide higher surface texture, so a surface paved with this material type has higher skid resistance. It can improve the surface durability because larger aggregate sizes increase the thickness of the pavement. This type of microsurfacing is appropriate for high speed or heavily trafficked roads. It can also be used for rut filling and minor surface reprofiling.

Normally, only types II and III are used for microsurfacing, with type I being used for slurry seal.

21.2.2.2 Emulsion

The emulsions used in slurry seal are traditionally slow setting and can be anionic or cationic. Slurry surfacing using quick setting cationic emulsion CQS-1h (ASTM D2397/D2397M-13 (ASTM, 2013)) can be opened to traffic more quickly than is the case with traditional slurry seals. Only polymer modified, quick setting cationic emulsions can be used in microsurfacing. In China it is named BCR and defined in Technical Guidelines for Micro-surfacing and Slurry Seal (RIOH, 2006b). In Europe, the emulsions should comply with the requirements of BS EN 13808:2013 (BSI, 2013). Latex is mostly used as a polymer modifier for microsurfacing emulsions. Particular polymers can promote adhesion between emulsion and aggregate, and also increase the cohesion of the binder in service, with the result that the microsurfacing is not easily detached under heavy traffic.

21.2.2.3 Mineral filler

The mineral filler is usually Portland cement or hydrated lime, which is added to the mixture to control its consistency, setting and curing rate. Portland cement is normally used up to 2% of total aggregate for slurry seal and up to 3% for microsurfacing. The rate of hydrated lime is normally 0.3–0.5% of the weight of total aggregate. Mineral filler should be considered as part of the aggregate.
21.2.2.4 Water
Water used in slurry surfacing should be clean and free of salt, silt etc. The proportion of water influences the mix time and the consistency of the slurry mixture.

21.2.2.5 Additives
Additives are often used to extend the mix time of slurry mixtures when ambient temperatures are high. Sometimes, additives that accelerate curing are used when ambient temperatures are low. Additives are usually surfactants (substances that reduce surface tension).

21.2.3 Mixture design and specifications
The objectives of mixture design are to assess the compatibility of components and the consistency of the slurry mixture, and to determine the amount of each component. Appropriate consistency should result in the slurry surfacing having a uniform appearance and exhibiting suitable roughness.

21.2.3.1 Trial mixtures
Trial mixtures are used to determine
- the appropriate combination of aggregate and emulsion
- the type and content of mineral filler
- the approximate water content
- the type and content of additive, if appropriate.

In the trial, all preliminary materials are mixed, allowing the mix time to be assessed. During this process, the consistency, breaking and curing characteristics of the slurry mixture are checked and adjusted as necessary. The consistency of a slurry mixture with conventional slow setting emulsion can be measured using the applicable standard (e.g. ASTM D3910-11 (ASTM, 2011), BS EN 12274-3 (BSI, 2002b) or, in China, Technical Guidelines for Micro-Surfacing and Slurry Seal (RIOH, 2006b). The minimum mix time is 180 s for slurry seal and 120 s for microsurfacing. These minimum values are necessary to ensure that the slurry mixture can be successfully spread on the road surface.

21.2.3.2 Wet stripping
This test is used to determine the water resistance of the slurry mixture quickly and simply, and to screen the raw material. It is used in the USA and China. The test process may make reference to ISSA TB-114 (Wet Stripping Test for Cured Slurry Seal Mix) (ISSA, 2012).

21.2.3.3 Cohesion of mixture
The set and cure characteristics of slurries can be assessed by measuring the torsion resistance. For microsurfacing the torsion resistance should be up to
2.0 Nm within 1 h. The test method is controlled by standards such as ASTM D6372-05 (ASTM, 2010b), BS EN 12274–4 (BSI, 2003a) or in China, Technical Guidelines for Micro-Surfacing and Slurry Seal (RIOH, 2006b).

21.2.3.4 Wet track abrasion
This test dictates the minimum binder content. The test measures the loss of weight of the slurry mixture specimen after having been soaked for 1 hour and abraded for a specified time by a rubber pad or wheel. The higher the binder content of the slurry mixture, the lower the abrasion loss of the specimen. If the soak time of the specimen is 6 days, the abrasion loss dictates the water resistance of the slurry mixture. The test method is controlled by standards such as ASTM D3910-11 (ASTM, 2011) and D6372-05 (ASTM, 2010b), BS EN 12274–5 (BSI, 2003b) or in China, Technical Guidelines for Micro-Surfacing and Slurry Seal. Figure 21.15 shows an example of a wet track abrasion machine.

When microsurfacing is used on heavily trafficked roads, two other tests are necessary.

- **Loaded wheel test.** This test determines the maximum permissible binder content if bleeding of the surface under heavy traffic is to be avoided.

![Figure 21.15 A wet track abrasion machine](image_url)
The test measures the amount of sand that adheres to the compacted microsurfacing specimen under a specified load. The higher the binder content, the greater is the amount of adhered sand. The test method is controlled by standards such as ASTM D6372-05 (ASTM, 2010b) or in China, Technical Guidelines for Micro-Surfacing and Slurry Seal (RIOH, 2006b). Figure 21.16 shows a loaded wheel test machine.

Aggregate filler–bitumen compatibility. This test assesses the moisture sensitivity of the slurry mixture. Cylindrical specimens are soaked for 6 days after which the amount of water absorption is measured. Thereafter, the specimens are placed in the abrasion machine and the samples are abraded for 3 h. The loss of mass due to abrasion is measured. Finally, the specimens are placed in boiling water for 30 minutes after which they are weighed. The ratio of the mass of boiled specimens after being boiled to the mass before being boiled is an indication of the integrity of the samples. The detailed method may refer to ASTM D6372-05 (ASTM, 2010b) or, in China, Technical Guidelines for Micro-Surfacing and Slurry Seal (RIOH, 2006b). A further standard that may apply is the BSI test method set out in BS EN 12274-7 (BSI, 2006), which is similar to the ASTM D6372-05 and the method used in China.

21.2.3.5 Specifications
The International Slurry Surfacing Association has developed two guides and a series of technical bulletins. The two slurry surfacing guides are

- ISSA A105, Recommended Performance Guidelines for Emulsified Asphalt Slurry Seal Surfaces (ISSA, 2010a)
ASTM has two standard practice documents that address materials, design and construction of slurry seal and microsurfacing. They are

- ASTM D3910-11, Standard practices for design, testing, and construction of slurry seal (ASTM, 2011)

CEN also developed two standards on slurry surfacing

- BS EN 12273, Slurry surfacing. Requirements (BSI, 2008b)


### 21.2.4 Construction

The surface that is to be treated should be thoroughly cleaned before applying the slurry surfacing. Any loose material on the surface can cause the slurry surfacing to peel off. Any existing defects such as potholes, cracks etc. should be treated in advance of applying the slurry surfacing. A tack coat or bond coat may be required. Slurry surfacing cannot be applied at low temperatures or in wet conditions. Normally the air temperature should exceed 10°C.

A slurry/micropaver is a special continuous flow machine for laying slurry surfacing or microsurfacing. These machines contain an aggregate bin, a bitumen emulsion tank, a water tank, an additive tank, a mineral filler feed, a paddle mixer and a spreader box. All materials can be metered and transported into the paddle mixer, which feeds a spreader box towed behind the machine. These machines are capable of applying up to 8000 m²/day. Figure 21.17 depicts a slurry/micropaver.

A ‘scratch course’ may be required for pavements having minor ruts or depressions. It is applied using the standard spreader box with a steel or stiff rubber screed. If the ruts are deep, they should be filled in advance using a rut filling box. Scratch courses and areas where ruts have been filled should be allowed to cure under traffic for at least 24 h before the surface course is applied. The surface course on flat pavements is applied using a standard spreader box. Figure 21.18 illustrates rut filling and paving a surface course.
**Figure 21.17** Slurry/micropaver

**Figure 21.18** Rut filling and paving a surface course
It is very important that the consistency of a slurry mixture should be correct when it is applied. The water content has a significant effect on the consistency of the slurry. If the water content is too low, the slurry will be stiff, making uniform application difficult to achieve. A low water content may result in the slurry mixture breaking in the spreader box. If the slurry mixture contains too much water the surface texture of the finished surface will be reduced.

Light compaction with a pneumatic tyred roller may be required depending on the traffic condition. Rolling is applied as soon as the material has set sufficiently to support the compaction plant, and normally consists of two full passes. Figure 21.19 illustrates rolling a slurry surface.

21.2.5 Defects
Defects that must be avoided during installation include

- poor joints
- drag marks
- washboarding
- flush surface
- ravelling
- bleeding
- delamination.
The cause(s) may be due to shortcomings of operatives, the paving machine, raw materials, mixture design etc.

21.2.5.1 Poor joints
Poor joints can be caused by excessive overlap or areas left uncovered on longitudinal or transverse joints. Operatives should be careful to avoid these issues. Figure 21.20 shows acceptable joints.

21.2.5.2 Drag marks
Drag marks on a slurry surface are of two types. One is caused by oversized particles or broken slurry mixtures that have lodged under the strike off plate and left furrows on the surface as they are pulled along with the paving machine. The other is caused by the strike off plate having some small particles of aggregate or hardened slurry stuck to it. Small particles may stick to the strike off plate when the spreader box is lifted up off the road when paving has stopped. Another possibility is that some slurry has hardened and has stuck to the strike off plate during the application process of a slurry having insufficient mixing time. Figure 21.21 illustrates two types of drag marks. The drag marks caused by oversize particles can be repaired by judicious use of a squeegee.
21.2.5.3 Washboarding

Figure 21.22 illustrates the washboarding effect. It occurs when the slurry mixture is too stiff and the spreader box is set up incorrectly. This results in the spreader box bumping during installation and causing the washboarding effect. Washboarding can be eliminated by use of a secondary strike off plate. It can be avoided by adjusting the rubbers on the primary strike off plate, making the slurry mixture slower setting. In addition, adding weight to the back of the spreader box may also be effective.
21.2.5.4 Flush surface
If the slurry has too much water or emulsion, the total volume of water and emulsion is greater than the voids of the aggregate, and thus the aggregate is submerged by the surplus water. Figure 21.23 shows an example of a flush surface. Due to the emulsion being too hot, high ambient temperature or the use of dirty aggregate, it may be difficult to spread the slurry. In such circumstances operatives may be tempted to add more water, and this may result in a flush surface.

21.2.5.5 Ravelling
Ravelling occurs after opening to traffic. This can be due to the slurry curing more slowly than expected, possibly due to low ambient temperatures or high humidity. Either could result in the slurry having insufficient resistance to shear when exposed to traffic. In addition, aggregate that is not clean may also delay setting of the slurry. Ravelling may also occur as a result of excessive rainfall.

21.2.5.6 Bleeding
Bleeding may occur after reopening to traffic or during the first summer. The cause may be a high binder content, poor adhesion between aggregate and binder, or a softening point of the binder that is too low for heavy traffic.
21.2.5.7 Delamination
This is separation of the slurry and the layer underneath at the interface. Following any such debonding, the slurry may break up as a result of traffic. Figure 21.24 illustrates this defect. Such failures occur due to poor bond between the subsurface material and the slurry. This may arise because the subsurface is polluted by soil, oil etc., which has not been cleaned sufficiently before application of the slurry. Other reasons why this condition may occur are the existing surface is too old or has excessive ravelling, or the slurry itself may be too stiff. Finally, it has been suggested that the slurry and the subsurface may contract at different rates during inordinately cold periods of weather, causing the bond between the two materials to break.

21.3 High friction surfaces
In 1965, a study carried out by the Greater London Council (GLC) suggested that 70% of all road accidents occurred at or within 15 m of conflict locations such as road junctions and pedestrian crossings (Hatherly and Lamb, 1970). This caused Les Hatherly, then the chief engineer of the GLC, to approach Shell, asking whether it would be possible to produce a high friction surfacing suitable for such sites. Shell developed a bitumen extended epoxy resin system and, following various successful road trials, the result was Shellgrip.
In the intervening period, alternative systems have been developed, including polyurethane resin, acrylic resin and thermoplastic rosin ester materials. A resin is generally a manufactured material or a natural secretion from certain plants, whereas rosin is a hard residue from the distillation of turpentine. A number of these systems are discussed below.

- **Bitumen extended epoxy resins.** These are produced by blending two components, one containing a resin and the other a hardener. When combined, they react chemically to form a very strong three-dimensional structure. A purpose designed machine applies the material onto the road followed by the application of calcined bauxite. The curing time is usually between 2 and 4 h. The excess aggregate is swept off the surface and the road can be opened to traffic.

- **Epoxy resin.** Such systems are a comparatively recent development. They have the advantages that they set very rapidly and adhere well to most surfaces, including concrete.

- **Polyurethane resin.** These resins are normally two part or three part binder systems with good adhesion to most surfaces. Some systems require a primer to be applied prior to application of the resin. They are normally applied by hand.

- **Acrylic resins.** These are fast setting two component systems that adhere well to most surfaces. They are transparent, which makes them ideal for pigmentation.

- **Thermoplastic rosin esters.** These are normally blended with calcined bauxite and heated, and hand screeded onto the road surface. The material cures rapidly and may be pigmented.

The UK’s national road construction and maintenance specification recognises the benefits of these materials and permits the use of systems covered by highway authorities product approval scheme (HAPAS) certification (Highways Agency, 2008).

Figure 21.25 shows a photograph of a cold-applied high friction surface.

### 21.4 Grouted macadams

Grouted macadams have evolved from a specialist treatment to a mainstream surfacing option. They are proprietary products that do not require CE marking. In the UK, the Department for Transport is seeking to include grouted macadams in the new highways maintenance efficiency programme (HMEP – an initiative funded by the Department for Transport to support the highways sector to transform highway services) local roads standard maintenance specification (HMEP, 2014) in recognition of their growing popularity. Currently, agrément certification is used to distinguish between different variants and to ensure consistency of manufacture and installation.
Grouted macadams offer an innovative solution to problems with which conventional surfacing materials struggle to cope.

### 21.4.1 Development

Grouted macadams originated in France in the 1950s. The materials were designed from a simple idea: to alter the characteristics of a new running surface to meet a specific need. Their use in Europe is widespread but, although these products have had a presence in the UK since the 1960s, their use has been relatively limited there. However, that is now changing, and grouted macadams are beginning to emerge as a serious alternative to conventional materials.

### 21.4.2 Laying

There are two forms of grouted macadam:

- cementitious grouted macadam
- asphaltic grouted macadam.

These are named according to the grout used within them, which is either cement or bitumen based. The cement grout adds strength and resistance to contaminants, whereas the bitumen grout provides an impervious seal.
The grouted macadam is laid using a two stage process. A design mix open graded bituminous ‘receiving course’ is laid by a conventional paving machine, generally to a depth of 40 mm, and compacted. The specified grout is then applied to the surface of the material and allowed to percolate into the voids to the required depth, providing the surface course with its chosen characteristics.

### 21.4.3 Cementitious grouted macadam
Cementitious grouted macadams are used in areas of high stress or where fuel, salt and leachate contamination is an issue. They are effectively a hybrid between asphalt and concrete. The most well known of these are Hardicrete, which was first awarded an agrément certificate in 1988 (certificate number 88/1969), and Hardipave, which was awarded a HAPAS certificate in 2006 (certificate number 06/H120).

Their use is widespread throughout the public and private sectors

- bus lanes, laybys and stations
- industrial yards
- coastal promenades
- roundabouts
- junctions
- waste and amenity
- ports and container terminals
- airfields.

The laying of cementitious grouted macadam is shown in Figure 21.26.

### 21.4.4 Asphaltic grouted macadam
Asphaltic grouted macadams use a bituminous grout instead of cement in order to seal the surface and improve flexibility. The sealed surface prevents the ingress of water, while the grout lying beneath the surface of the aggregate gives the product an increased bitumen content, which has a significant effect in slowing the rate of oxidation (discussed in section 20.1.1), and therefore extends product life. The earliest asphaltic grouted macadam, Milepave, gained HAPAS accreditation in 2006. Asphaltic grouted macadams have proved very successful when surfacing on top of concrete, by protecting the structure beneath. Preventing water ingress, they are designed to prevent erosion of the substrate and subsequent slab movement or breakage. In addition, the improved flexibility helps to delay crack formation or subsequent deterioration due to cracking.

The laying of asphaltic grouted macadam is shown in Figure 21.27.
Figure 21.26 Laying cementitious grouted macadam: (a) laying the receiving course; (b) application of cementitious grout; (c) cross section through the finished surface. (Courtesy of Miles Macadam Ltd.)

Figure 21.27 Laying asphaltic grouted macadam: (a) laying the receiving course; (b) application of asphalt grout; (c) finished overlay on concrete; (d) finished carriageway. (Courtesy of Miles Macadam Ltd.)
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Alternative asphalts

Some asphalts are different from those commonly employed on highway construction or maintenance. These mixtures do not always comply with particular specifications but nevertheless are useful in helping to solve specific problems. Examples of these alternative asphalts discussed in this chapter are

- foamed asphalts
- warm mix asphalts
- asphalts made using Shell Thiopave® (also referred to as ‘Thiopave asphalt’. THIOPAVE is a registered trade mark owned by Shell Brands International AG)
- recycling of asphalt pavements
- modification of bitumen by the addition of rubber.

In this chapter, these asphalts are described and explained.

22.1 Asphalts produced using foamed bitumen

Asphalts are extensively used for pavement construction, primarily due to their pressure dissipation and waterproofing properties. In order to mix bitumen with road building aggregates, the viscosity of the bitumen has to be reduced significantly. Traditionally, this was achieved by heating the bitumen and mixing it with heated aggregates, which act like a binding agent to produce an asphalt. Other methods of reducing the bitumen viscosity include dissolving the bitumen in solvents, the use of emulsifying agents and the use of foamed bitumen.

Foaming of bitumen is a means of temporarily reducing the binder viscosity and increasing the binder volume of a bitumen. This technique was first
introduced at the end of the 1950s. Professor Csanyi at the Engineering Experiment Station of Iowa State University studied the potential of foamed bitumen in cold asphalt applications for full depth reclamation projects (Csanyi, 1957). At that time, steam was injected under pressure into hot bitumen through specially designed nozzles, yielding bitumen in the form of a foam. The process reduced the viscosity and surface energy in the foamed bitumen, enabling intimate coating when mixed with moist aggregate at ambient temperatures (Csanyi, 1957). This process was subsequently found to be impractical due to the complexity of the equipment and the difficulties in accurately metering the steam. Mobil Oil refined the foaming technology in the 1960s, and developed an expansion chamber in which cold water containing 1–5% by weight of bitumen is injected under pressure into hot bitumen to produce foam (Bowering and Martin, 1976; Maccarrone et al., 1994, 1995).

Foamed bitumen is typically produced by adding small amounts of water (approximately 2–3% by weight of binder) to hot bitumen. This typically involves a process in which water is injected into the hot bitumen in an expansion chamber, resulting in foaming. The liquid bitumen thus expands by a factor of 15–20 times its original volume, causing foam, as shown in Figure 22.1 (Wirtgen Group, 2002).

The basis of the bitumen foaming process is as follows. The moment that a cold water droplet (at ambient temperature) makes contact with the bitumen at 170–180°C, the following chain of events occurs (Jenkins, 2000).

- The bitumen exchanges energy with the surface of the water droplet, heating the droplet to a temperature of 100°C and cooling the bitumen.

![Figure 22.1 Mechanism of the bitumen foaming process](image-url)
The transferred energy of the bitumen exceeds the latent heat of steam, resulting in explosive expansion and the generation of steam. Steam bubbles are forced into the continuous phase of bitumen under pressure, in the expansion chamber.

With emission from the spray nozzle, the encapsulated steam expands until a thin film of slightly cooler bitumen holds the bubble intact through its surface tension.

During expansion, the surface tension of the bitumen film counteracts the ever diminishing steam pressure until a state of equilibrium is reached.

Due to the low thermal conductivity of bitumen and water, the bubble can remain stable for a period of time, usually measurable in seconds.

This process occurs for a multitude of bitumen bubbles that occur, resulting in the material described as ‘foamed bitumen’. The intensity and effectiveness of the foaming process is governed by the details of the operation, and is related to the applicable physical conditions such as the applied air pressure and bitumen temperature. The rheology of the foamed bitumen can vary and is difficult to predict. The physical properties of the foamed bitumen depend on various factors, including binder type, grade and modification, amount of water used, type of foaming technology used and temperature.

The physical properties of the bitumen are temporarily altered when the injected water comes into contact with the hot bitumen (at about 160–170°C) resulting in vapours that are trapped in thousands of tiny bitumen globules (Fu, 2009). The introduction of water can be effected by using a direct water injection system through a foaming nozzle, by adding water-bearing chemical additives such as zeolites, or by using moist aggregate (Chowdhury and Button, 2008; D’Angelo et al., 2008). The foam dissipates in less than a minute, and the bitumen retains its original physical properties. When the colloidal mass (the bitumen) cools at the ambient temperature, the steam in the bubbles condenses, resulting in bubbles that collapse and cause the foam eventually to ‘break or decay’ (Jenkins et al., 1999a). The collapse of foamed bitumen with time is referred to as ‘decay’, also described as ‘breaking’.

**22.1.1 Foam characterisation**
The rapid evaporation of water produces a very large volume of foam that slowly reduces in volume with time. The foamed bitumens used in base layer stabilisation applications were typically characterised using the following three parameters (Jenkins, 2000)

- expansion ratio
- half life (lifetime)
- foam index.
However, there is no test method available to measure these three parameters precisely. The expansion ratio of the foam is defined as the ratio between the maximum volume achieved during the foamed state and the final volume of the binder once the foam has dispersed (Wirtgen Group, 1998). The half life is defined as the time between the moment the foam achieves maximum volume and the time when it disperses to half of the maximum volume. Half life is measured in seconds, and it depends strongly on the binder/mixture temperature (i.e. higher temperature mixtures have a shorter half life) (Jenkins et al., 1999b). An example of the relationship between expansion ratio and water content, and also half life and water content, is shown in Figure 22.2.

Typically, for foamed bitumen the expansion ratio increases with foaming temperature and water content. In addition, the half life decreases with increasing foaming temperature and water content (Kim and Lee, 2006). Finally, a decrease in the half life causes the thinning of the foam film, as well as a reduction in the viscosity. As the viscosity decreases, the surface tension of the bitumen film decreases as well, and the steam pressure within the bubbles exceeds the surface tension of the bitumen and water bubbles, resulting in subsequent collapse of the bubbles (Hande, 2013).

The foam index is a measure of the area under the breaking or decay curve (i.e. the change in expansion ratio with time). It reflects the stored energy in the foam for a particular bitumen when foamed at a known temperature with water at a predetermined application rate (Jenkins, 2000). The decay curve defines the rate at which the foam collapses, and gives an indication of the
time available for mixing. An important factor in foaming is the design of the nozzle and the injection pressure necessary to obtain an effective water droplet spray in contact with the hot bitumen. The foaming characteristics of specific bitumens are further influenced by the following.

- The temperature of the bitumen – for most bitumens, the foaming characteristics improve with higher temperatures.
- The expansion ratio – as more water is added, the expansion ratio increases, while the half-life decreases. The water helps in creating the foam but the foam can collapse quickly due to the rapid escape of steam.
- Addition of compounds – some compounds (e.g. some silicones) can be effective anti-foaming agents. In contrast, other chemical compounds can increase the half life of the foam from seconds to minutes.

22.1.2 Foamed asphalts

‘Foamed asphalt’ refers to a mixture of mineral aggregates and foamed bitumen. Due to the foaming process, the surface area of the bitumen increases, resulting in a considerable reduction in its viscosity. This reduction in binder viscosity, together with the accompanying increase in volume, means that foamed bitumen effectively coats cold or wet aggregates. This contrasts with traditional asphalts, which can only be manufactured with hot and dry aggregates. Foamed asphalts can be produced in situ or in an asphalt production plant. Such plants can be fixed, free standing or mobile.

22.1.3 Mixing of foamed asphalt

In this section, typical examples of the use of foamed bitumen in the paving industry are discussed.

22.1.3.1 Cold mix asphalt

Foamed asphalts produced using cold or moist aggregates are widely used for stabilising granular materials. Foamed asphalt stabilisation can be effected through cold in-place recycling of traditional asphalts or by way of cold central plant recycling (Bonvallet, 2001; Brosseau et al., 1997; Khweir et al., 2001). Foamed asphalt stabilised base is made up of combinations of recycled materials, including reclaimed asphalt pavement (RAP), recycled concrete and/or graded aggregate base with a foamed asphalt binder, to produce a partially stabilised base material. The foamed asphalt content is usually too low to coat all the aggregate particles fully contrary to what happens in conventional asphalts. During the mixing process, the bitumen ‘bubbles’ burst into tiny ‘patches’ that adhere, mainly, to the fine aggregate fraction. The resulting mortar of fine aggregate and bitumen binds the coarse aggregate particles after compaction, leading to increased
cohesion and stiffness of the aggregate assemblage (Schwartz and Khosravifar, 2013). The foaming characteristics, the moisture content and aggregate grading, together with adequate compaction, are key factors in maximising the performance of aggregates stabilised with foamed bitumen.

