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Dr Arthur Lyons, author of texts on building materials, was formerly head of quality, principal lecturer and teacher fellow for building materials at the Leicester School of Architecture, Faculty of Art and Design, De Montfort University, Leicester, UK. He was educated at Trinity Hall Cambridge, Warwick and Leicester Universities in the fields of natural sciences and polymer science, and has a postgraduate diploma in architectural building conservation. He was a lecturer in building materials within schools of architecture and surveying for 35 years. In recognition of his services to architects and architecture, Arthur Lyons was honoured with life membership of the Leicestershire and Rutland Society of Architects and he is a Fellow of the Higher Education Academy. He retains his active interest in architecture through liaison with the local society of architects and the Leicester School of Architecture of De Montfort University. In addition to this text, Arthur Lyons has written chapters in Metric Handbook—Planning and Design Data, 3rd edition, 2008, Architectural Press; ICE Manual of Construction Materials, 2009, Institution of Civil Engineers and Construction Materials Reference Book, 2010, Butterworth-Heinemann.
Materials for Architects and Builders is written as an introductory text to inform students at undergraduate degree and national diploma level of the relevant visual and physical properties of the widest range of building materials. The fourth edition has been significantly enhanced by the addition of more colour images, illustrating the materials and, in many cases, their use in buildings of architectural merit. The text embraces the broad environmental issues with sections on energy saving and recycled materials. An additional chapter on sustainability reflects the current debate on climate change and governmental action to reduce carbon emissions and ameliorate global warming. There are eighteen chapters covering the wide range of materials under standard headings. Each chapter describes the manufacture, salient properties and typical uses of the various materials, with the aim of ensuring their appropriate application within awareness of their ecological impact.

European Standards are taking over from the previous British Standards, and for most key materials, the European Norms have now been published. Generally, this has led to an increase in the number of relevant standards for building materials. However, in some cases, both the British and European Standards are current and are therefore included in the text and references.

New and rediscovered old materials, where they are becoming well integrated into standard building processes, are described, together with innovative products yet to receive general acceptance. Other materials no longer in use are generally disregarded, except where increased concern for environmental issues has created renewed interest. The use of chemical terminology is kept to the minimum required to understand each subject area, and is only significantly used within the context of the structure of plastics. Tabulated data is restricted to an informative level appropriate to student use. An extensive bibliography and listed sources of technical information are provided at the end of each chapter to facilitate direct reference where necessary.

The text is well illustrated with over 270 line drawings and colour photographs, showing the production, appearance and appropriate use of materials, but it is not intended to describe construction details as these are illustrated in the standard texts on building construction. Environmental concerns including energy-conscious design and the effects of fire are automatically considered as part of the broader understanding of the various materials.

The text is essential reading for honours and foundation degree, BTEC and advanced GNVQ students of architecture, building, surveying and construction, and those studying within the broad range of built environment subjects, who wish to understand the principles relating to the appropriate use of construction materials.

Arthur Lyons
January 2010
I acknowledge the support of the Leicester School of Architecture, Faculty of Art and Design, De Montfort University, Leicester, and Professor Brian Ford, School of the Built Environment, University of Nottingham. I wish to thank my wife, Susan, for her participation and support during the production of this work, and also my daughters Claire and Elizabeth for their constant encouragement. I am indebted to the numerous manufacturers of building materials for their trade literature and for permissions to reproduce their published data and diagrams. I am grateful to building owners, architectural practices and their photographers for the inclusion of the photographs; and to Her Majesty’s Stationery Office, the Building Research Establishment, the British Standards Institute and trade associations for the inclusion of their material.

I should like to thank the following organisations for giving permission to use illustrations:

Aircrete Products Association (Fig. 2.3); Angle Ring Company Ltd. (Fig. 5.10); Architectural Ceramics (Figs. 8.6, 8.9 and 8.10); British Cement Association (Figs. 3.5, 3.9 and 3.23); British Fenestration Rating Council (Fig. 7.17); British Standards Institute (Figs. 2.8 and 5.28). Permission to reproduce extracts from BS EN 771 Part 1: 2003 and BS 6915: 2001 is granted by BSI. British Standards can be obtained in PDF or hardcopy formats from the BSI online shop: www.bsigroup.com/shop or by contacting BSI Customer Services for hardcopies only: Tel: +44 (0) 20 8996 9001; Email: cservices@bsigroup.com; Building Research Establishment (Figs. 2.3, 4.15 and 9.18)—photographs from GBG 58, Digest 476 and IP 10/01, reproduced by permission of BRE; Baggeridge Brick Plc. (Figs. 1.22, 1.24 and 1.25); Construction Resources (Fig. 4.39); Copper Development Association (Figs. 5.23–5.25); Corus (Figs. 5.2, 5.4–5.6, 5.11, 5.13 and 5.16); EH Smith (Fig. 8.8); Glass Block Technology (Fig. 7.5); Hanson Brick Ltd. (Fig. 1.5); Ibstock Brick Ltd. (Figs. 1.3, 1.9, 1.10, 1.12, 1.15, 1.21 and 2.9); Imperial Chemical Industries Plc. (Fig. 15.4); James & Son Ltd. (Fig. 11.8); KME UK Ltd. (Fig. 5.25); Lead Contractors Association (Figs. 5.27 and 5.29); Lead Sheet Association (Fig. 5.26); Lignacite Ltd. (Fig. 2.7); Make Architects (Fig. 4.1); Marshalls Plc. (Fig. 2.14); Metal Cladding and Roofing Manufacturers Association (Fig. 5.15); Metra Non-ferrous Metals Ltd. and Rheinzinc (Fig. 5.31); Monodraught (Figs. 16.6 and 16.7); NCS — Natural Color System®—property of Scandinavian Colour Institute AB, Stockholm 2009. References to NCS® in this publication are used with permission of the Scandinavian Colour Institute AB (Fig. 15.3); Natural Stone Products Ltd. (Fig. 9.13); Pilkington Plc. (Figs. 7.6, 7.9, 7.11, 7.24 and 7.25)—images are reproduced by permission of Pilkington Plc; Pyrobel (Fig. 7.15); Ruberoid Building Products (Fig. 6.3); Securiglass Company Ltd. (Fig. 7.13); Smith of Derby (Fig. 11.2); Solar Century—www.solarcentury.com (Figs. 16.2 and 16.3); Stancliffe Stone (Figs. 9.3, 9.4 and 9.8); Steel Construction Institute (Figs. 5.7 and 5.12); Stone Federation of Great Britain (Fig. 9.6); TRADA Technology Ltd. (Figs. 4.15 and 4.20); Trent Concrete Ltd. (Figs. 1.23, 3.20, 3.21, 9.21, 11.5 and 11.6) and Zinc Development Association (Fig. 5.31).

The text uses the generic names for building materials and components wherever possible. However, in a few cases, products are so specific that registered trade names are necessarily used. In these cases the trade names are italicised in the text.
Specific information relating to the materials described in each chapter is given at the end of the appropriate section; however, the following are sources of general information relating to construction materials.

- Building Regulations 2000 and 2006, including current amendments and approved documents
- RIBA Office Library and Barbour Index
- Building Research Establishment (BRE) publications
- Trade association publications
- Trade exhibitions
- Trade literature
- Architecture and built environment journals
- British Board of Agrément certificates
- British Standards
- European Standards
- Eurocodes
- Institution of Civil Engineers—Manual of Construction Materials, 2009

European Standards (EN) have been published for a wide range of materials. A full European Standard, known in the UK as BS EN, is mandatory and overrules any conflicting previous British Standard which must be withdrawn. Prior to full publication, the draft European Standards are coded pr EN and are available for comment, but not implementation. BRE Information Paper IP 3/99 (1999) identifies the issues relating to the adoption in the UK of the structural Eurocodes.

The Building Research Establishment (BRE) publishes informative and authoritative material on a wide range of subjects relating to construction. Trade associations and manufacturers produce promotional literature and websites relating to their particular area of interest within the building industry. Architecture and building journals give news of innovations and illustrate their realisation in quality construction. Much literature has recently been presented, including from governmental organisations, in respect of the need to reduce energy consumption within the built environment sector to ameliorate the effects of global warming and climate change.

Information for this text has been obtained from a wide selection of sources to produce a student text with an overview of the production, nature and properties of the diverse range of building materials. New individual products and modifications to existing products frequently enter the market; some materials become unavailable. Detailed information and particularly current technical data relating to any specific product for specification purposes must therefore be obtained directly from the manufacturers or suppliers and cross-checked against current standards and regulations.
ABBREVIATIONS

General

AAC autoclaved aerated concrete
ABS acrylonitrile butadiene styrene
AC aggressive chemical (environment)
ACEC aggressive chemical environment for concrete
APAO atactic poly α-olefin
APM additional protective measures
APP atactic polypropylene
AR alkali-resistant
ASR alkali–silica reaction
BER building emission rate
BFRC British Fenestration Rating Council
BRE Building Research Establishment
BREEAM BRE environmental assessment method
BS British Standard
CAD computer-aided design
CCA chromated copper arsenate
CFCs chlorofluorocarbons
CG cellular glass
CIGS copper indium gallium selenide
CIS copper indium selenide
CL cross-laminated (timber)
CMYK cyan magenta yellow black
COSHH control of substances hazardous to health
CPE chlorinated polyethylene
CPVC chlorinated polyvinyl chloride
CS calcium silicate
CSA Canadian standards association
CSM chlorosulphonated polyethylene
DC design chemical (class)
DC direct current
DD draft for development
DER dwelling emission rate
DPC damp-proof course
DPM damp-proof membrane
DR (CR) dezincification-resistant
DRF durability of reaction to fire
DS design sulphate (class)
DSA design stage assessment
DZR dezincification-resistant
EN Euronorm
ENV Euronorm pre-standard
EP expanded perlite
EPC energy performance certificate
EPDM ethylene propylene diene monomer
EPR ethylene propylene rubber
EPS expanded polystyrene
ETFEX ethylene tetrafluoroethylene copolymer
EV exfoliated vermiculite
EVA ethylene vinyl acetate
EVOH ethyl vinyl alcohol copolymer
FEF flexible elastomeric foam
FPA flexible polypropylene alloy
FRP fibre reinforced polymer
FSC forest stewardship council
GGBS ground granulated blastfurnace slag
GRC glass-fibre reinforced cement
GRG glass-fibre reinforced gypsum
GRP glass-fibre reinforced plastic or polyester
GS general structural (timber)
HAC high alumina cement
HACC high alumina cement concrete
HB hardboard
HCFCs hydrochlorofluorocarbons
HD high density
HDPE high-density polythene
HIP home information pack
HL hydraulic lime
HLS hue lightness saturation
ICB insulating corkboard
ICF  insulating concrete formwork
IGU  insulating glass unit
LA  low alkali (cement)
LD  low density
LDPE  low-density polythene
LED  light-emitting diode
LRV  light reflectance value
LVL  laminated veneer lumber
MAF  movement accommodation factor
MB  mediumboard
MDF  medium density fibreboard
MF  melamine formaldehyde
MMC  modern methods of construction
MPa  mega Pascal
MTCS  Malaysian timber certification scheme
MW  mineral wool
NA  National Annex (British Standard)
NAOBL  National windspeed database
NCS  Natural Color System®
NHL  non-hydraulic lime
ODP  ozone depletion potential
OPC  ordinary Portland cement
OSB  oriented strand board
PAS  publicly available specification
PBAC  polystyrene-bead aggregate cement
PC  polycarbonate
PCM  phase change materials
PCR  post-construction review
PE  polyethylene
PEF  polyethylene foam
PEFC  programme for the endorsement of forest certification
PET  polyethylene terephthalate
PEX  crosslinked polyethylene
PF  phenolic foam/phenol formaldehyde
PFA  pulvrised fuel ash
PHA  partially halogenated alkane
PIB  polyisobutylene
PIR  polyisocyanurate foam
PMMA  polymethyl methacrylate
PP  polypropylene
pr EN  draft Euronorm
PS  polystyrene
PTFE  polytetrafluoroethylene
PUR  rigid polyurethane foam
PV  photovoltaic
PVA  polyvinyl acetate
PVB  polyvinyl butyral
PVC  polyvinyl chloride (plasticised)
PVC-U  polyvinyl chloride (unplasticised)
PVC-UE  extruded polyvinyl chloride
PVDF  polyvinylidene fluoride
RBDM  reinforced bitumen membrane
RGB  red green blue
SAP  standard assessment procedure
SB  softboard
SBEM  simplified building energy model
SBS  styrene butadiene styrene
SAP  standard assessment procedure
SCC  self-compacting concrete
SFI  sustainable forest initiative
Sg  specific gravity
SIP  structural insulated panel
SS  special structural (timber)
ST  standard (concrete mix)
SUDS  sustainable urban drainage system
T  tolerance (class)
TER  target emission rate
TFS  thin film silicon
TH  temperate hardwood
THF  tetrahydro furan
TMT  thermally modified timber
TPE  thermoplastic elastomer
TPO  thermoplastic polyolefin
TRADA  timber research and development association
TRM  total relative movement
UF  urea formaldehyde
UHPC  ultra high performance concrete
UV  ultra-violet (light)
VET  vinyl ethylene terpolymer
VOC  volatile organic compounds
WER  window energy rating
WF  wood fibre
WPC  wood plastic composite
WW  wood wool
XPS  extruded polystyrene

**Units**

- dB  decibel
- GPa  giga pascal (1 GPa = 1000 MPa)
- MPa  mega pascal (1 MPa = 1 N/mm²)
- μm  micron (10⁻⁶ m)
- nm  nanometre (10⁻⁹ m)

**Chemical symbols**

- Al  aluminium
- As  arsenic
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**Cement notation**

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<tr>
<td>C₃S</td>
<td>tricalcium silicate</td>
</tr>
<tr>
<td>C₃A</td>
<td>tricalcium aluminate</td>
</tr>
<tr>
<td>C₄AF</td>
<td>tetracalciumaluminoferite</td>
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Introduction

Originally, bricks were hand-moulded from moist clay and then sun-baked, as is still the practice in certain arid climates. The firing of clay bricks dates back well over 5000 years, and is now a sophisticated and highly controlled manufacturing process; yet the principle of burning clay, to convert it from its natural plastic state into a dimensionally stable, durable, low-maintenance ceramic material, remains unchanged.

The quarrying of clay and brick manufacture are high-energy processes, which involve the emission of considerable quantities of carbon dioxide and other pollutants including sulphur dioxide. The extraction of clay also has long-term environmental effects, although in some areas former clay pits have now been converted to bird sanctuaries or put to recreational use. However, well-constructed brickwork has a long life with low maintenance, and although the use of Portland cement mortar prevents the recycling of individual bricks, the crushed material is frequently recycled as aggregate in further construction.

The elegant cathedral at Evry near Paris (Fig. 1.1), designed by Mario Botta, illustrates the modern use of brickwork. The cathedral of Saint Corbinian, built with 670,000 bricks, was dedicated in 1997. The building exhibits fine detailing both internally and externally. Externally the cylindrical form rises to a circle of trees. Internally the altar is surmounted by a corbelled structure leading one’s view upwards to the central rooflight. Three-dimensional internal brickwork is finely detailed to generate the desired acoustic response.

Clay bricks

The wide range of clays suitable for brick making in the UK gives diversity to the products available. This variety is further increased by the effects of blending clays, the various forming processes, the application of surface finishes and the adjustment of firing conditions. Earlier this century most areas had their own brickworks with characteristic products; however, ease of road transportation and continuing amalgamations within the industry have left a reduced number of major producers and only a few small independent works. Most UK bricks are defined as high density (HD) fired-clay masonry units with a gross dry density greater than 1000 kg/m$^3$. The European standard (BS EN 771-1: 2003) also refers to low density (LD) fired-clay masonry units, and these blocks are described in Chapter 2.

The main constituents of brick-making clays are silica (sand) and alumina, but with varying quantities of chalk, lime, iron oxide and other minor constituents such as fireclay, according to their source. The largest UK manufacturer uses the Lower Oxford clays of Bedfordshire, Buckinghamshire and Cambridgeshire to produce the Fletton brick. This clay contains some carbonaceous content that reduces the amount of fuel required to burn the bricks, lowering cost and producing a rather porous structure. Other particularly characteristic bricks are the strongly coloured Staffordshire Blues and Accrington Reds from clays containing high iron content and the yellow London stocks from the Essex and Kent chalky clays with lower iron content.

The European standard (BS EN 771-1: 2003) also refers to low density (LD) fired-clay masonry units, and these blocks are described in Chapter 2.
SIZE

Within Europe, the dimensions of clay masonry units (BS EN 771-1: 2003) have not been standardised, but in the UK, the standard metric brick referred to in the National Annex (informative) to BS EN 771-1: 2003 is $215 \times 102.5 \times 65 \text{ mm}$, although this size is not a specified requirement. Dimensions are given in the order, length, width and height, respectively. These UK dimensions match those in BS 4729: 2005, which relates to special shapes and sizes of bricks. The standard brick weighs between 2 and 4 kg, and is easily held in one hand. The length (215 mm) is equal to twice its width (102.5 mm) plus one standard 10 mm joint and three times its height (65 mm) plus two standard joints (Fig. 1.2).

The building industry modular co-ordination system (BS 6750: 1986) is based on the module (M) of 100 mm and multimodules of 3M, 6M, 12M, 15M, 30M and 60M. For metric brickwork, the base unit is 3M or 300 mm. Thus four courses of 65 mm brickwork with joints give a vertical height of 300 mm, and four stretchers with joints co-ordinate to 900 mm.

Table 1.1 illustrates the two types of dimensional tolerance limits set for clay masonry units including the metric brick, which relate to the square root of the work size dimension. Measurements are based on a random sample of ten bricks. The calculation based on the use of the square root of work size ensures that the dimensional tolerance limits are appropriate for the wide range in size of clay masonry units used within the European Union (BS EN 771-1: 2003).

Tolerances

Mean value

Tolerance limits are set for the difference between the stated work size (e.g. 215, 102.5 and 65 mm) and the measured mean from the samples, for each of the three brick dimensions (length, width and height).
Fig. 1.2 Brick and co-ordinating sizes

These are categorised as T1, T2 and Tm where Tm is a tolerance quoted by the manufacturer.

| T1 | ±0.40 √(work size dimension) mm or 3 mm if greater |
| T2 | ±0.25 √(work size dimension) mm or 2 mm if greater |
| Tm | deviation in mm declared by the manufacturer |

**Range**
The maximum range of size for any dimension is designated by categories R1, R2 and Rm.

| R1 | 0.6 √(work size dimension) mm |
| R2 | 0.3 √(work size dimension) mm |
| Rm | range in mm declared by the manufacturer |

**Table 1.1 Tolerances on brick sizes**

<table>
<thead>
<tr>
<th>Brick (work) dimensions (mm)</th>
<th>Maximum deviation (±) of mean from declared work dimension (mm)</th>
<th>Maximum range of size within sample of ten bricks (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>T1</td>
<td>T2</td>
</tr>
<tr>
<td>Length</td>
<td>215</td>
<td>6</td>
</tr>
<tr>
<td>Width</td>
<td>102.5</td>
<td>4</td>
</tr>
<tr>
<td>Height</td>
<td>65</td>
<td>3</td>
</tr>
</tbody>
</table>

*Note: Limits for Tm and Rm are as declared by the manufacturer (these may be wider or closer than the other categories).*
There is no direct correlation between the limits on mean value (T) and those for the range (R); thus, a brick conforming to category T2 may be within the wider range R1. Category R2 bricks may only be required for very tight dimensional control, as in short runs of brickwork.

**Alternative sizes**

The metric standard evolved from the slightly larger Imperial sizes, which varied significantly, but typically were $9 \times 4\frac{3}{8} \times 2\frac{7}{8}$ in (229 $\times$ 112 $\times$ 73 mm) or $8\frac{5}{8} \times 4\frac{1}{8} \times 2\frac{5}{8}$ in (219 $\times$ 105 $\times$ 67 mm). Some manufacturers offer a range of bricks to full Imperial dimensions, alternatively to an appropriate height (e.g. 50, 68, 70, 73, 76 or 80 mm) for bonding in to Imperial brickwork for restoration and conservation work.

The 1970s also saw the introduction of metric modular bricks with co-ordination sizes of either 200 or 300 mm in length, 100 mm in width and either 75 or 100 mm in height. The popularity of these bricks has now declined but they did give the architect opportunities for increasing or reducing horizontal emphasis and scale within the context of traditional brickwork.

A recent development has been the production of longer bricks to offer enhanced horizontality to brick facades (Fig. 1.3). Lengths up to 440 mm are available in a range of colours and textures for normal, quarter or third bonding, alternatively stack bonding in non-load-bearing situations. Sizes include $240 \times 115 \times 50$ mm and $290, 327$ and $440 \times 102 \times 50$ and 65 mm.