Mixtures with a low fines content will not mix well with foamed bitumen because the bitumen will not disperse properly. The expansion ratio of the foam for producing such cold mixtures should be high enough to coat as much of the aggregate as possible after the introduction of the foam into the mixer. The type and size of the mixer is important in ensuring the production of a homogeneous asphalt. The half life of the foam determines the distribution of the foam into the aggregate in these cold mixtures (Schwartz and Khosravifar, 2013). If the half life is too short, the bitumen reverts to a high viscosity liquid, with the risk of lumps of bitumen or pieces of mortar/mastic being formed (Kim et al., 2007). A longer half life may produce a mixture that has good workability when using cold or moist aggregates. For such mixtures, the lifetime of the bitumen foam can range from 20 s up to a few minutes. Long half lives are obtained by introducing additives into the mixture. Optimum values of expansion ratio and foam half life are dictated by the type of application. A key determinant is the residence time in the paddle mixer.

### 22.1.3.2 Warm mix asphalt

Warm mix asphalt technology originated in Europe. Warm mix asphalts are produced and placed at lower temperatures than is the case with conventional asphalts (Koenders et al., 2000). The use of foamed bitumen in asphalts manufactured using warm mix technology to construct pavements has increased (Prowell et al., 2007). However, there is no specification or test method to evaluate the quality of foam generated by different foaming techniques. Recently, an automated repeatable test procedure called the ‘asphalt foam collapse test method’ has been developed (Hande, 2013). This test can be used to measure the reduction in the height of foam bitumen over time by way of an image analysis process. The system permits calculation of parameters of the foam binder quality. In addition, two new parameters, bubble size distribution and surface area index, are introduced as quality parameters. It is reported that these quality parameters are promising candidates for evaluating the workability and the effectiveness of coatings, as well as the performance of asphalt pavements. In order to produce foamed asphalts, the bitumen has to be incorporated in the aggregates while still in its foamed state. The binder contents typically used are based on the mixture design, and are determined as the percentage (by weight) required for the mixture to have optimum properties. Some plant modifications are...
It is necessary to incorporate these technologies. In addition, the mixture design process needs to be altered to include the steps of water injection and foaming action. Furthermore, the laboratory foam device can be used to determine the ‘foamability’ of the bitumen (Cazciiu et al., 2008). Using this device, the asphalt mixture design can be carried out in the laboratory. The procedure adopted for the foaming process using laboratory foaming devices is very similar to that adopted in a large scale production plant facility. In the laboratory, foamed asphalts can be produced as shown in Figure 22.3. Foamed bitumen is poured over warm aggregates at a temperature of 100–135°C in the mixing bucket (Figure 22.3(a)), and the warm aggregates are mixed with the foamed bitumen using a bucket mixer (Figure 22.3(b)). The foamed warm mix asphalt produced after the mixing process is shown in Figure 22.3(c).

A major technical concern in using this technology is that, in mixtures produced at reduced temperatures, the moisture may not escape completely and may thus be trapped inside the mixture after compaction. Whether or not this occurs depends on the prevailing environmental conditions. The trapped moisture in the asphalt can cause decreased adhesion between the bitumen and the aggregates, and lower the cohesion of the bitumen itself.
resulting in stripping, ravelling and other forms of pavement distress (Caro et al., 2008; Kiggundu and Roberts, 1988). It is reported that the rate of moisture dissipation in US high performance grade asphalts manufactured with stiffer binders was found to be slower than is the case in US low performance grade asphalts made with soft binders. Also, the size distribution of moisture bubbles in the binders varies with different bitumen grades (Kutay and Ozturk, 2012a). Furthermore, the results have revealed that the water content and air pressure have a significant influence on the expansion ratio, the half life and the flow index. Low water content and low pressure would result in foams with smaller bubbles than those in foams made with high water content and pressure. This would eventually affect the aggregate coating. Traditional mixing procedures that are adequate for the production of hot mixed and/or cold mixed asphalts have to be adapted in order to accommodate the different characteristics of foamed bitumen. Key parameters to consider are as follows.

- In mixing with hot bitumen or hot bitumen emulsions, the actual mixing has to ensure good coverage, resulting in increased mixing time and energy demand.
- Foam properties and mixing times must be set to ensure a homogeneous mixture.
- Good distribution of the bitumen in the mixture is obtained very quickly as a result of high expansion of the bitumen foam.
- Volume expansion must be such that the bitumen is distributed evenly in the asphalt.

22.1.4 Application and use

The use of foamed bitumen in base stabilisation and in situ recycling in base layers has attracted significant attention throughout the world (Schwartz and Khosravifar, 2013). The process is adaptable, allowing the treatment of virgin materials and recycled aggregates without heating and drying, thus saving energy compared with traditional hot mix production. Mixed material can be stockpiled for a period before being used, reducing the amount of waste. Although foamed bitumen is suitable for stabilisation and recycling work, the resultant mixtures still cannot be objectively regarded as being equivalent to traditional asphalts. In general terms, most cold products are currently inferior to traditional hot asphalt products.

If this is to change, three factors will influence the possibility of a move from traditional asphalts to cold mixed mixtures

- cost
- market demand
- performance.
In 2012, the US National Asphalt Pavement Association (NAPA) stated that the plant foaming of bitumen using water injection systems is a technology that enjoys widespread use in the USA for producing warm mix asphalt mixtures, with more than 88% of the market share as compared with other warm mix asphalt technologies used in various road projects (NAPA, 2012). The foaming process has gained popularity among asphalt producers due to the improved workability and compactibility of asphalts produced at lower temperatures. This technology can be used to add a new coating of bituminous binder to secondary aggregates, including RAP, creating a fresh asphalt and reducing demand for the extraction of virgin aggregate. When warm mix asphalt is used, there is a potential for the asphalt producer to save money on fuel costs, and the contractor is able to obtain better compaction (typically 1–2% higher in situ density) (Dale, 2012). Improved density translates into longer pavement life. Many paving contractors have reported that warm mix asphalts require lower compactive effort (fewer rollers/passes) than would be the case with traditional asphalts (Dale, 2012).

### 22.2 Warm mix asphalts

Warm mix asphalt technologies allow asphalt manufacturers to reduce production temperatures. In addition, warm mix asphalts can be compacted effectively at lower temperatures; typically, temperature reductions of 10–40°C have been reported (Bennert et al., 2011; Estakhri et al., 2010; Jenkins et al., 1999b; Romier et al., 2006). Such reductions in temperatures have the obvious benefit of reducing fuel consumption and decreasing the production of greenhouse gas emissions. World Bank estimates suggest that for every 10°C decrease in production temperature, savings of nearly 1 litre of fuel oil and 1 kg of carbon dioxide emissions are realised per tonne of asphalt produced (Krambeck, 2009). In addition, the engineering benefits include better compaction, the ability to haul over longer distances and extension of the paving season due to being able to lay asphalts at lower temperatures (Koenders et al., 2000).

Numerous warm mix asphalt techniques have been developed with the goal of either reducing the effective viscosity of the binder or providing superior workability at lower temperatures than is the case with a traditional asphalt. Asphalt technologies can be classified on the basis of the amount of temperature reduction. Warm asphalt mixtures are separated from half-warm asphalt mixtures according to the mix temperature (D’Angelo et al., 2008). There are many commercially available products that promise temperatures 10–40°C below those used with conventional asphalts, including some suggesting temperatures slightly above 100°C, with other technologies stating temperatures below the boiling point of water (Zaumanis, 2010). They are categorised as
Another way to classify warm mix asphalts is based on the technology that is used to produce the asphalt (NAPA, 2012). Warm mix asphalt technologies are typically classified into three main categories according to the use of:

- organic or wax additives
- chemical additives
- foaming techniques.

Foaming processes and organic or wax additives are used to achieve the temperature reduction by reducing the viscosity of the binder. These processes typically show a decrease in the viscosity of the binder above the melting point of the wax, making it possible to produce asphalts at lower temperatures (Hill, 2011). After crystallisation, these additives have the effect of increasing the stiffness of the binder and thus, in turn, improving the resistance of the asphalt to permanent deformation (Hanz et al., 2011). Chemical additives typically include a variety of chemical packages, which may be a combination of emulsification agents, surfactants, polymers and additives, to improve coating, mixture workability and compaction, as well as adhesion promoters (anti-stripping agents) (Hill, 2011). These chemical additives are typically used either in the form of an emulsion or are added to the bitumen during the asphalt production process and then mixed with hot or warm aggregates (D’Angelo et al., 2008). These materials affect the surface bonding between the asphalt binder and the aggregate, and are very likely to improve the fracture resistance of the mixture, and have the potential to improve the resistance to deformation (Xiao et al., 2012). Foaming processes and additives use water to foam the asphalt binder and reduce its viscosity prior to or during the mixing period. This particular category comprises the largest variety of warm mix asphalt methods available in the market, but there is the potential for such mixtures to show moisture damage and rutting due to the addition of water to the mixture during the foaming process (Hanz et al., 2011; Kvasnak et al., 2010; Xiao et al., 2011).

To date, more than 30 different warm mix asphalt technologies have been marketed and are available in North America. Depending on which process/technology is being used, the additional cost per tonne for warm mix asphalt with additives is now typically in the range 0–4% (Dale, 2012). At present, most of the US state departments of transportation and all the US Federal Lands Division have technical specifications and/or contractual language in place that would allow warm mix asphalt technology for use on federal aid or federal lands projects. Warm mix asphalt technology has gained popularity in most of US state departments of transportation and,
from 2007, these organisations have road projects constructed using warm mix asphalt techniques. Some of the current challenges for implementing warm mix asphalt technologies are as follows.

- The technologies require changes to asphalt production plants for the incorporation of additives or foam into the mixture.
- Changes are needed in the asphalt mixture production operation.
- Greater quality control and adoption of best practices are needed for the mixture producers to achieve all the benefits of using warm mix asphalt technology, including fuel savings and lower emissions.

It is estimated that the use of warm mix asphalt in the paving industry in the USA will produce energy cost savings of US$3.6 billion by 2020 (NAPA, 2012).

Some warm mix asphalt techniques alter the physical and chemical properties of the asphalt that is produced, and this may cause changes in the long term behaviour of the material. Several researchers have evaluated the performance of warm mix asphalts with respect to various modes of pavement distress. One major concern is the increased susceptibility of warm mix asphalts to deformation (Vargas-Nordcbeck and Timm, 2012). It is possible that the asphalt binder in warm mix asphalt may not harden sufficiently at relatively lower production temperatures, and hence may develop higher post-construction densification or distortion under early age traffic. Conceptually, warm mix asphalt technology has the potential to be detrimental at early age and beneficial in the long term in-service performance. Another concern is that warm mix asphalts may have an increased propensity to moisture induced damage (Bonaquist, 2011; WMTWG, 2007; Xiao et al., 2009). In warm mix asphalts, aggregates are heated to relatively low production temperatures, and therefore may not thoroughly dry before they are mixed with the asphalt binder, thereby reducing the amount of bitumen absorbed in the aggregate. However, results from numerous field trials indicate that a variety of warm mix asphalt additives have required production temperatures 15–40°C lower than those used with traditional asphalts, with no detrimental effect on short term performance (EAPA, 2010). An examination of the field performance of a number of warm mix asphalt projects evaluated by the Texas Transportation Institute suggested that warm mix asphalt pavement test sections were found to be equivalent to comparable control sections made using conventional asphalts (Estakhri et al., 2010). X-ray computed tomography (a technology that uses computer processed X-rays to produce images of specific areas of the scanned object) applied to the laid warm mix asphalts has indicated that the in situ density of the mixture or air void distribution with depth in the material may be even more uniform than is the case with hot mix asphalts. The ability to produce asphalts using this technology has been proved through ongoing implementation in various
paving projects across the USA and in Europe. This may eventually become the standard practice for asphalt production (WMTWG, 2007). In future, environmental aspects of projects will become even more important, and the use of warm mix asphalt may prove to be one means of achieving a lower carbon footprint (EAPA, 2010).

22.3 Shell Thiopave®, a sulfur-based asphalt modifier for enhanced asphalt road applications

Shell’s technology using Shell Thiopave enables a proportion of the bitumen to be replaced in conventional asphalt. It is produced at lower temperature, equivalent to other warm mixtures, resulting in a modified asphalt with enhanced structural properties.

22.3.1 Introduction and background

The concept of using sulfur as a bitumen extender/modifier has been around for three-quarters of a century. Bencowitz and Boe (1938) demonstrated that sulfur is able to modify the bitumen properties, and sulfur-extended asphalts were used in the paving industry in the 1970s (Kandhal, 1982; Kennepohl et al., 1975). At that time, numerous projects around the world used hot liquid sulfur added to the bitumen, in an in-line blending process, prior to the introduction of the bitumen to the mixture. Pavement surveys have validated the performance of these sulfur-extended asphalts as, at the very least, comparable, and often improved, relative to conventional asphalts (Beatty et al., 1987). However, while hydrogen sulfide and sulfur dioxide emissions met the regulatory standards of the time, the odour and irritation were an ongoing source of worker concern.

A process to produce sulfur in pellet form was developed in the late 1990s, and this facilitated the handling of elemental sulfur in a solid form. The process was used to develop sulfur-extended asphalt modifier (SEAM), which is the previous Shell technology incorporating plasticisers in the sulfur pellets (Deme et al., 2004). This represented a major step forward because it enabled the sulfur compound to be added directly into the mixer while manufacturing the asphalt. SEAM significantly reduced emissions and odour compared with the addition of sulfur in liquid form. This technology has been developed further into Thiopave (Figure 22.4). Compared with conventional asphalts, the use of Thiopave enables asphalt production at lower temperatures (125–140°C), with a target temperature of 130°C, which reduces off-gassing while benefiting energy consumption at the hot mix plant.

22.3.2 Asphalt modification principle

22.3.2.1 Role of Thiopave

Thiopave comprises a range of sulfur-based and complementary non-sulfur-based additive components. Individual Thiopave components may differ in
type and appearance depending on the proposed usage. Although Thiopave pellets (i.e. pellets of Thiopave) act as an asphalt modifier, there are situations in which an additional workability additive is required. The workability additive improves the ability of the bitumen to coat the aggregates at lower production temperatures, and also lowers the temperature at which compaction can be effected. This also helps to limit the generation of hydrogen sulfide, reduces the energy used in production and serves as an adhesion promoter between the bitumen and the aggregate.

The addition of the pellets into the bitumen modifies the bitumen properties (Strickland et al., 2007, 2008). Bitumen and sulfur combine at a temperature above the melting point of the pellets. Part of the sulfur is chemically combined with the bitumen and acts as an extender. This portion of sulfur is dissolved in the bitumen, modifying the bitumen properties by lowering the viscosity and increasing the ductility. Above a certain concentration, the remaining sulfur predominantly stays as free sulfur, which crystallises when the mixture cools. Depending on the quantity of pellets added, the crystallisation gives different levels of strengthening, with sulfur crystallisation acting as a structuring agent in the asphalt. An example of the structure of the modified asphalt obtained is shown in Figure 22.5.
22.3.2.2 Target applications
Thiopave is normally used in dense asphalts for binder courses and bases. However, in some countries it has been used in dense surface courses. It should not be used in tunnels, underground car parks or any places where there is a confined space, in case of fire. The modified asphalt is not prone to burning, and in this it is no different to conventional asphalt, but some sulfur compounds, mainly sulfur dioxide and hydrogen sulfide, may be emitted if the asphalt is exposed to a naked flame.

22.3.2.3 Design of Thiopave asphalt
The combination of Thiopave and bitumen results in a fraction of the bitumen being replaced by sulfur in the asphalt. Because the specific gravity of sulfur is virtually twice that of bitumen, in order to maintain the same volume of binder as in the conventional asphalt, the quantity of additive needed to replace the bitumen is almost doubled. In order to produce a paving mixture with enhanced structural properties, a bitumen/Thiopave mass ratio of 60w%/40w% (i.e. 60% bitumen/40% Thiopave by weight) is normally recommended, whereas for general applications the proportion should be limited to 70w%/30w% (i.e. 70% bitumen/30% Thiopave by weight) to retain maximum flexibility. However, these ratios are merely indicative, and the ideal proportion depends on the asphalt type, aggregate and bitumen.
quality. Therefore, it is recommended that an asphalt design using Thiopave be performed to verify the properties of a proposed mixture.

**22.3.3 Thiopave asphalt properties**

**22.3.3.1 Workability and compaction**

The blending of Thiopave and bitumen during manufacture of the asphalt results in a binder that has lower viscosity than the bitumen itself. Above its melting point (>120°C), Thiopave becomes liquid, with very low viscosity, and partly combines with the bitumen, coating the aggregates as a conventional binder would do but at a lower temperature.

Asphalt laying and compaction is carried out using the same equipment as would be employed with conventional asphalts, although the Thiopave asphalt is generally more workable. During paving, compaction starts at around 110°C although, as Thiopave includes a compaction additive, this results in compaction being effective at much lower temperatures. This technology has been accepted for use in warm mix asphalts in the USA (Powell and Taylor, 2012).

**22.3.3.2 Water sensitivity**

Asphalts made with Thiopave pellets generally exhibit the same resistance to water ingress as conventional asphalts. However, some specific studies have assessed the effect of water on both loose asphalts and compacted specimens, and found contradictory results. The immersion of loose asphalts in hot water (60°C) indicated no significant difference in binder film detachment for a period of up to 72 h, and indicated no difference in the effect of water saturation for either mixture type (Strickland et al., 2007). However, with a particular limestone aggregate, the retained Marshall stability and the tensile strength ratio were found to have reduced by 10% for asphalts made with modified sulfur pellets compared with conventional asphalts. Typically, with these aggregates, the use of anti-stripping agents such as amine additives has been found to improve the water resistance properties significantly. There is no generic rule for water sensitivity with the new sulfur technology, and this demonstrates that when a new asphalt is designed it is always good engineering practice to examine its resistance to water ingress.

**22.3.3.3 Deformation resistance**

The specific structure formed in the binder and the air voids in the asphalt enhance the stiffness of the asphalt, without significantly impairing flexibility. As the nature of the crystallites does not change in the pavement temperature range, roads constructed using Thiopave asphalts exhibit superior resistance to deformation compared to those constructed using conventional asphalts. This behaviour was demonstrated in the rutting experiments carried out by
Deme et al. (2004) utilising the asphalt pavement analyzer (APA – discussed in section 16.2.1), as shown in Figure 22.6, all made with a standard performance grade PG 58-28 bitumen (performance grade is the US framework classification for bitumen specifications). The conventional asphalts were much more susceptible to deformation than the same mixture when 40w% of the binder was Thiopave. As shown in Figure 22.6, gravel aggregate mixture, which shows the greatest deformation in a traditional asphalt, exhibits much less deformation when the asphalt is manufactured using modified sulfur pellets. In both cases, asphalt mixtures containing Thiopave demonstrate an improved resistance to deformation, and, to some extent, the inclusion of Thiopave has a positive impact when using lower quality aggregate sources.

22.3.3.4 Stiffness modulus
The stiffness modulus of an asphalt is generally increased when the concentration of Thiopave in the binder exceeds 25–30% by mass. This increase is significantly higher at elevated temperatures and long loading times than at low temperature and short loading times. This means that the modified asphalt is less sensitive to temperature and loading time than is the case with conventional asphalts. Figure 22.7 illustrates this with two asphalt master curves built at a reference temperature of 10°C using the principle of loading time or frequency–temperature superimposition. The asphalts compared are

- conventional asphalt made with a 70/100 pen grade bitumen
- the same asphalt formulation modified with Thiopave at 40% mass in the binder, achieving the same binder volume as for the asphalt above.

Master curves are commonly used when considering matters related to asphalt rheology. Figure 22.7 shows a representation of the stiffness of the two asphalts described above as a function of the temperature and loading time applied. Temperature and loading time are translated into a value of reduced frequency. Accordingly, Figure 22.7 enables the performance of

<table>
<thead>
<tr>
<th>Gravel aggregate</th>
<th>Crushed rock aggregate</th>
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<tr>
<td>Conventional asphalt</td>
<td>Asphalt with modified sulfur pellets</td>
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<td>Conventional asphalt</td>
<td>Asphalt with modified sulfur pellets</td>
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Figure 22.6 Rutting behaviour of different asphalts at 58°C
both asphalts to be compared against a key structural parameter (i.e. deformation). The left hand side of the figure represents high temperature and low speed of traffic (i.e. conditions in which the amount of deformation increases). The right hand side represents low temperature and high traffic speed. The higher the curve the less will be the amount of deformation. What Figure 22.7 demonstrates is that the mixture containing Thiopave is less susceptible to deformation, particularly when conditions are likely to lead to significant deformation.

22.3.3.5 Resistance to fatigue cracking
The resistance to fatigue cracking, as is the case for most asphalts, depends on the test conditions and the asphalt composition. Given similar binder volumes in a conventional and Thiopave asphalt, the fatigue resistance of the modified asphalt would be slightly lower in an imposed strain test because it has much higher stiffness, resulting in higher stresses being generated within the asphalt during fatigue testing. A decrease of 10% of the acceptable strain level of the asphalt has been observed in some cases, although some other studies have shown very similar behaviour in a two-point bending test at imposed strain (Cocurullo et al., 2014). Conversely, at imposed stress, the modified mixture always behaves better, with much higher allowable stresses. As an example, the lower fatigue resistance in the constant strain test mode can be overcome by increasing the amount of binder, as can be seen
in Figure 22.8. The figure shows the fatigue lines as the test results of a four-point bending test carried out at 20°C and 10 Hz. The conventional asphalt had 4.6% bitumen PG 70-10 (this is a US asphalt manufactured with a bitumen PG 70-10). The modified asphalt used the same PG 70-10 bitumen but with 6.6% binder, of which 60% was bitumen and 40% was Thiopave.

22.3.3.6 Low temperature properties

The low temperature performance of asphalts made with Thiopave can be assessed using the thermal stress restrained specimen test (TSRST), in accordance with the AASHTO TP 10 test method (AASHTO, 1993), with some modifications to the specimen size. The temperature conditioning in the TSRST does not attempt to simulate the in situ temperature cycling condition, which causes low temperature fatigue cracking. Moreover, it monotonically decreases temperature until the specimen fails in tension (single event thermal cracking). However, some studies have shown that the low temperature behaviour of asphalt concrete pavements can be predicted using the TSRST (Bouldin et al., 2000; Hannale et al., 1994).

Similar studies were carried out at Shell laboratories in 2008 indicating that the presence of Thiopave in the binder does not detrimentally affect the low temperature performance of the mixture compared with hot mix asphalt, as shown in Figure 22.9. The development of the induced stress in the specimens, with the temperature decrease, was very similar for both types of asphalt, although the modified asphalt exhibited a slightly higher stress
and a lower fracture temperature. The conventional asphalt was made with 70/100 pen grade bitumen, and the sulfur modified mixture was made with the same bitumen and 40% Thiopave pellets by mass. Both asphalts had the same total volume of binder.

Figure 22.10 shows the fracture temperature and the induced stress for both a traditional asphalt and Thiopave asphalt at 30% and 40% Thiopave pellets
by mass in the binder (data represent the mean value of three tests per asphalt formulation). It can be seen that the sulfur modified specimens behave the same as, or even slightly better than, traditional asphalt in terms of cold fracture temperature and induced stress. This suggests that the bitumen dictates the cold temperature behaviour (Strickland et al., 2008).

22.3.4 Road applications
22.3.4.1 Production of Thiopave asphalt
Thiopave is suitable for use in most batch and drum mix plants, with the exception of parallel flow drum plants, for which the drum temperature profile needs some investigation to ascertain their suitability. It is supplied as a solid pellet and added at ambient temperature towards the end of the asphalt manufacturing process. The high shear rate environment during mixing disperses Thiopave throughout the asphalt without the need to extend the mixing time. The target mixing temperature for asphalt containing Thiopave is 130°C, although the recommended range is 125–140°C. This represents a reduction of 20–40°C compared to asphalts made with unmodified bitumen.

Depending on the configuration of the asphalt plant, some modifications of the plant may be required to manage hazards associated with handling sulfur. These modifications are identified as a part of the plant assessment, which must be carried out for each individual plant.

22.3.4.2 Laying and compaction of Thiopave asphalts
Laying and compacting Thiopave asphalts is, in general, no different to that for a conventional asphalt. However, to avoid unwanted emissions during paving, the screed heater on the paver needs to be set to a maximum temperature of 130°C. Fuming typically occurs during rest breaks, when material under the screed may be heated for a prolonged period of time, but it may also occur due to localised heating if the screed heater is set at too high a temperature.

22.3.5 Road performance of asphalts containing Thiopave
More than 100 projects have been carried out worldwide using Thiopave. In most cases, the product brings benefits to the asphalt in terms of improved resistance to deformation and superior stiffness.

In China, in 2006, a 12 km long section of highway was built in Yunan Province (Getun Highway). The Shell technology was chosen to meet the strict specifications forming part of the Marshall mixture design, including a specified minimum level of resistance to deformation. Resistance to deformation was given particular attention because it is one of the main issues on this highway. The base was manufactured using 60% bitumen and
40% Thiopave pellets, while a 65%/35% ratio was used for the surface course. To date, the traffic has been assessed as 4500 vehicles/day, including around 400 heavy trucks. The value of a standard axle load used for pavement design in China is 100 kN. Regular inspections confirm that the section is performing well (Figure 22.11).