**MANUFACTURE OF CLAY BRICKS**

There are five main processes in the manufacture of clay bricks:

- extraction of the raw material;
- forming processes;
- drying;
- firing;
- packaging and distribution.

**Extraction of the raw material**

The process begins with the extraction of the raw material from the quarry and its transportation to the works, by conveyor belt or road transport. Topsoil and unsuitable overburden are removed first and used for site reclamation after the usable clay is removed.
The raw material is screened to remove any rocks, then ground into fine powder by a series of crushers and rollers with further screening to remove any over-size particles. Small quantities of pigments or other clays may be blended in at this stage to produce various colour effects; for example, manganese dioxide will produce an almost black brick and fireclay gives a teak brown effect. Occasionally, coke breeze is added into the clay as a source of fuel for the firing process. Finally, depending on the subsequent brick forming process, up to 25% water may be added to give the required plasticity.

Forming processes

Handmade bricks
The handmade process involves the throwing of a suitably sized clot of wet clay into a wooden mould on a bench. The surplus clay is struck off with a framed wire and the green brick removed. The bricks produced are irregular in shape with soft arrises and interestingly folded surfaces. Two variations of the process are pallet moulding and slop moulding.

In pallet moulding, a stock board, the size of the bed face of the brick, is fixed to the bench. The mould fits loosely over the stock board, and is adjusted in height to give appropriate thickness to the green brick. The mould and board are sanded to ease removal of the green brick which, is produced with a frog or depression on one face. In the case of slop moulding, the stock mould is placed directly on the bench, and is usually wetted rather than sanded to allow removal of the green brick which, unlike the pallet moulded brick, is smooth on both bed faces (Fig. 1.4).

Soft mud process
The handmade process has now been largely automated, with the clay being mechanically thrown into pre-sanded moulds; the excess clay is then removed and the bricks released from the mould. These soft mud process bricks retain much of the individuality associated with true handmade bricks, but at a lower cost.

Pressed bricks
In the semi-dry process used for Fletton bricks the appropriate quantity of clay is subjected to a sequence of four pressings within steel moulds to produce the green brick. These bricks usually have a deep frog on one bed face. For facing bricks, texturing on both headers and one stretcher may be applied by a series of rollers. A water spray to moisten the surface, followed by a blast of a sand/pigment mixture, produces the sand-faced finish.

With clays that require a slightly higher water content for moulding, the stiff plastic process is used in which brick-size clots of clay are forced into the moulds. A single press is then required to form the
Extruding wire-cut bricks

Engineering bricks made by this process often have shallow frogs on both bed faces. In all cases the size of the mould is calculated to allow for the anticipated drying and firing shrinkage.

Extruded wire-cut bricks

In this process clay with a water content of up to 25% is fed into a screw extruder which consolidates the clay and extracts the air. The clay is forced through a die and forms a continuous column with dimensions equal to the length and width of a green brick (Fig. 1.5). The surface may then be textured or sanded, before the clay column is cut into brick units by a series of wires. The bed faces of wire-cut bricks often show the drag marks where the wires have cut through the extruded clay. Perforated wire-cut bricks are produced by the incorporation of rods or tines between the screw extruder and the die. The perforations save clay and allow for a more uniform drying and firing of the bricks without significant loss of strength. Thermal performance is not significantly improved by the incorporation of voids.

Drying

To prevent cracking and distortion during the firing process, green bricks produced from wet clays must be allowed to dry out and shrink. Shrinkage is typically 10% on each dimension depending upon the moisture content. The green bricks, laid in an open chequerwork pattern to ensure a uniform loss of moisture, are stacked in, or passed through, drying chambers which are warmed with the waste heat from the firing process. Drying temperatures and humidity levels are carefully controlled to ensure shrinkage without distortion.

Firing

Both intermittent and continuous kilns are used for firing bricks. The former is a batch process in which the single kiln is loaded, fired, cooled and unloaded. In continuous kilns, the firing process is always active; either the green bricks are moved through a fixed firing zone, or the fire is gradually moved around a series of interconnected chambers to the unfired bricks. Both continuous systems are more energy efficient than the intermittent processes. Generally, for large-scale production, the continuous tunnel kiln (Fig. 1.6) and the Hoffman kiln (Fig. 1.7) are used. Clamps and intermittent gas-fired kilns are used for the more specialised products. Depending on the composition of the clay and the nature of the desired product, firing temperatures are set to sinter or vitrify the clay. Colour variations called kiss-marks occur where bricks were in contact with each other within the kiln and are particularly noticeable on Flettons.

Tunnel kiln

In the tunnel kiln process the bricks are loaded 10 to 14 high on kiln cars which are moved progressively.

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Fig. 1.5 Extruding wire-cut bricks

Fig. 1.6 Tunnel kiln
through the preheating, firing and cooling zones. A carefully controlled temperature profile within the kiln and an appropriate kiln car speed ensure that the green bricks are correctly fired with the minimum use of fuel, usually natural gas. The maximum firing temperature within the range 940°C to 1200°C depends on the clay, but is normally around 1050°C, with an average kiln time of three days. The oxygen content within the atmosphere of the kiln will affect the colour of the brick products. Typically a high temperature and low oxygen content are used in the manufacture of blue bricks. A higher oxygen content will turn any iron oxide within the clay red.

**Hoffman kiln**
Introduced in 1858, the Hoffman kiln is a continuous kiln in which the fire is transferred around a series of chambers which can be interconnected by the opening of dampers. There may be 12, 16 or 24 chambers, although 16 is usual. The chambers are filled with typically 100,000 green bricks. The chambers in front of the fire, as it moves around, are preheated, and then firing takes place (960°C–1000°C), followed by cooling, unloading and resetting of the next load. The sequence moves on one chamber per day, with three days of burning. The usual fuel is natural gas, although low-grade coal and landfill methane are used by some manufacturers.

**Intermittent gas-fired kilns**
Intermittent gas-fired kilns are frequently used for firing smaller loads, particularly specials. In one system, green bricks are stacked onto a concrete base and a mobile kiln is lowered over the bricks for the firing process. The firing conditions can be accurately controlled to match those within continuous kilns.

**Clamps**
The basis of clamp firing is the inclusion of coke breeze into the clay, which then acts as the major source of energy during the firing process. In the traditional process, alternate layers of unfired bricks and additional coke breeze are stacked up and then sealed over with waste bricks and clay. The clamp is then ignited with kindling material and allowed to burn for two to five weeks. After firing, the bricks are hand selected because of their variability from under- to over-fired. Currently some handmade bricks are manufactured in gas-fired clamps which give a fully controlled firing process but still produce bricks with the characteristic dark patches on their surfaces due to the burnt breeze content.

**Packaging and distribution**
Damaged or cracked bricks are removed prior to packing. Most bricks are now banded and shrink-wrapped into packs of between 300 and 500, for easy transportation by forklift truck and specialist road vehicles. Special shapes are frequently shrink-wrapped onto wooden pallets.

**SPECIFICATION OF CLAY BRICKS**
To specify a particular brick it is necessary to define certain key criteria which relate to form, durability and appearance. The European Standard BS EN 771-1: 2003 requires an extensive minimum description for masonry units, including the European Standard number and date (e.g. BS EN 771-1: 2003), the type of unit (e.g. high density - HD), dimensions and tolerances from mean value, configuration (e.g. a solid or frogged brick), compressive strength and freeze/thaw resistance. Also, depending on the particular end use, additional description may be required. This may, as appropriate, include dry density, dimensional tolerance range, water absorption, thermal properties, active soluble salt content, moisture movement, reaction to fire and vapour permeability.

Within the building industry the classification usually also includes some traditional descriptions:

- place of origin and particular name (e.g. Staffordshire smooth blue);
- clay composition (e.g. Gault, Weald or Lower Oxford Clay, Etruria Marl, Keuper Marl [Mercian Mudstones] or shale);
- variety — typical use (e.g. Class A engineering, common or facing);
- type — form and manufacturing process (e.g. solid, frogged, wire cut);
- appearance — colour and surface texture (e.g. coral red rustic).

**Table 1.2  Properties of clay engineering bricks**

<table>
<thead>
<tr>
<th>Physical property</th>
<th>Clay engineering bricks</th>
</tr>
</thead>
<tbody>
<tr>
<td>Defined properties</td>
<td></td>
</tr>
<tr>
<td>Minimum compressive strength (MPa)</td>
<td>≥ 125</td>
</tr>
<tr>
<td>Maximum water absorption (% by mass)</td>
<td>&lt; 4.5</td>
</tr>
<tr>
<td>Freeze/thaw resistance class</td>
<td>F2</td>
</tr>
<tr>
<td>Active soluble salt content class</td>
<td>S2</td>
</tr>
</tbody>
</table>

Note:
The water absorption limits for all clay bricks used for damp-proof courses for buildings (DPC1) and external works (DPC2) are included in the table.

**Type**

Type refers to the form of the brick and defines whether it is solid, frogged, cellular, perforated or of a special shape (Figs. 1.8 and 1.9). Bricks may be frogged on one or both bed faces; perforations may be few and large or many and small. Cellular bricks have cavities closed at one end. Keyed bricks are used to give a good bond to plaster or cement rendering. Because of the wide range of variation within brick types, the manufacturer is required to give details of the orientation and percentage of perforations in all cases.

For maximum strength, weather resistance and sound insulation, bricks should be laid with the frogs uppermost so that they are completely filled with mortar; with double-frogged bricks the deeper frog should be uppermost. However, for cheapness, speed

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**Fig. 1.8 Brick types**
and possibly minimisation of the dead weight of construction, frogged bricks are frequently laid frog-down. Inevitably this leads to a resultant reduction in their load-bearing capacity.

**Standard specials**

Increasingly, *specials* (special shapes) are being used to enhance the architectural quality of brickwork. British
Fig. 1.9 (continued)
Standard BS 4729: 2005 illustrates the range of standard specials, which normally can be made to order to match standard bricks (Fig. 1.9).

Designation of standard specials:
- Angle and cant bricks
- Bullnose bricks
- Copings and cappings
- Plinth bricks
- Arch bricks
- Radial bricks
- Soldier bricks
- Cuboid bricks
- Bonding bricks
- Brick slips

Manufacturers also frequently make purpose-made specials (special specials) to the particular requirements of the architect or builder. Inevitably, delivery on specials takes longer than for ordinary bricks, and their separate firing frequently leads to some colour variation between the specials and the standard bricks, even where the clay used is identical. The more complex specials are handmade, usually in specially shaped stock moulds, although some can be made by modifying standard bricks before firing. The range of shapes includes copings and cappings (for parapets and freestanding walls), bullnose (for corner details, e.g. window and door reveals), plinths (for corbelling details and cills), cants (for turning angles), arches and brick slips (to mask reinforced concrete lintels etc.). Special bricks are also manufactured by cutting standard bricks, then, if necessary, bonding the pieces with epoxy resins. This has the advantage of ensuring an exact colour match to the standard bricks. Many brick slips, dog leg and arch voussoir sets (bricks to create an arch) are produced by this method. Specialist brick cutting companies also supply bricks for conservation projects as in the refurbishment of the elegant facade of St Pancras International train station in London.

APPEARANCE

The colour range of bricks manufactured in the UK is extensive. The colours range from the light buffs, greys and yellows through pastel pink to strong reds, blues, browns and deep blue/black, depending mainly on the clay and the firing conditions, but also on the addition of pigments to the clay or the application of a sand facing. Colours may be uniform, varied over the surface of individual bricks or varied from brick to brick. The brick forms vary from precise to those with rounded arrises; textures range from smooth and sanded to textured and deeply folded, depending on the forming process (Fig. 1.10).
Fig. 1.10 Typical range of clay brick textures and colours. *Photographs: Courtesy of Ibstock Brick Ltd.*
In view of the variability of bricks from batch to batch it is essential that they should be well mixed, preferably at the factory before palleting, or failing this, on site. If this is not done sufficiently, accidental colour banding will appear as the brickwork proceeds. Sand-faced bricks are liable to surface damage on handling, which exposes the underlying colour of the brick. Chipping of the arrises on bricks with through colour is visually less detrimental. Where rainwater run-off is an important factor, e.g. on cills and copings, smooth rather than heavily rusticated bricks should be used, as the latter would saturate and stain. Handmade bricks with deep surface folds (Fig. 1.11) should be laid frog up so that the creases or smiles tend to shed the rain-water from the face of the brickwork.

Glazed bricks, available in a wide range of intense colours, are sometimes used for their strong aesthetic effect (Fig. 1.12) or resistance to graffiti. They are either manufactured in a two-stage process, which involves the initial firing of the green brick to the biscuit stage, followed by the application of a slip glaze and a second firing. In the alternative one-stage process, the glaze is applied before a single firing. Manufacturers offer a standard range of colours or the option to specify from the RAL colour range. Bricks may be fully or partially glazed and special shapes are normally available to order.

The visual acceptability of facing bricks and the quality of the bricklaying would normally be assessed on site by the construction of a reference panel of at least 1m² to the standard PAS 70: 2003, using randomly selected bricks with examples of any colour banding, the proposed bonding, mortar and jointing. All subsequent brick deliveries and constructed brickwork should then be checked against the reference panel.

**DURABILITY**

**Frost resistance**

Bricks are classified into one of the three categories, F2, F1 and F0 according to their frost resistance within a standardised freezing test (Table 1.3). Only category F2 bricks are totally resistant to repeated freezing and thawing when in a saturated condition. Category F1 bricks are durable, except when subjected to repeated freezing and thawing under saturated

<table>
<thead>
<tr>
<th>Durability designation</th>
<th>Freeze/thaw resistance</th>
</tr>
</thead>
<tbody>
<tr>
<td>F2</td>
<td>Masonry subjected to severe exposure</td>
</tr>
<tr>
<td>F1</td>
<td>Masonry subjected to moderate exposure</td>
</tr>
<tr>
<td>F0</td>
<td>Masonry subjected to passive exposure</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Active soluble salts content</th>
</tr>
</thead>
<tbody>
<tr>
<td>S2</td>
</tr>
<tr>
<td>S1</td>
</tr>
<tr>
<td>S0</td>
</tr>
</tbody>
</table>

---

**Fig. 1.11** Handmade Hoskins Bromley Red bricks. *Photograph: Arthur Lyons*
conditions. Therefore, category F1 bricks should not be used in highly exposed situations such as below damp-proof courses, for parapets or brick-on-edge copings, but they are suitable for external walls which are protected from saturation by appropriate detailing. Category F0 bricks must only be used where they are subject to passive exposure, as when protected by cladding or used internally.

Fig. 1.12 Glazed bricks. Selection of colours and Atlantic House, London. Architects: Proun Architects. Photographs: Courtesy of Ibstock Brick Ltd.
Soluble salt content

The soluble salt content of bricks is defined by three categories: low (S2), normal (S1) and no limits (S0) (Table 1.3). Both the S2 and S1 categories have defined maximum limits for sodium/potassium and magnesium salt contents. The soluble salts derive from the original clay or from the products of combustion during the firing process. Soluble salts can cause efflorescence and soluble sulphates may migrate from the bricks into the mortar or any rendering, causing it to expand and deteriorate by sulphate attack. If used in an exposed situation, S1 and S0 category bricks should be bonded with sulphate-resisting cement mortar.

Efflorescence

Efflorescence sometimes appears as a white deposit on the surface of new brickwork (Fig. 1.13). It is caused by moisture carrying salts from inside the bricks and mortar to the surface where the water evaporates leaving the crystalline salts. Under most conditions it disappears without deleterious effect within one year. In exposed brickwork that is constantly subjected to a cycle of wetting and drying, efflorescence can occur at any time; further, a build-up and expansion of crystalline salts under the surface (crypto-efflorescence) may cause the face of the brickwork to crumble or spall.

Staining

The surface of brickwork may be stained by cement during the building process, or by lime leaching out of the fresh mortar (Fig. 1.14). In either case the excess should be brushed and washed off, without saturating the brickwork.

PHYSICAL PROPERTIES

Compressive strength

High density (HD) clay bricks are available with a range of compressive strengths from around 5 MPa to well over 100 MPa. The criteria for general use, damp-proof courses and engineering use are set out in Table 1.2 (page 8).

To determine the crushing strength of bricks, both bed faces are ground down until flat and parallel. The bricks are then crushed without filling the voids or frogs. Where frogs are to be laid upwards and filled in the construction, the crushing strength (MPa) is based on the net bearing area. Where frogs or voids are not to be filled, the crushing strength is based on the full gross area of the bed face.

Water absorption and suction

The level of water absorption is critical when bricks are to be used for damp-proof courses, or as engineering bricks. Appropriate limits are shown in Table 1.2,
although generally absorption ranges from 1% to 35%. Suction rates are now quoted by most brick manufacturers, as high values can adversely affect the bricklaying process. Bricks with high suction rates absorb water rapidly from the mortar, making it insufficiently plastic to allow for repositioning of the bricks as the work proceeds. Generally low or medium suction rates (1.0–2.0 kg/m² per min) are advantageous. In warm weather, high-suction-rate bricks may be wetted in clean water before laying, but any excess water will cause the brick to float on the mortar bed and also increase the risk of subsequent efflorescence and staining.

**Moisture and thermal movement**

After the firing process, bricks absorb moisture from the atmosphere and expand irreversibly, up to a maximum of 0.1%. It is therefore recommended that bricks should not be used for at least two weeks after firing (although it is now recognised that this irreversible process may continue at a decreasing rate for 20 years). Subsequent moisture and thermal movements are largely reversible, and movement joints allowing for a 1 mm movement per 1 m of brickwork should be allowed, typically at 10–12 m centres and at a maximum of 15 m, in restrained walls. Unrestrained or lightly restrained walls should have movement joints at 7–8 m centres. Horizontal movement joints should be at approximately 12 m intervals, as the vertical movement is of the same order as movement in the horizontal direction. In external unreinforced non-load-bearing walls a maximum spacing of 15 m between vertical movement joints is permitted (NA to BS EN 1996-2: 2006).

For many buildings the necessary movement joints can be made inconspicuous by careful detailing or featured as part of the design. Appropriate locations for movement joints would be where differing structural forms adjoin, such as abutments between walls and columns or where the height or thickness of a wall changes; alternatively, at design details such as brickwork returns, re-entrant corners or the recesses for downpipes. In expansion joints, fillers such as cellular polythene, polyurethane or foam rubber should be used, as these are easily compressible. Pointing should be with a flexible sealing compound such as two-part polysulphide.
Typical reversible moisture movement = 0.02%
Typical reversible thermal movement = 0.03%
Thermal movement = $5\times10^{-6}$ deg C $^{-1}$

**Thermal conductivity**

The thermal conductivity of brickwork is dependent on its density and moisture content, but generally clay bricks are poor thermal insulators. Brick manufacturers quote thermal conductivities at a standard 5% moisture content for exposed brickwork, and may also give the 1% moisture content figure for protected brickwork.

Using bricks with an average thermal conductivity of 0.96 W/m K, a typical partially filled cavity system is:

- 102.5 mm fairfaced brickwork
- 50 mm clear cavity
- 50 mm foil-faced rigid polyurethane insulation ($\lambda = 0.023$ W/m K)
- 100 mm lightweight blockwork ($\lambda = 0.15$ W/m K)
- 12.5 mm plasterboard on dabs

Giving a $U$-value of 0.27 W/m$^2$ K.

The thermal conductivity of clay bricks at 5% moisture content typically ranges from 0.65 to 1.95 W/m K.

**Fire resistance**

Clay brickwork generally offers excellent fire resistance by retaining its stability, integrity and insulating properties. The standard (BS 5628-3: 2005) indicates that 100 mm and 200 mm of load-bearing solid clay brick masonry will give 120 minutes and 360 minutes of fire resistance, respectively. Bricks with less than 1% organic material are automatically categorised as Euroclass A1 with respect to reaction to fire.

**Acoustic properties**

Good-quality brickwork is an effective barrier to airborne sound, provided that there are no voids through the mortar for the passage of sound. All masonry joints should be sealed and bricks laid with filled frogs to achieve the necessary mass per unit area and avoid air pathways.

At the junction between a cavity blockwork separating wall and an external brick and blockwork wall, if the external cavity is not fully filled with thermal insulation, then the separating wall cavity must be closed with a flexible cavity stop to reduce sound transmission sufficiently to comply with the Building Regulations Part E performance requirements.

Impact sound absorption by brickwork over the normal frequency range is fairly low and further decreased by the application of dense plaster or paint. However, the application of acoustic plasters or the addition of an independent panel of plasterboard backed by absorbent material improves impact sound insulation.

**QUALITY CONTROL**

To meet the consistent standards of quality required by clients, many brick manufacturers are now operating quality assurance systems. These require manufacturers to document all their operational procedures and set out standards to which products must adhere. Quality is controlled by a combination of an internal self-monitoring system and two to four independent spot-check reviews per year. Both the content of the technical literature and the products themselves are subjected to this scrutiny.