In Qatar, air temperatures around 50°C are not uncommon, which means that pavement rutting can be a significant problem on roads that carry high volumes of heavy commercial vehicles, a situation that is typical of construction related traffic in the region. The use of Thiopave was chosen as one of six technologies to meet these challenging conditions, and Thiopave was used in a 270 mm thick base (Figure 22.12). The road provides access to a sand processing plant carrying around 2000 heavy trucks per day. After three years of service, the section of the asphalt made with Thiopave is performing well, and no major distress has been observed. The road is still being monitored.

In India, in 2010, Thiopave was used on National Highway 3 near Nashik, Maharashtra (Figure 22.13). It was included in the dense bitumen macadam (DBM) as type I and II at 60 mm for each layer thickness (DBM type I and type II are standard asphalts in India). Three sections were paved

- Thiopave in both DBM layers
- Thiopave in DBM type I and conventional in DBM type II
- conventional asphalt in both layers.

Regular inspections of the road site are undertaken, including visual observations, coring of the layers and deflection measurements using a Benkelman beam. The fourth evaluation was carried out in March 2013 and confirmed the previous evaluations.

- The three sections are in good condition.
The indirect tensile stiffness modulus tests carried out on cores taken from the road showed that the Thiopave asphalts have higher stiffness at all temperatures (20, 30 and 40°C) than the cores of conventional asphalt.

Figure 22.12 Laying and compacting Thiopave asphalt in Qatar in September 2010

Figure 22.13 Laying Thiopave asphalt in India in 2010
The deflection on the full section of the Thiopave asphalt was reduced by 26% compared with the deflection found on the section constructed using conventional asphalt.

The deflection measured on the hybrid section was 18% less than that on the section constructed using conventional asphalts.

Overall, the Central Road Research Institute of India concluded that the results were encouraging, and that a further structure and functional evaluation would be sufficient to confirm the advantages of Thiopave asphalts over conventional asphalts.

In the USA, in 2009, Thiopave asphalt performance was assessed against conventional asphalt in a comparative full scale experimental test section at the National Center for Asphalt Technology (NCAT) pavement test track (Figure 22.14). The two sections were designed as follows:

- Surface course 32 mm thick
- Binder course 70 mm thick
- Base 76 mm thick over a 152 mm thick aggregate base.

The section of the Thiopave asphalt consisted of a binder course manufactured with 40% Thiopave by mass in the binder, a base manufactured with 30% Thiopave by mass in the binder, and a surface course manufactured with unmodified bitumen. The test sections were regularly monitored in terms of traffic, ride quality, rutting and fatigue cracking. At the conclusion of monitoring in 2012, 10.14 million equivalent single axle loads (ESALs) had been carried by both sections. It was observed that there was no cracking in any
of the sections, and that the ride quality was excellent. The deformation performance was below the rutting failure threshold enforced at the test track. The two sections were left in place for further trafficking. In 2013, after an additional 5 million ESALs, the ride quality was still good for both sections. A small amount of cracking had begun to appear, although more cracking was found on the control section.

22.4 Recycling of asphalt pavements

Recycling in general has a long history and has always been an alternative source of materials when resources are scarce. There is proof from archaeological studies that, around 400 BC, humans had started to reuse broken tools and pottery (Wikipedia, 2014). The main reasons for the commercialisation of recycled asphalt are (Nordic Road Forum, 2012) to

- reduce the consumption of virgin materials
- cut energy use
- minimise air and water pollution.

Recycling has been defined as the ‘use of waste materials for manufacturing new products’. RAP is by no means a waste material. On the contrary, it can be a high value component (Beer et al., 2007). However, it is important to distinguish between recycling of asphalts containing bitumen and recycling of asphalts containing tar. The fumes of the latter are carcinogenic; hence recycling of asphalt containing tar can only be undertaken on cold material (e.g. in combination with an emulsion). The tar also has to be coated to avoid leaching and water pollution. This section applies only to recycling of asphalts containing bitumen, not tar.

For asphalts containing bitumen, the constraints as described above do not apply. The only limitation is the proportion of RAP that can be included in an asphalt. The dose depends on

- the layer in which RAP is to be included
- the type of road (motorway, country road etc.) in which RAP is to be included
- the moisture content of the RAP
- RAP quality
- RAP homogeneity
- RAP management
- asphalt plant capabilities
- local legislation/specification
- the contract with the road owner.

In general, the value of RAP increases with a higher proportion of bitumen. Accordingly, RAP originating in a surface course is likely to have a higher value than RAP originating from a base. Another relevant factor in
determining the value is whether it was used in a low trafficked or heavily trafficked road, with the latter probably having a higher value because it is likely that it contained a higher proportion of bitumen. As the constitution of RAP is variable, clearly the process of adding RAP to newly manufactured asphalts has, also, to be variable. Using RAP in the surface course of a heavily trafficked carriageway is technically demanding and, of course, comes at a cost (Riebesehl, 2007).

A precondition for recycling asphalt is that the performance of asphalt containing RAP is at least as good as the performance of virgin asphalt (EAPA, 2008). Figure 22.15 shows the factors that apply to the use of RAP in different layers of a pavement, with the greener areas of the arrow being the most financially beneficial (Federal Highways Administration, 2011; Riebesehl and Beer, 2006).

As the proportion of RAP in a mixture increases so too does the amount of labour required. This is because the higher the amount of RAP, the more effort is required in respect of quality control, screening, fractionating etc. Thus, it is relatively easy to include 20% RAP in a base but much more effort is required to include 40% RAP successfully in a surface course.

### 22.4.1 Production and storage of reclaimed asphalt

Reclaimed asphalt is obtained (Deutscher Asphaltverband, 2011)

- by milling asphalt
- by breaking and lifting up asphalt in slabs (demolition asphalt)
from waste at asphalt production plants
as returned unwanted loads from the laying site.

Demolition asphalt needs to be crushed and further processed before it can be used as reclaimed asphalt.

Milled asphalt results from milling out (originally described as ‘planing’) existing asphalt pavements, collecting the material as it is milled, and transporting it to an asphalt plant. At present, cold milling machines can process the full depth of a pavement in a single pass. However, in some cases, it may be worthwhile milling different layers separately. Very often, the upper layers contain aggregates or binders with special characteristics, such as high polished stone value aggregate or polymers in the binder.

RAP can be stockpiled (Figure 22.16). However, the quality and consistency of the stockpile has to be known. A disadvantage of stockpiling is that the RAP is not separated into different sizes. As a consequence, the amount of RAP that can be added to a mixture is limited.

In order to maximise the amount of RAP in an asphalt, the RAP should be separated into different sizes. Each size must be stored separately in a facility that prevents water getting into the RAP and thus limiting its use (Figure 22.17).

22.4.2 Quality and suitability
Quality control and consistency of the RAP are key factors in the successful recycling of asphalts. There are a variety of test methods used to determine
the quality and suitability of RAP. Applicable local regulations must be obeyed. An unanswered question about RAP usage is how often an asphalt can be recycled. The answer is certainly not an infinite number of times. Research and practice demonstrate that RAP can be used more than once but then the dosage has to be limited.

The use of recycled asphalt shingles in asphalt is problematic. Usually these shingles contain special binders (oxidised grades, hard grades) that are not suitable for asphalt. Using such binders may result in premature failure of the pavement, as they can embrittle the mixture.

Asphalt containing tar as binder must not be used without special arrangements.

22.4.3 Recycling techniques
There are two main ways of recycling asphalt

- in an asphalt production plant
- in situ (FGSV, 2002).

22.4.3.1 Recycling in an asphalt production plant
This is discussed in detail in several sections in Chapter 14.

22.4.3.2 Recycling in situ
The main processes for recycling in situ are reshaping, repaving and remixing. A common feature of these techniques is that the existing pavement is heated in situ to make it flexible. In some processes, fresh material is mixed with the existing asphalt. Heating has to be carried out very carefully in order to avoid thermal ageing of the binder.

Reshape
In the reshape process (Figure 22.18), the top layer is carefully heated up, using infrared heaters, and scarified. The warm asphalt is then mixed, laid
with a screed and compacted. The aggregate in the existing layer must not be crushed. The asphalt itself remains unchanged, as no additional components are introduced into the existing mixture. Reshaping can be used to rehabilitate evenness of the transverse and longitudinal profile. The process is only feasible on layers that have a good foundation, as stability cannot be improved by reshaping.

Repave
In this process, the existing layer is reshaped, as explained above, and a new top layer is laid on the hot reshaped surface. The new material for the surface course (mixed at an asphalt production plant) is transported by a conveyor belt to a second screed, which lays the asphalt on top of the reshaped surface. Both layers are compacted at the same time. As both layers are hot, a good bond is achieved.

Remix
In the remix process (Figure 22.19), the existing top layer is carefully heated, and lifted by a special purpose machine, where fresh asphalt is added. The machine produces a homogeneous mixture of the existing and fresh asphalt. The resulting mixture is laid and compacted using the special purpose machine.

An enhanced version of the remix process is available in which an additional thin layer is placed on top of the asphalt that has been remixed as described above. This can be achieved using the same machine (remix plus) or a paver (remix compact) (Figure 22.20).
Industry guidance

The use of RAP is only practical when the resulting material performs at least as well as conventional asphalt. This has to be demonstrated in the laboratory and in the field.

Recycling of asphalt pavements requires a RAP management system. This includes, among other things:

- a facility for the dry storage of RAP
- the capability of segregating RAP into its constituent aggregate sizes
- a system for close monitoring of RAP quality
- the ability to monitor and evaluate the properties of the bitumen in the RAP (binder from recycled asphalt shingles contains hard or blown bitumen, and adding these components to an asphalt can cause premature failure of the pavement).

The work necessary to ensure the successful use of RAP is a function of the role of the affected layer. In the case of a surface course or a binder course on a heavily trafficked arterial route, the effort required would be quite significant.

Depending on the target proportion of RAP to be included in the asphalt, modifications to the asphalt production plant may be required.

The use of RAP may be economically and environmentally attractive but it comes at a cost, as the quality of the resultant final asphalt must be fit for the intended purpose. This has to be taken into account not only by asphalt producers but also by construction companies and road authorities.

### 22.5 Modification of bitumen by the addition of rubber

Polybutadiene, polyisoprene, natural rubber, butyl rubber, chloroprene and random styrene–butadiene–rubber, among other rubbers, have all been used to modify bitumen. The effect is mainly to increase the viscosity of the bitumen. Rubbers are also used in a vulcanised (cross-linked) state, (e.g. particles of crumb rubber reclaimed from old scrap tyres, to modify and improve the performance of conventional bitumen, while providing an outlet for the disposal of waste tyres) (Gillen, 2007; Oliver, 1999; WRAP, 2008).
In the USA, some states, including Arizona, California, Florida, Louisiana and Texas, have been predominantly using rubberised asphalt products for several decades. It was reported that, together, these states recycled over 35.6 million tyres in asphalt paving applications from 1995 to 2001 (Rubber Pavements Association, 2002). Approximately 1500 tyres are used to pave one lane mile of highway, an initiative that saves precious landfill capacity. The principal source of raw material for rubber modified binders is tyre rubber, which is a composite of natural rubber, synthetic rubber and carbon black (Chesner et al., 2005). Materials like natural rubber provide the elastic properties, while the synthetic rubber improves the thermal stability properties of the compound and the addition of carbon black improves the binder’s durability.

Crumb rubber (CR), which may also be described as ‘ground rubber’ if it complies with a particular grading, is typically used to produce CR modified binders (CRMB). CR is produced by shredding and grinding reclaimed old scrap tyres from trucks, passenger cars or both, into very small particles (Kandhal, 2006). The tyre source affects the type of rubber present: truck tyres contain a greater proportion of natural latex than do passenger car tyres (Bandini, 2011). It should be noted that natural latex has been found to be more reactive with bitumen than ground tyre rubber. Some state agencies also specify the maximum ratio of car tyre rubber to truck tyre rubber used for binder modification. This is to ensure that an adequate amount of natural rubber is incorporated in the blended material. In the grinding process, most of the steel wires and reinforcing fibres or fluff from the waste recycled tyres are removed. The finer the CR is ground, the more it will react with bitumen; however, additional grinding adds to the cost of the CR material.

In the tyre recycling industry, the fine grinding technique used for scrap tyres is based on either the ambient method or the cryogenic method. The ambient method consists of grinding the rubber at or slightly above ambient temperature using a granulator or a cracker mill (Caltrans, 2014). Finer CR particles (i.e. <0.4 mm) can be produced using a micro-milling process. The CR particles produced using this grinding method are predominantly used for producing CRMB in the paving industry.

Cryogenic tyre grinding consists of using liquid nitrogen to freeze the scrap tyre until it becomes brittle, and then using a hammer mill to crack the frozen rubber into smaller particles. The resulting material is composed of clean, flat particles that generally have a smooth surface texture and a low surface area, and are considered to be less reactive with asphalt binders.

In the USA, most of the south eastern states have changed their bitumen specification to allow asphalt binder suppliers and mixture producers to use
either ambient or cryogenic rubber for producing CRMB to meet the performance grading requirement (Willis et al., 2012) (in the USA, bitumens are classified by their PG). Figure 22.21, shows scanning electron microscopy images of CRMB produced by ambient and cryogenic methods using different sizes of rubber particles. The CR is often sieved and separated into categories based on grading to meet the requirements of a particular application or agency. In the US market, three types of CR particle size are typically available:

- type 1: 2 mm coarse
- type 2: 1.1–0.8 mm
- type 3: 0.6 mm.

The size designation indicates the first sieve with an upper range specification of 5–10% of material retained (Bandini, 2011). The use of CR in asphalts and pavement rehabilitation treatments such as fog seal, crack seal and chip seal can be achieved using two different processes: the ‘wet process’ and the ‘dry process’. In addition, under the wet process, two types of production processes are used to produce rubber modified binders, one process requiring agitation and the other involving no agitation. Each of these is described below.

22.5.1 Wet process with agitation
In the wet process, the CR and other additives, such as extender oil, natural rubber and polymers (as required), are mixed thoroughly with the bitumen, which is maintained at temperatures in the range 180–225°C prior to manufacture of the asphalt. Extender oils such as naphthene or aromatic oil are typically used to promote the reaction between the binder and the CR. Extender oil assists with the dispersion by chemically suspending the rubber particles in the binder. In the US literature, this mixture is often referred to as
‘asphalt rubber’ (AR). This technology was developed in the early 1960s by Charles H. McDonald, while working as an engineer with the City of Phoenix, Arizona. The manufacture of AR can be done on site, and hence it is also called ‘field blend asphalt rubber’. AR is currently being used in California, Arizona and Louisiana. California has been a major user of AR since the 1980s, with around 30% of all asphalts produced there incorporating AR.

For this technology, in order to meet the ASTM D6114/D6114M-09 specification (ASTM, 2009), the rubber content used should typically be at least 15% (by weight) of the total blend to provide acceptable properties in the material when laid. Use of higher rubber contents (18% and 22% by weight) is often reported or specified (Hanson et al., 1996). When using such a high proportion of rubber in the asphalt, the rubber particles are typically agitated sufficiently in the hot bitumen until there is a reaction between the bitumen and rubber particles. This ‘reaction’ is not a chemical process but rather a dispersion process that includes the absorption of aromatic oils from the bitumen into the polymer chain of the rubber. The rubber particles swell to three to five times their original size as they absorb oils, which causes the viscosity of the AR modified binder to increase during the first hour or so (Kandhal, 2006). The rate of reaction is affected by the following properties:

- the temperature of the binder (higher temperatures cause a quicker reaction)
- the surface characteristics of the rubber particle (rougher surfaces react quicker)
- the size of the rubber particle (smaller particles swell quicker but to a lesser extent)
- the period for which the rubber and bitumen are kept at the reaction temperature (a longer time gives a greater reaction).

During the blending process the rubber particles swell, and this changes the proportion of the ground rubber in the binder, which can partially break down. It is reported that the particle size reduction increases with mixing time, and decreases with increasing original rubber particle size (ARTS, 2010). After the ‘reaction’ process and associated swelling has stopped, the rotational viscosity of the blend levels off. The targeted viscosity at the job site for the product would be a minimum of 1.5 Pa·s at 190°C after the interaction period is completed. Because it possesses a short storage life, the AR modified binder has to be agitated constantly in order to maintain the rubber particles in suspension and ensure that they are distributed evenly in the storage tanks. Hence, it is preferable that mixing takes place adjacent to the production plant.
For AR modified binders, a minimum targeted rotational viscosity of 1.5 Pa·s at 190°C is necessary in order to increase binder contents significantly above those of conventional asphalts without excessive ‘drain-down’ (that portion of the bitumen that separates and flows downwards through the mixture) (Caltrans, 2014). If the AR modified binder is maintained at high temperature for a prolonged period of time (as little as 6 h), the rubber particles start to degrade, causing the viscosity of the binder to decrease. ASTM D6114/D6114M-09 (ASTM, 2009) specifies that, among other characteristics, the rubber particles used to produce AR modified binder should contain less than 0.75% moisture (by weight), no visible non-ferrous metal particles, no more than 0.01% ferrous metal particles (by weight) and should be free flowing.

22.5.2 Wet process – no agitation
Terminal blending is another type of wet process in which the CR is blended with the hot bitumen at the refinery or at the bitumen storage and distribution terminal (Bandini, 2011). In practice, terminal blended rubber modified binders are typically produced using rubber contents lower than 15% by weight. At present, this technology is gaining popularity with asphalt producers in the USA, as the process does not require agitation with paddles or augers when stored in tanks in order to keep the rubber particles in suspension. Previously, this was one of the major concerns in the industry when handling modified binders with higher rubber contents. Some of the methods or the processing conditions reported and in use to improve the storage stability of the CRMBs without agitation include (Perez-Lepe et al., 2003)

- the addition of substances to create a chemical stabilisation of the blends
- mixing rubberised binder blends in the presence of a low percentage of polymers
- the addition of a compatibiliser (see the definition below) to activate the CR, or with sulfur to improve cross-linking
- blowing oxygen gases through CR modified (CRM) binder blends
- pre-treatment of rubber particles with hydrogen peroxide (H₂O₂)
- lowering the storage temperature of CRM blends
- increasing the surface area of the rubber particles
- using fine CR (reducing particle size)
- high shear blending/mixing rate (see the note below)
- ensuring that the CR is fully digested into the binder, so that the resulting solubility for such binders is >97%.

In general, compatibilisers (a type of polymeric additive) are used in the production of CRM binders. Compatibilisers typically cause the CR to interact completely with the binder, thereby improving the rheological properties
of the binder. The compatibiliser will typically consist of a reactive component such as one or more glycidyl groups. The effect of any of these reactive functional groups is beneficial in improving the solubility properties and also in aiding the dispersion of CR in bitumen.

The bitumen is heated in a tank to an elevated temperature, and CR is introduced into the tank and is digested into the bitumen. During this process, samples will be collected to test for solubility (i.e. to determine how many residual CR particles remain in the bitumen). This test is carried out during the manufacturing process to ensure that the rubber particles are completely digested in the binder. Most manufacturers use a high shear blending process to make sure that the ground tyre rubber is completely digested. The solubility of the finished product is generally above 97%.

Laboratory studies have shown that adding 10% rubber (passing 0.6 mm sieve) is equivalent to changing from a PG 67-22 binder to a PG 76-22 binder (Willis et al., 2012). ('76' is the average 7 day maximum pavement design temperature (°C) and ‘22’ is the minimum pavement design temperature (°C).) It is reported that the longer mixing process causes the CR particles to partially melt and separate, creating small CR ‘chips’ in the binder. These smaller CR ‘chips’ homogeneously mix with the binder to give a polymer-like structure, which improves their engineering properties (e.g. fatigue resistance) (Kutay and Ozturk, 2012b). Furthermore, increasing the rubber content in the binder would also increase the critical low temperature grade of the modified material. These research data suggested and provided support for US state agencies to move to a performance grade specification for CRMBs instead of specifying raw materials. With the development of performance related properties and the establishment of binder compatibility of rubber modified binders manufactured using the terminal blending process, many states have introduced new specifications for CRMBs, and these are often referred to as PG 76-22TR (tyre rubber) or PG 82-22RM (rubber modified).

In order to meet the PG specifications, terminal blended binder suppliers typically use a minimum CR content of 7% and the size of the rubber particle used would be smaller than 0.6 mm. In addition, they may also use 0.5–4% styrene–butadiene–styrene (SBS) polymer, depending on the performance grade required. Terminal blended products are provided by binder suppliers, and generally come with a certificate of compliance stating the material meets performance grade specifications. Terminal blended rubber modified binders are typically used in open graded and gap graded surface courses in Texas, Florida, Louisiana, New York, Arizona, California and Nevada (Asphalt Institute, 2008). The two main distinctions between the types of wet process methods are
differences in the viscosities of the final product
whether or not agitation is required to maintain relatively uniform
distribution of the rubber particles in suspension in storage.

22.5.3 Dry process

CR modifier can also be used as a substitute for a percentage of the fine aggregate in an asphalt concrete grading rather than as part of the binder. This approach, called the ‘dry process’, was first developed in Sweden in the 1960s. In this process, the tyre rubber is ground to a size similar to that of some of the aggregate fractions, and introduced into the mixing zone (paddle mixer), clear of direct contact with any heating flames (Bandini, 2011). In this process, CR is introduced with the aggregate fraction before adding the binder. The CR acts as an ‘elastic mineral aggregate’ and replaces part of the aggregate fraction. Typically, the rubber percentage used in the dry process varies from 3% to 5% (by weight of the aggregate) (Hicks et al., 2013). It should be noted that, while the bitumen in the dry process is considered to be unmodified, the rubber tends to absorb some of the lighter fractions from the bitumen during the mixing process. However, the properties of the resulting mixture would be modified (Amirkhanian, 2001). The designer must allow for this reaction to take place by slightly increasing the binder content, otherwise there will be a high risk of ravelling in service. The dry process can be used in dense, open graded and gap graded asphalts to accommodate the rubber particles in the aggregate grading, but cannot be used for cold mix applications such as chip seals and surface treatments. The mixture design should take into account the lower specific gravity of the CR compared with that of conventional aggregates (Caltrans, 2014).

The dry process can be implemented in batch asphalt plants and drum mix asphalt plants. The granulated or crumb rubber is usually packed and stored in sacks at the plant. Good control of the feeding of rubber and the temperature are critical in the mixing process, as these variables affect the performance of the resulting mixture and also the performance of the pavement into which the material is incorporated. There have been reports in the literature that batch plants require a dry mix cycle to ensure that the heated aggregate is mixed with the CR before applying the binder. For the mixing process, a hopper typically used for introducing RAP into a drum mixer can be used to introduce CR modifier in the same way (Colucci et al., 1994). Due to the limited reaction time with this process, only the surface of the coarse rubber particle reacts with the binder, which creates a binder–rubber interface that bonds the two materials together. The applicable mixture design process includes a quantity of finer rubber particles for partial reaction with the binder, with a further quantity that acts, in effect, as coarse aggregate.
The fine rubber particles produce the swelling that occurs in the mixture, while the coarse CR particles retain their shape and rigidity to function as aggregate. Some of the major concerns and challenges when using this technology include the following.

- The dry process requires more CR modifier material per tonne of asphalt than does the wet process to achieve the desired properties. This has the effect of increasing the cost of the mixture compared with that of material resulting from the wet process (Colucci et al., 1994).
- The additional binder content needed to produce sufficient rubber–binder reaction on the surface of the rubber particles also carries with it a higher cost.
- These mixtures also appear to be more temperature sensitive, as vertical expansion and extensive cracking have been observed in laboratory studies following 6 h of conditioning at 60°C (Rahman, 2004).
- Visual inspection studies have shown that plucking of rubber particles from the asphalt specimens after moisture conditioning is predominant, especially for mixtures having a rubber content of 5% or more.
- Surface ravelling may also occur due to a lack of bonding between the rubber and binder, meaning that the surface aggregate is not retained.

The above are some of the major reasons why wet process technology is favoured over dry process systems.

22.5.4 Applications of CRMB

The USA and India are currently the leading users of CR for the modification of asphalt. China, Canada and some European countries are also now taking a serious look at this technology. Although initially limited to trial roads, the use of CRMB now covers a wide range of applications, including:

- anti-reflective cracking systems in asphalts
- thin overlay systems on concrete or asphalt, including acting as a stress absorbing membrane interlayer
- stress absorbing membranes
- waterproofing treatment of pavements, typically using chip seal technology beneath the top layer or on the base of the pavement
- bridge deck waterproofing
- noise reduction pavement
- anti-icing pavement (Way et al., 2010).

Thus CRMB can be found in and around the surfacing courses (i.e. the binder course and surface course of new pavements). Typical concentrations of rubber particles in bitumen are in the range 7–15%. High rubber particle content (>20%) allows the formation of a physical network that partially prevents the sedimentation of the particles. In all cases, when dispersed into
bitumen, the CR particles highly influence its rheology and increase its viscosity, which is then sometimes detrimental to the workability.