**Unfired clay bricks**

Unfired clay bricks for internal non-load-bearing applications are produced from clays which would be less suitable for standard fired bricks. Earth bricks require only low energy input for drying and have high potential recyclability. They have the advantage of inhibiting condensation and regulating the relative humidity of the internal environment. Earth bricks should be laid with clay or moderately hydraulic lime mortar and finished with a breathable finish such as clay or lime plaster, alternatively lime wash or permeable paint. Products include vertically perforated bricks with keyed or smooth finishes to sizes $220 \times 105 \times 67$ mm and $220 \times 105 \times 133$ mm (Fig. 1.15). Compressive strengths are in the range 2.9 to 3.8 MPa.

**Reclaimed clay bricks**

Reclaimed bricks are often selected for aesthetic reasons, but their appearance is not a guarantee of durability. In particular, frost resistance is uncertain
and it is not valid to test a sample to classify the whole consignment. Imperial sizes may vary considerably and some material may be contaminated with sulphates, or be liable to efflorescence from absorbed soluble salts. However, the strength and water absorption properties of reclaimed bricks are usually appropriate for domestic-scale construction. Reclaimed paving bricks that have previously been exposed to frost will normally be durable, but walling bricks may not be durable when used as pavers.

Recently some buildings have been constructed using lime mortar, specifically with a view to the potential recycling of the metric bricks at the end of the useful life of the construction. Lime mortar is significantly easier to clean from bricks than modern Portland cement mortar.

Of the estimated 2500 million bricks demolished each year in the UK only about 5% are reclaimed and nearly half are crushed and used as hardcore fill.

**Brickwork**

**CLAY BRICKWORK**

The bonding, mortar colour and joint profile have a significant visual effect on brickwork. The overall effect can be to emphasise as a feature or reduce to a minimum the impact of the bonding mortar on the bricks. Additionally, the use of polychromatic brickwork with complementary or contrasting colours for quoins, reveals, banding and even graphic designs can have a dramatic effect on the appearance of a building. The three-dimensional effects of decorative dentil courses and projecting corbelled features offer the designer further opportunities to exploit the effects of light and shade. Normally, a projection of 10–15 mm is sufficient for the visual effect without causing increased susceptibility to staining or frost damage. Curved brickwork constructed in stretcher bond shows faceting and the overhang effect, which is particularly accentuated in oblique light. With small-radius curvatures, the necessary change of bonding pattern to header bond can also be a visual feature, as an alternative to the use of curved-radius bricks.

The *Gothic Revival* exterior of the Queens Building, De Montfort University, Leicester (Fig. 1.16), illustrates the visual effects of polychromatic brickwork and voussoir specials. The energy-efficient building maximises use of natural lighting, heating and ventilation, using massive masonry walls to reduce peak temperatures. The mortar, which matches the external coral-red brickwork, reduces the visual impact of the individual bricks, giving the effect of planes rather than walls. This is relieved by the colour and shadow effects of the polychromatic and corbelled features, which are incorporated in the ventilation grilles and towers. The special bricks, cill details and banding are picked out in a deeper cadmium red and silver buff to contrast with the characteristic Leicestershire red-brick colouring.

**Mortars**

The mortar in brickwork is required to give a bearing for the bricks and act as a sealant between them. Mortars should be weaker than the individual bricks, to ensure that any subsequent movement does not cause visible cracking of the bricks, although too weak a mix would adversely affect durability of the brickwork. Mortar mixes are based on blends
of either cement/lime/sand, masonry cement/sand or cement/sand with plasticiser. Suitable alternatives to Portland cement are listed in Table 1.4 to PD 6678: 2005. When the mix is gauged by volume, an allowance has to be made for bulking of damp sand. The five mix classes are shown in Table 1.5. A typical 1:1:6 (cement : lime : sand) mix (M4—designation (iii)) would generally be appropriate and durable for low-rise construction, but for calculated structural brickwork or for increased resistance to frost in exposed situations a greater-strength mortar may be required. For designed mortars to BS EN 998-2: 2005, the specification is performance-based in relation to the compressive strength M in MPa at 28 days. For prescribed mortars a combination of the constituent proportions and the 28-day compressive strength is required. In the repointing of old brickwork it is particularly important to match the porosity of the brick to the water-retention characteristics of the mortar. This prevents excessive loss of water from the mortar before hydration occurs, which may then cause the pointing to crumble.

The use of lime mortar, as in the Building Research Establishment environmental building in Garston, Watford, will allow for the ultimate reuse of the bricks at the end of the building’s life cycle. The recycling of bricks is not possible, except as rubble, when strong Portland cement mortar is used.

Sands for mortars are normally graded to BS EN 13139: 2002 into categories designated by a pair of sieve sizes d/D which define the lower and upper size limits in mm, respectively. The majority of the particle size distribution should lie between the stated limits. The preferred grades are 0/1 mm, 0/2 mm, 0/4 mm, 0/8 mm, 2/4 mm and 2/8 mm. Typically between

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**Table 1.4** Appropriate cements for mortars (PD 6678: 2005)

<table>
<thead>
<tr>
<th>Cement</th>
<th>Standard</th>
<th>Cement notation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Portland cement</td>
<td>BS EN 197-1</td>
<td>CEM I</td>
</tr>
<tr>
<td>Sulphate-resisting Portland cement</td>
<td>BS 4027</td>
<td></td>
</tr>
<tr>
<td>Portland slag cement</td>
<td>BS EN 197-1</td>
<td>CEM II/A-S or II/B-S</td>
</tr>
<tr>
<td>Portland fly ash cement</td>
<td>BS EN 197-1</td>
<td>CEM II/A-V or II/B-V</td>
</tr>
<tr>
<td>Portland limestone cement</td>
<td>BS EN 197-1</td>
<td>CEM II/A–LL(L)</td>
</tr>
<tr>
<td>Masonry cement (inorganic filler other than lime)</td>
<td>BS EN 413-1</td>
<td>Class MC</td>
</tr>
<tr>
<td>Masonry cement (lime)</td>
<td>BS EN 413-1</td>
<td>Class MC</td>
</tr>
</tbody>
</table>
Mortar classes to BS EN 998-2: 2003 equate to compressive strengths at 28 days in MPa.
M1 class is listed in BS EN 998-2: 2003, but not in BS 5628-3: 2005 or NA to BS EN 1996-1-1: 2005.
Designations to BS 5628-3: 2005.

In all cavity brickwork, wall ties manufactured from galvanised steel, stainless steel or polypropylene to BS EN 845-1: 2003 should be incorporated (Fig. 1.18). They should be installed according to BS 5628-3: 2005 drip down and level or sloping down (≤25°) towards the outer leaf. Wall ties are categorised as Types 1 to 4 to BS 5628 according to their strength and appropriate use, and lengths are dependent on the cavity width (Table 1.6). The Building Regulations refer only to types A and B. Type A are the normal butterfly ties, and Type B are double triangle ties. Stainless steel ties are specified for all housing. An estimated minimum service life of 60 years is recommended by the publication DD 140-2: 1987. Low thermal conductivity wall ties are available, manufactured in pultruded composite basalt fibre and epoxy resin.

Where mortar bed-joints do not co-ordinate between masonry leaves, slope-tolerant cavity wall ties must be used. In partially filled cavities, the wall ties should clip the insulation cavity batts to the inner leaf. In all cases the cavity, insulation and ties should be kept clear of mortar droppings and other residues by using a protective board. Asymmetric wall ties are used for fixing masonry to timber or thin-joint aircrete blockwork. Movement-tolerant wall ties bend, or slide within a slot system fixed to one leaf of the masonry. Screw-in wall ties are available for tying new masonry to existing walls.

Coloured mortars
Mortar colour has a profound effect on the overall appearance of the brickwork, as with stretcher bond and a standard 10 mm joint, the mortar accounts for 17% of the brickwork surface area. A wide range of light-fast coloured mortars is available which can be used to match or contrast with the bricks, thus

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### Table 1.5 Mortar mix designations and classes

<table>
<thead>
<tr>
<th>Mortar designation</th>
<th>Mortar class</th>
<th>Cement:lime:sand</th>
<th>Masonry cement:sand</th>
<th>Cement:sand with plasticiser</th>
</tr>
</thead>
<tbody>
<tr>
<td>(i) M12</td>
<td>1:0:3</td>
<td></td>
<td>1:½:3</td>
<td></td>
</tr>
<tr>
<td>(iii) M4</td>
<td>1:1:5</td>
<td></td>
<td>1:1:6</td>
<td>1:4 – 1:5</td>
</tr>
<tr>
<td>(iv) M2</td>
<td>1:2:8</td>
<td></td>
<td>1:2:9</td>
<td>1:5½ – 1:6½</td>
</tr>
<tr>
<td>(v) M1</td>
<td>1:3:10</td>
<td></td>
<td>1:3:12</td>
<td>1:6½ – 1:7</td>
</tr>
</tbody>
</table>

Notes:
Mortar classes to BS EN 998-2: 2003 equate to compressive strengths at 28 days in MPa.
M1 class is listed in BS EN 998-2: 2003, but not in BS 5628-3: 2005 or NA to BS EN 1996-1-1: 2005.
Designations to BS 5628-3: 2005.

85 and 99% of the sand should pass through the larger sieve limit, and between 0 and 20% should pass through the smaller sieve size limit. The grades with more fines (63 micron or less) require more cement to achieve the same strength and durability as the equivalent mortars mixed with a lower fines content.

Ideally, brickwork should be designed to ensure the minimal cutting of bricks and built with a uniform joint width and vertical alignment of the joints (perpends). During construction, brickwork should be kept clean and protected from rain and frost. This reduces the risk of frost damage, patchiness and efflorescence. Brickwork may be rendered externally or plastered internally if sufficient mechanical key is provided by appropriate jointing or the use of keyed bricks. For repointing existing brickwork, it is necessary to match carefully the mortar sand and use lime mortar where it was used in the original construction.

Bonding

Figure 1.17 illustrates the effects of bonding. The stretcher bond is standard for cavity walls and normally a half-lap bond is used, but an increase in horizontal emphasis can be achieved by the less standard quarter or third bond. In conservation work it may be necessary to use half bricks (snap headers) to match the appearance of bonding in solid brick walls. For one-brick-thick walls more variations are possible; most typical are the English and Flemish bonds. The equivalent English and Flemish garden wall bonds, which have more stretchers, are primarily used for one-brick-thick walls where the reduced number of headers makes it easier to build both sides fairfaced. Panels of herringbone brickwork (raking bond), or dog tooth and dentil courses as in Victorian brickwork, can generate interesting features.
Fig. 1.17 Brick bonding

- Stretcher bond
- English bond
- Flemish bond
- English garden wall bond
- Flemish garden wall bond

Fig. 1.18 Wall ties

- Butterfly tie
- Double-triangle tie
- Pressed tie
- Wire tie
- Vertical-twist tie
- Spiral tie
- Insulation-retaining wall ties
highlighting the bricks as units or creating a unity within the brickwork. The coloured mortars contain inert pigments, which are factory-blended to a tight specification to ensure close colour matching between batches. Occasionally, black mortars may bloom due to lime migration to the surface. Coloured mortars can be used creatively to enhance the visual impact of the brickwork and even create designs on sections of otherwise monochromatic brickwork. The quantity of pigment should not exceed 10% by weight of the cement.

Mortar colours may also be modified by the use of stains after curing; however, such applications only penetrate 2 mm into the surface, and therefore tend to be used more for remedial work. Through-body colours are generally more durable than surface applications.

**Joint profiles**

The standard range of joint profiles is illustrated in Fig. 1.19. It is important that the main criterion should be the shedding of water to prevent excessive saturation of the masonry, which could then deteriorate. Normally the brickwork is jointed as the construction proceeds. This is the cheapest and best method as it gives the least disturbance to the mortar bed. Pointing involves the raking out of the green mortar to a depth of 13–20 mm, followed by refilling the joint with fresh mortar. This is only appropriate when the desired visual effect cannot be obtained directly by jointing; for example, when a complex pattern of coloured mortar joints is required for aesthetic reasons.

The square recessed (raked) joints articulate the brickwork by featuring the joint, but these should only be used with durable, (F2, S2) high-absorption bricks under sheltered conditions; furthermore, the recess should be limited to a maximum depth of 6 mm. The struck or weathered joint also accentuates the light and shade of the brickwork, while, as a tooled joint, offering good weather resistance in all grades of exposure. If the visual effect of the joint is to be diminished, the flush joint may be used, but the curved recessed (bucket handle) joint, which is compressed by tooling, offers better appearance and weathering properties. No mortar should be allowed to smear the brickwork, as it is difficult to remove subsequently without the use of dilute acid or pressure jets of water.

**Reinforced brickwork**

Reinforcement may be introduced vertically or horizontally into brickwork (Fig. 1.20). Bed-joint

### Table 1.6 Wall ties to BS 5628-1: 2005

<table>
<thead>
<tr>
<th>Types</th>
<th>Least leaf thickness (mm)</th>
<th>Nominal cavity width (mm)</th>
<th>Tie length (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Types 1, 2, 3 or 75</td>
<td>≤75</td>
<td>200</td>
<td></td>
</tr>
<tr>
<td>4 depending on</td>
<td>90</td>
<td>76–100</td>
<td>225</td>
</tr>
<tr>
<td>design loading and cavity width</td>
<td>90</td>
<td>101–125</td>
<td>250</td>
</tr>
<tr>
<td>90</td>
<td>126–150</td>
<td>275</td>
<td></td>
</tr>
<tr>
<td>90</td>
<td>151–175</td>
<td>300</td>
<td></td>
</tr>
<tr>
<td>90</td>
<td>176–300</td>
<td>≤50 mm embedding in each leaf</td>
<td></td>
</tr>
</tbody>
</table>

**Notes:**

Type 1—Masonry: Heavy duty. Suitable for most masonry construction except where large differential movements are anticipated.

Type 2—Masonry: General purpose. Suitable for construction up to 15 m above ground level, when built of double leaves of similar thicknesses in the range 90–150 mm. Maximum basic wind speed of 31 m/sec.

Type 3—Masonry: Basic. As Type 2 but for lower basic winds speeds of 25 m/sec maximum.

Type 4 — Masonry: Light duty. Suitable for dwellings up to 10 m in height, when built of double leaves of similar stiffness and thicknesses in the range 90–150 mm. Suitable for most internal separating cavity walls.

All Specifications are subject to verification by calculation.

A minimum embedded length of 50 mm is required in each leaf.

Wall ties in leaves of 90 mm or more should be placed at a minimum density of 2.5 ties per square metre. Thinner walls require increased provision.
reinforcement, usually austenitic stainless steel, should be completely surrounded by mortar with a minimum cover of 15 mm. For continuity in long walls, sections of reinforcement should be sufficiently end lapped. Vertical reinforcement is possible in the cavity or in pocket-type walls, where the void spaces are formed in the brickwork, then reinforcement and concrete is introduced after the masonry is completed. Care should be taken in the use of vibrators to compact the concrete within new masonry.

Decorative brickwork

Tile-bricks replicate the visual effect of hung tiles. A profiled edge on each brick overhangs the brick course below creating the tiled effect. The bricks are laid in 10 mm mortar which is cut back 15 mm from the front face, using only a 1 mm vertical joint so that the tiles appear to butt joint. A range of colours with smooth and sand-faced finishes is available with matching specials for corner returns. The brickwork must be completed in courses ahead of the blockwork to ensure that internal vertical joints are sealed. The tile-bricks are typically used for gable-ends on housing.

Profiled bricks (Fig. 1.21), which co-ordinate with standard brickwork, can create features with enhanced shadow effects. Shapes include sawtooth, spheres, pyramids and wave effects in a variety of colours and finishes to create feature panels or to be interspaced within standard brickwork.

Plaques, motifs, murals and sophisticated sculptures (Fig. 1.22) can be manufactured to individual designs both for new buildings and for the renovation or refurbishment of Victorian terracotta. The designs are carved as a bas-relief in soft solid through-colour brickwork or moulded in the unfired clay in relatively small units and joined on site with a matching mortar. For repetitive units, the clay is shaped in an appropriate wooden mould. Relief depths of 10–30 mm give shadow and contrast sufficient for most sculptural effects to be seen, although the viewing distances and angles must be considered. For large brickwork sculptures, the whole unit may be built in green bricks, with allowances made for the mortar joints and drying contraction. The design is then carved, numbered, dismantled, fired and reassembled on site.

Thin-bed masonry

The use of thin-bed masonry, with joints of between 2 and 6 mm, significantly reduces the visual effect of the mortar joints from 17% in 10 mm joint standard brickwork, to only 8% in 4 mm joints. This effect is further enhanced by the use of glue-mortars which
are applied to create a recessed joint. Thus the joint becomes only a shade line and the visual effect of the wall is totally determined by the colour and texture of the bricks. Because the glue-mortar is stronger than traditional mortar and has tensile properties, the brickwork patterns are not constrained to standard stretcher bonding. The glue-mortar is applied in two lines to both the horizontal and vertical joints, and therefore solid or perforated bricks rather than frogged bricks are most appropriate. Thin-bed masonry wall ties and special aramid bed-joint reinforcement are used as appropriate. The system offers the creative designer significant alternative aesthetic opportunities.

Preassembled brickwork

The use of preassembled brickwork supported on reinforced concrete or steel frames offers the builder a potentially higher level of quality control and increased speed of construction on site (Fig. 1.23). It also offers the scope to create complex details and forms, such as long low arches, that would be expensive or impossible in traditional brick construction. Specialist manufacturers produce large complete brick-clad precast concrete panels with either whole bricks or brick slips.
Typically the rear faces of brick slips are drilled at an angle, then stainless steel rods inserted and fixed with resin adhesive. The brick slips are laid out with spacers within the panel mould, prior to the addition of steel reinforcement and concrete. Finally the brick slips are pointed up giving the appearance of normal brickwork.

Factory-made brickwork panels of solid wall dimensions and single-storey height, incorporating openings and ducts for windows, doors and electrical installations, enable fast erection on site for domestic-scale construction. One manufacturer supplies bay window units constructed on steel frames for easy installation on site. The panels can either be pre-pointed or left unfinished for subsequent pointing on site.

**Brick cladding systems**

A significant revolution for brick-faced building has been the development of brick slip and brick tile cladding systems, designed to have the appearance and durability of traditional brickwork, but with a significantly reduced construction time. In one system, external walls are constructed with 215 mm aerated concrete blockwork and faced with an extruded polystyrene insulation panel to which 16 mm brick slips are applied onto the pre-formed grid, giving the appearance of standard external leaf brickwork. The polystyrene grid panels have an overlap to ensure horizontal joints are watertight and are tongued and grooved to interlock vertically. Adhesive is applied to the polystyrene and the brick slips are pushed into place with the appropriate horizontal spacing. Mortar is applied with either a pointing gun or a mortar bag and tooled to the required joint profile. With the use of highly insulating blocks, this type of construction can achieve $U$-values as low as 0.27 W/m$^2$.

An alternative system uses a plastic-coated galvanised-steel profile fixed to the structural wall (Fig. 1.24). The specially shaped brick tiles then clip into the steel system with appropriate vertical joint spacing. Mortar (typically a 1 : 1 : 6 mix) is applied with a pointing gun and smoothed off to the required profile, usually bucket-handle. A range of special tiles is manufactured to produce dados, plinths, cills and external returns, giving the appearance of traditional brickwork. Because the brickwork is non-structural, a range of bond patterns including stack, quarter and diagonal is optional. This type of pre-fabrication offers the potential for increased off-site construction work, and some manufacturers supply pre-formed brick-tile panels ready for fixing on site.

Another use for 20 mm brick slip systems is the refurbishment of poor-quality brickwork or previously
rendered blockwork, giving an enhancement to the aesthetic of fairfaced brickwork.

**CLAY BRICK PAVING**

Many clay brick manufacturers produce a range of plain and chamfered paving bricks together with a matching range of paver accessories. Bricks for flexible paving are frequently nibbed to set the spacing correctly. The material offers a human scale to large areas of hard landscape, especially if creative use is made of pattern and colour. Typical patterns (Fig. 1.25) include herringbone, running bond, stack bond, basket-weave and the use of borders and bands. However, it should be noted that not all paver sizes co-ordinate for herringbone and basket-weave designs. Profiled brick designs include decorative diamond and chocolate-bar patterns, and pedestrian-management texturing. The paving bricks may be laid on a hard base with mortar joints or alternatively on a flexible base with fine sand brushed between the pavers. Edge restraint is necessary to prevent lateral spread of the units.

The British Standard (BS EN 1344: 2002) stipulates minimum paver thicknesses of 40 mm and 30 mm for flexible and rigid construction, respectively. However, 50 mm pavers are generally used for flexible laying and 60 mm pavers are necessary when subjected to substantial vehicular traffic (BS 7533-1: 2001). Table 1.7 shows the standard sizes. Clay pavers are classified on the basis of freeze/thaw resistance. Pavers with designation FP0 are unsuitable for saturated freezing conditions, while pavers designated FP100 may be used under freeze/thaw conditions. The Standard BS EN 1344: 2002 classifies pavers into five categories (T0 to T4) of transverse breaking strength, with the lowest category T0 being appropriate only for rigid construction. Slip resistance for the unpolished pavers is categorised

<table>
<thead>
<tr>
<th>Table 1.7 Standard work sizes for pavers</th>
</tr>
</thead>
<tbody>
<tr>
<td>Length (mm)</td>
</tr>
<tr>
<td>215</td>
</tr>
<tr>
<td>215</td>
</tr>
<tr>
<td>210</td>
</tr>
<tr>
<td>210</td>
</tr>
<tr>
<td>200</td>
</tr>
<tr>
<td>200</td>
</tr>
</tbody>
</table>
Fig. 1.25 Typical range of clay pavers and hard landscape at Birmingham. Photographs: Courtesy of Baggeridge Brick Plc.
Fig. 1.25 (continued)
Calcium silicate bricks

Calcium silicate bricks, also known as sandlime or flintlime bricks, were first produced commercially in Germany in 1894, and then in the UK in 1905. Initially their use was confined to common brick applications, but in the 1950s, their durability for foundations was exploited. Research into mix design and the development of improved manufacturing processes subsequently led to the production of a full range of load-bearing-strength classes and attractive facings. Calcium silicate bricks are competitively priced and have about 3% of the UK brick market.

SIZE

The work size for calcium silicate bricks is $215 \times 102.5 \times 65$ mm, the same as for clay bricks, with a co-ordinating size of $225 \times 112.5 \times 75$ mm, allowing for 10 mm mortar joints. Generally, calcium silicate bricks are more accurate in form and size than fired clay bricks, which inevitably distort in the manufacturing process. The dimensional tolerances for calcium silicate bricks defined in the standard (BS EN 771-2: 2003) are generally $\pm 2$ mm on each dimension, except for thin-layer mortar when a maximum of only $\pm 1$ mm tolerance is permitted on the height. Actual tolerances are usually less than one millimetre. Modular and bricks of Imperial thickness can be made to order.

MANUFACTURE OF CALCIUM SILICATE BRICKS (SANDLIME AND FLINTLIME BRICKS)

The raw materials are silica sand (approximately 90%), hydrated lime, crushed flint, colouring pigments and water. (If quicklime is used, it is fully hydrated before the bricks are pressed, to prevent expansion under the steam treatment.) A mixture of sand, lime and water is used to manufacture the natural white sandlime brick. The addition of colouring pigments or crushed-flint aggregate to the standard components or the application of texturing to the brick surface gives the full product range of smooth, rustic and textured finishes.

The appropriately proportioned blend is pressed into brick units, stacked on bogies, moved into
the autoclave and subjected to steam pressure (0.8–1.3 MPa) for 4 to 15 hours at 180°C (Fig. 1.26). This causes the hydrated lime to react chemically with the surface of the sand particles, enveloping them with hydrated calcium silicates which fill much of the void spaces between the sand particles. Subsequently the calcium silicates react slowly with carbon dioxide from the atmosphere to produce calcium carbonate, with a gradual increase in the strength of the bricks.

APPEARANCE

The manufacturing process results in accurate shapes and dimensions, and with the untextured calcium silicate bricks, a smooth finish. The colour range is extensive, from white and pastel shades through to deep reds, blues, browns, greens and yellows. The visual effect on the brickwork tends to be that of precision. The bricks tend to be more brittle than clay bricks and are therefore more susceptible to damage on their arrises.

SPECIFICATION OF CALCIUM SILICATE BRICKS

Types

Both solid and frogged calcium silicate bricks are available. Manufacturers produce a wide range of matching specials to BS 4729: 2005; special specials to clients’ requirements and brick slips for facing reinforced concrete.

Durability

Calcium silicate bricks have good frost resistance, but should not be exposed repeatedly to strong salt solutions, acids or industrial effluent containing magnesium or ammonium sulphates. The bricks have a negligible salt content and therefore efflorescence, and sulphate attack on the mortar, cannot arise from within the bricks. The bricks are themselves resistant to sulphate attack and can therefore be used below ground with a suitable sulphate-resisting cement mortar. However, calcium silicate bricks should not be used as pavers where winter salting can be expected.

PHYSICAL PROPERTIES

Compressive strength

The British Standard BS EN 771-2: 2003 defines the range of compressive strength classes, as shown in Table 1.8.
Table 1.8  Minimum compressive strength for calcium silicate bricks

<table>
<thead>
<tr>
<th>Compressive strength class</th>
<th>Normalised compressive strength (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>5.0</td>
</tr>
<tr>
<td>7.5</td>
<td>7.5</td>
</tr>
<tr>
<td>10</td>
<td>10.0</td>
</tr>
<tr>
<td>15</td>
<td>15.0</td>
</tr>
<tr>
<td>20</td>
<td>20.0</td>
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<tr>
<td>25</td>
<td>25.0</td>
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<td>35</td>
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<tr>
<td>40</td>
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</tr>
<tr>
<td>45</td>
<td>45.0</td>
</tr>
<tr>
<td>50</td>
<td>50.0</td>
</tr>
<tr>
<td>60</td>
<td>60.0</td>
</tr>
<tr>
<td>75</td>
<td>75.0</td>
</tr>
</tbody>
</table>

**Weight**

Most standard calcium silicate bricks weigh between 2.4 and 3.0 kg, but densities can range from below 500 to above 2200 kg/m³.

**Water absorption**

Water absorption is usually in the range 8–15% by weight.

**Moisture and thermal movement**

Unlike clay bricks, which expand after firing, calcium silicate bricks contract. This shrinkage is increased if the bricks become wet before use; therefore, site protection of brick stacks from saturation is essential. Similarly, unfinished brickwork should be protected from both saturation and freezing during construction. Reversible moisture movement is greater for calcium silicate bricks than for clay bricks, so expansion joints must be provided at intervals between 7.5 and 9.0 m. Such movement joints should not be bridged by rigid materials. Generally a weak mortar mix should be used (e.g. 1 : 2 : 9 cement : lime : sand), except below damp-proof course level (DPC) and for copings, to prevent visible cracking of either the mortar or the bricks.

Typical reversible moisture movement = ±0.05%
Typical reversible thermal movement = ±0.05%
Thermal movement = \(8–14 \times 10^{-6} \text{ deg C}^{-1}\)

**Thermal conductivity**

The thermal conductivities are equivalent to those of clay bricks of similar densities.

The thermal conductivity of calcium silicate brick ranges from 0.6 W/m K (Class 20) to 1.3 W/m K (Class 40).

**Fire resistance**

The fire resistance of calcium silicate bricks is similar to that of clay bricks, with solid 100 mm calcium silicate brickwork giving 120 minutes and 200 mm giving 360 minutes’ fire resistance, according to BS 5628-3: 2005. The standard illustrates only marginal differences in fire resistance between calcium silicate and clay bricks. Calcium silicate bricks (with less than 1% organic material) are designated Euroclass A1 with respect to reaction to fire.

**Acoustic properties**

Acoustic properties are related to mass and are therefore the same as for clay bricks of equivalent density.

**CALCIUM SILICATE BRICKWORK**

Most design considerations are the same for either clay or calcium silicate brick. However, calcium silicate bricks are particularly popular for their light-reflecting properties, for example in light wells or atria. Their smooth crisp appearance with a non-abrasive surface is particularly appropriate for some interior finishes and also forms an appropriate base for painted finishes. The use of complementary coloured mortars enhances the aesthetic effect when using strongly coloured bricks. Their dimensional accuracy gives some advantage in the bricklaying process, and cost is comparable to that of the equivalent clay bricks.

The interior of the Queens Building of De Montfort University, Leicester (Fig. 1.27), illustrates the effective use of calcium silicate brickwork in creating a light internal space. Incorporated within the ivory Flemish-bond brickwork are restrained bands of polychromatic features and robust articulation of obtuse-angle quoins. The accuracy of the brickwork emphasises the clarity of the internal form, reflecting the disciplines of engineering that the building houses.
Fig. 1.27 Polychromatic calcium silicate brickwork—Queens Building, De Montfort University, Leicester. Architects: Short Ford & Associates. Photographs: Lens-based media — De Montfort University
Concrete bricks

Developments in the use of iron oxide pigments have produced a wide range of colour-stable quality concrete-brick products. Currently concrete bricks are competitively priced and hold approximately 10% of the total brick market share.

SIZE

The British Standard BS 6073-2: 2008, which gives guidance on the application of the European Standard BS EN 771-3: 2003, does not refer to standard concrete sizes for bricks, but only aggregate concrete coursing units as set out in Table 1.9. However, the National Annex NA (informative) of the current BS EN 771-3: 2003 standard still refers to UK work sizes of bricks including the unit 215 × 103 × 65 mm as for clay bricks.

Due to their manufacturing process, concrete bricks can be made to close tolerances, so accurate alignment is easy to achieve on site. Half-brick walls can readily be built fairfaced on both sides.

MANUFACTURE OF CONCRETE BRICKS

Concrete bricks are manufactured from blended dense aggregates (e.g. crushed limestone and sand) together with cement under high pressure in steel moulds. Up to 8% of appropriately blended iron oxide pigments, depending on the tone and depth of colour required, are added to coat the cement particles which will then form the solid matrix with the aggregate. The use of coloured aggregates also increases the colour range. The accurate manufacturing process produces bricks that have clean arrises.

Table 1.9 Standard work sizes for aggregate concrete coursing units (BS 6073-2: 2008)

<table>
<thead>
<tr>
<th>Length (mm)</th>
<th>Width (mm)</th>
<th>Height (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>190</td>
<td>90</td>
<td>65</td>
</tr>
<tr>
<td>190</td>
<td>90</td>
<td>90</td>
</tr>
<tr>
<td>215</td>
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<td>65</td>
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<tr>
<td>290</td>
<td>90</td>
<td>90</td>
</tr>
<tr>
<td>440</td>
<td>90</td>
<td>65</td>
</tr>
<tr>
<td>440</td>
<td>100</td>
<td>65</td>
</tr>
<tr>
<td>440</td>
<td>90</td>
<td>140</td>
</tr>
<tr>
<td>440</td>
<td>100</td>
<td>140</td>
</tr>
</tbody>
</table>

APPEARANCE

A wide range of colours, including multicolours, is available, from red, buff and yellow to green and black. Surfaces range from smooth to simulated natural stone, including those characteristic of handmade and textured clay bricks. Because of the wide range of pigments used in the manufacturing process, it is possible to match effectively new concrete bricks to old and weathered clay bricks for the refurbishment or extension of old buildings.

SPECIFICATION OF CONCRETE BRICKS

Types

Concrete bricks may be solid, perforated or frogged, also common or facing according to the manufacturer. 'Facing masonry units' may or may not be left exposed externally, while 'exposed masonry units' offer visual qualities without further protection. Compressive strengths and densities are manufactured to client requirements. A normal range of specials to BS 4729: 2005 is produced, although as with clay and calcium silicate bricks, a longer delivery time must be anticipated. The manufacturer’s reference, the crushing strength, the dimensions and the brick type must be clearly identified with each package of concrete bricks. High-compressive-strength concrete units should be used below ground where significant sulphate levels are present according to the classification given in the BRE Special Digest 1, Concrete in aggressive ground (2005).

DURABILITY

Concrete bricks are resistant to frost and are therefore usable in all normal levels of exposure. Like all concrete products, they harden and increase in strength with age. As with calcium silicate bricks, they can be made free of soluble salts and thus free from efflorescence. Concrete bricks should not be used where industrial effluents or acids are present.

PHYSICAL PROPERTIES

Weight and compressive strength

The standard brick weighs typically 3.2 kg depending on its composition. The compressive strengths of concrete masonry units are specified according to Table 1.10, within the range 2.9 to 40.0 MPa (BS 6073-2: 2008).
Table 1.10  Typical strengths of concrete units (BS 6073-2: 2008)

| Compressive strength (MPa) | 2.9 | 3.6 | 7.3 | 8.7 | 10.4 | 17.5 | 22.5 | 30.0 | 40.0 |

Notes:
Strengths 10.4 to 40.0 MPa inclusive apply only to aggregate concrete units and not aircrete.

Water absorption

Water absorption is typically 8%, but engineering-quality bricks average less than 7% after 24 hours’ cold immersion, and are suitable for aggressive conditions such as retaining walls, below damp-proof course level and for inspection chambers.

Moisture and thermal movement

Concrete bricks have a typical drying shrinkage of 0.04%, with a maximum of 0.06%. Moisture and thermal movements are greater than those for calcium silicate bricks and movement joints should be at 5–6 m centres. Because of their moisture movement, prior to laying, concrete bricks should not be wetted to overcome excessive suction, but the water retentivity of the mortar should be adjusted accordingly. Brick stacks should be protected on site from rain, frost and snow.

Thermal conductivity

The thermal conductivities of concrete bricks are equivalent to those of clay and calcium silicate bricks of similar densities. Partially filled cavities, maintaining a clear cavity, are recommended to prevent water penetration to the inner leaf.

The thermal conductivity of concrete bricks ranges between 1.4 and 1.8 W/m K.

An appropriate level of thermal insulation for external walls can be achieved using concrete brickwork. A typical partial cavity fill system is:

102.5 mm concrete facing brick
50 mm clear cavity airspace
45 mm foil-faced rigid polyurethane insulation ($\lambda = 0.023$ W/m K)
115 mm high-performance lightweight blockwork ($\lambda = 0.11$ W/m K)
12.5 mm plasterboard on dabs

giving a $U$-value of approximately 0.27 W/m$^2$ K depending on the thermal conductivity of the concrete bricks used.

Fire resistance

The fire resistance of concrete bricks is of the same order as that of clay and calcium silicate bricks. Concrete bricks (with less than 1% organic material) are designated Euroclass A1 with respect to reaction to fire.

Acoustic properties

Dense concrete bricks are suitable for the reduction of airborne sound transmission. On a weight basis, they are equivalent to clay and calcium silicate bricks.

CONCRETE BRICKWORK

With the wide range of colour and texture options offered by concrete-brick manufacturers, it is frequently difficult to distinguish visually, except at close quarters, between concrete and clay brickwork. The visual effects of using coloured mortars and various jointing details are as for clay bricks, but for exposed situations the use of raked joints is not recommended.

Mortarless brick system

The non-structural cladding system is based on interlocking S-section concrete bricks, with appropriate special units for 45$^\circ$ and 90$^\circ$ internal and external angles also a PVC-U section as a base starter track. The visual effect is that of standard brickwork without the mortar joints. The system is suitable for cladding steel or timber frame buildings up to 18 m in height. It may also be used on conventional masonry or insulated concrete formwork (ICF) construction. The system has resistance to moisture penetration but is not airtight. A range of colours from buff, beige and brown through to red is available. The horizontal joints are slightly chamfered to give an impression of standard brickwork coursing.
References

FURTHER READING


STANDARDS


BS 4729: 2005 Clay and calcium silicate bricks of special shapes and sizes. Recommendations.

BS 5628 Code of practice for use of masonry:


Part 2: 2005 Structural use of reinforced and prestressed masonry.


BS 6073 Precast concrete masonry units:


BS 6100 Building and civil engineering vocabulary:

Part 0: 2002 Introduction.


BS 6515: 1984 Specification for polythene damp-proof courses for masonry.


BS 7533 Pavements constructed of clay, natural stone or concrete pavers.

BS 8000 Workmanship on building sites:


BS 8103 Structural design of low rise buildings:


BS 8208 Guide to assessment of suitability of external cavity walls for filling with thermal insulation:

Part 1: 1985 Existing traditional cavity construction.


BS EN 771 Specification for masonry units:


Part 2: 2003 Calcium silicate masonry units.


Part 4: 2003 Autoclaved aerated concrete masonry units.

BS EN 772 Methods of test for masonry units:


BS EN 845 Specification for ancillary components for masonry:

Part 1: 2003 Ties, tension straps, hangers and brackets.


BS EN 934-3: 2003 Admixtures for masonry mortar.


BS EN 1015 Methods of test for mortar for masonry.

BS EN 1015 Methods of test of masonry.

36 MATERIAls FOR ARCHITECTS AND BUILDERS

BS EN 1365-1: 1999 Fire resistance tests for loadbearing elements. Walls.
BS EN 1996 Eurocode 6: Design of masonry structures:
  Part 1.2: 2005 Structural fire design.
BS EN 13139: 2002 Aggregates for mortar.
PAS 70: 2003 HD clay bricks. Guide to appearance and site measured dimensions and tolerance.

BUILDING RESEARCH ESTABLISHMENT PUBLICATIONS

BRE Digests

BRE Digest 329: 2000 Installing wall ties in existing construction.
BRE Digest 441: 1999 Clay bricks and clay brick masonry (Parts 1 and 2).
BRE Digest 460: 2001 Bricks, blocks and masonry made from aggregate concrete (Parts 1 and 2).
BRE Digest 461: 2001 Corrosion of metal components in walls.

BRE Special digests

BRE SD1: 2005 Concrete in aggressive ground.
BRE SD4: 2007 Masonry walls and beam and block floors. U-values and building regulations.

BRE Good building guides

BRE GBG 62: 2004 Retro-installation of bed joint reinforcement in masonry.

BRE Information papers

BRE IP 10/93 Avoiding latent mortar defects in masonry.
BRE IP 10/99 Cleaning exterior masonry.
BRE IP 11/00 Ties for masonry walls. A decade of development.
BRE IP 3/01 Dynamic stiffness of wall ties used in masonry cavity walls. Measurement procedure.

BRICK DEVELOPMENT ASSOCIATION PUBLICATIONS

Design notes


Building note


Technical information paper


TeChniCal paper

Lilley, A.A., Flexible brick paving: application & design 1990: Highways & Transportation 10(37).

ADVIsory ORGANISATIONS

Brick Development Association, The Building Centre, 26 Store Street, London WC1E 7BT, UK (0207 323 7030).
Mortar Industry Association, Gillingham House, 38-44 Gillingham Street, London SW1V 1HU, UK (020 7963 8000).
Introduction

The variety of commercially available concrete blocks is extensive, from dense through to lightweight, offering a range of load-bearing strength, sound and thermal insulation properties. Where visual blockwork is required, either internally or externally, fairfaced blocks offer a selection of textures and colours at a different visual scale compared to that associated with traditional brickwork. Externally, visual concrete blockwork weathers well, provided adequate attention is given to the quality of the material and rainwater run-off detailing. Blockwork has considerable economic advantages over brickwork in respect of speed of construction, particularly as the lightweight blocks can be lifted in one hand.

Whilst clay blocks are used extensively for masonry construction on the continent of Europe, until recently there had been little demand from the building industry within the UK. However, both fired and unfired-clay blocks are now commercially available within the UK. Gypsum blocks may be used for internal non-load-bearing partitions and the internal insulation of walls. Inverted reinforced concrete T-beams with concrete or clay block infill is a standard form of domestic-scale floor construction.

Concrete paving blocks, which offer opportunities for creative hard landscaping with their diversity of form and colour, are widely used for town pedestrian precincts and individual house driveways. Concrete interlocking blocks with planting are used to create environmental walls.

Concrete blocks

TYPES AND SIZES

Concrete blocks are defined as solid, cellular or hollow, as illustrated in Fig. 2.1.

Concrete blocks are manufactured to various work face dimensions in an extensive range of widths, offering a wide choice of load-bearing capacity and level of insulation. Manufacturers work size dimensions should be indicated as length, width and height, in that order, to BS EN 771-3: 2003 and BS 6073-2: 2008. However, the National Annex NA (informative) to BS EN 771-3: 2003 and the British Standard particularly note that previous standard UK practice was to specify blocks by length, height and thickness.

The standard work face size, which co-ordinates to three courses of metric brickwork allowing for 10 mm mortar joints, is 440 × 215 mm (Fig. 2.2), but the other sizes in Table 2.1 are marketed for aesthetic and constructional reasons. For example, narrow bands of a different colour may be used as visual features within fairfaced blockwork, and foundation wall blocks are normally laid flat. The use of thin-joint masonry offers speedier construction, especially when using large format blocks (Fig. 2.3), which are approximately equivalent in size to two standard units. However, blocks heavier than 20 kg should not be lifted by a single person as this potentially can lead to injury. Within the 20 kg limit are 100 mm wide aircrrete blocks with face dimensions 610 × 375 mm for speedy construction using the thin-joint system.
The European Standard (BS EN 771-3: 2003) describes a wide range of aggregate concrete masonry units incorporating either dense or lightweight aggregates. Under the European Standard, the minimum description for concrete blocks includes the European Standard number and date (e.g. BS EN 771-3: 2003), the type of unit (e.g. common or facing), work size dimensions and tolerance category, configuration (e.g. solid or with voids) and compressive strength. Also, depending on the particular end use, additional description may be required. This may, as appropriate, include surface finish, net and gross dry density, co-ordinating size, thermal properties and moisture movement. Tolerance limits for regular-shaped blocks are defined at four levels in Table 2.2. Compressive strengths of concrete masonry units are classified as Category I or Category II. Category I units have a tighter control with only a 5% risk of the units not achieving the declared compressive strength.