22.5.5 Limitations of CRMB
Asphalt modification with ground tyre rubber provides the obvious environmental benefit of using scrap tyres. The initial costs of this technology are high and this is still an issue, but the results of life cycle cost analysis indicate that the use of CRMB is, in many cases, an economical option (Hicks et al., 2013). Notwithstanding, aspects of CRMB technology pose a number of concerns:

- a lack of availability of rubber tyre recycling processing facilities in the vicinity
- the cost of such facilities
- a shortage of quality CR
- the need to establish a Superpave mixture design procedure (an asphalt design procedure commonly used in the USA)
- inexperience in the asphalt industry of using this technology
- binder performance grading of AR modified binder
- mixtures are highly temperature sensitive
- compaction issues
- weather restrictions
- limited paving window
- higher mixing temperatures (180–195°C) are required, and these cause the generation of odours and fumes
- the fact that storing the CRMB at high temperature might accelerate the process of particle sedimentation to the bottom of the tank or truck, which results in inconsistent supply quality (Presti, 2013).

22.5.6 Technology gaining popularity in the USA
While environmental stewardship is important, some state agencies and/or contractors are investigating using CRMBs instead of polymers such as SBS in binder modification. If CRMBs can perform such that they are perceived to be equivalent to polymer modified binders, state agencies and contractors will have additional options in the event of another polymer shortage such as occurred in 2008 in the asphalt industry in the USA. Many state agencies are considering using or increasing the use of CR in paving applications, and are looking to other state agencies that have strong recycling programmes for guidance on best practice when handling and using recycling materials. Presently, many asphalt producers are looking at increasing the range of rubberised asphalt paving applications. They hope to do so by using different techniques, including the use of additives in the mixtures, which will enable a reduction in paving temperatures (by using warm mix
technologies) (Punith et al., 2012). This could allow mixtures to be placed at night and in cooler climates, or pelletised asphalt rubber (a patented technology) to be added to the heated aggregates and mixed thoroughly to produce polymerised asphalts (Amirkhanian and Kelly, 2012). These pellets are less than 20 mm in size and contain some form of stiffener (e.g. hydrated lime) and other chemicals.

Recent efforts have focused on using bio binders consisting of bio oil that has reacted with CR. Preliminary laboratory results have shown that such binders can produce a binder that is comparable to binders derived from crude petroleum (Peralta et al., 2012).

Shell Bitumen developed a patented technology (Shell Mexphalte RM) to improve the homogeneity and the stability of CRMB during transport and storage at the asphalt plant. It also limited the maximum mixing and laying temperature to 170°C, thus avoiding the odours and other emissions from rubber particles.

At present, CR serves as an asphalt modifier that can ease the asphalt industry’s dependence on the supply of polymers. Studies conducted at NCAT have shown that the use of CR in speciality mixtures such as open graded friction courses and stone mastic asphalts (described in the USA as ‘stone matrix asphalt’) can allow contractors to prevent drain down (that portion of the bitumen that separates and flows downwards through the mixture) without the use of additional fibres such as cellulose fibres. In addition to removing additional fibres from such mixtures, using CRMBs to modify the mixture can also eliminate the need to use polymer modified binders to avoid rutting. Two year field studies carried out at the NCAT test track, showed that the test sections made of CR modified mixes performed in a manner that is similar to SBS modified mixes in every aspect of pavement performance (West et al., 2012). Based on this research, many US states allow the use of CR as an alternative to traditional PG 76-22 for heavily trafficked roads.

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Chapter 23

Certification of bitumens and asphalts

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This chapter examines the certification of bitumen and asphalts (in the context of construction products) in Europe, where this practice is well established, although other similar systems are used elsewhere.

Certification is the term used to describe the procedures used to confirm defined characteristics of a product or process, usually undertaken by an organisation that is independent of the producer. This is referred to as a third party certification scheme, and is the most common type of certification. The third party is referred to as a certification body.

Other types of certification can exist in which a producer self-certifies products under its own quality system, or when the purchaser is involved in assessing the product (a second party scheme).

Certification is usually applied to either a management system or a product produced to a stated specification. Management systems are certified to BS EN ISO 9001:2008 (BSI, 2008a), which can be supplemented by additional requirements, such as those found in the national highway sector schemes. Product certification is a declaration of the properties or performance of the product in the terms defined by the appropriate standard, and a declaration of compliance with a stated standard.

Initially, certification was adopted voluntarily by some producers as a means of improving customer confidence in their products. Gradually, clients in the construction sector began to include quality management and certification requirements in their specifications, and this has continued until it is now a requirement of most specifications. In recent years, product certification has become a regulatory requirement for some products.

Certification is the responsibility of the producer, who selects a certification body with the relevant experience to certify the product or management
system. The producer will need to develop, document and implement a system prior to the initial assessment by the certification body.

The duration of the assessment and subsequent surveillance arrangements will depend on the scope of the quality system and the size of the organisation being assessed. It is usually a staged process involving examination of the system documentation and observing the implementation of the quality system.

Once the certification body has awarded the certification, it carries out routine surveillance, usually on an annual basis with reassessment every 3 years.

23.1 The regulatory framework

In the construction industry, the principal regulatory requirements for certification are found in the Construction Products Regulation 2011 (EC, 2011), which was implemented in the UK on 1 July 2013. This Regulation superseded the Construction Products Directive (EC, 1989) and the accompanying UK legislation.

The Construction Products Regulation builds upon the earlier Construction Products Directive and aims to break down technical barriers to trade in construction products within the European economic area.

The Regulation stipulates that construction works as a whole and in their separate parts must be fit for their intended use, taking into account, in particular, the health and safety of persons involved throughout the life cycle of the works.

It also requires that, subject to normal maintenance, construction works must satisfy defined basic requirements for construction works for an economically reasonable working life.

These basic requirements are found in Annex 1 of the Regulation

- mechanical resistance and stability
- safety in case of fire
- hygiene, health and the environment
- safety and accessibility in use
- protection against noise
- energy economy and heat retention
- sustainable use of natural resources.

These basic requirements apply to the works as a whole, not individual products. The characteristics of the products themselves must include at least one essential characteristic that relates to the basic requirements above. A product may have several essential characteristics relating to more than one of the basic requirements.
The Construction Products Regulation (EC, 2011) provides for four main elements

- a system of harmonised technical specifications
- an agreed system of conformity assessment for each product family
- a framework of notified bodies
- CE marking of products.

23.1.1 Technical specifications
The requirements for technical specifications are discussed in detail later in section 23.2.

23.1.2 Conformity assessment
The methods of conformity assessment are set out in Annex V of the Regulation, and six systems of assessment and verification of the constancy of performance (AVCP) are described with requirements for certification with differing levels of involvement of external bodies.

All the harmonised standards require that a system of factory production control (FPC) is put in place by the producer. An FPC system is similar to a BS EN ISO 9001:2008 system but includes requirements for inspection, sampling and testing during production. For the higher levels of AVCP, the FPC system must be certified against the relevant product standard, and some producers also choose to have their system certified for products with the lower levels of AVCP.

Asphalts, bitumens and surface treatments require AVCP system 2+. In the UK, aggregates with a polished stone value (PSV) of 58 or greater also require system 2+, and those with a PSV less than 58 require AVCP system 4.

The differences between these two systems of AVCP are shown in Table 23.1.

<table>
<thead>
<tr>
<th>Table 23.1</th>
<th>Tasks and responsibilities for AVCP systems 2+ and 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>AVCP system</td>
<td>2+</td>
</tr>
<tr>
<td>Tasks for the producer</td>
<td></td>
</tr>
<tr>
<td>Factory production control</td>
<td>Yes</td>
</tr>
<tr>
<td>Further testing of samples taken at a factory according to a prescribed test plan</td>
<td>Yes</td>
</tr>
<tr>
<td>Declaration of performance (initial type testing)</td>
<td>Yes</td>
</tr>
<tr>
<td>Tasks for third party notified accreditation body</td>
<td></td>
</tr>
<tr>
<td>Certification of factory production control</td>
<td>Yes</td>
</tr>
<tr>
<td>Surveillance of factory production control</td>
<td>Yes</td>
</tr>
</tbody>
</table>
23.1.3 Notified bodies
Notified bodies are the product certification bodies, FPC certification bodies and testing laboratories, which are considered to be competent to carry out the conformity assessment tasks described in the Construction Products Regulation (EC, 2011). These organisations are first approved by their respective member states to carry out certain designated tasks, and then notified to the European Commission (EC) and other member states.

In the UK, the notification of bodies under the Construction Products Regulation is carried out by the Department for Communities and Local Government, which is the UK notifying authority. It delegates the assessment and ongoing surveillance of notified bodies to the United Kingdom Accreditation Service (UKAS).

Notified bodies are required to participate in the group of notified bodies, with their European counterparts, to discuss practical implementation matters in order to achieve a consistent approach to the tasks.

Once a harmonised technical specification is available for its product, a manufacturer required to use a notified body for assessment can approach any such body in the European economic area that has been notified for the particular harmonised technical specification and task, according to the appropriate conformity assessment procedure. It does not have to use a body operating in the same country as the place of manufacture or where the product is to be used.

Details of notified bodies and their designated tasks can be found on the New Approach Notified and Designated Organisations Information System (Nando) website (Nando, 2014).

23.1.4 CE marking
CE marking on a product is the manufacturer’s declaration that the product complies with the basic requirements of all the directives or regulations that apply to it. It also indicates to the appropriate bodies that the product may be legally offered for sale in its country.

The CE mark can only be applied to products within the scope of a harmonised European standard or a European technical assessment. It indicates that a product is consistent with its declaration of performance (DoP) as made by the manufacturer. The requirements for the DoP are given in the relevant product standard and are normally based on those properties measured for FPC.

The mark is shown in Figure 23.1 and is required to be fixed to the product itself. Where this is not practical, it should be on the accompanying documentation. In addition, the documentation supplied with the product should include the manufacturer’s DoP. Typical examples of the CE mark certificate can be found in Annex ZA of a harmonised standard.
For those products that require notified body involvement in the CE marking (e.g. those subject to AVCP 2+), the number of the notified body must also be shown beneath the CE symbol.

### 23.1.5 Implications for specifiers and purchasers

In addition to the requirements for the manufacture and labelling of products, the Construction Products Regulation (EC, 2011) also contains requirements that relate to public bodies, their specifications and procurement practices.

Articles 8.4 and 8.5 of the Construction Products Regulation place obligations on member states to ensure that the use of construction products bearing the CE mark shall not be impeded by rules imposed by public bodies or private bodies acting as a public undertaking.

Those acting as such a body, either in a monopoly position or under a public mandate, should not specify the performance of products other than in accordance with the basic requirements covered by the harmonised section of the harmonised European standards or European technical assessment under which the CE mark is applied.

Article 8.6 requires that methods used by the member states in their requirements for construction works, as well as other national rules in relation to the essential characteristics of construction products, shall be in accordance with harmonised standards.

Public bodies also have other requirements with which they have to comply when specifying and procuring materials, and these are given in both European directives and UK legislation to implement European directives. At the time of writing (May 2014), a revision to the public procurement Directive (EC, 2004) is due to be published, and the relevant UK regulations will be updated accordingly in due course.

### 23.2 Standards

Certification is usually undertaken in relation to a standard. A standard is a document that provides requirements, specifications, guidelines or characteristics that can be used consistently to ensure that materials, products, processes and services are fit for their purpose.
Standards themselves are used by producers on a voluntary basis but there are circumstances in which their use is mandatory, if required by legislation.

While some clients or industries may develop their own standards, the majority of standards used are developed and published by the recognised standards bodies at national, European or international level.

### 23.2.1 National standards

Most countries have their own national standards body. The British Standards Institution (BSI) fulfils this role in the UK, with DIN and AFNOR, for example, operating in Germany and France, respectively (AFNOR, 2014; DIN, 2014). They work with government, industry and consumers to develop standards at national, European and international levels. They represent their country as members of the International Organization for Standardization (ISO) and the European Committee for Standardization (CEN).

British standards are developed through a committee structure, with representation from producers, consumers and government (local and national). Additional expertise can be co-opted as necessary. Other national standards bodies operate in a similar manner.

The committees are responsible for providing the national representatives to participate in European or international standards work, and also provide a national response when required in the development of these European or international standards.

National standards can be used as the basis for certification of products and systems.

Further information about the BSI and other national standards bodies, their processes and the standards they produce can be found on their websites.

### 23.2.2 European standards

CEN is an association that brings together the national standardisation bodies of 33 European countries. It is one of three European standardisation organisations (together with the European Committee for Electrotechnical Standardization (CENELEC) and the European Telecommunications Standards Institute (ETSI)) that have been officially recognised by the European Union and by the European Free Trade Association (EFTA) as being responsible for developing and defining voluntary standards at European level.

CEN operates with a structure of technical committees, each having responsibility for a defined technical area. The committees are subdivided into working groups and task groups.
The development of a European standard can be initiated by a proposal from any interested party, although most standardisation work is proposed through the national standardisation bodies.

If a standard is required to support legislation, the European standardisation organisations will receive a mandate from the EC or the secretariat of EFTA. This is a request to develop and adopt European standards in support of European policies and legislation.

Once CEN accepts a proposal to develop a standard, all member countries must put any national activity within the scope of the project on hold. This means that they do not initiate new projects, nor revise existing standards at national level. This is referred to as ‘standstill’.

The new standard is developed by the technical committees and working groups, and then submitted for public comment through the CEN enquiry process. Once this is completed and comments resolved, a final version is drafted, which is then submitted to the CEN national members for a weighted formal vote.

When approved, the standard is announced in the *Official Journal of the European Union* (OJEC, 2014) and subsequently published. It must be adopted and given the status of a national standard in all member countries within 2 years, and any national standards that would conflict with it must be withdrawn.

When a new standard is published, the national standards body is able to provide a national annex. Such an annex provides the opportunity for specific guidance to be given about the application of the standard in that country.

In the UK, if the BSI committee considers that more extensive guidance is required, then this is made available in the form of a BSI published document (PD). For example, PD 6691:2010 (BSI, 2010b) gives guidance on the use of BS EN 13108 Bituminous mixtures – Material specifications (BSI, 2006b, 2006c).

More information about CEN and European standards can be found on the CEN website (CEN, 2014).

### 23.2.3 Harmonised European standards

A harmonised standard is a European standard produced in response to a mandate to develop a European standard that provides solutions for compliance with a legal provision. The mandate provides guidelines with which these standards must comply to meet the basic requirements or other provisions of relevant EU harmonisation legislation.
Compliance with harmonised standards provides a presumption of conformity with the corresponding requirements of harmonisation legislation. Manufacturers, suppliers or conformity assessment bodies can use harmonised standards to demonstrate that products, services or processes comply with relevant EU legislation.

CE marking can only be applied to products manufactured to a harmonised standard.

Harmonised standards contain an Annex ZA, which is the part of the standard that is used for the purpose of CE marking of construction products, and transforms all or part of a European product standard into a harmonised European product standard.

The Annex ZA identifies those clauses of the standard (or refers to clauses in another standard) that cover the essential characteristics included in the mandate to which it responds, considered in the answer to the mandates prepared by the technical committee and accepted by the commission.

It also gives details of the AVCP system(s) to which the product needs to be submitted before the manufacturer is able to draw up the DoP and to affix the CE marking.

23.2.4 International standards
The ISO develops standards for international use. At the time of writing (May 2014), 161 countries are ISO members.

ISO standards are voluntary standards and do not inhibit any development or revision of national standards. In some instances they have been adopted and implemented as European standards (e.g. BS EN ISO 9001:2008 (BSI, 2008a)), and in other instances as British standards, when there is no conflict with European standards.

More information about ISO and international standards can be found on the ISO website (ISO, 2014).

23.3 Certification for products and systems outside the scope of CEN harmonised standards
There are several options available to enable products, systems and services outside the scope of CEN harmonised standards to gain certification. These include

- European technical assessments
- national highway sector schemes (NHSS)
- highways authorities product approval scheme (HAPAS)
- BSI Kitemark.
23.3.1 European technical assessments

The European technical assessment is defined in the Construction Products Regulation (EC, 2011) as a document providing information on the assessment of the performance of a construction product, in relation to its essential characteristics.

It is considered to be a harmonised technical specification for the purposes of the Regulation, and consequently provides a way for the manufacturer to CE mark a product in the following situations:

- The product is not or not fully covered by any harmonised technical specification such as European assessment documents (EADs) or harmonised European standards.
- No mandate has yet been issued by the EC for an appropriate standard.
- The product is covered by a EAD.

European technical assessments are carried out by technical approval bodies, which are members of the European Organisation for Technical Assessment (EOTA), using EADs. New EADs, where required, are developed by EOTA and technical approval bodies in conjunction with manufacturers. The EC is informed of the development work and comments on the final draft. After approval, publication of the EAD is announced in the OJEU.

The EOTA website contains further information about the procedures and available documents (EOTA, 2014).

23.3.2 National highways sector schemes

NHSS are developed in partnership with all sides of the highways industry, and their purpose is to interpret the international quality management standard (BS EN ISO 9001:2008 (BSI, 2008a)) for specific activities within the UK highways sector. These include:

- NHSS 13: The Supply and Application of Surface Treatments to Road Surfaces (UKAS, 2013)
- NHSS 14: The Quality Management of the Production of Asphalt Mixes (UKAS, 2014b)
- NHSS 15: The Supply of Paving Bitumens (UKAS, 2014c)

Each scheme is managed by technical advisory committees with industry and client representation. Training organisations, certification bodies and UKAS are also represented.

Consensus is reached on key elements of the schemes such as:

- Minimum levels of services or products.
- Minimum standards of workmanship.
testing
training and competency of the workforce
training and competency of auditors used by certification bodies.

Once an organisation has been successfully assessed, its BS EN ISO 9001:2008 registration is endorsed with the details of the relevant schemes. The organisation is then required to arrange for its details to be included on the schedule of registered suppliers for the NHSS, maintained by LANTRA. This is available for viewing on the NHSS schedule of suppliers website (LANTRA, 2014).

The use of registered suppliers, where an NHSS exists, is a mandatory requirement of the Specification for Highway Works (SHW), and is referenced by other specifiers (Highways Agency, 2014).

NHSS are mostly used in the UK but any competent organisation registered to BS EN ISO 9001:2008 can apply for assessment. At the time of writing, only UK certification bodies are able to provide assessments.

Since the introduction of NHSS in the mid-1990s, some or parts of the activities covered have become the subject of harmonised standards or are affected by the requirements of the Construction Products Regulation (EC, 2011). These schemes have been either withdrawn or amended to remove any potential conflicts, and now provide specific guidance on elements such as training and competency.

Details of the current schemes and copies of the scheme requirements can be found on the UKAS website (UKAS, 2014a).

23.3.3 The highway authorities product approval scheme

The highway authorities product approval scheme (HAPAS) was established in 1995 at the request of specifiers and producers to provide a national approval scheme for products that were not within the scope of existing standards or were innovatory.

It was intended to rationalise the ad hoc processes that many clients had instigated for proprietary products, and to provide a form of approval that would be transferrable between clients.

HAPAS is operated by the British Board of Agrément, and consists of assessment processes for a number of different product sectors. The scheme has operated principally in the UK but is open to any supplier that wishes to supply in the UK. It has been used on a very limited basis by specifiers in other countries.

The process for each product sector is described in guidelines produced by a specialist advisory group drawn from all sides of the highways industry.
These guidelines follow a standard approach to assessment, which typically includes

- Stage 1 – assessment of applicant’s data
- Stage 2 – assessment of quality assurance/factory production control
- Stage 3 – laboratory testing
- Stage 4 – system installation trial
- Stage 5 – system performance trial (if required)
- Stage 6 – certification.

Once certification has been achieved by an organisation, it is subject to annual surveillance and periodic reassessment. Changes made to the design of the product may also necessitate a reassessment, depending on the scale of the change.

HAPAS certification has been a requirement of the SHW for products such as thin surface course systems and high friction surfacing, but the introduction of the Construction Products Regulation (EC, 2011) allows alternative forms of certification to comply with the requirements in Series 100 of SHW.

Further details of HAPAS can be found on the British Board of Agrément website (British Board of Agrément, 2014).

### 23.3.4 BSI Kitemark

The BSI Kitemark (BSI, 2014b) is a registered product certification scheme developed many years ago and operated by the BSI in its role as a certification body. Although it originated in the UK, it is now recognised worldwide.

Certification of conformity with standards is provided through Kitemark schemes to many industry sectors, including the construction industry. The schemes involve an initial assessment of conformity to the relevant standard and an assessment of the quality management system operated by the supplier.

A programme of ongoing surveillance is undertaken, including routine testing of the product or service and assessment of production quality controls.

This type of scheme can provide improved customer confidence when products are not within the scope of CE marking.

The Kitemark scheme is fully described on the BSI website (BSI, 2014a).

### 23.4 Requirements for product certification

Product certification to meet the requirements of harmonised standards consists of two key elements

- a DoP
- implementation of FPC and certification if AVCP 2+ applies to the product.
The preparation of the DoP is the responsibility of the producer, and it must be renewed at least every 5 years. The tests on which it is based must also have been undertaken within the preceding 5 years. The tests required for the DoP are detailed in Annex ZA of the product standard, and are usually drawn from those tests included within the FPC regime.

Previously, under the Construction Products Directive, the DoP was referred to as the initial type test (ITT).

The testing schedules and frequencies are also included in the product standards.

For those products to which AVCP 2+ applies, the producer will need to involve a notified body in the initial inspection, certification and annual surveillance of the FPC. A reassessment of the FPC is undertaken every 3 years.

Some more details about the different product types are given below. References are given to the UK version of the published European standard.

### 23.4.1 Bitumen

Bitumen for paving applications is certified to the requirements of one of the following standards:

- BS EN 12591:2009 Bitumen and bituminous binders – Specifications for paving grade bitumens (BSI, 2009)
- BS EN 13808:2013 Bitumen and bituminous binders – Framework for specifying cationic bituminous emulsions (BSI, 2013a)
- BS EN 13924:2006 Bitumen and bituminous binders – Specifications for hard paving grade bitumens (BSI, 2006d)
- BS EN 14023:2010 Bitumen and bituminous binders – Specification framework for polymer modified bitumens (BSI, 2010a)

All these standards contain sections describing FPC, and an Annex ZA describing AVCP requirements and CE marking.

AVCP system 2+ applies to all these products.

### 23.4.2 Aggregate

Aggregate requirements for road surfacing are given in BS EN 13043:2002, Aggregates for bituminous mixtures and surface treatments for roads, airfields and other trafficked areas (BSI, 2002).

FPC and CE marking requirements are included but this standard contains two systems of AVCP, the one that applies being dependent on the properties of the aggregate.
The products are categorised in accordance with their PSV, and AVCP 2+ applies to those aggregates with a PSV of 58 or greater. AVCP 4 applies to the remainder. This is because, in the UK, coarse aggregates for use in skid-resistant surfacings (PSV ≥ 58) have been defined as high specification aggregates with an influence on road safety.

23.4.3 Asphalt
The specifications for asphalts are found in the BS EN 13108 series of standards, which are currently being reviewed and updated. The requirements for FPC and ITT (the previous term for a DoP) for all the mixtures are grouped together as

- BS EN 13108-20:2006 Bituminous mixtures – Material specifications – Part 20: Type testing (BSI, 2006b)

Part 20 of the standard identifies which test methods are used for each of the different types of asphalt, and Part 21 gives the requirements for the frequency of testing. Testing is undertaken on the constituent materials and also on the manufactured asphalt.

AVCP 2+ applies to asphalts, except where they are used in situations subject to reaction to fire regulations. Where this situation applies, the level of AVCP may be higher, and will need to be determined from Annex ZA.2, taking into account the implications of any national legislation.

The scope of BS EN 13108 is the manufacture of asphalts, and the CE marking applied relates to the material as it leaves the production facility, not the installed material.

23.4.4 Surface treatments
Both surface dressing and slurry surfacing are now within the scope of harmonised standards, and can therefore be CE marked. The relevant standards are

- BS EN 12271:2006 Surface dressing – requirements (BSI, 2006a)

Each of these standards contains requirements for FPC, including the required tests and frequencies. The testing requirements include the constituent materials, the equipment used and a method of visual assessment of defects after 1 year in service.

The declaration of performance is treated differently than in most other standards, as these products cannot be sampled from a production plant.
prior to installation and tested in a laboratory in the conventional way. Instead, the producer carries out a type approval installation trial (TAIT).

The trial is undertaken on a representative site at least 200 m long, selected by the producer. The installation is monitored by the producer and all the data required by the FPC are recorded. The design details, including the traffic volumes defined by road category, are also recorded. The performance is then assessed by the producer after 1 year in service using the visual assessment method in either BS EN 12272-2:2003 (BSI, 2003) or BS EN 12274-8:2005 (BSI, 2005) as appropriate and measurement of the macrotexture is carried out.

The results from these tests are recorded on a TAIT report, which provides the performance categories for the CE mark certificate.

AVCP 2+ is applied to both types of surface treatment.

References
British Board of Agrément (2014) HAPAS Certificates. See http://www.bbacerts.co.uk/productapproval/hapas [accessed 16/10/2014].


UKAS (United Kingdom Accreditation Service) (2013) NHSS 13: The Supply and Application of Surface Treatments to Road Surfaces. UKAS, Feltham, UK.


UKAS (2014b) NHSS 14: The Quality Management of the Production of Asphalt Mixes. UKAS, Feltham, UK.

UKAS (2014c) NHSS 15: The Supply of Paving Bitumens. UKAS, Feltham, UK.