The European Standard (BS EN 771-4: 2003) gives the specification for autoclaved aerated concrete (AAC) masonry units. The maximum size of units within the standard is 1500 mm length × 600 mm width × 1000 mm height. The tolerance limits on the dimensions are defined in Table 2.3, and are dependent on whether the units are to be erected with standard or thin-layer mortar joints. The standard manufacturer’s description for AAC masonry units must include the European Standard number and date (e.g. BS EN 771-4: 2003), dimensions and tolerances, compressive strength (Category I or Category II, as for concrete.
Table 2.1  Typical work sizes and strengths for concrete blocks to BS 6073: 2008

<table>
<thead>
<tr>
<th>Length (mm)</th>
<th>Width (mm)</th>
<th>Height (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aggregate concrete blocks Coursing blocks</td>
<td></td>
<td></td>
</tr>
<tr>
<td>190</td>
<td>90</td>
<td>65</td>
</tr>
<tr>
<td>190</td>
<td>90</td>
<td>90</td>
</tr>
<tr>
<td>215</td>
<td>100</td>
<td>65</td>
</tr>
<tr>
<td>290</td>
<td>90</td>
<td>90</td>
</tr>
<tr>
<td>440</td>
<td>90</td>
<td>65</td>
</tr>
<tr>
<td>440</td>
<td>100</td>
<td>65</td>
</tr>
<tr>
<td>440</td>
<td>90</td>
<td>140</td>
</tr>
<tr>
<td>440</td>
<td>100</td>
<td>140</td>
</tr>
<tr>
<td>Standard blocks</td>
<td></td>
<td></td>
</tr>
<tr>
<td>390</td>
<td>90</td>
<td>190</td>
</tr>
<tr>
<td>390</td>
<td>100</td>
<td>190</td>
</tr>
<tr>
<td>390</td>
<td>140</td>
<td>190</td>
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<tr>
<td>390</td>
<td>190</td>
<td>190</td>
</tr>
<tr>
<td>440</td>
<td>75</td>
<td>215</td>
</tr>
<tr>
<td>440</td>
<td>90</td>
<td>215</td>
</tr>
<tr>
<td>440</td>
<td>100</td>
<td>215</td>
</tr>
<tr>
<td>440</td>
<td>140</td>
<td>215</td>
</tr>
<tr>
<td>440</td>
<td>190</td>
<td>215</td>
</tr>
<tr>
<td>440</td>
<td>215</td>
<td>215</td>
</tr>
<tr>
<td>Aircrete concrete blocks Coursing blocks</td>
<td></td>
<td></td>
</tr>
<tr>
<td>215</td>
<td>90–150</td>
<td>65</td>
</tr>
<tr>
<td>215</td>
<td>90–150</td>
<td>70</td>
</tr>
<tr>
<td>Standard blocks</td>
<td></td>
<td></td>
</tr>
<tr>
<td>440</td>
<td>50–350</td>
<td>215</td>
</tr>
<tr>
<td>610</td>
<td>50–350</td>
<td>215</td>
</tr>
<tr>
<td>620</td>
<td>50–350</td>
<td>215</td>
</tr>
</tbody>
</table>

Notes:
Not all sizes are produced by all manufacturers but other sizes may be available.
Widths quoted by certain manufacturers include: 50, 70, 75, 100, 115, 125, 130, 140, 150, 190, 200, 215, 255, 265, 275, 300 and 355 mm.
Foundation blocks with face sizes of typically 255 x 290, 255 x 300, 310 x 350, 400 x 215, 440 x 140 or 440 x 215 mm are laid flat.
Beam and block floor rectangular units are usually 100 mm thick, with face dimensions of 440 x 215, 440 x 350, 440 x 540, 610 x 350 or 620 x 215 mm.
Common crushing strengths to BS 6073-2: 2008 are 2.9, 3.6, 7.3, 8.7, 10.4, 17.5, 22.5, 30.0 and 40.0 MPa, but some manufacturers supply additional intermediate strengths (e.g. 4.2 MPa).

Fig. 2.3 Thin-joint masonry using large format blocks. Photograph reproduced from GBG 58 by permission of BRE and courtesy of Aircrete Products Association

units) and dry density. Further description for specific purposes may include durability, configuration (e.g. perforations or tongued and grooved jointing system) and intended use.

Table 2.2  Limit of tolerances on aggregate concrete block sizes

<table>
<thead>
<tr>
<th>Tolerance category</th>
<th>D1</th>
<th>D2</th>
<th>D3</th>
<th>D4</th>
</tr>
</thead>
<tbody>
<tr>
<td>Length (mm)</td>
<td>+3</td>
<td>+1</td>
<td>+1</td>
<td>+1</td>
</tr>
<tr>
<td>Width (mm)</td>
<td>−5</td>
<td>−3</td>
<td>−3</td>
<td>−3</td>
</tr>
<tr>
<td>Height (mm)</td>
<td>+3</td>
<td>±2</td>
<td>±1.5</td>
<td>±1.0</td>
</tr>
</tbody>
</table>

Notes:
BS 6073: 2008 states that tolerance categories D3 and D4 are intended for use with thin-layer mortar joint systems. Therefore, most units used within the UK conform to tolerance categories D1 and D2.
Closer tolerances may be declared by the manufacturer.
Table 2.3  Limit of tolerances on autoclaved aerated concrete block sizes

<table>
<thead>
<tr>
<th>Standard joints of general purpose and lightweight mortar</th>
<th>Thin-layer mortar joints</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>GPLM</strong></td>
<td><strong>TLMA</strong></td>
</tr>
<tr>
<td>Length (mm)</td>
<td>−5 to +3</td>
</tr>
<tr>
<td>Width (mm)</td>
<td>±3</td>
</tr>
<tr>
<td>Height (mm)</td>
<td>−5 to +3</td>
</tr>
</tbody>
</table>

Notes:
For autoclaved aerated concrete units of category TLMB, the maximum deviation from flatness of bed faces and plane parallelism of bed faces is ≤1.0 mm in each case.
Closer tolerances may be declared by the manufacturer.

MANUFACTURE

Dense concrete blocks, which may be hollow, cellular or solid in form, are manufactured from natural dense aggregates including crushed granite, limestone and gravel. Medium and lightweight concrete blocks are manufactured incorporating a wide range of aggregates including expanded clay, expanded blastfurnace slag, sintered ash and pumice. Concrete is cast into moulds, vibrated and cured. Most aerated (aircrete or autoclaved aerated concrete) blocks are formed by the addition of aluminium powder to a fine mix of sand, lime, fly ash (pulverised fuel ash) and Portland cement. The hydrogen gas generated by the dissolution of the metal powder produces a non-interconnecting cellular structure. The process is accelerated by pressure steam curing in an autoclave (Fig. 2.4). For some products, additional insulation is afforded by the filling of voids in the cellular blocks or by bonding on a layer of extruded polystyrene, polyurethane or foil-faced phenolic foam (Fig. 2.1). Standard blocks, typically natural grey or buff in colour, are usually shrinkwrapped for delivery. Different grades of blocks are usually identified by scratch marks or colour codes.

PROPERTIES

Density and strength

The British Standard BS 6073-2: 2008 lists common compressive strengths of 2.9, 3.6, 7.3, 8.7, 10.4, 17.5, 22.5, 30.0, and 40.0 MPa for the range of aircrete and aggregate concrete blocks. However, the majority of concrete blocks fall in the range from 2.8 to 30 MPa, with associated densities of 420–2200 kg/m³ and thermal conductivities from 0.10 to 1.5 W/m K at 3% moisture content (Table 2.4). Drying shrinkages are typically in the range 0.03–0.05%.

Durability

Dense concrete blocks and certain aerated lightweight blocks are resistant to freeze/thaw conditions below damp-proof course (DPC) level. However, some lightweight concrete blocks, with less than 7 MPa crushing strength should not be used below DPC level, except for the inner skin of cavity construction.

Fixability

Aerated and lightweight concrete blocks offer a good background for fixings. For light loads, nails to a depth of 50 mm are sufficient. For heavier loads, wall plugs and proprietary fixings are necessary. These fixings should avoid the edges of the blocks.

Fig. 2.4 Manufacture of aerated blocks
Table 2.4  Typical relationship between density and thermal conductivity for concrete blocks

<table>
<thead>
<tr>
<th>Nominal density (kg/m³)</th>
<th>2200</th>
<th>2000</th>
<th>1800</th>
<th>1600</th>
<th>1400</th>
<th>1200</th>
<th>1000</th>
<th>900</th>
<th>800</th>
<th>750</th>
<th>700</th>
<th>600</th>
<th>500</th>
<th>460</th>
<th>420</th>
</tr>
</thead>
<tbody>
<tr>
<td>Typical thermal conductivity (W/m K)</td>
<td>1.5</td>
<td>1.2</td>
<td>0.83</td>
<td>0.63</td>
<td>0.47</td>
<td>0.36</td>
<td>0.27</td>
<td>0.24</td>
<td>0.20</td>
<td>0.19</td>
<td>0.17</td>
<td>0.15</td>
<td>0.12</td>
<td>0.11</td>
<td>0.10</td>
</tr>
</tbody>
</table>

Notes: Blocks of differing compositions may vary significantly from these average figures and manufacturers’ data should be used.

Thermal insulation

The Building Regulations Approved Document Part L (2006 Edition) requires new dwellings (Part L1A) and other new building types (Part L2A) to be compliant with an overall energy and carbon performance, the Target Emission Rate (TER) based on the whole building (Chapter 7, page 246). Individual $U$-values for elements are therefore not set, except for extensions on existing dwellings (Part L1B) and other existing buildings (Part L2B) where an indicative $U$-value of 0.30 W/m² K is the standard for new exposed walls. The limiting area-weighted $U$-value standard for wall elements in new buildings is 0.35 W/m² K, but to achieve the Target Emission Rate overall, most buildings will require wall $U$-values within the range 0.27–0.30 W/m² K.

The following material combinations achieve a $U$-value of 0.27 W/m² K (Fig. 2.5).

- Partially filled cavity
  - 102.5 mm fairfaced brickwork outer leaf
  - 50 mm clear cavity
  - 50 mm foil-faced polyurethane foam ($\lambda = 0.023$ W/m K)
  - 100 mm lightweight blocks ($\lambda = 0.15$ W/m K)
  - 12.5 mm plasterboard on dabs ($\lambda = 0.16$ W/m K).

- Fully filled cavity
  - 102.5 mm fairfaced brickwork outer leaf
  - 100 mm full-fill cavity of blown mineral wool ($\lambda = 0.038$ W/m K)
  - 100 mm lightweight blocks ($\lambda = 0.15$ W/m K)
  - 13 mm dense plaster.

Similarly, a $U$-value of 0.27 W/m² K can be achieved with 100 mm external fairfaced blockwork as an alternative to fairfaced brickwork, provided that the...
necessary additional thermal resistance is furnished by slightly increased cavity insulation. The thin-joint mortar system for inner leaf blockwork gives slightly enhanced $U$-values compared to the equivalent standard 10 mm joint blockwork construction.

Rendered solid wall construction can also achieve a $U$-value of 0.27 W/m$^2$ K (Fig. 2.5).

**Solid wall**

- 16 mm external render
- 215 mm high-performance lightweight blocks ($\lambda = 0.11$ W/m K)
- 50 mm lining of 9.5 mm plasterboard ($\lambda = 0.16$ W/m K) backed with 40 mm phenolic foam insulation ($\lambda = 0.023$ W/m K).

The following material combinations achieve a $U$-value of 0.20 W/m$^2$ K. These tighter specifications are required to achieve higher ratings in respect of the Code for Sustainable Homes.

**Partially filled cavity**

- 102.5 mm fairfaced brickwork outer leaf
- 50 mm clear cavity
- 75 mm foil-faced polyurethane foam ($\lambda = 0.022$ W/m K)
- 100 mm lightweight blocks ($\lambda = 0.15$ W/m K)
- 12.5 mm plasterboard on dabs ($\lambda = 0.16$ W/m K).

**Fully filled cavity**

- 102.5 mm fairfaced brickwork outer leaf
- 150 mm full-fill cavity of blown mineral wool ($\lambda = 0.038$ W/m K)
- 100 mm lightweight blocks ($\lambda = 0.15$ W/m K)
- 13 mm dense plaster.

**Solid wall**

- 16 mm external render
- 70 mm phenolic foam insulation ($\lambda = 0.023$ W/m K)
- 215 mm high-performance lightweight blocks ($\lambda = 0.11$ W/m K)
- 12.5 mm plasterboard on dabs ($\lambda = 0.16$ W/m K).

For domestic construction the appropriate Robust Details should be used to ensure compliance with thermal and sound requirements of the Building Regulations.

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**Phase change material blocks**

Phase change materials (PCMs) incorporated into aerated concrete blocks offer some additional thermal stability to the internal environment by absorbing excessive summer heat, which is then released during the cooler periods. This phase change at 26°C effectively increases the thermal capacity of the lightweight blocks. One manufacturer colour codes the phase change material blocks green for easy identification. The phase change material is described in Chapter 12 (page 337).

**Fire resistance**

Concrete block construction offers good fire resistance. Solid unplastered 90 mm blocks can give up to 60 minutes’ fire protection when used as load-bearing walls; certain 150 mm and most 215 mm solid blocks can achieve 360 minutes’ protection. Dense, lightweight and autoclaved aerated concrete blocks with less than 1% organic material are automatically categorised as Euroclass A1 with respect to reaction to fire.

**Sound insulation**

The Building Regulations 2000 Approved Document E (2003) recognises the need to provide adequate sound insulation both between and within dwellings also between rooms in hostels, hotels and residential accommodation. The regulations require minimum airborne sound insulation of $45R_w$ dB for separating walls and $40R_w$ dB for internal bedroom or WC walls. The passage of airborne sound depends on the density and porosity of the material. The use of Robust Details or Pre-Completion Testing is required to demonstrate compliance. The following alternative systems should perform to the required airborne insulation standard for separating walls of new build dwellings.

- 12.5 mm plasterboard on dabs
- 8 mm render
- 100 mm dense (1600–2200 kg/m$^3$) or lightweight (1350–1600 kg/m$^3$) blockwork
- 75 mm clear cavity only linked by appropriate wall ties
- 100 mm dense (1600–2200 kg/m$^3$) or lightweight (1350–1600 kg/m$^3$) blockwork
- 8 mm render
- 12.5 mm plasterboard on dabs.

These alternatives perform to the required standard only if there are no air leaks within the construction,
all joints are filled, the cavities are kept clear except for the approved wall ties and any chasing out on opposite sides of the construction is staggered. Vertical chases should, in any case, not be deeper than one third of the block thickness. Horizontal chases should be restricted to not more than one sixth of the block thickness, due to the potential loss of structural strength.

**Sound absorption**

The majority of standard concrete blocks with hard surfaces are highly reflective to sound, thus creating long reverberation times within building enclosures. Acoustic absorbing concrete blocks are manufactured with a slot on the exposed face which admits sound into the central cavity (Fig. 2.1). Since the void space is lined with sound-absorbing fibrous filler, incident sound is dissipated rather than reflected, significantly reducing reverberation effects. Acoustic control blocks in fairfaced concrete are suitable for use in swimming pools, sports halls, industrial buildings and auditoria.

**SPECIALS**

Most manufacturers of blocks produce a range of specials to match their standard ranges. Quoins, cavity closers, splayed cills, flush or projecting copings, lintel units, bullnose ends and radius blocks are generally available, and other specials can be made to order (Fig. 2.6). The use of specials in fairfaced blockwork can greatly enhance visual qualities. Matching full-length lintels may incorporate dummy joints and should bear on to full, not cut, blocks.

**FAIRFACED BLOCKS**

Fairfaced concrete blocks are available in a wide range of colours from white, through buff, sandstone, yellow, to pink, blue, green and black. Frequently the colour is all through, although some blocks have an applied surface colour. Most blocks are uniform in colour, but there is some variability with, for example, flecked finishes. Textures range from polished, smooth and weathered (sand- or shot-blasted) to striated and split face (Fig. 2.7), the latter intended to give a random variability associated more with natural stone.

Glazed masonry units are manufactured by the application of a thermosetting material to one or more faces of lightweight concrete blocks, which are then heat-treated to cure the finish. The glazed blocks are available in an extensive range of durable bright colours and are suitable for interior or exterior use.

Where required, profiled blocks to individual designs can be glazed by this system. Most manufacturers produce a range of specials to co-ordinate with their standard fairfaced blocks, although, as with special bricks, they may be manufactured from a different batch of mix, and this may give rise to slight variations. In specific cases, such as individual lintel blocks, specials are made by cutting standard blocks to ensure exact colour matching.

**Clay blocks**

**FIRED-CLAY BLOCKS**

Masonry clay honeycomb-insulating blocks can be used as a single skin for external load-bearing construction as an alternative to standard cavity construction. These fired-clay honeycomb blocks combine structural strength, insulation and, when externally rendered, moisture protection. The internal surface
is normally finished directly with gypsum plaster. Blocks for monolithic construction are 260 mm long × 240 mm high and either 300 or 365 mm wide giving wall $U$-values of 0.36 and 0.30 W/m² K, respectively, when rendered and plastered. For internal walls, blocks are 400 mm long and range in widths from 100 to 125 and 150 mm. Horizontal joints require 10 mm of a lightweight mortar, but the vertical joint edges, if tongued and grooved, remain dry. The British Standard (BS EN 771-1: 2003) illustrates a selection of high density (HD) vertically perforated units and a range of low density (LD) fired-clay masonry units. The LD units may be vertically or horizontally perforated, with butt jointing, mortar pockets or a tongue and groove system (Fig. 2.8). Special blocks are available for corners, lintels, door and window openings, but individual blocks can also be cut.

Fairfaced fired-clay blocks, as illustrated in Fig. 2.9, offer an alternative to traditional brickwork. They are manufactured to natural, riven or textured finish in a range of colours including terracotta red, ochre, buff and blue, and also to high gloss or satin finish in strong or pastel shades. Where used as infill, rather than load-bearing, alternative bonding is possible including stack bond. Typical work sizes, depending on the manufacturer, are 215 × 215 mm, 327 × 215 mm, 327 × 140 mm, 440 × 215 mm, 390 × 240 mm, 390 × 190 mm and 490 × 190 mm with widths of 90 and 102 mm. A standard 10 mm mortar joint is appropriate, which may match or contrast with the block colour.

**UNFIRED-CLAY BLOCKS**

Unfired blocks manufactured from clay and sometimes incorporating straw may be used for non-load-bearing partition walls. Blocks (typically 500 × 250 mm and 450 × 225 mm × 100 mm wide) may be tongued and grooved or square edged, but only the horizontal joints require fixing with a thin layer of cellulose-based adhesive or clay mortar. Blocks are easily cut to create architectural features, and are usually finished with a skim coat of clay plaster, although they may be painted directly. Internal walls are sufficiently strong to support shelving and other fixtures. Unfired-clay block walls are recyclable or biodegradable and have the advantage of absorbing odours and stabilising internal humidity and temperature by their natural absorption and release of moisture and heat. A 100 mm thick wall gives a 45 dB sound reduction and 90 minutes’ fire resistance. (The thermal conductivity of perforated unfired-clay blocks is typically 0.24 W/m K.)

**Gypsum blocks**

Gypsum blocks are available with densities ranging from 600 to 1500 kg/m³ and thicknesses from 50 to
Fig. 2.8  (a) Low-density and high-density units. Permission to reproduce extracts from BS EN 771-1: 2003 is granted by the British Standards Institute. (b) Hollow clay blocks in Greece. Photograph: Arthur Lyons
100 mm. The maximum thickness to BS EN 12859: 2008 is 150 mm. The preferred face dimensions are 666 mm in length and 500 mm in height, with a maximum length of 1000 mm. Gypsum blocks are classified by density and water absorption.

Gypsum blocks - density class:
- Low density (L) 600–800 kg/m³
- Medium density (M) 800–1100 kg/m³
- High density (H) 1100–1500 kg/m³

Gypsum blocks - water absorption class:
- H3 >5%
- H2 ≤5%
- H1 ≤2.5%

The standard BS EN 15318: 2007 details the sound insulation properties of gypsum block partitions in relation to wall thickness and block density. Gypsum blocks may be used as non-load-bearing partitions and internal insulation of walls. They are assembled with gypsum-based adhesives as specified in BS EN 12860: 2001.

**Blockwork**

**FAIRFACED CONCRETE BLOCKWORK**

Within fairfaced blockwork, an appropriate choice of size is important to both co-ordination and visual scale. Whilst blocks can be cut with a masonry cutter, the addition of small pieces of block, or the widening of perpends over the 10 mm standard, is unacceptable. The insertion of a thin jumper course at floor or lintel height may be a useful feature in adjusting the coursing. Curved blockwork may be constructed from standard blocks, the permissible curvature being dependent on the block size. The oversail between alternate courses should not normally exceed 4 mm in fairfaced work. If the internal radius is exposed, then the perpends can be maintained at 10 mm with uncut blocks, but if the external radius is exposed, the blocks will require cutting on a splay. For tighter curves specials will be required.
Thin-joint blockwork may be constructed with mortar joints of only 2–3 mm, provided that the aircrete or equivalent blocks have been manufactured to fine tolerances and on-site workmanship is good. The special rapid-setting mortar sets typically within 60 minutes and the full bond strength is achieved after only 2 hours, allowing more courses to be laid each day. In the case of brick and block cavity construction, the inner leaf is built first, providing a weatherproof enclosure as quickly as possible. The outer skin of brickwork can subsequently be built up, using wall ties fixed to the face, either screwed or hammered into the completed blockwork. Bed joints in thin-layer mortar blockwork do not necessarily co-ordinate with those of the brickwork, so conventional cavity wall ties can only be used if they are slope-tolerant.

Usually, inner leaf construction commences with a line of 440 × 215 mm standard height blocks, with normal bedding mortar to compensate for variations in the foundation level, followed by the larger 440 or 620 mm × 430 mm high blocks, which should weigh less than 20 kg for repeated lifting by one operative. Heavier blocks require mechanical lifting or two-person handling. Thin-joint mortars, consisting of polymer-modified 1 : 2 cement : sand mix with water-retaining and workability admixtures, are factory pre-mixed and require only the addition of water, preferably mixed in with an electrically powered plasterer’s whisk. The mortar is applied manually with a serrated scoop of the appropriate width or through a pumped system to achieve uniformity. Work should only proceed at temperatures above 5°C.

The main advantages of thin-joint systems over traditional 10 mm joint blockwork are:

- increased productivity allowing storey-height inner leaves to be completed in one day;
- up to 10% improved thermal performance due to reduced thermal bridging by the mortar;
- improved airtightness of the construction;
- the accuracy of the wall, which allows internal thin-coat sprayed plaster finishes to be used;
- higher quality of construction and less wastage of mortar.

The acoustic properties of thin-joint mortar walls differ slightly from those of walls constructed with 10 mm mortar joints. Resistance to low-frequency noise is slightly enhanced, whilst resistance to high-frequency sound is slightly reduced.

Completed thin-joint blockwork acts as a monolithic slab, which, if unrestrained, may crack at the weaker points, such as near openings. To avoid this, the block units should be laid dry to avoid shrinkage and bed joint reinforcement (1.5 mm thick GRP mesh) should be appropriately positioned. Larger structures require movement joints at 6 m centres.

Certain extruded multi-perforated clay and calcium silicate blocks are designed for use with thin mortar bed joints and dry interlocking vertical joints. One system of clay blocks requires only horizontal proprietary adhesive joints of 1 mm applied with a special roller tool, as the units are ground to exact dimensions after firing. Blocks may be used for inner and/or outer leaf cavity construction or for solid walls. Whilst this reduces the initial construction time, exposed sides of the units subsequently require plaster or cement render to minimise heat loss by air leakage. Typical block sizes are 300 × 224 mm and 248 × 249 mm with widths of 100, 140, 190 and 365 mm.
BOND

A running half-block bond is standard, but this may be reduced to a quarter bond for aesthetic reasons. Blockwork may incorporate banding of concrete bricks, but because of differences in thermal and moisture movement, it is inadvisable to mix clay bricks with concrete blocks. Horizontal and vertical stack bonds and more sophisticated variations, such as basket-weave bond, may be used for infill panels within framed structures (Fig. 2.10). Such panels will require reinforcement within alternate horizontal bed joints, to compensate for the lack of normal bonding.

REINFORCEMENT

Blockwork will require bed-joint reinforcement above and below openings where it is inappropriate to divide the blockwork up into panels, with movement joints at the ends of the lintels. Bed-joint reinforcement would be inserted into two bed joints above and below such openings (Fig. 2.11). Cover to bed reinforcement should be at least 25 mm on the external faces and 13 mm on the internal faces. Combined vertical and horizontal reinforcement may be incorporated into hollow blockwork in accordance with BS 5628-2: 2005, where demanded by the calculated stresses. Typical situations would be within retaining basement walls, and large infill panels to a framed structure.

MOVEMENT CONTROL

Concrete blockwork is subject to greater movements than equivalent brickwork masonry. Therefore, the location and form of the movement joints requires greater design-detail consideration, to ensure that inevitable movements are directed to the required locations and do not cause unsightly stepped cracking or fracture of individual blocks. Ideally, such movement joints should be located at intersecting walls, or other points of structural discontinuity, such as columns. Additionally, movement joints are required at changes in thickness, height or loading of walls, above and below wall openings, and adjacent to movement joints in the adjoining structure (Fig. 2.12). External unreinforced non-load-bearing concrete masonry walls with a length to height ratio of 3:1 or less must be separated into a series of panels with vertical movement-control joints at approximately 9 m centres or more frequently for masonry walls with a length to height ratio of more than 3:1 (NA to BS EN 1996-2: 2006).

Wall ties should allow for differential movement between the leaves in cavity construction and should be spaced at 900 mm horizontally and 450 mm vertically, for 50–75 mm cavities.

MORTARS

The mortar must always be weaker than the blocks to allow for movement.
The usual mixes for standard 10 mm joints are by volume:

- cement/lime/sand 1:1:5 to 1:1:6
- cement/sand + plasticiser 1:5 to 1:6
- masonry cement/sand 1:4 to 1:5

Below DPC level a stronger mix is required and sulphate-resisting cement may be necessary depending on soil conditions.

- cement/sand 1:4
- cement/lime/sand 1:1.5:4.5

Where high-strength blockwork is required, stronger mortars may be necessary. Mortar joints should

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**Fig. 2.11** Reinforced blockwork

**Fig. 2.12** Blockwork movement joints

Below DPC level a stronger mix is required and sulphate-resisting cement may be necessary depending on soil conditions.

- cement/sand 1:4
- cement/lime/sand 1:1.5:4.5

Where high-strength blockwork is required, stronger mortars may be necessary. Mortar joints should
be slightly concave, rather than flush. Bucket handle and weathered or struck joints are suitable for external use, but recessed joints should only be used internally. Coloured mortars should be ready mixed or carefully gauged to prevent colour variations. Contraction joints should be finished with a bond breaker of polythene tape and flexible sealant. For expansion joints, a flexible filler is required, e.g. bitumen-impregnated fibreboard with a polythene foam strip and flexible sealant. Where blockwork is to be rendered, the mortar should be raked back to a depth of 10 mm for additional key. Masonry should not be built when the temperature is at or below 3°C and falling or unless it is at least 1°C and rising (BS 5628-3: 2005).

FINISHES

Internal finishes

Plaster should be applied normally in two coats to 13 mm. Blocks intended for plastering have a textured surface to give a good key. Dry lining may be fixed with battens or directly with adhesive dabs to the blockwork. Blockwork to be tiled should be first rendered with a cement/sand mix. Fairfaced blockwork may be left plain or painted. Where standard blocks are to be painted, the appropriate grade should be used.

Unfired-clay blocks should be finished with breathable materials, such as clay or lime plaster, clay boards, limewash or highly vapour-permeable paint.

External finishes

External boarding or hanging tiles should be fixed to battens, separated from the blockwork with a breather membrane. For external rendering a spatterdash coat should be applied initially on dense blockwork, followed by two coats of cement/lime/sand render. The first 10 mm coat should be the stronger mix (e.g. 1:1:6); the 5 mm second coat must be weaker (e.g. 1:2:9). Cement/sand mixes are not recommended as they are more susceptible to cracking and crazing than mixes incorporating lime. The render should terminate at damp-proof course level with a drip or similar weathering detail.

FOUNDATIONS

Foundation blocks laid flat offer an alternative to trench fill or cavity masonry. Portland cement blocks should not be used for foundations where sulphate-resisting cement mortar is specified, unless they are classified as suitable for the particular sulphate conditions. Sulphate and other chemically adverse ground conditions are classified in the BRE Special Digest 1 (2005) from DS1 (Design Sulphate Class 1) to the most aggressive, DS5. Foundation blocks can be of either dense or appropriate lightweight concrete, the latter providing enhanced floor edge insulation. Interlocking foundation blocks, with a tongue and groove vertical joint, slot together with only bed-joint mortar being required. A handhold makes manipulating these blocks on site much easier than lifting standard rectangular blocks. Thicknesses within the range 255–355 mm are standard.

Beam and block flooring

Beam and block flooring offers an alternative system to traditional solid ground floors within domestic-scale construction (Fig. 2.13), and may also be used for upper storeys. Systems are described in BS EN 15037 Part 1: 2008 and Parts 2 and 3: 2009. Beams may be inverted T or I in form, alternatively incorporate partially exposed lattice girder reinforcement to be covered within a concrete topping. The infill may be standard 100 mm concrete blocks with a minimum transverse crushing strength of 3.5 MPa. Insulation will be required to achieve a $U$-value between 0.20
and 0.25 W/m² K. For first and subsequent floors, the infill can be full-depth solid or hollow concrete blocks or hollow clay blocks which may require a cast in situ structural topping to comply with Building Regulations.

The following material combination achieves a U-value of 0.20 W/m² K:

- 18 mm particleboard (λ = 0.13 W/m K)
- 100 mm continuous insulation (λ = 0.030 W/m K)
100 mm concrete block ($\lambda = 0.46 \text{ W/m K}$)
dense concrete inverted T-beam at 515 mm centres
($\lambda = 1.65 \text{ W/m K}$)
under-floor ventilated space.

Landscape blockwork

**BLOCK PAVING**

Concrete block paving units are manufactured to a wide range of designs as illustrated in Fig. 2.14. Blocks may be of standard brick form ($200 \times 100 \text{ mm}$) to thicknesses of 60, 80 or 100 mm depending on the anticipated loading. Alternative designs include tumbled blocks, which emulate granite setts, and various interlocking forms giving designs based on polygonal and curvilinear forms. Colours range from red, brindle, buff, brown, charcoal and grey through to silver and white, with smooth, textured or simulated stone finishes. For most designs, a range of kerb blocks, drainage channels, edging and other accessory units are available. Concrete paving blocks are usually laid on a compacted sub-base with 50 mm of sharp sand. Blocks are frequently nibbed to create a narrow joint to be filled with kiln-dried sand. For the wider joints that occur between the simulated stone setts a coarser grit can be used to prevent loss by wind erosion.

Physical properties, including water absorption, freeze/thaw resistance, abrasion resistance and tolerances on size, are categorised in BS EN 1338: 2003. Guidance on the design and construction of pavements for a range of applications is given in BS 7533 Parts 1–13 inclusive. The types of tactile paving surfaces – blister, rib and groove – are defined and coded in the standard DD CEN/TS 15209: 2008.

Sustainable urban drainage systems (SUDS) are designed to reduce the environmental impact of impermeable hard landscaping surfaces which create rapid rainwater run-off. With permeable surfaces, including nibbed blocks, the rainwater permeates through the spacing and is dispersed by natural drainage into the underlying soil, or may be collected through rainwater harvesting systems for further use.

Where the appearance of grass is required, but with the traffic-bearing properties of a concrete block pavement, a selection of porous (hollow) blocks is available which can be filled with soil and seeded to give the required effect. Different block depths and sub-bases can be specified according to the anticipated traffic loading. Sulphate-resisting blocks are available if dictated by the soil conditions.

**EARTH-RETAINING BLOCKWORK**

A range of precast-cellular concrete-interlocking blocks is manufactured for the construction of dry-bed retaining walls. Soil is placed in the pockets of each successive course to allow for planting. The rear is backfilled with granular material to allow for drainage. The size of the block determines the maximum construction height, but over 20 m can be achieved with very deep units. A face angle of $15^\circ$ to $22^\circ$ is typical to ensure stability, but other gradients are possible with the appropriate block systems. Limited wall curvature is possible without cutting the standard blocks. The systems are used both for earth retention and to form acoustic barriers.

**References**

**FURTHER READING**


**STANDARDS**

BS 5628 Code of practice for use of masonry:
Part 2: 2005 Structural use of reinforced and prestressed masonry.
BS 5977 Lintels:
BS 6073 Precast concrete masonry units:
BS 6100 Glossary of building and civil engineering terms:
  Part 0: 2002 Introduction.
BS 7533 Pavements constructed with clay, natural stone or concrete pavers:
BS 8000 Workmanship on building sites:
BS 8103 Structural design of low-rise buildings:
BS EN 413-1: 2008 Masonry cement. Composition, specifications and conformity criteria.
BS EN 771 Specification for masonry units:
  Part 4: 2003 Autoclaved aerated concrete masonry units.
  Part 5: 2003 Manufactured stone masonry units.
BS EN 772 Methods of test for masonry units:
BS EN 845 Specification for ancillary components for masonry:
  Part 1: 2003 Ties, tension straps, hangers and brackets.
BS EN 934 Admixtures for concrete, mortar and grout:
  Part 2: 2001 Concrete admixtures. Definitions, requirements, conformity, marking and labelling.
BS EN 1996 Eurocode 6: Design of masonry structures:
  Part 1.2: 2005 Structural fire design.
BS EN 12859: 2008 Gypsum blocks. Definitions, requirements and test methods.
BS EN 13139: 2002 Aggregates for mortar.
BS EN 15037 Precast concrete products. Beam-and-block floor systems:
  Part 2: 2009 Concrete blocks.
DD CEN/TS 15209: 2008 Tactile paving surface indicators produced from concrete, clay and stone.
BS EN 15254 Extended application of results from fire resistance tests. Nonloadbearing walls:
BS EN 15318: 2007 Design and application of gypsum blocks.

BUILDING RESEARCH ESTABLISHMENT PUBLICATIONS

BRE Special digests
BRE SD1: 2005 Concrete in aggressive ground.
BRE SD4: 2007 Masonry walls and beam and block floors. U-values and building regulations.

BRE Digests
BRE Digest 432: 1998 Aircrete: thin joint mortar system.
BRE Digest 460: 2001 Bricks, blocks and masonry made from aggregate concrete (Parts 1 and 2).
BRE Digest 461: 2001 Corrosion of metal components in walls.
BRE Digest 468: 2002 AAC ‘aircrete’ blocks and masonry.

BRE Good building guides
BRE GBG 44: 2000 Insulating masonry cavity walls (Parts 1 and 2).
BRE GBG 50: 2002 Insulating solid masonry walls.
BRE GBG 58: 2003 Thin layer masonry mortar.
BRE GBG 62: 2004 Retro-installation of bed joint reinforcement in masonry.
BRE GBG 67: 2006 Achieving airtightness (Parts 1, 2, and 3).
BRE GBG 68: 2006 Installing thermal insulation (Parts 1 and 2).

BRE Information papers
BRE IP 1/99 Untied cavity party walls. Structural performance when using AAC blockwork.
BRE IP 7/05 Aircrete tongue and grooved block masonry.
BRE IP 1/06 Assessing the effects of thermal bridging at junctions and around openings.
BRE IP 8/08 Determining the minimal thermal resistance of cavity closers.

ADVISORY ORGANISATIONS
Aircrete Products Association, 4th floor, 60 Charles Street, Leicester LE1 1FB, UK (0116 253 6161).
British Concrete Masonry Association, Grove Crescent House, 18 Grove Place, Bedford MK40 3JJ, UK (01234 353745).
Concrete Block Association, 60 Charles Street, Leicester LE1 1FB, UK (0116 253 6161).
Concrete Society, Riverside House, 4 Meadows Business Park, Station Approach, Blackwater, Camberley, Surrey GU14 9AB, UK (01276 607140).
Mortar Industry Association, Gillingham House, 38-44 Gillingham Street, London SW1V 1HU, UK (020 7963 8000).
**Introduction**

In the broadest sense, the term *cement* refers to materials which act as adhesives. However, in this context, its use is restricted to that of a binding agent for sand, stone and other aggregates within the manufacture of mortar and concrete. Hydraulic cements and limes set and harden by internal chemical reactions when mixed with water. Non-hydraulic materials will harden slowly by absorption of carbon dioxide from the air.

Lime was used as a binding agent for brick and stone by the ancient civilisations throughout the world. The concept was brought to Britain in the first century AD by the Romans, who used the material to produce lime mortar. Outside Britain, the Romans frequently mixed lime with volcanic ashes, such as pozzolana from Pozzuoli in Italy, to convert non-hydraulic lime into a hydraulic cement suitable for use in constructing aqueducts, baths and other buildings. However, in Britain, lime was usually mixed with artificial pozzolanas, for example crushed burnt clay products, such as pottery, brick and tile. In the eighteenth century, so-called *Roman cement* was manufactured by burning the *cement stone* (argillaceous or clayey limestone), collected from the coast around Sheppey and Essex.

In 1824, Joseph Aspdin was granted his famous patent for the manufacture of *Portland cement* from limestone and clay. Limestone powder and clay were mixed into a water slurry which was then evaporated by heat in *slip pans*. The dry mixture was broken into small lumps, calcined in a kiln to drive off the carbon dioxide, burnt to clinker and finally ground into a fine powder for use. The name Portland was used to enhance the prestige of the new concrete material by relating it to Portland stone which, in some degree, it resembled. Early manufacture of Portland cement was by intermittent processes within bottle, and later chamber, kilns. The introduction of the rotating furnace in 1877 offered a continuous burning process with consequent reductions in fuel and labour costs. The early rotating kilns formed the basis for the development of the various production systems that exist now. Year 1989 was the peak of production when 18 million tonnes of cement was manufactured within the UK. About half of this was required by the ready-mixed concrete industry; the remainder was divided roughly equally between concrete-product factories and bagged cement for general use.

**Lime**

**MANUFACTURE OF LIME**

Lime is manufactured by calcining natural calcium carbonate, typically hard-rock carboniferous limestone. The mineral is quarried, crushed, ground, washed and screened to the required size range. The limestone is burnt at approximately 950°C in either horizontal rotary kilns or vertical shaft kilns which drive off the carbon dioxide to produce the lime products. Limestone and chalk produce non-hydraulic or air limes whilst limestone/clay mixtures produce hydraulic limes. Subsequent treatments produce the product range given in Fig. 3.1. Air or non-hydraulic limes are classified by BS EN 459-1: 2001 according to their calcium and magnesium oxide contents, and hydraulic limes are classified according to their compressive strengths (Table 3.1).
Types of building lime (BS EN 459-1: 2001)

<table>
<thead>
<tr>
<th>Designation</th>
<th>Notation</th>
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<tbody>
<tr>
<td>Calcium lime 90</td>
<td>CL 90</td>
</tr>
<tr>
<td>Calcium lime 80</td>
<td>CL 80</td>
</tr>
<tr>
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<td>CL 70</td>
</tr>
<tr>
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<td>DL 85</td>
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<tr>
<td>Natural hydraulic lime 5</td>
<td>NHL 5</td>
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</tbody>
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Notes:
Air or non-hydraulic limes are also classified according to their delivery conditions—quicklime (Q), hydrated lime (S), semi-hydrated dolomitic lime [calcium hydroxide and magnesium oxide] (S1) and completely hydrated dolomitic lime [calcium hydroxide and magnesium hydroxide] (S2).

NON-HYDRAULIC OR AIR LIMES

Limestone and chalk when burnt produce quicklime or lump lime (calcium oxide). Quicklimes include calcium limes (CL) and dolomitic limes (DL) depending on the composition of the starting mineral as defined by BS EN 459-1: 2001. The dolomitic limes contain significant quantities of magnesium oxide.

$$\text{CaCO}_3 \xrightarrow{950^\circ C} \text{CaO} + \text{CO}_2$$

Slaking of lime

Slaking—the addition of water to quicklime—is a highly exothermic reaction. The controlled addition of water to quicklime produces hydrated lime (S) (mainly calcium hydroxide) as a dry powder.

$$\text{CaO} + \text{H}_2\text{O} \rightarrow \text{Ca(OH)}_2$$

It is suitable for use within mortars or in the manufacture of certain aerated concrete blocks. Generally, the addition of lime to cement mortar, render or plaster increases its water-retention properties, thus retaining workability, particularly when the material is applied to absorbent substrates such as porous brick. Lime also increases the cohesion of mortar mixes allowing them to spread more easily. Hydrated lime absorbs moisture and carbon dioxide from the air, and should therefore be stored in a cool, draught-free building and used whilst still fresh.