UKAS (2014d) NHSS 16: The Laying of Asphalt Mixes. UKAS, Feltham, UK.
Other important uses of bitumens and asphalts

The majority of bitumen and asphalt usage is in road construction. However, bitumens and asphalts are also used in a range of other important applications. This chapter considers some of these applications:

- bitumen in roofing applications
- bituminous adhesives for pavement markers application
- airfield pavements
- conventional and high speed railway applications
- bridges
- recreational areas
- motor racing tracks including Formula 1 tracks
- vehicle testing circuits
- hydraulic applications
- coloured surface courses and surface treatments
- asphalt kerbs.

The UK’s Mineral Products Association (MPA) has published a number of useful application guides, including one for decorative and coloured surfacing...
(MPA, 2009a), asphalt in construction of games and sport areas (MPA, 2009b), miscellaneous uses of asphalt (MPA, 2009c) and airfields (MPA, 2009d). They can all be found on the MPA’s website (MPA, 2014). Additionally, two books published by Shell provide substantial detail on industrial (Morgan and Mulder, 1995) and hydraulic applications (Scönian, 1999).

24.1 Bitumen in roofing applications
The bitumen industry has three primary areas for its products

- paving
- roofing
- industrial applications.

Roofing products consume approximately 10–15% of the market share of asphalts and bitumens. The roofing sector of the bitumen market is divided between residential steep-slope shingles and commercial modified waterproof membranes. This section addresses bitumen roofing applications in both residential and industrial construction.

24.1.1 Roofing asphalt flux manufacture
Roofing flux is a vacuum residue produced during the distillation of crude oil. It is called roofing flux because it is used to produce harder grades of oxidised bitumen (via a bitumen blowing unit) for use in roofing products. Over 2000 different crudes are produced worldwide, and they are all different in terms of both their physical and chemical properties. The most critical aspect of roofing flux is the source of the crude oil; less than 20% of crudes make acceptable material. The main sources are Middle East and North and South American crudes.

24.1.2 Atmospheric distillation
Crude oil is a very complex mixture of hydrocarbons differing in molecular weights and boiling ranges. Manufacturing of residue involves several steps of physical separation and chemical treatment. The first step is fractional atmospheric distillation, where the crude is heated to temperatures in the range 350–380°C. The column bottom stream is called atmospheric or ‘long’ residue. It requires further vacuum flashing before it can be used as roofing flux. The process is illustrated in Figure 2.1.

24.1.3 Vacuum flashing
The atmospheric residue needs further distillation at reduced pressure in a high vacuum unit. Typical process conditions are 10–100 mmHg pressure with a temperature in the range 350–425°C. The vacuum column removes the non-condensable gas that enters the vacuum unit with the feed. The process is illustrated in Figure 2.2.
To prevent cracking of the heavy components, the residence time at temperatures above 350°C has to be as short as possible. The bottom stream obtained by vacuum distillation of the long residue is called the vacuum or ‘short’ residue (also known as vacuum tower bottom (VTB)). This VTB residue can be used as roofing flux if the crude source is suitable.

### 24.1.4 Roofing products

The two general categories of roofing products are steep slope and low slope. Steep slope roofings are coverings installed on slopes exceeding 14°, and mostly occur on residential properties. Low slope roofing includes waterproofing systems installed on slopes that are 14° or less. This type of roofing is usually used for commercial buildings. Steep slope roofing is the dominant product, representing over 75% of the total roofing market. Data on the usage of the available types are shown in Table 24.1 (ARMA et al., 2011). As can be seen from Table 24.1, in the USA most steep slope roofing consists of shingles, an example of which is shown in Figure 24.1. Note that built-up systems are those where several layers of roof felt are laminated together with bitumen.

Asphalt shingles are composed of the following components:

- fibreglass felt
- coating asphalt (to provide weather resistance and adhesion)
- mineral filler
- mineral granules.

### Table 24.1 North American and European bitumen roofing production by roof type (2006)

(© 2011 Asphalt Roofing Manufacturers Association)

<table>
<thead>
<tr>
<th></th>
<th>Steep slope roofing: $\text{m}^2 \times 10^6$</th>
<th>Low slope roofing: $\text{m}^2 \times 10^6$</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>North America</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Shingles</td>
<td>3403</td>
<td>0</td>
</tr>
<tr>
<td>Built-up roofing</td>
<td>0</td>
<td>298</td>
</tr>
<tr>
<td>Bitumen membranes and underlay</td>
<td>1418</td>
<td>0</td>
</tr>
<tr>
<td>Polymer modified bitumen roofing</td>
<td>0</td>
<td>309</td>
</tr>
<tr>
<td>Total</td>
<td>4821</td>
<td>607</td>
</tr>
<tr>
<td>Market share</td>
<td>89%</td>
<td>11%</td>
</tr>
<tr>
<td><strong>Europe</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Shingles</td>
<td>40</td>
<td>0</td>
</tr>
<tr>
<td>Built-up roofing</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>Bitumen membranes and underlay</td>
<td>14</td>
<td>273</td>
</tr>
<tr>
<td>Polymer modified bitumen roofing</td>
<td>0</td>
<td>610</td>
</tr>
<tr>
<td>Total</td>
<td>54</td>
<td>885</td>
</tr>
<tr>
<td>Market share</td>
<td>6%</td>
<td>94%</td>
</tr>
</tbody>
</table>
24.1.5 Roofing coating specification
The American Society for Testing and Materials (ASTM) produces standard specifications for coating bitumens used in the roofing market in ASTM D312:00 (ASTM, 2000) (Table 24.2).

24.1.6 Roofing shingles specification
Standard specifications for asphalt shingles made from glass felt and surfaced with mineral granules are given in ASTM D3462/D3462M-10a (ASTM, 2010d). The document requires that several properties are met by

<table>
<thead>
<tr>
<th>Test</th>
<th>Minimum/maximum</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Type I</td>
</tr>
<tr>
<td>Softening point: °F</td>
<td>135/151</td>
</tr>
<tr>
<td>Softening point: °C</td>
<td>57/66</td>
</tr>
<tr>
<td>Penetration (25°C, 100 g, 5 s): dmm</td>
<td>18/60</td>
</tr>
<tr>
<td>Penetration (0°C, 200 g, 60 s): dmm</td>
<td>3+</td>
</tr>
<tr>
<td>Penetration (46°C, 50 g, 5 s): dmm</td>
<td>90/180</td>
</tr>
<tr>
<td>Ductility (25°C, 5 cm/min): cm/min</td>
<td>10.0+</td>
</tr>
<tr>
<td>Solubility in TCE: %</td>
<td>99.0</td>
</tr>
<tr>
<td>Flash point: °C</td>
<td>288+</td>
</tr>
</tbody>
</table>

TCE, trichloroethylene.
bitumen shingles, as shown in Table 24.3. As can be seen in Table 24.3, two principal properties – penetration and softening point – are required that relate solely to the coating bitumen used in the shingle manufacture. These two properties provide some indication of the stiffness of the bitumen at intermediate and high temperatures.

### Table 24.3 Standard specification for roofing shingles (ASTM D3462/D3462M-10a (ASTM, 2010d))

<table>
<thead>
<tr>
<th>Property</th>
<th>Min.</th>
<th>Max.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Loss of volatile matter: %</td>
<td>–</td>
<td>1.5</td>
</tr>
<tr>
<td>Sliding of granular surfacing: in.</td>
<td>–</td>
<td>1/16</td>
</tr>
<tr>
<td>Tear strength: g</td>
<td>16.7</td>
<td>–</td>
</tr>
<tr>
<td>Fastener pull-through resistance (73°F): lbf</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Average of single-layer specimens</td>
<td>20</td>
<td>–</td>
</tr>
<tr>
<td>Average of multi-layer specimens</td>
<td>30</td>
<td>–</td>
</tr>
<tr>
<td>Fastener pull-through resistances (32°F): lbf</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Average of single-layer specimens</td>
<td>23</td>
<td>–</td>
</tr>
<tr>
<td>Average of multi-layer specimens</td>
<td>40</td>
<td>–</td>
</tr>
<tr>
<td>Wind resistance</td>
<td>Class A</td>
<td>–</td>
</tr>
<tr>
<td>Fire resistance</td>
<td>Class A</td>
<td>–</td>
</tr>
<tr>
<td>Penetration of bitumen (without mineral stabiliser, 77°F): dmm</td>
<td>15</td>
<td>–</td>
</tr>
<tr>
<td>Pliability (73°F, all directions)</td>
<td>4 of 5 shall pass</td>
<td></td>
</tr>
<tr>
<td>Asphalt softening point (without mineral stabiliser, R&amp;B): °F</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Unmodified</td>
<td>190</td>
<td>235</td>
</tr>
<tr>
<td>Polymer modified</td>
<td>190</td>
<td>320</td>
</tr>
<tr>
<td>Weight of displaced granules: g</td>
<td>–</td>
<td>1.0</td>
</tr>
</tbody>
</table>

R&B, ring and ball test.

24.1.7 **Roofing flux specification**

There is no standard specification for roofing flux, either from the ASTM or from the industry itself. Thus, flux specifications vary between producers, manufacturers and suppliers. Roofing industry companies have different requirements for flux quality depending on their specific needs.

The most critical criteria for roofing flux, as related to roofing manufacture, are the properties of the crude oil used in making the roofing flux, as the end product is the oxidised coating material resulting from the air blowing process. The chemistry of the crude oil contributes to the resulting performance. Even if a flux meets the required physical properties, it does not follow that it has the required coating properties. Therefore, in most cases, a new flux has to pass an approval process before it can be used in the manufacture of roofing products.

As there is no well established relationship between the flux and the coating performance, the flux requirement (not a specification) can only come from a pre-approved crude oil resource or from a pre-approved supply. Assay of the crude oil does not indicate whether it will result in a good roofing flux or a
good shingle coating. Roofing manufacturers need good oxidised coating performance rather than good flux properties.

However, there are some basic requirements that can be used for flux: for example, for paving grade bitumens the flash point (Cleveland open cup (COC) method) needs to be a minimum of 550°F (250°C); and, for reasons of safety, some roofing manufacturers require the closed cup flash point (Pensky–Martens closed cup (PMCC) test) to be a minimum of 450°F (232°C). For a roofing flux made from a pre-approved crude oil, the viscosity range can be used to specify the properties. This can be the absolute viscosity at 60°C, or the kinematic viscosity at 135°C. The stain index is also critical, because it represents the volatile oil content and relates to the discolouration performance of the coating. The requirement for the stain index varies between different manufacturers (Tables 24.4 and 24.5).

Table 24.4 Physical properties of four bitumens typically used in North American roofing made from crude source A (ARMA et al., 2011)

<table>
<thead>
<tr>
<th>Test and units</th>
<th>Test methoda</th>
<th>Straight run bitumen</th>
<th>Oxidised bitumen</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Polymer modified bitumen base</td>
<td>Roofing flux</td>
</tr>
<tr>
<td>Softening point: °C</td>
<td>ASTM D36</td>
<td>41</td>
<td>41</td>
</tr>
<tr>
<td>Penetration (25 °C): dmm</td>
<td>ASTM D5</td>
<td>170+</td>
<td>220+</td>
</tr>
<tr>
<td>Viscosity (60 °C): poise</td>
<td>ASTM D4402</td>
<td>140</td>
<td>140</td>
</tr>
<tr>
<td>Loss on RTFO (163 °C): %</td>
<td>ASTM D2872</td>
<td>0.1880</td>
<td>0.0418</td>
</tr>
<tr>
<td>Flash point (COC): °C</td>
<td>ASTM D92</td>
<td>313</td>
<td>313</td>
</tr>
<tr>
<td>Stain index</td>
<td>ASTM D2746</td>
<td>b</td>
<td></td>
</tr>
<tr>
<td>Weathering</td>
<td>ASTM D4798</td>
<td>b</td>
<td></td>
</tr>
</tbody>
</table>


b Varies by manufacturer

Table 24.5 Physical properties of four bitumens typically used in North American roofing made from crude source B (ARMA et al., 2011)

<table>
<thead>
<tr>
<th>Test and units</th>
<th>Test methoda</th>
<th>Straight run bitumen</th>
<th>Oxidised bitumen</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Polymer modified bitumen base</td>
<td>Roofing flux</td>
</tr>
<tr>
<td>Softening point: °C</td>
<td>ASTM D36</td>
<td>42</td>
<td>37</td>
</tr>
<tr>
<td>Penetration (25 °C): dmm</td>
<td>ASTM D5</td>
<td>170+</td>
<td>300+</td>
</tr>
<tr>
<td>Viscosity (60 °C): poise</td>
<td>ASTM D4402</td>
<td>150</td>
<td>100</td>
</tr>
<tr>
<td>Loss on RTFO (163 °C): %</td>
<td>ASTM D2872</td>
<td>0.066</td>
<td>0.165</td>
</tr>
<tr>
<td>Flash point (COC): °C</td>
<td>ASTM D92</td>
<td>320</td>
<td>318</td>
</tr>
<tr>
<td>Stain index</td>
<td>ASTM D2746</td>
<td>b</td>
<td></td>
</tr>
<tr>
<td>Weathering</td>
<td>ASTM D4798</td>
<td>b</td>
<td></td>
</tr>
</tbody>
</table>


b Varies by manufacturer
24.1.8 Product approval

Shingle manufacturers provide guarantees on their products with periods ranging from 15 years’ to a lifetime warranty. The durability of the coating (filled and unfilled) is critical. The accelerated xenon arc weathering test (ASTM, 2011) is the traditional method used to evaluate the long term ageing performance. The coating panel must show less than 10% of failure after a 2000 h cycle of exposure. The time needed to pass the weathering test is usually 3–4 months.

The coating material also needs to meet certain softening point/penetration relationship criteria, without the inclusion of any additives. Once the coating has passed the testing phases, a plant trial is carried out to evaluate the material’s performance in the process. A trial is typically one to three truck loads (approximately 25–75 tonnes). Following a trial, it usually takes around a month to complete the product evaluation.

24.1.9 Air blowing of roofing flux

Oxidation requires a complex series of reaction processes in order to produce the bitumen coating product from a roofing flux. The blowing process can be carried out in batch or continuous mode. The batch production mode is the one most commonly used in the USA. The reaction process introduces oxygen into the system; the temperature is typically in the range 480–500°F (218–260°C). A series of chemical thermal reactions takes place during production. There are mainly two types of reaction:

- dehydrogenation, which introduces oxygen to hydrogen at the molecular level
- polymerisation, which converts most aromatic compounds to a condensed asphaltene structure.

It has been found that all the oxygen taken up by the bitumen can be accounted for by the formation of hydroxyl, carbonyl, acid and ester groups (ARMA et al., 2011).

The air blowing time depends on the properties of the flux, and the final oxidised product requirements. As a general rule, the softer the flux is at the start, the longer it will take to achieve the final product. This is illustrated in Figure 24.2: the higher the softening point to be achieved, the longer the time that is needed.

In terms of its chemical composition, the flux changes during the oxidation process, the proportion of saturates tends to remain the same, some aromatics convert to resins, some resins convert to asphaltenes and the asphaltenes themselves remain unchanged. The net result is that there is an increase in asphaltene content and a dramatic drop in aromatic content (Table 24.6).
After air blowing (Figure 2.4), the end product (typically called ‘bitumen coating’ in Europe and ‘asphalt coating’ in the USA) exhibits decreased penetration, an increased softening point, higher viscosity, improved weathering resistance and good temperature susceptibility.

24.1.10 Asphalt low slope roofing products
Low slope roofs have slopes of 14° or less. Most low slope roofs are used in commercial or industrial buildings. Bitumen is used in two different low slope roof systems: polymer modified bitumen membranes and built-up roofing (BUR) products. In the USA, these two systems account for 35–40% of the low slope market. The total share of low slope bitumen systems is even higher in western Europe, where polymer modified bitumen systems dominate the low slope bitumen market (Figure 24.3).

24.1.11 Polymer modified membrane systems
Polymer modified bitumen membranes are often used in commercial roofing, including low slope roofing applications.

The most common base bitumens are European paving grades 160/220 and 100/150 pen, depending on the market and customer requirements.

| Saturates: % | 15.0 | 14.5 |
| Aromatics: % | 38.5 | 27.2 |
| Resins: % | 35.5 | 27.2 |
| Asphaltenes: % | 11.0 | 31.1 |

Table 24.6 Typical changes in SARA between the flux and the coating
Polymers typically used in this application are atactic polypropylene (APP) and styrene–butadiene–styrene (SBS). Different polymers require different bases. In most instances, a base with the highest saturate content would be suitable with an APP, whereas a base with a higher aromatic content would be appropriate for use with an SBS application.

An APP modified bitumen membrane felt typically contains
- bitumen 52–63%
- APP 15–25%
- filler 20–30%,

while an SBS modified membrane felt typically contains
- bitumen 60–70%
- SBS 5–15%
- filler 20–35%.

Specifications for APP modified bituminous sheet materials are given in ASTM D6223-02 (ASTM, 2009e), and specifications for SBS modified bituminous sheet materials using glass fibre reinforcement can be found in D6163-00 (ASTM, 2008) (Table 24.7).

**24.1.12 Future trends for roofing bitumen shingles and coating evaluation**

Although rheological testing of paving grade bitumens has become much more widespread over the last two decades as a result of the Strategic Highway Research Program (SHRP), the conventional empirical tests (e.g. penetration, ring and ball (R&B) softening point and viscosity) remain the
dominant tests and requirements for oxidised coating and BUR products in
the bitumen roofing industry in North America. The rapid development of the
dynamic shear rheometer (DSR), including the 4 mm parallel plate measure-
ment for low temperature testing and the master curve application for paving
grade bitumen, and the belief that the results can be linked to bitumen per-
formance, has created an interest in employing these techniques for use in
the roofing industry. More detailed information about rheological testing of
bitumen is given in Chapter 7.

Due to proprietary issues in the roofing industry, some individual roofing
companies have undertaken research based on rheological properties using
rotational rheology tests. This is an attempt to link the rheological properties
of the bitumen to the performance of coatings and shingles, and character-
isation of roofing fluxes. However, little has been done to produce a unified
or standard specification for bitumen fluxes or coatings. For the last 10 years,
the Asphalt Institute has, in cooperation with the roofing industry, been very
actively involved in research based on rheological properties (Asphalt Insti-
tute, 2014). The purpose of this research is to explore the possibility of devel-
oping a new standard based on the use of rheological parameters as a
measure of oxidised bitumen coating, which could be linked to shingle per-
formance with a view to replacing the conventional empirical testing. For
example, a useful bitumen relationship has been developed based on the
complex modulus ($G^*$) and penetration (for more details see Chapter 7)

$$\log(G^*; 25^\circ C, f=0.4 \text{ Hz}) = 2.923 - 1.9 \log(\text{penetration})$$

<table>
<thead>
<tr>
<th>Source information for APP roofing felt requirementsa</th>
<th>UEAtcb</th>
<th>Manufacturer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Country</td>
<td></td>
<td>A</td>
</tr>
<tr>
<td>Penetration (25°C, 0.1 mm; ASTM D5)</td>
<td></td>
<td>France</td>
</tr>
<tr>
<td>Penetration (50°C, 0.1 mm; ASTM D5)</td>
<td></td>
<td>&lt;80</td>
</tr>
<tr>
<td>Penetration (60°C, 0.1 mm; ASTM D5)</td>
<td></td>
<td>90–120</td>
</tr>
<tr>
<td>Dynamic viscosity (180°C; ASTM D2170): cP</td>
<td></td>
<td>4000–6000</td>
</tr>
<tr>
<td>Softening point (R&amp;B; ASTM D36): °C</td>
<td>120</td>
<td>140–155</td>
</tr>
<tr>
<td>Flexibility at low temperature (ASTM C711-14): °C</td>
<td></td>
<td>−5</td>
</tr>
<tr>
<td>Penetration (50°C, 0.1 mm; ASTM D5)</td>
<td></td>
<td>70–100</td>
</tr>
<tr>
<td>Dynamic viscosity (180°C; ASTM D2170): cP</td>
<td></td>
<td>3500–5500</td>
</tr>
<tr>
<td>Softening point (R&amp;B; ASTM D36): °C</td>
<td>140–155</td>
<td>150</td>
</tr>
<tr>
<td>Flexibility at low temperature (without damage): °C</td>
<td>0</td>
<td>−20</td>
</tr>
</tbody>
</table>

R&B, ring and ball test.
b UEAtc: European Union for technical approval in construction.
The current consensus is to use the rotational rheology test to characterise the oxidised bitumen coating material at intermediate and higher temperatures:

- DSR at 25°C, 8 mm plate, 1% strain at a frequency of 2.5 rad/s
- DSR temperature sweep tests from 90°C to 110°C in 10°C increments.

Similar studies of predicting ageing behaviour using rheological testing have been carried out on paving grade binders (Farrar et al., 2013; Mike, 2013; Qin et al., 2014), and these suggest that a master curve can be generated on the basis of the temperature sweep test.

The rheological parameter ($G^{*}2.5$ – the complex modulus multiplied by 2.5) was chosen to create a correlation with conventional physical properties such as penetration and softening point (Mike, 2013) (Table 24.8). The data suggest that there is a linear relationship in the penetration range 13–25 with a variability of 13.5%. Figure 24.4 shows the relationship between $G^{*}$ (complex modulus) and the softening point for oxidised coating materials.

The relationship from all data is

$$ \log G^{*}2.5 = -1.6281 \times \log(\text{pen}) + 8.6582 $$

Penetration = 12 dmm

![Figure 24.4](image-url) Relationship between the rheological parameter $G^{*}2.5$ (Pa) and the softening point (°C)
The proposed use of the rheological parameter \(G^*2.5\) criterion to replace the softening point and penetration tests is as follows.

- **Proposed procedure to replace softening point**
  - perform a temperature sweep using a DSR
  - 25 mm parallel plate, 10% strain, 1 rad/s
  - start at 90°C, increase to 110°C in 10°C increments
  - plot \(\eta^*(1 \text{ rad/s})\) versus temperature on a semi-logarithmic graph
  - calculate \(T_c\) when \(\eta^*(1 \text{ rad/s}) = 1200 \text{ Pa.s.}\)

- **Proposed procedure to replace penetration at 25°C**
  - perform single point temperature test using a DSR
  - 8 mm parallel plate, 1% strain, 2.5 rad/s, 25°C
  - determine \(G^*2.5\) and compare with the recommended specification value.

Work is underway to evaluate the proposed criteria further.

### 24.2 Bituminous adhesive for use with pavement markers

Flexible bituminous pavement marker adhesive is a hot-melt thermoplastic bituminous material used for bonding raised pavement markers and recessed pavement markers to the pavement. There are two types of bituminous marker adhesive:

- **standard bituminous marker adhesive**, which consists of an asphalt base with a homogeneously mixed mineral filler
- **flexible bituminous marker adhesive**, which consists of a highly polymer modified asphalt – this type normally contains ground tyre rubber (GTR) as one modifier.

Both adhesives must be suitable for bonding ceramic and plastic markers to cementitious, asphalt concrete and surface dressed road surfaces, and be applicable when road surface and marker temperatures are in the range 4–71°C. Prior to application, adhesives are heated in either an air or an oil-jacketed melter up to a temperature of around 220°C, and must be able to tolerate these elevated temperatures without deteriorating.

#### 24.2.1 Manufacture of bituminous marker adhesive

Standard bituminous marker adhesive (BMA) can be made from a relatively soft bitumen base made harder through light distillation, an oxidation process...
or the addition of a hard component. Alternatively, a hard bitumen base could be employed at the outset, modified with additives and filler. More often than not, the formula and production process are the intellectual property of the manufacturer (DMS, 2008).

Flexible BMA is a polymer modified marker adhesive. It is normally a GTR-based modified product. In order to achieve other requirements such as ductility, other elastic polymers may be needed in the formulation. Companies that make these products in the USA include:

- Crafo (Crafco, 2008)
- Martin Asphalt (Martin Asphalt, 2014)
- Henry.

Each company produces both standard and flexible BMA in their own brands. Both standard and flexible BMA need fillers added to them to comply with the specific standards listed in Tables 24.9 and 24.10. The filler content can be varied, depending on the neat base. Normally, the filler content is in the range 40–75%.

### Table 24.9 Adhesive properties

<table>
<thead>
<tr>
<th>Property</th>
<th>Material</th>
<th>Test methoda</th>
<th>Standard adhesive</th>
</tr>
</thead>
<tbody>
<tr>
<td>Softening point: °C</td>
<td>ASTM D36</td>
<td></td>
<td>93</td>
</tr>
<tr>
<td>Penetration (100 g, 5 s, 25°C, 0.1 mm)</td>
<td>ASTM D5</td>
<td></td>
<td>10</td>
</tr>
<tr>
<td>Flow (5 h, 70°C (158°F)): mm</td>
<td></td>
<td></td>
<td>5</td>
</tr>
<tr>
<td>Heat stability flow (5 h, 70°C (158°F)): mm</td>
<td></td>
<td></td>
<td>5</td>
</tr>
<tr>
<td>Viscosity (10 rpm, 204°C (400°F)): Pa·s</td>
<td></td>
<td></td>
<td>7.5</td>
</tr>
<tr>
<td>Flash point (COC): °C</td>
<td>ASTM D92</td>
<td></td>
<td>288</td>
</tr>
<tr>
<td>Ductility (5 cm/min, 25°C): cm</td>
<td>ASTM D113</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Flexibility (1 in. mandrel, 90° bend, 10 s)</td>
<td>ASTM D3111</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

a ASTM D36/D36M-14e1 (ASTM, 2014b); ASTM D5 (ASTM, 2006); ASTM D92 (ASTM, 2012b); ASTM D113 (ASTM, 1999); ASTM D3111 (ASTM, 2010c).

### Table 24.10 Properties of flexible bituminous marker adhesives

<table>
<thead>
<tr>
<th>Specification</th>
<th>Test methoda</th>
<th>Requirements</th>
<th>Typical results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Softening point: °C</td>
<td>ASTM D36</td>
<td>93 min.</td>
<td>102</td>
</tr>
<tr>
<td>Penetration (100 g, 5 s, 25°C, 0.1 mm)</td>
<td>ASTM D5</td>
<td>30 max.</td>
<td>22</td>
</tr>
<tr>
<td>Rotational viscosity (191°C): Pa·s</td>
<td>ASTM D4402</td>
<td>2–6</td>
<td>3.8</td>
</tr>
<tr>
<td>Flow (70°C): mm</td>
<td>ASTM D5329</td>
<td>5 max.</td>
<td>0</td>
</tr>
<tr>
<td>Flexibility (−7°C)</td>
<td>ASTM D3111</td>
<td>No breaks or cracks</td>
<td>pass</td>
</tr>
<tr>
<td>Ductility (25°C, 5 cm/min): cm</td>
<td>ASTM D113</td>
<td>15 min.</td>
<td>30</td>
</tr>
<tr>
<td>Ductility (4°C, 1 cm/min): cm</td>
<td>ASTM D113</td>
<td>5 min.</td>
<td>10</td>
</tr>
<tr>
<td>Flash point: °C</td>
<td>ASTM D92</td>
<td>288 min.</td>
<td>315</td>
</tr>
</tbody>
</table>

a ASTM D36/D36M-14e1 (ASTM, 2014b); ASTM D5 (ASTM, 2006); ASTM D4402/D4402M-13 (ASTM, 2013b); ASTM D5329-09 (ASTM, 2009d); ASTM D3111 (ASTM, 2010c); ASTM D113 (ASTM, 1999); ASTM D92 (ASTM, 2012b).
24.2.2 Requirements and specifications for bituminous marker adhesives

24.2.2.1 Adhesive properties
The adhesive must be smooth and homogeneous and contain no visible particles, and must comply with the requirements shown in Table 24.9.

24.2.2.2 Properties of base bitumen
This applies to standard bituminous adhesive only. The filler-free bitumen, obtained from the extraction and Abson recovery process, must have the properties shown in Table 24.11.

24.3 Airfield pavements
Uninterrupted flow of air traffic and safe traffic conditions are essential for all airport owners and their customers. The combination of the large number of airports and the increase in air travel encouraged by low cost flights means that construction works are ongoing at many airports. This is necessary for reasons of expansion and maintenance to keep the taxiways and runways in a serviceable condition.

In airports, both asphalt pavements and concrete pavements are used. Consideration of a number of factors related to cost, performance and environment will assist in explaining the increasing use of asphalt pavements on airfields.

In matters of cost, flexible pavements have the following characteristics.

- Initial construction costs for flexible pavements are generally lower than those for concrete pavements.
- Less time is required for curing before the facility can be opened to traffic.
- Although, in general, maintenance costs for asphalt pavements can be slightly higher than those for concrete pavements, repair of flexible pavements is far easier and quicker than repair of concrete pavements, meaning less downtime for airport operation, a facet that may completely reverse the maintenance cost argument.
- Whole life costs of an asphalt pavement are generally lower than those of a concrete pavement.

<table>
<thead>
<tr>
<th>Property</th>
<th>Minimum</th>
<th>Maximum</th>
<th>Test method</th>
</tr>
</thead>
<tbody>
<tr>
<td>Penetration (100 g, 5 s, 25°C): dmm</td>
<td>25</td>
<td></td>
<td>ASTM D5</td>
</tr>
<tr>
<td>Viscosity (135°C): Pa s</td>
<td>1.2</td>
<td></td>
<td>ASTM D2171</td>
</tr>
<tr>
<td>Viscosity ratio (135°C)</td>
<td></td>
<td>2.2</td>
<td>ASTM D1754</td>
</tr>
</tbody>
</table>

* ASTM D5 (ASTM, 2006); ASTM D2171 (ASTM, 2010a); ASTM D1754 (ASTM, 2009b).
In relation to performance and environmental issues, asphalt pavements can exhibit the following characteristics in comparison to concrete pavements:

- Lower noise levels
- Less cracking, and thus less ravelling or loss of aggregate, which is an important safety aspect for aircraft
- Lower spray and aquaplaning risks, particularly with negatively textured asphalts
- Higher friction when wet
- Vastly superior surface regularity
- Recycling of existing asphalt in new construction is easy and already common practice
- Asphalt pavements require about 20% less energy to produce and construct than other pavements (EAPA, 2014).

The use of flexible pavements manufactured with polymer modified bitumen has become much more common in airfield works. Traditional asphalt mixture design and the design of flexible pavements do not take account of the improved characteristics of asphalts that are modified in this manner.

Many UK airports now use open graded mixtures for the surface course. This material, known as the ‘friction course’, allows water to permeate through the layer, thus maximising the frictional characteristics of the running surface. An impervious binder course, placed below the surface course, and laid to appropriate falls, causes the rainwater to flow away from the pavement. On airfields carrying aircraft that are relatively light, the use of the standard road mixtures may well be quite adequate.

### 24.3.1 Design of flexible airfield pavements

Airfield and highway flexible pavements have many similarities but also some significant differences. The key distinctions between highways and airfields are the types and frequencies of loads that are experienced when the pavement is in service. Airfield pavements tend to experience far fewer load repetitions over their design lives than do highway pavements. For highways, the number of heavy goods vehicles is the primary characteristic used to specify pavement structure and materials. This spectrum of traffic has been quantified by the use of standard axles in Europe, and equivalent single axle loads (ESALs) in the USA and other areas where US practice is followed as the controlling factor for both pavement design and selection of asphalts. For airfields, the pavement is generally designed and specified based on a particular design of aircraft. Depending on whether the airport is a small general aviation airport or a large commercial airport, the design aircraft can be as small as a Cessna Skyhawk, which has...
a gross weight of approximately 1.36 tonnes, or an Airbus A380, which has a maximum take off weight of 590 tonnes (Burns Cooley Dennis, Inc., 2009).

Another factor related to loads is tyre pressure. Small aircraft can have tyre pressures similar to cars, while some military fighter jets can have tyre pressures in excess of 20 bar.

Another difference between airfield and highway pavements is the traffic patterns. For highways, the traffic is generally channelised and falls within narrow wheel tracks on the road. Traffic patterns on airfields can vary from channelised moving (taxiways) to channelised stacked (ends of runways or taxiways) to evenly distributed and random (aprons) to occasional (runway edges) to almost never (shoulders and blast pads).

The number of loads or passes on airfields is, in general, much lower than that found on roads carrying high volumes of traffic but the loads and tyre pressures on aircraft can be significantly higher. This has important consequences for the pavement design and the choice of materials. Figure 24.5 illustrates the significant difference in loading and tyre pressures on roads and airfields.

The high loads on airfields will produce critical stresses and strains in the subgrade and lower layers of the pavements if not properly designed. The extent to which critical stresses and strains are produced in the upper layers of the pavement is dependent on the wheel configuration of the aircraft.

Figure 24.5 Loading and tyre pressures on road and airfield
The traffic situation and loading severity varies greatly at different locations in an airfield, with the ends of runways and taxiways being subject to the greatest concentrations of traffic. In particular, the design of taxiways that are used for departing traffic is critical. Runways are more critical with respect to cracking and adhesion but have a lesser priority from a design perspective.

24.3.1.1 Design methods
The International Civil Aviation Organization (ICAO) classifies the bearing capacity of airfield pavements in terms of a ‘pavement classification number’ (PCN). Similarly, the severity of loading from aircraft is expressed in terms of an ‘aircraft classification number’ (ACN). Several design procedures can be used to select materials, layer thickness, mixture type etc. that will provide a pavement structure with PCN values that exceed the ACN values required by the ICAO.

Some of the most commonly used design procedures for flexible pavements on military and civilian airfields are

- FAA method (Federal Aviation Administration, USA)
- LCN method (UK load classification number system)
- the French method (ICAO French practice)
- Canadian Ministry of Transportation method
- Corps of Engineers method (CBR method)
- Asphalt Institute method
- Shell Pavement Design Method (SPDM).

Most of these systems are empirical, being based on the assessment of the California bearing ratio (CBR). Exceptions are the Asphalt Institute method and SPDM, which are analytical pavement design methods based on calculations and criteria for stresses and strains in the various pavement layers.

These design systems recommend a minimum thickness of asphalt pavement. The methods listed above have few or no specific requirements for the asphalt layers. For this reason, theoretical models for pavement responses are often used as a supplementary tool to optimise the pavement construction and selection of materials. In relation thereto, analytical software such as BISAR is often used. BISAR was designed to calculate and analyse the stresses and strains within the layers of a pavement, and this program was used in drawing up the SPDM design charts that were issued in 1978. More information on SPDM can be found in sections 13.9 and 13.10.

24.3.2 Requirements for airfield pavements
Airfield pavements must perform certain functions in line with agreed international standards. The consequences of a pavement that performs poorly...
may be dangerous and very costly. Detached pieces of aggregate may be ingested by jet engines or strike aircraft moving at speed, causing serious and possibly fatal foreign object damage (FOD). As a consequence, maintenance on airfield pavements is often focused on minimising the risk of FOD.

The most important requirements for asphalt pavements on airfields are adequate

- wet friction for landing aircraft
- resistance to ravelling and cracking to minimise the risk of FOD
- surface regularity
- surface water drainage
- resistance to rutting and fatigue
- pavement bearing capacity
- resistance to deterioration due to spilled fuel (especially on military airfields), de-icing chemicals and other chemicals
- sufficient bonding strength between asphalt layers to avoid slippage cracking.

The asphalt/bitumen solutions used to meet specific airport requirements will vary substantially from location to location depending on several factors such as climate, traffic intensity, types of aircraft etc. Asphalts offer great flexibility in their composition and physical characteristics, and may be specially designed to meet the needs at individual airports. Bitumen, obviously a very important component in relation to the properties of the asphalt, can be selected in different stiffness grades, being specially designed to meet climatic challenges, good resistance to damage resulting from fuel spillage, and improved rutting, low temperature cracking and fatigue properties. Conventional bitumen cannot always meet these demands. Thus, the use of polymer modified bitumen is, in most cases, necessary to meet the specific needs of an airport pavement.

Asphalt concrete with specification requirements for Marshall stability, flow and void content has normally been used for airfields in the past. The surfaces of these pavements were usually grooved to provide sufficient macrotexture and to create drainage channels, leading to improved frictional characteristics. However, the introduction of stone mastic asphalt and porous asphalt on airfields has resulted in improved wet friction properties. Grooves, which often lead to premature ravelling, are unnecessary when stone mastic asphalt or porous asphalt is used as the surface course. The use of an optimised and adequate asphalt composition may offer improved pavement service life. Polymer modified bitumens based on SBS are generally used in dense asphalts, stone mastic asphalt and porous asphalts (e.g. Shell Cariphalte® PG76 and Shell Cariphalte® PG82). These bitumens provide the best possible aggregate adhesion to prevent cracking and ravelling, while at
the same time providing the pavement with an improved anti-rutting performance.

The development of new asphalts has been followed by improvements in mechanical tests for bitumen and asphalts, as well as in specifications and mixture design procedures. Specifications for some recent airport works have been based on various sets of special binder specifications (Dubai Airport in the UAE and Muscat Airport in Oman, Kuala Lumpur in Malaysia, Gardermoen Airport in Norway, Schiphol Airport in the Netherlands, Arianda Airport in Sweden and various airports in Greenland). At Dubai Airport, Muscat Airport and Schiphol Airport, specifications for rutting and crack resistance were used based on the wheel tracking test, dynamic creep test and indirect tensile test, while the binder for Gardermoen Airport was specified using SHRP specifications. Superpave Mix Design was implemented at Muscat Airport. These developments in tests and test criteria for airports have led to new asphalt and bitumen solutions that will further encourage clients to choose asphalts for airfield applications.

The nature of the loading on airfield pavements may produce critical shear stresses in the interface between asphalt layers, and result in slippage cracking if the bonding strength is insufficient. For main airfields with severe traffic, it is advisable to use bond coats (bond coats are always manufactured with polymer modified bitumen) with an application rate resulting in 300 g/m² of residual binder.

Some airports at which Shell Bitumen supplied the bitumen (as at 2014) are listed in Table 24.12.

### 24.4 Conventional and high speed railway applications
#### 24.4.1 Conventional railway

In the USA, Japan and many European countries, asphalts have been tested and used in the construction of railway track beds for many years. One of the earliest known applications dates back to 1894, when an asphalt base, approximately 150 mm thick surfaced with a 30 mm thick surface course, was laid as a tramway bed in Visalia, California. In Japan, asphalt has been used in railway construction since the late 1950s, and it has been the subject of numerous tests and commercial applications in Europe.

In railway construction, asphalts are used in one of four ways (Beecken, 1986):

- to stabilise the aggregate ballast
- as an asphalt layer below the aggregate ballast
- as a direct base for the track sleepers
- as a direct base for the rails.
Table 24.12 Some airports at which Shell Bitumen supplied the bitumen (as at 2014)

<table>
<thead>
<tr>
<th>Country</th>
<th>Name of job/city or state</th>
<th>Product group</th>
<th>Year</th>
</tr>
</thead>
<tbody>
<tr>
<td>Germany</td>
<td>Heringsdorf (public)</td>
<td>Cariphalte</td>
<td>1995</td>
</tr>
<tr>
<td>Germany</td>
<td>Parchim (public)</td>
<td>Cariphalte</td>
<td>1996</td>
</tr>
<tr>
<td>Norway</td>
<td>Bardufoss Airport</td>
<td>Cariphalte</td>
<td>1996</td>
</tr>
<tr>
<td>Germany</td>
<td>Spandahlem (military)</td>
<td>Cariphalte</td>
<td>1998</td>
</tr>
<tr>
<td>Northern Ireland</td>
<td>St Angelo Airport</td>
<td>Cariphalte</td>
<td>1999</td>
</tr>
<tr>
<td>Germany</td>
<td>Baden-Baden (public)</td>
<td>Cariphalte</td>
<td>2000</td>
</tr>
<tr>
<td>UK</td>
<td>East Midlands Airport</td>
<td>Cariphalte</td>
<td>2000</td>
</tr>
<tr>
<td>France</td>
<td>Merignac Airport/Bordeaux</td>
<td>Mexphalte</td>
<td>2001</td>
</tr>
<tr>
<td>France</td>
<td>Toulouse Blagnac Airport/Toulouse</td>
<td>Mexphalte</td>
<td>2001</td>
</tr>
<tr>
<td>Germany</td>
<td>Hamburg (public)</td>
<td>Cariphalte</td>
<td>2001</td>
</tr>
<tr>
<td>Philippines</td>
<td>Diosdado Macapagal International Airport</td>
<td>Cariphalte</td>
<td>2001</td>
</tr>
<tr>
<td>Brunei</td>
<td>Brunei Airport</td>
<td>Cariphalte</td>
<td>2002</td>
</tr>
<tr>
<td>Philippines</td>
<td>Subic Bay International Airport/ Zambales</td>
<td>Cariphalte</td>
<td>2002</td>
</tr>
<tr>
<td>South Africa</td>
<td>Cape Town Airport</td>
<td>Cariphalte</td>
<td>2002</td>
</tr>
<tr>
<td>Argentina</td>
<td>Ezeiza International Airport</td>
<td>Cariphalte</td>
<td>2003</td>
</tr>
<tr>
<td>Germany</td>
<td>Berlin (public)</td>
<td>Cariphalte</td>
<td>2003</td>
</tr>
<tr>
<td>Greece</td>
<td>Alexandroupoli Airport</td>
<td>Cariphalte</td>
<td>2003</td>
</tr>
<tr>
<td>Australia</td>
<td>Sydney Airport Maintenance/NSW</td>
<td>Cariphalte Fuelsafe</td>
<td>2004</td>
</tr>
<tr>
<td>Belgium</td>
<td>Zaventem Airport/Brussels</td>
<td>Cariphalte</td>
<td>2004</td>
</tr>
<tr>
<td>Germany</td>
<td>Fraport Airport/Frankfurt (public)</td>
<td>Cariphalte</td>
<td>2004</td>
</tr>
<tr>
<td>Germany</td>
<td>Finkenwerder Hamburg (private Airbus)</td>
<td>Cariphalte</td>
<td>2004</td>
</tr>
<tr>
<td>Thailand</td>
<td>Bangkok International Airport</td>
<td>Cariphalte Fuelsafe</td>
<td>2004</td>
</tr>
<tr>
<td>Thailand</td>
<td>Udorn Thaini Airport</td>
<td>Cariphalte</td>
<td>2004</td>
</tr>
<tr>
<td>UK</td>
<td>Heathrow Airport</td>
<td>Cariphalte Fuelsafe</td>
<td>2005</td>
</tr>
<tr>
<td>Cambodia</td>
<td>Siem Reap Airport</td>
<td>Cariphalte Fuelsafe</td>
<td>2006</td>
</tr>
<tr>
<td>Cambodia</td>
<td>Phnom Penh Airport</td>
<td>Cariphalte Fuelsafe</td>
<td>2008</td>
</tr>
<tr>
<td>Scotland</td>
<td>Edinburgh Airport</td>
<td>Cariphalte</td>
<td>2008</td>
</tr>
<tr>
<td>UAE</td>
<td>Al Bateen Air Base/Abu Dhabi</td>
<td>Cariphalte PG76-22</td>
<td>2010</td>
</tr>
<tr>
<td>UAE</td>
<td>Qusaiwera Air Base/Abu Dhabi</td>
<td>Cariphalte Fuelsafe</td>
<td>2012</td>
</tr>
<tr>
<td>UAE</td>
<td>Sweihan Air Base/Abu Dhabi</td>
<td>Cariphalte PG76-22</td>
<td>2013</td>
</tr>
<tr>
<td>Netherlands</td>
<td>Schiphol Airport</td>
<td>Cariphalte</td>
<td>1993–2004</td>
</tr>
<tr>
<td>Norway</td>
<td>Gardermoen Airport repairs</td>
<td>Cariphalte</td>
<td>2000–2004</td>
</tr>
<tr>
<td>Greenland</td>
<td>Thule (military)</td>
<td>Cariphalte</td>
<td>2004–2005</td>
</tr>
<tr>
<td>UAE</td>
<td>Dubai International Airport</td>
<td>Cariphalte, Cariphalte Fuelsafe</td>
<td>2005–2013</td>
</tr>
<tr>
<td>Hong Kong</td>
<td>HKIA</td>
<td>Cariphalte Fuelsafe</td>
<td>2006–2007</td>
</tr>
<tr>
<td>Singapore</td>
<td>Changi Airport</td>
<td>Cariphalte</td>
<td>2006–2007</td>
</tr>
<tr>
<td>UAE</td>
<td>Abu Dhabi International Airport</td>
<td>Cariphalte PG76-10</td>
<td>2007–2010</td>
</tr>
<tr>
<td>UAE</td>
<td>Jebel Ali/Al Maktoum International Airport</td>
<td>Cariphalte, Cariphalte</td>
<td>2007–2013</td>
</tr>
<tr>
<td>Sri Lanka</td>
<td>Hambantota International Airport</td>
<td>Cariphalte PG76-22</td>
<td>2011–2013</td>
</tr>
<tr>
<td>Oman</td>
<td>Muscat International Airport</td>
<td>Cariphalte PG76-22</td>
<td>2011–2014</td>
</tr>
<tr>
<td>Oman</td>
<td>Salalah International Airport</td>
<td>Cariphalte PG76-22</td>
<td>2012–2014</td>
</tr>
</tbody>
</table>

24.4.2 Cement asphalt mortar used in high speed railway application

Recent years have seen moves towards improving the capacity of railways and the speeds of trains. China, in particular, has invested heavily in high

722
speed railway technology, having carried out a number of major projects in the main cities.

Traditional rail tracks tend to require significant maintenance, and often have weak subgrades and drainage deficiencies. High speed railways overcome this by using a simple but advanced structure consisting of a continuous slab of concrete base (like a highway structure) with the rails supported on a bearing board, which in turn are supported on a layer of cement asphalt mortar. Such a system, shown in Figure 24.6, allows for adjustment of levels while buffering the high speed heavy loading. Figure 24.7 shows a

Figure 24.6 Cross section of ballastless slab track. (Liu and Liu, 2009)

Figure 24.7 Ballastless high speed track in China
photograph of such a system, a ballastless high speed track in China (Liu and Liu, 2009; Ministry of Railways, 2008).

24.4.2.1 Composition of cement asphalt mortar
Cement asphalt mortar consists of a bitumen emulsion, fine aggregate and additives. Additives include defoamers, superplasticisers and aluminium powder, which are added to give better control of the performance of the emulsion in the field. It is very common to use slow setting modified bitumen emulsion, which will tolerate longer application times without final curing. Both cationic and anionic modified bitumen emulsions have been used to make the mortar in high speed railway projects. Examples where modified bitumen emulsion systems have been used are two high speed railway projects in China: the Harbin–Dalian railway, in which a cationic modified emulsion base was employed, and the Beijing–Shanghai railway, in which an anionic modified emulsion base was employed.

24.4.2.2 Materials specification
Polymer modified cationic bitumen emulsion
The properties of the cationic polymer modified bitumen emulsion plays an important role in successful cement asphalt mortar performance. Typical requirements are shown in Table 24.13.

<table>
<thead>
<tr>
<th>Table 24.13 Polymer modified cationic bitumen emulsion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Test item</td>
</tr>
<tr>
<td>--------------------------------------------------------</td>
</tr>
<tr>
<td>Sieve test (1.18 mm)</td>
</tr>
<tr>
<td>Appearance</td>
</tr>
<tr>
<td>Particle charge</td>
</tr>
<tr>
<td>Viscosity (Engela, 25(^\circ)C)</td>
</tr>
<tr>
<td>Storage stability (1 day, 25(^\circ)C)</td>
</tr>
<tr>
<td>Storage stability (5 days, 25(^\circ)C)</td>
</tr>
<tr>
<td>Miscibility</td>
</tr>
<tr>
<td>Discharge temperature (^\circ)C</td>
</tr>
<tr>
<td>Low temperature storage stability (^\circ)C (1 day, –5(^\circ)C)</td>
</tr>
<tr>
<td>Residual properties</td>
</tr>
<tr>
<td>Residual content %wt</td>
</tr>
<tr>
<td>Penetration (25(^\circ)C, 100 g, 5 s)</td>
</tr>
<tr>
<td>Softening point (^\circ)C</td>
</tr>
<tr>
<td>Solubility</td>
</tr>
<tr>
<td>Ductility (15(^\circ)C)</td>
</tr>
<tr>
<td>Ductility (5(^\circ)C)</td>
</tr>
</tbody>
</table>

\(^a\) ASTM D36/D36M-14e1 (ASTM, 2014b); ASTM D5 (ASTM, 2006); ASTM D113 (ASTM, 1999); ASTM D244-09 (ASTM, 2009c).

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Typical requirements for polymer modified anionic bitumen emulsion are shown in Table 24.14. Important requirements for the emulsified bitumen include binder content, cement miscibility, mixing temperature, water blending ratio, mixing time and workability.

Fine aggregate quality requirements
The source of the fine aggregate ranges from river gravel to crushed rock. The maximum particle size of aggregate is 1.18 mm. The finished aggregate should meet the quality requirements shown in Table 24.15.

24.4.2.3 Cement asphalt mortar performance requirements
The workability of cement asphalt mortar is difficult to control for the following reasons:

- It has a relatively high bitumen content
- The working temperature
- The mixing conditions
- The temperature of the mixture.