Lime putty

Lime putty is produced by slaking quicklime with an excess of water for a period of several weeks until a creamy texture is produced. Alternatively, it can be made by stirring hydrated lime into water, followed by conditioning for at least 24 hours. However, the traditional direct slaking of quicklime produces finer particle sizes in the slurry, and the best lime putty is produced by maturing for at least six months. Lime putty may be blended with Portland cement in mortars where its water-retention properties are greater than that afforded by hydrated lime. Additionally, lime putty, often mixed with sand to form coarse stuff, is used directly as a pure lime mortar, particularly in restoration and conservation work. It sets, not by reaction with sand and water, but only by carbonation, and is therefore described as non-hydraulic. Lime wash, as a traditional surface coating, is made by the addition of sufficient water to lime putty to produce a thin creamy consistency.

Carbonation

Lime hardens by the absorption of carbon dioxide from the air, which gradually reconverts calcium oxide back to calcium carbonate.

$$\text{CaO} + \text{CO}_2 \xrightarrow{\text{slow}} \text{CaCO}_3$$

The carbonation process is slow, as it is controlled by the diffusion of carbon dioxide into the bulk of the material. When sand or stone dust aggregate is added to the lime putty to form a mortar or render, the increased porosity allows greater access of carbon dioxide and a
speedier carbonation process. The maximum size of aggregate mixed into lime mortars should not exceed half the mortar-joint width.

**HYDRAULIC LIMES**

Hydraulic limes are manufactured from chalk or limestone containing various proportions of clay impurities. The materials produced have some of the properties of Portland cement, and partially harden through hydration processes, rather than solely through carbonation, as happens with non-hydraulic pure calcium oxide lime. Hydraulic limes rich in clay impurities are more hydraulic and set more rapidly than those with only a low silica and alumina content. Natural hydraulic limes are traditionally categorised as *feebly*, *moderately* or *eminently hydraulic* depending on their clay content, which is in the ranges 0–8%, 8–18% and 18–25%, respectively. These traditional grades equate approximately to the 28 day compressive strengths of 2, 3.5 and 5 MPa, respectively, for NHL2, NHL3.5 and NHL5. Eminently hydraulic lime mortar is used for masonry in exposed situations, moderately hydraulic lime mortar is used for most normal masonry applications and feebly hydraulic lime mortar is appropriate for conservation work and solid wall construction. Grey semi-hydraulic lime is still produced within the UK in small quantities from chalk containing a proportion of clay. It is used with very soft bricks and for conservation work. Natural hydraulic limes (NHLs) are produced by burning chalk and limestone but hydraulic limes (HLs) are produced by blending the constituents in appropriate proportions.

Hydraulic lime, usually imported from France, is mainly used for the restoration of historic buildings, where the use of modern materials would be inappropriate. It is gauged with sand only, giving a mix which develops an initial set within a few hours, but which hardens over an extended period of time. The workable render or mortar mixes adhere well and, because the material is flexible, the risks of cracking and poor adhesion are reduced. The dried mortar is off-white in colour and contains very little alkali, which in Portland cement mortars can cause staining, particularly on limestone. Hydraulic lime may be used for interior lime washes, and also for fixing glass bricks where a flexible binding agent with minimum shrinkage is required. Unlike hydrated lime, hydraulic lime is little affected by exposure to dry air during storage.

**LIME MORTAR**

Various advantages of lime-based mortars over Portland cement mortars are reported. The production of building lime consumes less energy, thus reducing greenhouse emissions compared to the equivalent manufacture of Portland cement. The subsequent carbonation process removes CO₂ from the atmosphere. Lime-based mortars remain sufficiently flexible to allow thermal and moisture movement, but additionally, due to the presence of uncarbonated lime, any minor cracks are subsequently healed by the action of rainwater. The recycling of bricks and blocks is easier due to the lower adherence of the mortar. Lime mortar construction is more breathable than Portland cement masonry and lime mortars are more resistant to sulphate attack than standard Portland cement mixes due to their lower tricalcium aluminate content.

Typical lime mortars are within the range 1 : 2 and 1 : 3, lime : aggregate ratio. A 1 : 2 lime : sand mix made with NHL3.5 lime equates approximately to the Standard BS 5628-1 designation (iii) class M4 mortar and a 1 : 3 mix equates roughly to a designation (iv) class M2 Portland cement mix. A well-graded sharp sand should be used. Because of the slow carbonation process, masonry lifts are limited, and the mortar must be allowed some setting time to prevent its expulsion from the joints. Little hardening occurs at temperatures below 5°C.

The use of lime mortars will eventually be incorporated into an annex for use with the Standard BS 5628-1: 2005.

**HEMP LIME**

Hemp is grown, particularly in France, for its fibre which is used in the manufacture of certain grades of paper. The remaining 75% of the hemp stalks, known as hemp hurd, is a lightweight absorbent material which has the appearance of fine wood chips. When mixed with hydraulic lime it produces a cement mixture which sets within a few hours and gradually ‘petrifies’ to a lightweight solid due to the high silica content of hemp hurd. The material mix can be poured and tamped or sprayed as required and formwork may be removed after 24 hours or less. The set material, sometimes referred to as ‘hempcrete’, which has good thermal insulation properties and a texture similar to that of cork, has been used for the construction of floors, walls using plywood formwork, and also blocks for blockwork. The material is also used as a solid infill
Fig. 3.2 Manufacture of Portland cement—the wet process

for timber-frame construction. In this case the combination of the moisture-absorbing properties of the hemp with the nature of lime, affords some protection to the timber-framing which it encloses. The moisture-absorbing properties of hemp lime give rise to higher thermal efficiencies than are calculated for the material based on conventional thermal transmittance data. To build a conventional house of hemp lime would require approximately 40 m$^3$ of the material containing 7–10 tonnes of hemp, produced from about 1 ha of land.

EXTERNAL LIME RENDERING

External lime rendering is usually applied in a two- or three-coat system, to give an overall thickness of up to 30 mm. In exposed situations, hydraulic lime is used and the thicker initial coat may be reinforced with horsehair. The final coat can be trowelled to receive a painted finish; alternatively, pebble dash or rough cast may be applied.

Cement

MANUFACTURE OF PORTLAND CEMENT

Portland cement is manufactured from calcium carbonate (chalk or limestone) by expulsion of the carbon dioxide, and sintering, at the point of incipient fusion, the resulting calcium oxide (lime) with the clay and iron oxide. Depending on the raw materials used and their water content at extraction, four key variations in the manufacturing process have been developed. These are the wet, semi-wet, semi-dry and the dry processes.

Wet process

The wet process (Fig. 3.2), which was the precursor to the other developments, is still used in some areas for processing chalk and marl clay. Clay is mixed with water to form a slurry, when any excess sand is removed by settlement. An equivalent slurry is prepared from the chalk which is then blended with the clay slurry, screened to remove any coarse material and stored in large slurry tanks. After final blending, the slurry is fed into the top of large slowly rotating kilns. The kilns, which are refractory brick-lined steel cylinders up to 200 m long, are fired to approximately 1450$^\circ$C, usually with pulverised coal. The slurry is dried, calcined and finally sintered to hard grey/black lumps of cement clinker.

A major development in energy conservation has been the elimination or reduction in the slurry water content required in the manufacturing process, as this consumed large quantities of heat energy during its evaporation.

Semi-wet process

In the semi-wet process, chalk is broken down in water and blended into a marl clay slurry. The 40% water content within the slurry is reduced to 19% in a filter press; the resulting filter-cake is nodularised by
extrusion onto a travelling preheater grate or reduced in a crusher/dryer to pellets. Heating to between 900°C and 1100°C in tower cyclones precalculnes the chalk; the mix is then transferred to a short kiln at 1450°C for the clinkering process.

**Semi-dry process**

In the semi-dry process, dry shale and limestone powders are blended. About 12% water is added to nodularise the blend, which is then precalcinced and clinkered as in the semi-wet process.

**Dry process**

In the dry process (Fig. 3.3) limestone, shale and sand (typically 80%, 17% and 3%, respectively) are milled to fine powders, then blended to produce the dry meal, which is stored in silos. The meal is passed through a series of cyclones, initially using recovered kiln gases to preheat it to 750°C, then with added fuel to precalcine at 900°C, prior to passage into a fast-rotating 60 m kiln for clinkering at 1450°C. In all processes an intimately mixed feedstock to the kiln is essential for maintaining quality control of the product. Most plants operate primarily with powdered coal, but additionally other fuels including petroleum coke, waste tyre chips, smokeless fuel plant residues or reclaimed spoil heap coal are used when available. Oil, natural gas and landfill gas have also been used when economically viable. The grey/black clinker manufactured by all processes is cooled with full heat recovery and ground up with 5% added gypsum (calcium sulphate) retarder to prevent excessively rapid flash setting of the cement.

The older cement grinding mills are open circuit allowing one pass of the clinker, which produces a wide range of particle size. This product is typically used for concrete production. The newer cement mills are closed circuit, with air separators to extract fine materials and with recycling of the oversize particles for regrinding. This product is frequently used in the ready-mixed market, as it can be controlled to produce cement with higher later strength. To reduce grinding costs, manufacturers accept load shedding and use off-peak electrical supplies where possible. The Portland cement is stored in silos prior to transportation in bulk, by road or rail, or in palletised packs. The standard bag is 25 kg for reasons of health and safety.

With the dry processing and additional increases in energy efficiency, a tonne of pulverised coal can now produce in excess of six tonnes of cement clinker compared to only three tonnes with the traditional wet process. Because the cement industry is so large, the combined output of carbon dioxide to the atmosphere, from fuel and the necessary decarbonation of the limestone or chalk, represents about 2% of the carbon dioxide emissions in Britain and up to 7.5% worldwide. According to data from the 2005 British Geological Survey, the UK has agreed to reduce its primary energy consumption by 25.6% per tonne of cement produced by 2010 from the baseline of 1990. Emissions of oxides of sulphur from the fuel are low as these gases are trapped into the cement clinker; however, the escape of oxides of nitrogen and dust, largely trapped by electrostatic precipitators, can only be controlled by constantly improving process technologies. On the basis of the final production of concrete, cement manufacture releases considerably less carbon dioxide per tonne than does primary steel manufacture; however, the relative masses for equivalent construction and the recycling potential of each should also be
Table 3.2 Typical composition of starting materials for Portland cement manufacture

<table>
<thead>
<tr>
<th>Component</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lime</td>
<td>68%</td>
</tr>
<tr>
<td>Silica</td>
<td>22%</td>
</tr>
<tr>
<td>Alumina</td>
<td>5%</td>
</tr>
<tr>
<td>Iron oxide</td>
<td>3%</td>
</tr>
<tr>
<td>Other oxides</td>
<td>2%</td>
</tr>
</tbody>
</table>

considered. Currently, some three quarters of European cement production is based on the dry process.

COMPOSITION OF PORTLAND CEMENT

The starting materials for Portland cement are chalk or limestone and clay, which consist mainly of lime, silica, alumina and iron oxide. Table 3.2 illustrates a typical composition.

Minor constituents, including magnesium oxide, sulphur trioxide, sodium and potassium oxides, amount to approximately 2%. (The presence of the alkali oxides in small proportions can be the cause of the alkali–silica reaction, which leads to cracking of concrete when certain silica-containing aggregates are used.) During the clinkering process, these compounds react together to produce the four key components of Portland cement (Table 3.3).

The relative proportions of these major components significantly affect the ultimate properties of the cements and are therefore adjusted in the manufacturing process to produce the required product range. Typical compositions of Portland cements are shown in Table 3.4.

A small reduction in the lime content within the initial mix will greatly reduce the proportion of the tricalcium silicate and produce an equivalent large increase in the dicalcium silicate component of the product. The cement produced will harden more slowly, with a slower evolution of heat. As tricalcium aluminate is vulnerable to attack by soluble sulphates it is the proportion of this component that is reduced in the manufacture of sulphate-resisting cement.

Under the British Standard BS EN 197-1: 2000, except in the case of sulphate-resisting cement, up to 5% of minor additional constituents may be added to cement. These fillers must be materials that do not increase the water requirements of the cement, reduce the durability of the mortar or concrete produced, or cause increased corrosion to any steel reinforcement. In the UK, typical fillers include limestone powder and dry meal or partially calcined material from the cement manufacturing processes.

SETTING AND HARDENING OF PORTLAND CEMENTS

Portland cement is hydraulic; when mixed with water it forms a paste, which sets and hardens as a result of various chemical reactions between the cementitious compounds and water. Setting and hardening

Table 3.3 Major constituents of Portland cement and their specific properties

<table>
<thead>
<tr>
<th>Compound</th>
<th>Chemical formula</th>
<th>Cement notation</th>
<th>Properties</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tricalcium silicate</td>
<td>3CaO·SiO₂</td>
<td>C₃S</td>
<td>Rapid hardening giving early strength and fast evolution of heat</td>
</tr>
<tr>
<td>Dicalcium silicate</td>
<td>2CaO·SiO₂</td>
<td>C₂S</td>
<td>Slow hardening giving slow development of strength and slow evolution of heat</td>
</tr>
<tr>
<td>Tricalcium aluminate</td>
<td>3CaO·Al₂O₃</td>
<td>C₃A</td>
<td>Quick setting which is retarded by gypsum. Rapid hardening and fast evolution of heat but lower final strength; vulnerable to sulphate attack</td>
</tr>
<tr>
<td>Tetracalcium aluminoferrite</td>
<td>4CaO·Al₂O₃·Fe₂O₃</td>
<td>C₄AF</td>
<td>Slow hardening; causes grey colour in cement</td>
</tr>
</tbody>
</table>

Table 3.4 Typical compositions of Portland cements

<table>
<thead>
<tr>
<th>Cement</th>
<th>Class</th>
<th>Composition</th>
<th>Fineness (m²/kg)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>%C₃S</td>
<td>%C₂S</td>
</tr>
<tr>
<td>Portland cement</td>
<td>42.5</td>
<td>55</td>
<td>20</td>
</tr>
<tr>
<td>52.5</td>
<td></td>
<td>55</td>
<td>20</td>
</tr>
<tr>
<td>White Portland cement</td>
<td>62.5</td>
<td>65</td>
<td>20</td>
</tr>
<tr>
<td>Sulphate-resisting Portland cement</td>
<td>42.5</td>
<td>60</td>
<td>15</td>
</tr>
</tbody>
</table>
are not dependent on drying out; indeed, Portland cement will harden under water. Only a small proportion of the added water is actually required for the chemical hydration of the cementitious constituents to hydrated calcium silicates. The additional water is needed to ensure the workability of the mix when aggregates are added, so that concrete, for example, can be successfully placed within formwork containing steel reinforcement. Water in excess of that required for hydration will ultimately evaporate leaving capillary pores in the concrete and mortar products. Typically, an increase in void space of 1% reduces crushing strength by 6%. It is therefore necessary to control carefully the water content of the mix by reference to the water/cement ratio. A minimum water/cement ratio of 0.23 is required to hydrate all the cement, although, as the cement powder is hydrated it expands, and thus a ratio of 0.36 represents the point at which cement gel fills all the water space. However, a water/cement of 0.42 more realistically represents the minimum water content to achieve full hydration without the necessity for further water to be absorbed during the curing process.

The setting and hardening processes should be distinguished. Setting is the stiffening of the cement paste which commences as soon as the cement is mixed with water. Because the major cementitious constituents set at different rates it is convenient to refer to initial set and final set. Typically, initial set, or the formation of a plastic gel, occurs after 1 hour and final set, or the formation of a rigid gel, within 10 hours. The setting process is controlled by the quantity of gypsum added to the cement in the final stages of production. Hardening is the gradual gain in strength of the set cement paste. It is a process which continues, albeit at a decreasing rate, over periods of days, months and years. The rate of hardening is governed partially by the particle-size distribution of the cement powder. Finely ground cement hydrates more rapidly, and therefore begins to set and harden more quickly. Furthermore, the relative proportions of tricalcium silicate and dicalcium silicate have a significant effect on the rate of hardening as indicated in Table 3.3.

During hydration, any sodium and potassium salts within the Portland cement are released into the pore water of the concrete, giving rise to a highly alkaline matrix. This effectively inhibits corrosion of any reinforcing steel embedded within the concrete, but if active silica is present in any of the aggregates, it may react to form an alkali–silica gel which absorbs water, swells and causes cracking of the concrete. This alkali–silica reaction can, however, be effectively prevented by limiting the total alkali content in the cement to less than 3 kg/m³. (Cement manufacturers normally specify alkali content in terms of equivalent percentage of sodium oxide.)

**TYPES OF CEMENT**

Cements are classified primarily on the main constituents such as Portland cement or blastfurnace cement. (In addition, there may be minor constituents up to 5% and also additives up to 1% by weight.)

The standard BS EN 197-1: 2000 lists five main types of cement:

- CEM I Portland cement
- CEM II Portland-composite cement
- CEM III Blastfurnace cement
- CEM IV Pozzolanic cement
- CEM V Composite cement

Within these five main types of cement, a wide range of permitted additional constituents, including silica fume, natural or industrial pozzolanas, calcareous or siliceous fly ash and burnt shale, may be incorporated. The full range of products is listed in Table 3.5, but not all are commercially available within the UK. Current standard production in addition to Portland cement includes Portland-limestone cement, Portland-fly ash cement and low early strength blastfurnace cement, but these composite cements account for only about 5% of the UK market. Calcium aluminate cement (also known as high alumina cement, HAC) has a totally different formulation compared with the range of Portland cements based on calcium silicates.

**STRENGTH CLASSES OF CEMENT**

The standard strength classes of cement are based on the 28 day compressive strength of mortar prisms, made and tested to the requirements of BS EN 196-1: 2005. The test uses specimens which are 40 × 40 × 160 mm, cast from a mix of 3 parts of CEN (European Committee for Standardisation) standard sand, 1 part of cement and 0.5 part of water. The sample is vibrated and cured for the appropriate time, then broken in halves and compression tested across the 40 mm face. Three specimens are used to determine a mean value from the six pieces.

Each cement strength class (32.5, 42.5 and 52.5) has sub-classes associated with the high early (R) and the
<table>
<thead>
<tr>
<th>Cement</th>
<th>Type</th>
<th>Notation</th>
<th>Portland cement clinker content (%)</th>
<th>Additional main constituent (%)</th>
<th>Sulphate resistance group</th>
</tr>
</thead>
<tbody>
<tr>
<td>Portland cement</td>
<td>I</td>
<td>CEM I</td>
<td>95–100</td>
<td>0</td>
<td>A</td>
</tr>
<tr>
<td>Portland slag cement</td>
<td>II</td>
<td>CEM II/A-S</td>
<td>80–94</td>
<td>6–20</td>
<td>A</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CEM II/B-S</td>
<td>65–79</td>
<td>21–35</td>
<td>A</td>
</tr>
<tr>
<td>Portland silica fume cement</td>
<td>II</td>
<td>CEM II/A-D</td>
<td>90–94</td>
<td>6–10</td>
<td>A</td>
</tr>
<tr>
<td>Portland pozzolana cement</td>
<td>II</td>
<td>CEM II/A-P</td>
<td>80–94</td>
<td>6–20</td>
<td>A</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CEM II/B-P</td>
<td>65–79</td>
<td>21–35</td>
<td>A</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CEM II/A-Q</td>
<td>80–94</td>
<td>6–20</td>
<td>A</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CEM II/B-Q</td>
<td>65–79</td>
<td>21–35</td>
<td>A</td>
</tr>
<tr>
<td>Portland fly ash cement</td>
<td>II</td>
<td>CEM II/A-V</td>
<td>80–94</td>
<td>6–20</td>
<td>A</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CEM II/B-V</td>
<td>65–79</td>
<td>21–35</td>
<td>A for class 21–24%</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>D for class ≥25%</td>
</tr>
<tr>
<td>Portland burnt shale cement</td>
<td>II</td>
<td>CEM II/A-T</td>
<td>80–94</td>
<td>6–20</td>
<td>A</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CEM II/B-T</td>
<td>65–79</td>
<td>21–35</td>
<td>A</td>
</tr>
<tr>
<td>Portland limestone cement</td>
<td>II</td>
<td>CEM II/A-L</td>
<td>80–94</td>
<td>6–20</td>
<td>B for class 32.5</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CEM II/B-L</td>
<td>65–79</td>
<td>21–35</td>
<td>C for class ≥42.5</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CEM II/A-LL</td>
<td>80–94</td>
<td>6–20</td>
<td>B for class 32.5</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CEM II/B-LL</td>
<td>65–79</td>
<td>21–35</td>
<td>C for class ≥42.5</td>
</tr>
<tr>
<td>Portland composite cement</td>
<td>II</td>
<td>CEM II/A-M</td>
<td>80–94</td>
<td>6–20</td>
<td>A</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CEM II/B-M</td>
<td>65–79</td>
<td>21–35</td>
<td>A</td>
</tr>
<tr>
<td>Blastfurnace cement</td>
<td>III</td>
<td>CEM III/A</td>
<td>35–64</td>
<td>36–65</td>
<td>A, or D for strictly controlled levels of tricalcium aluminate</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CEM III/B</td>
<td>20–34</td>
<td>66–80</td>
<td>A, or F for strictly controlled levels of tricalcium aluminate</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CEM III/C</td>
<td>5–19</td>
<td>81–95</td>
<td></td>
</tr>
<tr>
<td>Pozzolanic cement</td>
<td>IV</td>
<td>CEM IV/A</td>
<td>65–89</td>
<td>11–35</td>
<td>E</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CEM IV/B</td>
<td>45–64</td>
<td>36–55</td>
<td>E (BS EN 14216)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>VLH IV/B</td>
<td>45–64</td>
<td>36–55</td>
<td>E (BS EN 14216)</td>
</tr>
<tr>
<td>Composite</td>
<td>V</td>
<td>CEM V/A</td>
<td>40–64</td>
<td>36–60</td>
<td>G (BS 4027: 1996)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CEM V/B</td>
<td>20–39</td>
<td>61–80</td>
<td></td>
</tr>
</tbody>
</table>

**Notes:**
- The code letters used in the European Standard are:
  - D silica fume, F filler, K Portland cement clinker, L/LL limestone, M mixed, P natural pozzolana, Q industrial pozzolana, S granulated blastfurnace slag, T burnt shale, V siliceous fly ash, W calcareous fly ash.
  - (Limestone LL has a total organic carbon content limit of 0.2%; limestone L has a total organic carbon content limit of 0.5%.)
  - Grouping with respect to sulphate resistance A (low) to G (high) resistance (BRE Special Digest 1: 2005).
  - VLH refers to very low heat cements (BS EN 14216: 2004).
ordinary (N) development of early strength (Table 3.6). The strength classes and sub-classes give production standards for cements, but do not specify how a particular mix of cement, aggregate and admixtures will perform as a concrete; this needs to be determined by separate testing.