### Table 24.14 Polymer modified anionic bitumen emulsion

<table>
<thead>
<tr>
<th>Test item</th>
<th>Unit</th>
<th>Test method</th>
<th>Specification</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sieve test (1.18 mm)</td>
<td>%</td>
<td>ASTM D244</td>
<td>&lt;0.1</td>
</tr>
<tr>
<td>Appearance</td>
<td>–</td>
<td>–</td>
<td>Brown, smooth, no impurity</td>
</tr>
<tr>
<td>Particle charge</td>
<td>–</td>
<td>ASTM D244</td>
<td>Negative</td>
</tr>
<tr>
<td>Viscosity (Saybolt Furol, 25°C)</td>
<td>s</td>
<td>ASTM D244</td>
<td>&gt;20</td>
</tr>
<tr>
<td>Storage stability (1 day, 25°C)</td>
<td>%</td>
<td>ASTM D244</td>
<td>&lt;1.0</td>
</tr>
<tr>
<td>Storage stability (5 days, 25°C)</td>
<td>%</td>
<td>ASTM D244</td>
<td>&lt;5.0</td>
</tr>
<tr>
<td>Brookfield viscosity (40°C)</td>
<td>MPa s</td>
<td>ASTM D2171</td>
<td>≤1000</td>
</tr>
<tr>
<td>Miscibility</td>
<td></td>
<td>ASTM D244</td>
<td>Pass</td>
</tr>
<tr>
<td>Discharge temperature</td>
<td>°C</td>
<td>–</td>
<td>5–35</td>
</tr>
<tr>
<td>Low temperature storage stability</td>
<td>%</td>
<td>ASTM D244</td>
<td>No coagulation</td>
</tr>
<tr>
<td>(1 day, −5°C)</td>
<td>μm</td>
<td></td>
<td>≤7</td>
</tr>
<tr>
<td>Average particle size</td>
<td>μm</td>
<td></td>
<td>≤5</td>
</tr>
<tr>
<td>Mode</td>
<td>μm</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Residual properties</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Residual content</td>
<td>% wt</td>
<td>ASTM D244</td>
<td>≥60</td>
</tr>
<tr>
<td>Penetration (25°C, 100 g, 5 s)</td>
<td>0.1 mm</td>
<td>ASTM D5</td>
<td>40–120</td>
</tr>
<tr>
<td>Softening point</td>
<td>°C</td>
<td>ASTM D36</td>
<td>≥42</td>
</tr>
<tr>
<td>Solubility</td>
<td>%</td>
<td>ASTM D244</td>
<td>≥97</td>
</tr>
<tr>
<td>Ductility (25°C)</td>
<td>cm</td>
<td>ASTM D113</td>
<td>≥100</td>
</tr>
<tr>
<td>Ductility (5°C)</td>
<td>cm</td>
<td>ASTM D113</td>
<td>≥20</td>
</tr>
</tbody>
</table>

* ASTM D36/D36M-14e1 (ASTM, 2014b); ASTM D5 (ASTM, 2006); ASTM D113 (ASTM, 1999); ASTM D2171 (ASTM, 2010a); ASTM D244-09 (ASTM, 2009c).
In most cases, a successful cement asphalt mortar should meet the performance requirements set out in Table 24.16 for cationic types or Table 24.17 for anionic types.

### 24.5 Bridges

In many countries, steel and concrete bridges have asphalt surfacing. One of the main reasons for using asphalt is to protect the steel and concrete structures from the ingress of water and the deleterious effects of de-icing salts. In general, the asphalt bridge pavement system can be split into four different layers:

- A sealing/bonding layer (primer)

#### Table 24.15 Fine aggregate quality requirements

<table>
<thead>
<tr>
<th>Item</th>
<th>Unit</th>
<th>Requirements</th>
<th>Method(a)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fineness modulus</td>
<td>–</td>
<td>1.4–1.8</td>
<td>ASTM C117</td>
</tr>
<tr>
<td>Apparent density (g/cm^3)</td>
<td></td>
<td>≥2.55</td>
<td>ASTM C29</td>
</tr>
<tr>
<td>Water absorption (%)</td>
<td></td>
<td>&lt;3.0</td>
<td></td>
</tr>
<tr>
<td>Dust content (%)</td>
<td></td>
<td>&lt;1.0</td>
<td></td>
</tr>
<tr>
<td>Deleterious materials (%)</td>
<td></td>
<td>&lt;2.0</td>
<td></td>
</tr>
<tr>
<td>Compound colorimetry (–)</td>
<td></td>
<td>Lighter than standard</td>
<td></td>
</tr>
<tr>
<td>Oxidised compound (%)</td>
<td></td>
<td>&lt;0.02</td>
<td></td>
</tr>
<tr>
<td>Sulfide content ((SO_3)%)</td>
<td></td>
<td>≤0.5</td>
<td></td>
</tr>
<tr>
<td>Potential alkali reactivity of aggregates (%) (\text{(mortar-bar method)})</td>
<td></td>
<td>≤0.1</td>
<td>ASTM C289</td>
</tr>
</tbody>
</table>

\(a\) ASTM C117-13 (ASTM, 2013a); ASTM C29 (ASTM, 2009a); ASTM C127 (ASTM, 2012a); ASTM C289 (ASTM, 2007a).

#### Table 24.16 Cationic emulsified cement asphalt mortar requirements

<table>
<thead>
<tr>
<th>No.</th>
<th>Item</th>
<th>Unit</th>
<th>Requirements</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>CAM temperature (\text{°C})</td>
<td></td>
<td>5–40</td>
</tr>
<tr>
<td>2</td>
<td>Flow (s)</td>
<td></td>
<td>18–26</td>
</tr>
<tr>
<td>3</td>
<td>Workability (\text{minute})</td>
<td></td>
<td>≥30</td>
</tr>
<tr>
<td>4</td>
<td>Air content (%)</td>
<td></td>
<td>8–12</td>
</tr>
<tr>
<td>5</td>
<td>Apparent density (kg/m^3)</td>
<td></td>
<td>≥1,300</td>
</tr>
<tr>
<td>6</td>
<td>Compressive strength (\text{MPa})</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>1 day</td>
<td></td>
<td>≥0.1</td>
</tr>
<tr>
<td></td>
<td>7 days</td>
<td></td>
<td>≥0.7</td>
</tr>
<tr>
<td></td>
<td>28 days</td>
<td></td>
<td>≥1.8</td>
</tr>
<tr>
<td>7</td>
<td>Elastic modulus (28 days) (\text{MPa})</td>
<td></td>
<td>100–300</td>
</tr>
<tr>
<td>8</td>
<td>Separation (%)</td>
<td></td>
<td>≤1%</td>
</tr>
<tr>
<td>9</td>
<td>Change of volume (%)</td>
<td></td>
<td>1.0–3.0</td>
</tr>
<tr>
<td>10</td>
<td>Bleeding (%)</td>
<td></td>
<td>0</td>
</tr>
<tr>
<td>11</td>
<td>Freeze/thaw loss (28 days) (\text{-})</td>
<td></td>
<td>Relative dynamic modulus ≥60%, mass loss &lt;5% after 300 freeze/thaw cycles</td>
</tr>
<tr>
<td>12</td>
<td>Weathering tests (28 days) (\text{-})</td>
<td></td>
<td>No peeling, cracking Compressive strength ratio</td>
</tr>
</tbody>
</table>
Other important uses of bitumens and asphalts

Asphalt layers on bridge decks must absorb traffic loads, and successfully transfer them to the supporting elements of the structure without the asphalts themselves deteriorating. The surface course asphalt should retain good surface regularity, be resistant to deformation and provide adequate skidding resistance for vehicles. All the asphalt layers must protect the bridge structure from surface water, which in winter contains de-icing salt that promotes corrosion. These functions are generally not fulfilled or only partially fulfilled by one material, and a functional division can be made for the layers forming the surface of the deck. Asphalts typically used on bridges include dense asphalt concrete, mastic asphalt and stone mastic asphalt.

In general terms, the bridge deck is covered by a bitumen based waterproofing layer, a sealing/bonding layer, a protective layer, a binder course and a surface course. The thickness of the bridge pavement varies, and depends on the type of deck. The sealing layer must cover the full surface of the bridge deck.

<table>
<thead>
<tr>
<th>No.</th>
<th>Item</th>
<th>Unit</th>
<th>Requirements</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Cement asphalt mortar temperature</td>
<td>°C</td>
<td>5–35</td>
</tr>
<tr>
<td>2</td>
<td>Spread and passing</td>
<td>–</td>
<td>ASTM D5: ≥280 mm, T 280 ≤16 s</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>ASTM D30: ≥280 mm, T 280 ≤22 s</td>
</tr>
<tr>
<td>3</td>
<td>Flow</td>
<td>s</td>
<td>80–120</td>
</tr>
<tr>
<td>4</td>
<td>Separation</td>
<td>%</td>
<td>&lt;3%</td>
</tr>
<tr>
<td>5</td>
<td>Air content</td>
<td>%</td>
<td>&lt;10%</td>
</tr>
<tr>
<td>6</td>
<td>Apparent density</td>
<td>kg/m³</td>
<td>≥1800</td>
</tr>
<tr>
<td>7</td>
<td>Volume change</td>
<td>%</td>
<td>0–2.0</td>
</tr>
<tr>
<td>8</td>
<td>Flexural strength</td>
<td>MPa</td>
<td></td>
</tr>
<tr>
<td></td>
<td>1 day</td>
<td></td>
<td>≥1</td>
</tr>
<tr>
<td></td>
<td>7 days</td>
<td></td>
<td>≥2</td>
</tr>
<tr>
<td></td>
<td>28 days</td>
<td></td>
<td>≥3</td>
</tr>
<tr>
<td>9</td>
<td>Compressive strength</td>
<td>MPa</td>
<td></td>
</tr>
<tr>
<td></td>
<td>1 day</td>
<td></td>
<td>≥2</td>
</tr>
<tr>
<td></td>
<td>7 days</td>
<td></td>
<td>≥10</td>
</tr>
<tr>
<td></td>
<td>28 days</td>
<td></td>
<td>≥15</td>
</tr>
<tr>
<td>10</td>
<td>Elastic modulus (28 days)</td>
<td>MPa</td>
<td>7000–10 000</td>
</tr>
<tr>
<td>11</td>
<td>Freeze/thaw (28 days)</td>
<td>–</td>
<td>No failure on surface</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Mass loss &lt;2000 g/m²</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Dynamic modulus ratio greater than 60%</td>
</tr>
<tr>
<td>12</td>
<td>Fatigue test (28 days)</td>
<td>–</td>
<td>No cracking after 10 000 cycles</td>
</tr>
</tbody>
</table>

Concrete decks may be primed with any one of a range of materials, including bitumen emulsion, polymer modified bitumen, epoxy resin and polyurethane.

In steel structures there may be large deformations in the bridge deck, and therefore fatigue resistance in the asphalt layers is more important for steel bridges than for concrete bridges. Different countries use different asphalts in a range of layer thicknesses.

Mastic asphalt is often used on steel bridge decks, and also as a waterproofing layer on both concrete and steel bridges. Mastic asphalt is thermoplastic, which means that it does not crack quickly but is sensitive to deformation. A modified stone mastic asphalt is an alternative to mastic asphalt.

The asphalt mixture design for concrete bridge decks is easier because of the rigid behaviour of the deck. The asphalt for the surface layers can be adjusted to be stiffer. The protection layer should be designed to be slightly softer than the surface layer. Generally, on smaller bridges the surface course is often the same as that used on the adjacent road. On larger bridges, different surface courses can be used such as stone mastic asphalt or thin surface course systems.

Proper and timely maintenance and rehabilitation is essential for good performance and durability. For repairing cracked bridge decks, it is necessary to ascertain whether the cracks are structural or non-structural. Major rehabilitation works may range from the installation of a protective surface system, to complete replacement of the bridge deck. Preventive maintenance applied at a cost effective time will preserve and extend the useful life of a bridge.

A comprehensive paper on the use of asphalt pavements on bridge decks was issued by the European Asphalt Pavement Association (EAPA, 2013).

### 24.6 Recreational areas

Asphalts are used extensively for recreational areas (e.g. tennis, netball and volleyball courts, five-a-side football pitches). They are used in two ways:

- as the final surfacing over an appropriate base, which may itself be asphalt
- as a firm base beneath a synthetic surfacing.

By choosing an appropriate mixture, a pleasing and durable finish requiring little maintenance over long periods can be obtained. Asphalts are generally black or dark grey, providing a good contrast for line paints. If a coloured surfacing is required, this can be achieved by using a special pigmentable binder or by painting the surface with a special proprietary paint.

Synthetic sports surfaces are very popular for athletic tracks, football pitches and other specialised applications. Asphalts have been found to be ideal for forming the base for these surfaces.
Impact absorbing asphalt (IAA) is a patented technology (European Union, 1995) consisting of a relatively conventional asphalt concrete with a large proportion of the aggregate (typically up to 30%) replaced by dry crumb rubber obtained from the recycling of truck tyres. It has many potential applications. One example is as a surface layer for play areas. Another is its use as a shock pad underneath artificial sports surfaces, where it provides the required impact absorption, longitudinal and transverse regularity.

The technology is based on the dry process developed in the late 1960s in Sweden and traded in Europe under the name Rubit (Green and Tolonen, 1977; Heitzman, 1992; Takallou and Hicks, 1988). Production of the material involves the addition of crumb rubber to hot aggregate prior to mixing with bitumen. The material is then mixed and maintained at a high temperature. It is laid and compacted using conventional paving techniques.

A number of recreational applications can be found on the website of the Asphalt Paving Association of Iowa (2014).

24.7 Motor racing tracks, including Formula 1 tracks
International motor racing circuits are generally surfaced with asphalt. The main requirements for the racing tracks are

- very good surface regularity
- a high value of skid resistance
- sufficient cohesion to resist the high tangential forces exerted by wide racing tyres.

This last property is normally achieved by using dense mixtures such as dense asphalt concrete. However, stone mastic asphalts have also been used as racing track surface courses. Cohesion of the asphalt is greatly improved by using polymer modified bitumens.

As much as the surfaces of racetracks and standard roads may be similar, the stresses acting on them could hardly be more different. On a normal road, the greatest stresses occur when, for instance, a 44 tonne heavy goods vehicle brakes. On race tracks, the effects are the exact opposite. The hot tyres of the race cars develop an adhesive quality similar to tough chewing gum and pull the stones upwards. A high value of strain is also created by the high speeds, which result in intense pressure in front of the cars and a vacuum behind them. The effect has been described as: ‘For the asphalt, this is as if someone was pounding it with a hammer at the front and hoovering it at the back’ (Formula 1, 2014, citing Tilke engineers and architects).

Shell has developed products in order to fulfil many advanced professional applications for clients worldwide. Shell has wide experience in supplying Shell Cariphalte Racetrack to a variety of racing circuit projects.
Table 24.18 lists a number of projects where Cariphalte Racetrack has been used.

### 24.8 Vehicle testing circuits

Many vehicle testing circuits have steep banked curves, and special techniques have been developed by specialist contractors to ensure that the finished pavement is fully compacted, meets the surface tolerances and has a high degree of surface regularity. Figure 24.8 shows a surfacing being laid on the Rockingham Speedway track in England. On this project, considerable skill and expertise was required to lay and compact materials to the required specification under the prevailing conditions.

### 24.9 Hydraulic applications

The waterproofing properties of bitumens and asphalts have been recognised since ancient times. The ancient civilisations of Ur, Egypt, Babylon, Assyria and the Indus all used the naturally occurring surface seepages of bitumen for waterproofing and building. The reservoir dam at Mohenjo Daro in the Indus basin is particularly well preserved, and demonstrates that, in this field, asphalt can claim a life of 5000 years (Menzies, 1988).

Bitumens and asphalts have been used effectively for a range of hydraulic applications such as canal lining, reservoir lining, sea wall construction, coastal groynes, dam construction and the lining of leisure lakes. In the UK, dams at Dungonnell, Colliford, Winscar, Marchlyn and Sulby incorporate bitumens. A further example of the use of bitumens in hydraulic applications is shown in Figure 24.9, which shows a photograph of the...
Figure 24.8 Surfacing the Rockingham Speedway track. (Courtesy of Colas Ltd.)

Figure 24.9 Bituminous grouting on the face of the Megget Dam
Megget Dam in the south of Scotland. Reservoirs at Shotton and Towey in Wales and Leamington in England also make significant use of bitumens for their waterproofing properties.

The two principal properties that make bitumen ideal for hydraulic applications are its impermeability and the fact that it is chemically inert. In combination with suitable aggregates in mixtures that are specifically designed for particular applications, bitumen can impart these properties to the structures in which the asphalt is used. In many of these structures it is essential to have impermeability under pressure, and asphalts with void contents below 3% can successfully contain water column pressures up to 200 m.

Reservoir and canal embankments often have a slope of 1 : 1.75 but can be as steep as 1 : 1.25. Thus, asphalts used in such applications must have sufficient stability to be laid and compacted on such slopes without cracking during application and in service. The material must be flexible enough to accommodate differential settlement of the substrate.

One great advantage of asphalt is that it can be laid on the dam or reservoir face in a continuous manner, thus eliminating joints. Any joint constitutes a discontinuity and, at the high pressures that exist at the base of dams and reservoirs, a joint provides the opportunity for the water to find an outlet. The Shell Bitumen Hydraulic Engineering Handbook provides substantial detail on hydraulic applications (Scönnian, 1999).

24.10 Coloured surface courses and surface treatments
The appearance of traditional asphalts, especially when finished with the normal contrasting white lines, is generally pleasing to the eye. However, there are some locations where a specific colour is desired (Table 24.19), and there are a number of ways in which this can be achieved:

- incorporation of coloured pigments in the mixture during manufacture
- application of suitably coloured chippings to the asphalt during laying
- application of a coloured surface treatment after laying
- use of a conventional bitumen with a coloured aggregate
- use of a suitably coloured aggregate with a translucent binder.

An example of the application of coloured asphalt in a recreational park is shown in Figure 24.10.

24.10.1 Application of coloured chippings
Hot rolled asphalt and fine graded asphalt concrete (previously called fine cold asphalt) can have decorative coloured chippings rolled into the surface during the compaction phase. Hot rolled asphalt is suitable for most traffic conditions but fine graded asphalt concrete is only appropriate for lightly trafficked applications such as private drives and pedestrian areas.
To provide a decorative finish to both these types of asphalt, pigmented bitumen coated or clean resin coated chippings can be applied during laying. The bitumen or resin coating promotes adhesion to the surface. The use of uncoated chippings is not recommended, as they are liable to become detached from the surface. However, in the case of fine graded

<table>
<thead>
<tr>
<th>Purpose of providing coloured surface</th>
<th>Examples of applications</th>
</tr>
</thead>
<tbody>
<tr>
<td>Increasing safety</td>
<td>Road crossings, obstacles, bus stops, turning pre-selections, zone 30 km, tunnels</td>
</tr>
<tr>
<td>Controlling and guiding traffic</td>
<td>Footpath, cycle lane, bus lane, parking area</td>
</tr>
<tr>
<td>Improve visibility/safety, save energy</td>
<td>Tunnels, in front of schools, fire department exits</td>
</tr>
<tr>
<td>Optical illusion (make objects bigger)</td>
<td>Football area</td>
</tr>
<tr>
<td>Increase attractiveness</td>
<td>Pleasure areas, park lanes, pleasure grounds</td>
</tr>
<tr>
<td>Increasing attractiveness by avoiding contrast</td>
<td>Historical sites, natural or historical environment, sport areas, zoos, prestige sites, square, footpaths</td>
</tr>
<tr>
<td>Improving comfort</td>
<td>Cycling, walking, skating</td>
</tr>
<tr>
<td>Improving comfort or performance, reduce pavement warming</td>
<td>Footpath – reduced heat radiation</td>
</tr>
<tr>
<td>Improved appearance</td>
<td>Heavily trafficked road – reduced rutting</td>
</tr>
<tr>
<td></td>
<td>Using light chippings in footpaths</td>
</tr>
</tbody>
</table>

Figure 24.10 Example of coloured asphalt used in a recreational park
macadam that is laid on areas subject to little traffic, a light application of uncoated chippings (e.g. white limestone chippings) produces a very attractive finish if they are uniformly applied and properly embedded. Many footways in Scotland are finished with a 30 mm thick layer of hot rolled asphalt sand carpet (i.e. hot rolled asphalt surface course without the coarse aggregate) into which white limestone chippings are added. Decorative chippings cannot be successfully rolled into the surface of high stone content mixtures such as dense asphalt concretes or high stone content hot rolled asphalts.

24.10.2 Application of coloured surface treatments
There are three surface treatments that can be applied to an asphalt surface course to provide a decorative finish

- pigmented slurry seals
- surface dressings
- application of a coloured paint.

Pigmented slurry seals are available as proprietary products in a range of colours. However, as they are applied in a very thin coat (approximately 3 mm thick) they are only suitable for pedestrian and lightly trafficked areas.

Surface dressings are suitable for most categories of road application (Roberts and Nicholls, 2008) but are less suitable for pedestrian situations. The final colour of the surface dressing will be that of the aggregate used.

Proprietary coloured paints are available for painting on top of conventional black surface courses. These are only suitable for pedestrian and games areas (e.g. tennis courts, street painting).

24.10.3 Coloured aggregate bound by a conventional bitumen
When conventional bitumen is used, the depth of colour achieved is dependent on

- the colour of the aggregate itself
- the thickness of the binder film on the aggregate
- the rate at which the binder on the road surface is exposed by traffic.

In medium and heavily trafficked situations, the natural aggregate colour will start showing through fairly quickly, with a year being typical. In lightly trafficked situations, where coloured surfacings are often specified, it may take significantly longer (3 years or more) for the aggregate colour to become apparent.

24.10.4 Coloured aggregate bound by a translucent binder
Several proprietary asphalt concretes are available in which the binder is a clear resin rather than bitumen. A range of coloured surfacings can be manufactured by selecting appropriately coloured aggregates. The major
advantage of such a system is that the colour is obtained immediately. For more information on this subject see section 8.3.2.

24.11 Asphalt kerbs

Extruded asphalt kerbs are very simple to construct. They are produced by extruding a hot rolled asphalt through specialised equipment directly onto a binder course or surface course, which has been treated with a tack coat or bond coat. Asphalt kerbs are unsuitable for the edge of carriageways carrying heavy traffic, as they may be damaged if hit by vehicles, particularly commercial vehicles. However, they do provide a convenient means of producing a raised edge on other paved areas (e.g. perimeter kerbing to drives or car parks). In the UK, BS 5931:1980 (BSI, 2014) gives guidance on the use of asphalt for producing kerbs in situ.

An extruded asphalt kerb on a UK motorway is shown in Figure 24.11.

References
Asphalt Institute (2014) Use of a DSR to evaluate effect of aging on roofing membrane. AI spring meeting, San Antonio, TX, USA.


ASTM (2009a) C29. Standard test method for bulk density ('unit weight') and voids in aggregate. ASTM, West Conshohocken, PA, USA.


ASTM (2009c) D244-09. Standard test methods and practices for emulsified asphalts. ASTM, West Conshohocken, PA, USA.


ASTM (2010d) D3462/D3462M-10a. Standard specification for asphalt shingles made from glass felt and surfaced with mineral granules. ASTM, West Conshohocken, PA, USA.


ASTM (2012a) C127. Standard test method for density, relative density (specific gravity), and absorption of coarse aggregate. ASTM, West Conshohocken, PA, USA.
ASTM (2012b) D92. Standard test method for flash and fire points by Cleveland open cup tester. ASTM, West Conshohocken, PA, USA.


Mike A (2013) Rheology. Asphalt Institute, Lexington, KY, USA. Spring meeting.


Appendices

Appendix 1
Physical constants of bitumens

A1.1 Specific gravity
The specific gravity of a bitumen is primarily dependent on the grade of the bitumen and the temperature (Pfeiffer, 1950). Typical values of specific gravity for a range of grades of bitumen are given in Table A1.1, and the effect of temperature on specific gravity is detailed in Table A1.2. The conversion from volume of bitumen in litres to weight in tonnes at various temperatures and values of specific gravity is detailed in Table A1.3.

<table>
<thead>
<tr>
<th>Grade of bitumen</th>
<th>Typical specific gravity at 25°C</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Penetration grades</strong></td>
<td></td>
</tr>
<tr>
<td>160/220</td>
<td>1.015–1.025</td>
</tr>
<tr>
<td>100/150</td>
<td>1.020–1.030</td>
</tr>
<tr>
<td>70/100</td>
<td>1.020–1.030</td>
</tr>
<tr>
<td>50/70</td>
<td>1.020–1.030</td>
</tr>
<tr>
<td>40/60</td>
<td>1.025–1.035</td>
</tr>
<tr>
<td>30/45</td>
<td>1.025–1.035</td>
</tr>
<tr>
<td>20/30</td>
<td>1.030–1.040</td>
</tr>
<tr>
<td><strong>Oxidised grades</strong></td>
<td></td>
</tr>
<tr>
<td>75/30</td>
<td>1.015–1.030</td>
</tr>
<tr>
<td>85/25</td>
<td>1.020–1.030</td>
</tr>
<tr>
<td>85/40</td>
<td>1.010–1.025</td>
</tr>
<tr>
<td>95/25</td>
<td>1.015–1.030</td>
</tr>
<tr>
<td>105/35</td>
<td>1.000–1.015</td>
</tr>
<tr>
<td>115/15</td>
<td>1.020–1.035</td>
</tr>
<tr>
<td><strong>Hard grades</strong></td>
<td></td>
</tr>
<tr>
<td>H 80/90</td>
<td>1.045–1.055</td>
</tr>
<tr>
<td>H 110/120</td>
<td>1.055–1.065</td>
</tr>
<tr>
<td><strong>Cut-back grades</strong></td>
<td></td>
</tr>
<tr>
<td>50 s</td>
<td>0.992–1.002</td>
</tr>
<tr>
<td>100 s</td>
<td>0.995–1.005</td>
</tr>
<tr>
<td>200 s</td>
<td>0.997–1.007</td>
</tr>
</tbody>
</table>
A1.2 Coefficient of thermal expansion

The coefficient of thermal expansion of bitumen is effectively independent of the grade, and is $6.1 \times 10^{-4}/\degree C$.

A1.3 Coefficient of thermal contraction

The coefficient of thermal contraction (Jones et al., 1968) of bitumen is effectively independent of the grade, and is $3.45 \times 10^{-4}/\degree C$.

| Table A1.2 Typical specific gravities of bitumen at various temperatures |
|-----------------------------|-----------------------------|-----------------------------|-----------------------------|-----------------------------|-----------------------------|-----------------------------|
| Temperature: °C | Specific gravity at 25°C | 1.00 l/tonne | 1.01 l/tonne | 1.02 l/tonne | 1.03 l/tonne | 1.04 l/tonne | 1.05 l/tonne |
| 15.5 | 1.006 | 1.016 | 1.026 | 1.036 | 1.046 | 1.056 |
| 25 | 1.000 | 1.010 | 1.020 | 1.030 | 1.040 | 1.050 |
| 45 | 0.988 | 0.998 | 1.008 | 1.018 | 1.028 | 1.038 |
| 60 | 0.979 | 0.989 | 0.999 | 1.009 | 1.019 | 1.029 |
| 90 | 0.961 | 0.971 | 0.981 | 0.991 | 1.001 | 1.011 |
| 100 | 0.955 | 0.965 | 0.975 | 0.985 | 0.995 | 1.005 |
| 110 | 0.949 | 0.959 | 0.969 | 0.979 | 0.989 | 0.999 |
| 120 | 0.943 | 0.953 | 0.963 | 0.973 | 0.983 | 0.993 |
| 130 | 0.937 | 0.947 | 0.957 | 0.967 | 0.977 | 0.987 |
| 140 | 0.931 | 0.941 | 0.951 | 0.961 | 0.971 | 0.981 |
| 150 | 0.925 | 0.935 | 0.945 | 0.955 | 0.965 | 0.975 |
| 160 | 0.919 | 0.929 | 0.939 | 0.949 | 0.959 | 0.969 |
| 170 | 0.913 | 0.923 | 0.933 | 0.943 | 0.953 | 0.963 |
| 180 | 0.907 | 0.917 | 0.927 | 0.937 | 0.947 | 0.957 |
| 190 | 0.901 | 0.911 | 0.921 | 0.931 | 0.941 | 0.951 |
| 200 | 0.895 | 0.905 | 0.915 | 0.925 | 0.935 | 0.945 |

| Table A1.3 Conversion factors linking volume to weight for bitumen at various temperatures and specific gravities |
|-----------------------------|-----------------------------|-----------------------------|-----------------------------|-----------------------------|-----------------------------|-----------------------------|
| Temperature: °C | Specific gravity at 25°C | 1.00 l/tonne | 1.01 l/tonne | 1.02 l/tonne | 1.03 l/tonne | 1.04 l/tonne | 1.05 l/tonne |
| 25 | 995 | 984 | 973 | 963 | 953 | 943 |
| 45 | 1010 | 999 | 988 | 978 | 968 | 958 |
| 60 | 1020 | 1009 | 998 | 988 | 978 | 968 |
| 90 | 1041 | 1030 | 1019 | 1009 | 999 | 989 |
| 100 | 1047 | 1036 | 1026 | 1015 | 1005 | 995 |
| 110 | 1054 | 1043 | 1032 | 1022 | 1011 | 1001 |
| 120 | 1060 | 1049 | 1038 | 1028 | 1017 | 1007 |
| 130 | 1068 | 1056 | 1045 | 1034 | 1024 | 1013 |
| 140 | 1072 | 1063 | 1052 | 1041 | 1031 | 1019 |
| 150 | 1081 | 1070 | 1058 | 1047 | 1036 | 1026 |
| 160 | 1088 | 1076 | 1065 | 1054 | 1043 | 1032 |
| 170 | 1095 | 1083 | 1072 | 1060 | 1049 | 1038 |
| 180 | 1103 | 1091 | 1079 | 1067 | 1056 | 1045 |
| 190 | 1110 | 1098 | 1086 | 1074 | 1063 | 1052 |
| 200 | 1117 | 1105 | 1093 | 1082 | 1070 | 1058 |
A1.4 Electrical properties

Electrical resistance

Bitumen has a very high resistance and is, therefore, an ideal insulating material. Hard grades of bitumen have slightly higher resistance than soft grades. However, these differences are insignificant. The resistance of all grades of bitumen falls with increasing temperature, as shown in Table A1.4.

The influence of fillers on electrical resistance is negligible unless conductive fillers such as graphite, coke or metal powders are used in significant quantities.

<table>
<thead>
<tr>
<th>Temperature: °C</th>
<th>Resistance: Ω/cm</th>
</tr>
</thead>
<tbody>
<tr>
<td>30</td>
<td>$10^{14}$</td>
</tr>
<tr>
<td>50</td>
<td>$10^{13}$</td>
</tr>
<tr>
<td>80</td>
<td>$10^{12}$</td>
</tr>
</tbody>
</table>

Dielectric strength

The dielectric strength is measured in kilovolts per millimetre, and depends on the conditions of measurement and the shape of the electrodes. Hard grades of bitumen tend to have a higher dielectric strength than soft grades. The dielectric strength of all grades decreases with increasing temperature, as shown in Table A1.5.

<table>
<thead>
<tr>
<th>Temperature: °C</th>
<th>Dielectric strength (flat electrodes): kV/mm</th>
</tr>
</thead>
<tbody>
<tr>
<td>20</td>
<td>20–30</td>
</tr>
<tr>
<td>50</td>
<td>10</td>
</tr>
<tr>
<td>60</td>
<td>5</td>
</tr>
</tbody>
</table>

Dielectric constant (permittivity)

The dielectric constant (or permittivity) of bitumen is about 2.7 at 25°C, rising to about 3.0 at 100°C. The dielectric losses in bitumen rise with increasing temperature but fall with increasing frequency. The Transport and Road Research Laboratory showed that the rate at which bitumen weathers under the combined effects of oxygen, rain, oil deposition from traffic and ultraviolet component of sunlight is related to the dielectric constant of the bitumen (Green, 1977).

A1.5 Thermal properties

Specific heat

The specific heat of bitumen is dependent on both the grade of the bitumen and its temperature. Values of specific heat vary from 1675 to 1800 J/kg/°K at
0°C. The specific heat is increased by 1.67–2.51 J/kg for every 1°C increase in temperature.

**Thermal conductivity**

Bitumen is a good thermal insulating material, and typically has a thermal conductivity of 0.15–0.17 W/m/°K.

**References**


Appendix 2
Conversion factors for viscosities

The viscosity of bitumen is usually measured using either capillary or cup viscometers (for more detail, see section 6.3). In capillary viscometers, viscosity is determined by timing the flow of bitumen through a glass tube at a given temperature. The product of flow time and a calibration factor gives the kinematic viscosity in centistokes (cSt) or mm²/s. In cup viscometers, the time is recorded in seconds for a standard volume of bitumen to flow out through the orifice in the bottom of the cup. The values given in Table A2.1 enable conversion of a cup viscosity to a kinematic viscosity for a number of different types of cup viscometer, and vice versa.

Table A2.1 Conversion factors for viscosities

<table>
<thead>
<tr>
<th>Known viscosity</th>
<th>Kinematic: mm²/s</th>
<th>Redwood I: s</th>
<th>Redwood II: s</th>
<th>Saybolt Universal: s</th>
<th>Saybolt Furol: s</th>
<th>Engler: °E</th>
<th>Standard tar viscometer: s</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kinematic: mm²/s</td>
<td>–</td>
<td>4.05</td>
<td>0.405</td>
<td>4.58</td>
<td>0.458</td>
<td>0.132</td>
<td>0.0025</td>
</tr>
<tr>
<td>Redwood I: s</td>
<td>0.247</td>
<td>–</td>
<td>0.1</td>
<td>1.13</td>
<td>0.113</td>
<td>0.0326</td>
<td>–</td>
</tr>
<tr>
<td>Redwood II: s</td>
<td>2.47</td>
<td>10</td>
<td>–</td>
<td>11.3</td>
<td>1.13</td>
<td>0.326</td>
<td>0.0062</td>
</tr>
<tr>
<td>Saybolt Universal: s</td>
<td>0.218</td>
<td>0.885</td>
<td>0.0885</td>
<td>–</td>
<td>0.1</td>
<td>0.0287</td>
<td>–</td>
</tr>
<tr>
<td>Saybolt Furol: s</td>
<td>2.18</td>
<td>8.85</td>
<td>0.885</td>
<td>10</td>
<td>–</td>
<td>0.287</td>
<td>0.0054</td>
</tr>
<tr>
<td>Engler: °E</td>
<td>7.58</td>
<td>30.7</td>
<td>3.07</td>
<td>34.81</td>
<td>3.48</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Standard tar viscometer (10 mm cup): s</td>
<td>400</td>
<td>–</td>
<td>162</td>
<td>–</td>
<td>183</td>
<td>528</td>
<td>–</td>
</tr>
</tbody>
</table>
Appendix 3
Blending charts and formulae

Bitumen can be blended with a wide variety of crude oil-based fractions for different applications. Volatile light fractions (e.g. white spirit) are used when rapid curing is required. Less volatile fractions, such as cut-back kerosene, are used for the manufacture of cut-back bitumens. Figures A3.1 and A3.2 show the effect of adding cut-back kerosene and diesel, respectively, to 70/100, 160/220 and 250/330 pen bitumen. Similarly, Figure A3.3

Figure A3.1 Typical blending curves for 70/100, 160/220 and 250/330 pen bitumen and cut-back kerosene
Figure A3.2 Typical blending curves for 70/100, 160/220 and 250/330 pen bitumen and diesel

Figure A3.3 Typical blending curve for 40/60 pen bitumen and diesel
shows the effect on penetration when diesel is added to 40/60 pen bitumen.

Bitumens are miscible with each other in all proportions. The penetration and softening point of a blend of two bitumens can be estimated using the blending charts shown in Figures A3.4 and A3.5. These charts should only be used for blends of bitumen of the same type (i.e. those having the same penetration index). They should not be used for blends of oxidised and penetration bitumens. As an alternative to these charts, the following formulae can be used to estimate the penetration and softening point of a blend

\[ \log P = \frac{A \log P_a + B \log P_b}{100} \]

where \( P \) is the penetration of the final blend, \( P_a \) is the penetration of component a, \( P_b \) is the penetration of component b, \( A \) is the percentage of component a in the blend, and \( B \) is the percentage of component b in the blend

\[ S = \frac{A S_a + B S_b}{100} \]

where \( S \) is the softening point of the final blend, \( S_a \) is the softening point of component a, \( S_b \) is the softening point of component b, \( A \) is the percentage of component a in the blend, and \( B \) is the percentage of component b in the blend.

**How to use the blending charts – Figures A3.4 and A3.5**

For Figure A3.4, mark the measured penetration value (at 25°C) for the two bitumens you wish to blend (bitumens A and B) on the left hand and right hand y axes of the graph. Draw a straight line joining these two points – this is the blend line. Select the desired penetration of the final product you wish to produce and draw a horizontal line from the left hand y axis to the point at which it intersects the blend line. Draw a vertical line down from the point of intersection to the x axis and read the percentage of bitumen A and the percentage of bitumen B from the x axis. This is the estimated blend ratio required to produce bitumen of the desired penetration from the two components.

An example. Bitumen A has penetration of 100 dmm and bitumen B has a penetration of 20 dmm. To produce a bitumen with a penetration of 50 dmm from these two components would require a blend of 60% of bitumen A and 40% of bitumen B.

For Figure A3.5 the process is exactly the same but softening point values are used.
Figure A3.4 Chart for estimating the penetration of a blend of two bitumens

Figure A3.5 Chart for estimating the softening point of a blend of two bitumens
Appendix 4
Calculation of bitumen film thickness in an asphalt

It is possible to estimate the surface area of a sample of aggregate by coating it with an oil and measuring the quantity of oil required for complete coating. Alternatively, the surface area of an aggregate can be calculated by assuming a specific aggregate particle shape. Hveem (ASTM, 1992) calculated surface area factors, assuming a spherical particle shape and a specific gravity of 2.65. Typical surface area factors are shown in Table A4.1.

<table>
<thead>
<tr>
<th>Sieve size: mm</th>
<th>Surface area factor: m²/kg</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.075</td>
<td>32.77</td>
</tr>
<tr>
<td>0.150</td>
<td>12.29</td>
</tr>
<tr>
<td>0.300</td>
<td>6.14</td>
</tr>
<tr>
<td>0.600</td>
<td>2.87</td>
</tr>
<tr>
<td>1.18</td>
<td>1.64</td>
</tr>
<tr>
<td>2.36</td>
<td>0.82</td>
</tr>
<tr>
<td>&gt;4.75</td>
<td>0.41</td>
</tr>
</tbody>
</table>

The theoretical bitumen film thickness is then calculated as

\[ T = \frac{b}{100 - b} \times \frac{1}{\rho_b} \times \frac{1}{SAF} \]

where \( T \) is the bitumen film thickness (mm), \( \rho_b \) is the density of bitumen (kg/m³), \( SAF \) is the surface area factor (m²/kg), and \( b \) is the bitumen content (%).

Another method, developed in France (Norme Française, 1991), gives an approximation of the binder film thickness using the formula

\[ T = \frac{b}{a \times \sqrt{\Sigma}} \]
where $T$ is the bitumen film thickness (mm), $b$ is the bitumen content (% by total mass of the mixture), and $a$ is a correction coefficient taking into account the density of the aggregate. The coefficient $a$ is given by

$$a = \frac{2650}{SG_a}$$

where $SG_a$ is the density of the aggregate (kg/m$^3$). $\Sigma$ is the specific surface area of the aggregate, and is given by

$$\Sigma = 0.25G + 2.3S + 12s + 135f$$

where $G$ is the proportion by mass of aggregate over 6.3 mm, $S$ is the proportion by mass of aggregate between 6.3 and 3.15 mm, $s$ is the proportion by mass of aggregate between 3.15 and 0.80 mm, and $f$ is the proportion by mass of aggregate smaller than 0.80 mm.

References
Appendix 5
Bitumen product standards used across the world

The specification systems used in bitumen product standards can be broadly described as either empirical or performance based. Those based on empirical testing such as penetration, softening point and viscosity have been used for many decades in a variety of climates. The performance based specification system (PG grading) used in the USA is well established in product standards.

Derivations, combinations and local adaptations of these standards are used depending on the regulatory environment, climate and historical factors. Even when regional product standards are used (e.g. the harmonised norms in Europe), differences still exist between countries.

Penetration and the ring and ball softening point test are used as the main characteristics in product standards throughout Europe, many parts of Asia (including China), Africa and Argentina. Viscosity based standards are used more commonly in Australia and New Zealand, and for the softer grades in Europe and South America. India is the latest country to move to viscosity grading. The VG grading used in India is slightly different from the Australian C grade, the VG grades having a higher requirement at 60°C.

In Argentina, although penetration and viscosity standards are valid, viscosity specifications are becoming more widely used for both soft and hard grades. In Columbia, penetration grades are used, and in other countries in Latin America viscosity standards are most common.

China is unique in that it employs ductility (at both normal temperature and low temperature) for conventional bitumen specifications.

The PG grading system is the primary method for specifying bitumen in the USA and Canada. Here, individual regulators and specifiers determine the grades to be used on state roads. Other countries, such as China, use PG grading for premium binders and for some large construction projects such as airfields, motor sport circuits, heavy duty expressways or highways exposed to extreme weather conditions. However, other performance based systems are (at the time of writing, May 2014) under development in many countries.
The map in Figure A5.1 provides a summary of the type of specification systems used throughout the world.

Penetration/softening point used in:
- Europe
- Many parts of Asia
- Africa
- Argentina
- Brazil

Viscosity used in:
- Australia
- New Zealand
- Europe (soft grades)
- South America: Chile, Mexico and Argentina
- India

PG grading used in:
- USA and Canada
- Other locations: mainly for premium binders (e.g. China)

Figure A5.1 The type of specification systems used throughout the world

Table A5.1 lists the main bitumen product standards used in a selection of countries.

<table>
<thead>
<tr>
<th>Region/country</th>
<th>Regulatory body</th>
<th>Main product standards for bitumen used in asphalt</th>
</tr>
</thead>
<tbody>
<tr>
<td>Region/country</td>
<td>Regulatory body</td>
<td>Main product standards for bitumen used in asphalt</td>
</tr>
<tr>
<td>---------------------</td>
<td>----------------------------------------------------------------------------------</td>
<td>-----------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>USA and Canada</td>
<td>State department for transportation/local agency</td>
<td>AASHTO M 320-10. Standard specification for performance-graded asphalt binder. (American Association of State and Highway Transportation Officials, Washington, DC, USA)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>State and agency-specific standards</td>
</tr>
<tr>
<td></td>
<td>(Edited by) Research Institute of Highway Ministry of Transport</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Indian Roads Congress (IRC)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>IRC SP 53:2010. Guidelines on use of modified bitumen in road construction</td>
</tr>
<tr>
<td>Thailand</td>
<td>Department of Highways Standard</td>
<td>American Society for Testing and Materials (ASTM) and American Association of State and Highway Transportation Officials (AASHTO) for standard testing procedures</td>
</tr>
<tr>
<td></td>
<td>Specification for Highway Construction (DH-T)</td>
<td></td>
</tr>
<tr>
<td>Malaysia</td>
<td>Standards &amp; Industrial Research Institute of Malaysia (SIRIM)</td>
<td>MS 124:1996. Specification for penetration grade of bitumen for use in pavement construction</td>
</tr>
<tr>
<td></td>
<td>Jabatan Kerja Raya (JKR)</td>
<td>JKR/SPJ/2008-S4. Standard specification for road works</td>
</tr>
<tr>
<td>Philippines</td>
<td>Department of Public Works &amp; Highways (DPWH)</td>
<td>Standard specifications for highways, bridges and airports</td>
</tr>
</tbody>
</table>
Useful links for more information


Appendix 6
Asphalt product standards used across the world

A wide variety of asphalt product standards exist across different countries and regions and construction sectors. These are controlled, reviewed and amended by approved regulatory bodies.

Table A6.1 lists the main asphalt standards used in a selection of countries.

<table>
<thead>
<tr>
<th>Region/country</th>
<th>Regulatory body</th>
<th>Main standards for asphalt in road construction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Member countries of the EU</td>
<td>European Committee for Standardization (CEN)</td>
<td>EN 13108 series (there are several parts in the series specifying various types of asphalt)</td>
</tr>
<tr>
<td>USA and Canada</td>
<td>State department for transport/local agency</td>
<td>State and agency specific standards</td>
</tr>
<tr>
<td>India</td>
<td>Indian Roads Congress (IRC)</td>
<td>MORT&amp;H:05. Specifications for road and bridge works, 5th revision (2013)</td>
</tr>
<tr>
<td>Argentina</td>
<td>Direccion Nacional de Vialidad</td>
<td>Specifications for road construction (1998)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Asphalt Commission recommendations are applied for specific public and private contracts</td>
</tr>
<tr>
<td>Australia</td>
<td>Austroad Test Method and Specification</td>
<td>AGPT/T190 2014. Specification framework for polymer modified binders</td>
</tr>
<tr>
<td>Region/country</td>
<td>Regulatory body</td>
<td>Main standards for asphalt in road construction</td>
</tr>
<tr>
<td>----------------</td>
<td>-----------------------------------------------</td>
<td>---------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Malaysia</td>
<td>Standards &amp; Industrial Research Institute of Malaysia (SIRIM)</td>
<td>MS 124:1996. Specification for penetration grade of bitumen for use in pavement construction</td>
</tr>
<tr>
<td></td>
<td>Jabatan Kerja Raya (JKR)</td>
<td>JKR/SPJ/2008-S4. Standard specification for road works</td>
</tr>
<tr>
<td>Philippines</td>
<td>Department of Public Works &amp; Highways (DPWH)</td>
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Appendix 7
Reporting asphalt compositional analysis results

The composition of an asphalt is a basic factor that critically impacts its properties such as stiffness, fatigue resistance and durability etc. This explains why the choice of individual constituents and their proportions in the asphalt mixture design (see Chapter 12) is so important. Compositional analyses of asphalts are taken routinely by asphalt producers in order to verify their compliance with applicable standards. Clients often also carry out compositional analyses to give confidence that the asphalts used in their projects meet specified requirements.

The requirements of the asphalt composition (i.e. type and size distribution of the aggregates, and the type and content of the bitumen) have been covered in the relevant sections of this book. This appendix describes how the results of the asphalt compositional analysis are reported in different parts of the world.

In general, there are two aspects to asphalt compositional analysis: aggregate particle size distribution and bitumen content. The former, also known as the aggregate ‘grading’ or ‘gradation’, is typically determined by a sieve analysis that involves the use of a set of increasingly finer sieves to separate the aggregate into gradually smaller particle sizes. The mass of material retained by each sieve is then determined as a proportion of the total aggregate weight and reported as the cumulative quantity (by percentage) passing each sieve.

Bitumen content is most commonly reported as a percentage by mass of the total mixture but it can also be reported as a percentage by mass of the dry aggregate only. Both methods are acceptable provided that the method to be employed is clear, in order to avoid any misunderstanding. The choice of method depends largely on the road authorities or agencies concerned, and some allow both methods to be used. Table A7.1 indicates which method is used in different parts of the world.
The two percentages can be easily converted as follows

\[ P_{b(\text{agg})} = \frac{100 \times P_b}{100 - P_b} \]

or

\[ P_b = \frac{100 \times P_{b(\text{agg})}}{100 + P_{b(\text{agg})}} \]

where \( P_{b(\text{agg})} \) is the bitumen content as a percentage by mass of the dry aggregates, and \( P_b \) is the bitumen content as a percentage by mass of the mixture.

In a small number of specifications, it is required that the effective bitumen content is reported rather than the total bitumen content. This will involve additional effort to determine and deduct the binder quantity that is absorbed by the aggregates, in accordance with the following formula

\[ P_{be} = P_b - \left( \frac{P_{ba}}{100} \right) P_s \]

where \( P_{be} \) is the effective bitumen content as a percentage by mass of the total mixture, \( P_b \) is the bitumen content as a percentage by mass of the total mixture, \( P_{ba} \) is the absorbed bitumen as a percentage by mass of the dry aggregate, and \( P_s \) is the aggregate content as a percentage by mass of the total mixture.

---

**Table A7.1** Method of reporting bitumen content by country

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<th>Bitumen content reported as a percentage of the dry aggregate</th>
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<td>Malaysia</td>
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<td>Singapore</td>
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<td>Australia</td>
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In the asphalt mixture design process, the volumetric properties of the mixture and its components need to be determined. These, in turn, require the determination of the specific gravities of the constituent ingredients in addition to their masses. The percentage of bitumen by volume of mixture is given by

\[ P_{b\text{(vol)}} = \left( \frac{G_{mb}}{G_b} \right) P_b \]

where \( P_{b\text{(vol)}} \) is the bitumen content as a percentage by volume of the total mixture, \( P_b \) is the bitumen content as a percentage by mass of the total mixture, \( G_{mb} \) is the bulk specific gravity of the mixture, and \( G_b \) is the specific gravity of the bitumen.

However, for purposes of asphalt mixture design, the parameters that are commonly used are the volume of air voids, voids in the mineral aggregate and voids filled with bitumen (see Chapter 12). Hence, for the convenience of asphalt production control, the easier method of measuring and reporting the compositional analysis by the total mass is used.
# Appendix 8
## Volumetrics symbols

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<th>Description</th>
<th>Symbol</th>
<th>UK/EU*</th>
<th>Description</th>
<th>Symbol</th>
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<td></td>
<td>Volume of voids in mineral aggregate</td>
<td>$V_{ma}$</td>
<td></td>
<td>Volume of voids content in the mineral aggregate</td>
<td>VMA</td>
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<td></td>
<td>Bulk volume of compacted mix</td>
<td>$V_{mb}$</td>
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<tr>
<td></td>
<td>Voidless volume of paving mix</td>
<td>$V_{mnn}$</td>
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<td></td>
<td>Volume of voids filled with asphalt</td>
<td>$V_{fa}$</td>
<td></td>
<td>Volume of voids in the mineral aggregate filled with binder</td>
<td>VFB</td>
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<td></td>
<td>Volume of air voids</td>
<td>$V_a$</td>
<td></td>
<td>Volume of air voids</td>
<td>$V_m$</td>
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<tr>
<td></td>
<td>Volume of asphalt binder</td>
<td>$V_b$</td>
<td></td>
<td>Volume of binder</td>
<td>$V_{br}$</td>
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<td></td>
<td>Volume of absorbed asphalt binder</td>
<td>$V_{ba}$</td>
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<td></td>
<td>Volume of mineral aggregate (by bulk specific gravity)</td>
<td>$V_{sb}$</td>
<td></td>
<td>Volume of aggregate</td>
<td>$V_a$ or VIM</td>
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<td></td>
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<td>$V_{se}$</td>
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<td></td>
<td>Mass of air=0</td>
<td>$M_{air}$</td>
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* BS EN 12697-8:2003. Bituminous mixture – Test methods for hot mix asphalt – Part 8: Determination of air void characteristics of bituminous specimens. (This standard expresses volume as a percentage by volume.)
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Note: Page numbers in italics reference figures and tables separate from the corresponding text.

2L-PA see double layered porous asphalt
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AASHO see American Association of State Highway Officials
AASHTO see American Association of State Highway and Transportation Officials
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