The most commonly used cement within the UK (formerly ordinary Portland cement or OPC) is currently designated to the standard BS EN 197-1: 2000 as:

<table>
<thead>
<tr>
<th>Strength class</th>
<th>Compressive strength (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Early strength</td>
</tr>
<tr>
<td></td>
<td>2 day minimum</td>
</tr>
<tr>
<td>32.5N</td>
<td>16.0</td>
</tr>
<tr>
<td>32.5R</td>
<td>10</td>
</tr>
<tr>
<td>42.5N</td>
<td>20</td>
</tr>
<tr>
<td>42.5R</td>
<td>30</td>
</tr>
<tr>
<td>52.5N</td>
<td>30</td>
</tr>
<tr>
<td>52.5R</td>
<td>30</td>
</tr>
</tbody>
</table>

Notes:
The code letters in the standards are: (N) ordinary early strength development; (R) high early strength development.

<table>
<thead>
<tr>
<th>type of cement</th>
<th>strength class</th>
<th>ordinary early strength development</th>
</tr>
</thead>
<tbody>
<tr>
<td>CEM I 42.5 N</td>
<td>42.5</td>
<td></td>
</tr>
<tr>
<td>CEM I 42.5N</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

High early strength Portland cement is designated as:

<table>
<thead>
<tr>
<th>type of cement</th>
<th>strength class</th>
<th>high early strength development</th>
</tr>
</thead>
<tbody>
<tr>
<td>CEM I 42.5 R</td>
<td>42.5</td>
<td></td>
</tr>
<tr>
<td>CEM I 42.5R</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Low early strength low heat blastfurnace cement with a granulated blastfurnace slag content between 81% and 95% and a strength class of 32.5 is designated as:

<table>
<thead>
<tr>
<th>type of cement</th>
<th>strength class</th>
<th>low early strength development and low heat</th>
</tr>
</thead>
<tbody>
<tr>
<td>CEM III/C 32.5</td>
<td>32.5</td>
<td></td>
</tr>
<tr>
<td>CEM III/C 32.5LH</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Portland limestone cement with between 6% and 20% limestone of 0.5% total organic content, a strength class of 32.5 and normal early strength is designated as:

<table>
<thead>
<tr>
<th>type of cement</th>
<th>sub-type</th>
<th>strength class</th>
<th>ordinary early strength development</th>
</tr>
</thead>
<tbody>
<tr>
<td>CEM II/A-L 32.5N</td>
<td></td>
<td>32.5</td>
<td></td>
</tr>
</tbody>
</table>

Portland cements—strength classes 32.5, 42.5 and 52.5

The Portland cement classes 32.5, 42.5 and 52.5 correspond numerically to their lower characteristic strengths at 28 days. The 32.5 and 42.5 classes have upper characteristic strengths which are 20 MPa greater than the lower characteristic strengths, as designated by the class number. The class 52.5 has no upper strength limit. Statistically, the tested strengths must fall with no more than 5% of the tests below the lower limit or 10% of the tests above the upper limit. Thus class 42.5 Portland cement has a strength within the range 42.5–62.5 MPa, with a maximum of 5% of test results being below 42.5 MPa and a maximum of 10% of the test results being above 62.5 MPa.

Each class also has lower characteristic strength values at 2 days, except for class 32.5, which has a lower characteristic strength at 7 days. Where high early strength is required, for example to allow the early removal of formwork in the manufacture of precast concrete units, class 52.5 or class 42.5R is used. These Portland cements are more finely ground than class 42.5 to enable a faster hydration of the cement in the early stages. Class 32.5 cements for general-purpose and DIY use, frequently contain up to 1% additives to improve workability and frost resistance, together
Table 3.7 Very low heat special cements to BS EN 14216: 2004

<table>
<thead>
<tr>
<th>Cement</th>
<th>Type</th>
<th>Notation</th>
<th>Portland cement clinker content (%)</th>
<th>Additional main constituent (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blastfurnace cement</td>
<td>III</td>
<td>VLH III/B</td>
<td>20–34</td>
<td>66–80</td>
</tr>
<tr>
<td></td>
<td></td>
<td>VLH III/C</td>
<td>5–19</td>
<td>81–95</td>
</tr>
<tr>
<td>Pozzolanic cement</td>
<td>IV</td>
<td>VLH IV/A</td>
<td>65–89</td>
<td>11–35</td>
</tr>
<tr>
<td></td>
<td></td>
<td>VLH IV/B</td>
<td>45–64</td>
<td>36–55</td>
</tr>
<tr>
<td>Composite cement</td>
<td>V</td>
<td>VLH V/A</td>
<td>40–64</td>
<td>18–30 blastfurnace slag</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>18–30 pozzolana and fly ash</td>
</tr>
<tr>
<td></td>
<td></td>
<td>VLH/VB</td>
<td>20–38</td>
<td>31–50 blastfurnace slag</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>31–50 pozzolana and fly ash</td>
</tr>
</tbody>
</table>

with up to 5% minor additional constituents such as pulverised-fuel ash, granulated blastfurnace slag or limestone filler. Portland cement of strength class 42.5 accounts for approximately 90% of the total cement production within the UK.

White Portland cement
White Portland cement is manufactured from materials virtually free of iron oxide and other impurities, which impart the grey colour to Portland cement. Generally, china clay and limestone are used and the kiln is fired with natural gas or oil rather than pulverised coal. Iron-free mills are used for the grinding process to prevent colour contamination. Because of the specialist manufacturing processes, it is approximately twice the price of the equivalent grey product. To further enhance the whiteness up to 5% of white titanium oxide pigment may be added. The standard product is to strength class 52.5N. Typical applications include renderings, cast stone, pre-cast and in situ structural concrete and pointing.

Sulphate-resisting Portland cement
Sulphate-resisting Portland cement (BS 4027: 1996) is suitable for concrete and mortar in contact with soils and groundwater containing soluble sulphates up to the maximum levels (measured as sulphur trioxide) of 2% in soil or 0.5% in groundwater. In normal Portland cements the hydrated tricalcium aluminate component is vulnerable to attack by soluble sulphates, but in sulphate-resisting Portland cement this component is restricted to a maximum of 3.5%. For maximum durability a high-quality, dense, non-permeable concrete is required. Many sulphate-resisting cements are also defined as low alkali (LA) to BS 4027: 1996, containing less than 0.6% alkali (measured as sodium oxide). Thus durable concrete, without the risk of subsequent alkali–silica reaction, can be manufactured with alkali-reactive aggregates, using up to 500 kg/m³ of cement, provided no other alkalis are present.

Very low-heat special cements
Low-heat Portland cement (BS 1370: 1979) is appropriate for use in mass concrete, where the rapid internal evolution of heat could cause cracking. It contains a higher proportion of dicalcium silicate which hardens and evolves heat more slowly. The range of very low heat special cements, listed in Table 3.7, includes products based on blastfurnace, pozzolanic and composite cements (BS EN 14216: 2004). Very low heat special cements are appropriate for use only in massive constructions such as dams, but not for bridges or buildings.

Blended Portland cements
Blended Portland cements include not only masonry cement, with its specific end use, but also the wide range of additional materials classified within the European Standard EN 197-1: 2000.

Masonry cements
Portland cement mortar is unnecessarily strong and concentrates any differential movement within brickwork or blockwork into a few large cracks, which are unsightly and may increase the risk of rain penetration. Masonry cement produces a weaker mortar, which accommodates some differential movement, and ensures a distribution of hairline cracks within joints, thus preserving the integrity of the bricks and blocks. Masonry cements contain water-retaining mineral fillers, usually ground limestone, and air-entraining agents to give a higher workability than unblended Portland cement. They should not normally be blended with further admixtures but mixed with building sand in ratios between 1:4 and 1:6.5
Table 3.8  Strength classes of masonry cements to European Standard BS EN 413-1: 2004

<table>
<thead>
<tr>
<th>Strength classes</th>
<th>Compressive strength (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>7 day early strength</td>
</tr>
<tr>
<td>MC 5</td>
<td>≥7</td>
</tr>
<tr>
<td>MC 12.5</td>
<td>≥7</td>
</tr>
<tr>
<td>MC 12.5X</td>
<td>≥7</td>
</tr>
<tr>
<td>MC 22.5X</td>
<td>≥10</td>
</tr>
</tbody>
</table>

Notes:
Masonry cement is designated by MC; the letter X refers to cements which do not incorporate air-entraining agent.

depending on the degree of exposure of the brick or blockwork. The air entrained during mixing increases the durability and frost resistance of the hardened mortar. Masonry cement is also appropriate for use in renderings but not for floor screeds or concreting. It is therefore generally used as an alternative to Portland cement with hydrated lime or plasticiser. Inorganic pigments, except those containing carbon black, may be incorporated for visual effect. The strength classes for masonry cement are listed in Table 3.8.

Portland slag and blastfurnace cements
Granulated blastfurnace slag, formerly termed ground blastfurnace slag (GGBS), is a cementitious material, which in combination with Portland cement and appropriate aggregates, makes a durable concrete. The material is a by-product of the iron-making process within the steel industry. Iron ore, limestone and coke are fed continuously into blastfurnaces, where at 1500°C they melt into two layers. The molten iron sinks, leaving the blastfurnace slag floating on the surface, from where it is tapped off at intervals. The molten blastfurnace slag is rapidly cooled by water quenching in a granulator or pelletiser to produce a glassy product. After drying, the blastfurnace slag granules or pellets are ground to the fine off-white powder, granulated blastfurnace slag. The composition of the material is broadly similar to that of Portland cement as illustrated in Table 3.9.

Granulated blastfurnace slag may be intimately ground with Portland cement clinker in the cement mill, although usually it is mixed with Portland cement on site. The Standard (BS EN 15167: 2006) gives the constituent specification and the Standard BS EN 197-4: 2004 refers to blastfurnace cements with mixes from 36% to 80 and 95% of granulated blastfurnace slag, respectively (Table 3.10). By using 50% granulated blastfurnace slag as an alternative to Portland cement in a typical mix, emissions of carbon dioxide in the overall concrete production process are reduced by approximately 40%.

Concrete manufactured from a blend of Portland and granulated blastfurnace slag cements has a lower permeability than Portland cement alone; this enhances resistance to attack from sulphates and weak acids also to the ingress of chlorides which can cause rapid corrosion of steel reinforcement, for example in marine environments and near roads subjected to de-icing salts. Sulphate attack is also reduced by the decrease in tricalcium aluminate content. The more gradual hydration of granulated blastfurnace slag cement evolves less heat and more slowly than Portland cement alone; thus, a 70% granulated blastfurnace slag mix can be used for mass concrete, where otherwise a significant temperature rise could cause cracking. The slower evolution of heat is associated with a more gradual development of strength over the first 28 day period. However, the ultimate strength of the mature concrete is comparable to that of the equivalent Portland cement. The initial set with granulated blastfurnace slag blends is slower than for Portland cement alone, and the fresh concrete mixes are more plastic, giving better flow for placing and full compaction. The risk of alkali–silica reaction caused by reactive silica aggregates can be reduced by the use of granulated blastfurnace slag to reduce the active alkali content of the concrete mix to below the
Table 3.11  Strength classes of low early strength blastfurnace cements to British Standard BS EN 197-4: 2004

<table>
<thead>
<tr>
<th>Strength classes</th>
<th>Compressive strength (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>2 day early strength</td>
</tr>
<tr>
<td>32.5L</td>
<td>—</td>
</tr>
<tr>
<td>42.5L</td>
<td>—</td>
</tr>
<tr>
<td>52.5L</td>
<td>≥10</td>
</tr>
</tbody>
</table>

critical 3.0 kg/m^3 level. The classes for low early strength blastfurnace cements are listed in Table 3.11.

*Portland fly ash and pozzolanic cements*

Pozzolanic materials are natural or manufactured materials containing silica, which react with calcium hydroxide produced in the hydration of Portland cement to produce further cementitious products. Within the UK, natural volcanic pozzolanas are little used, but fly ash, formerly termed pulverised-fuel ash (PFA), the waste product from coal-fired electricity-generating stations, is used either factory mixed with Portland cement or blended in on site. Portland fly ash cement cures and evolves heat more slowly than Portland cement; it is therefore appropriate for use in mass concrete to reduce the risk of thermal cracking. Additions of up to 25% fly ash in Portland cement are often used; the concrete produced is darker than that with Portland cement alone. Concrete made with blends of 25–40% by weight of fly ash in Portland cement has good sulphate-resisting properties. However, in the presence of groundwater with high magnesium concentrations, sulphate-resisting Portland cement should be used. Fly ash concretes also have enhanced resistance to chloride ingress, which is frequently the cause of corrosion to steel reinforcement.

The fly ash produced in the UK by burning pulverised bituminous coal is siliceous, containing predominantly reactive silica and alumina. In addition to siliceous fly ash, the European Standard EN 197-1: 2000 does allow for the use of calcareous fly ash, which additionally contains active lime, giving some self-setting properties. The range of fly ash suitable for concrete is defined in the standard BS EN 450-1: 2005. Natural pozzolanas of volcanic origin and industrial pozzolanas from other industrial processes in Europe are used with Portland cement and are categorised under EN 197-1: 2000 as pozzolanic cements. Research is in progress to determine whether proportions greater than the currently permitted 40% of PFA in Portland cement are feasible, as this would significantly reduce manufacturing carbon dioxide emissions.

*Portland limestone cement*

The addition of up to 5% limestone filler to Portland cement has little effect on its properties. The addition of up to 25% limestone gives a performance similar to that of Portland cement with a proportionally lower cementitious content; thus, if equivalent durability to Portland cement is required, then cement contents must be increased. The two categories of limestone for Portland limestone cement are defined by their total organic carbon (TOC) content; LL refers to a maximum of 0.20% and L to a maximum of 0.50% by mass.

*Silica fume*

Silica fume or microsilica, a by-product from the manufacture of silicon and ferro-silicon, consists of ultrafine spheres of silica. The material, because of its high surface area, when blended as a minor addition to Portland cement, increases the rate of hydration, giving the concrete a high early strength and also a reduced permeability. This in turn produces greater resistance to chemical attack and abrasion. Silica fume may be added up to 5% as a filler, or in Portland silica fume cement to between 6% and 10%.

*Burnt shale*

Burnt shale is produced by heating oil shale to 800°C in a kiln. It is similar in nature to blastfurnace slag, containing mainly calcium silicate and calcium aluminate, but also silica, lime and calcium sulphate. It is weakly cementitious. The European Standard EN 197-1: 2000 allows for the use of burnt shale as a filler to 5%, or between 6% and 35% in Portland burnt shale cement.

*Fillers*

Fillers up to 5% by weight of the cement content may be added to cements to the standard EN 197-1: 2000. They should be materials which do not increase the water requirements of the cement. Fillers may be any of the permitted alternative main constituents (e.g. granulated blast furnace slag, pozzolanas, fly ash, burnt shale, silica fume or limestone), or other inorganic materials, provided that they are not already present as one of the main constituents. The most common fillers are limestone and either raw meal or partially calcined material from the cement-making process.
CEMENT ADMIXTURES

Admixtures may be defined as materials that are added in small quantities to mortars or concretes during mixing, in order to modify one or more of their physical or visual properties.

Plasticisers

Plasticisers, or water-reducing admixtures, are added to increase the workability of a mix, thus enabling easier placing and compaction. Where increased workability is not required, water reducers may be used to lower the water/cement ratio, giving typically a 15% increase in strength and better durability. The plasticisers, which are usually lignosulphonates or hydroxylated polymers, act by dispersing the cement grains. Some air entrainment may occur with the lignosulphonates, causing a 6% reduction in crushing strength for every 1% of air entrained.

Superplasticisers

Superplasticisers, such as sulphonated naphthalene or sulphonated melamine formaldehyde, when added to a normal 50 mm slump concrete produce a flowing, self-levelling or self-compacting concrete (SCC) which can be placed, even within congested reinforcement, without vibration. Alternatively, significantly reduced water contents can be used to produce early- and ultimately higher-strength concretes. As the effect of superplasticisers lasts for less than an hour, the admixture is usually added to ready-mixed concrete on site prior to discharge and placing. Standard concrete additives, fillers and steel or polypropylene fibres may be incorporated into self-compacting concrete which can be pumped or delivered by skip or chute. Good-quality off-the-form surface finishes can be achieved, especially with timber formwork. Self-leveling mixes for screeds between 3 and 20 mm thick can be adjusted to take light foot traffic after 3 to 24 hours. Renovation mixes, usually incorporating fibre-mat reinforcement, can be used over a range of existing floor surfaces to thicknesses usually in the range 4–30 mm.

Accelerators

Accelerators increase the rate of reaction between cement and water, thus increasing the rate of set and development of strength. This can be advantageous in precasting, where early removal of the formwork is required, and in cold weather when the heat generated speeds up the hardening processes and reduces the risk of frost damage. Only chloride-free accelerators, such as calcium formate, should be used in concrete, mortar or grout where metal will be embedded, because calcium chloride accelerators can cause extensive metallic corrosion. Accelerators producing a rapid set are not normally used within structural concrete.

Retarders

Retarders, typically phosphates or hydroxycarboxylic acids, decrease the rate of set, thus extending the time between initial mixing and final compaction, but they do not adversely affect 28 day strength. Retarders may be applied to formwork, to retard the surface concrete where an exposed aggregate finish is required by washing after the formwork is struck. Retarders are also frequently used in ready-mixed mortars to extend their workable life up to 36 hours. The mortars are usually delivered on site in date-marked containers of 0.3 m³ capacity.

Air-entraining admixtures

Air-entraining admixtures, typically wood resins or synthetic surfactants, stabilise the tiny air bubbles which become incorporated into concrete or mortar as it is mixed. The bubbles, which are between 0.05 and 0.5 mm in diameter, do not escape during transportation or vibration, improve the workability of the mix, reduce the risk of segregation and greatly enhance frost resistance. However, the incorporation of void space within concrete decreases its crushing strength by 6% for every 1% of air entrained; thus, for a typical 3% addition of entrained air, a reduction of 18% in crushing strength is produced. This is partially offset by the increase in plasticity, which generally produces a higher-quality surface and allows a lower water content to be used. The increased cohesion of air-entrained concrete may trap air against moulded vertical formwork reducing the quality of the surface.

Water-resisting admixtures

The water penetration through concrete can be reduced by incorporation of hydrophobic materials, such as stearates and oleates, which coat the surface of pores and, by surface tension effects, discourage the penetration of damp. The use of water-reducing admixtures also reduces water penetration by reducing
the water/cement ratio, thus decreasing pore size within the concrete. For mortars and renders a styrene–butadiene latex emulsion admixture can be used to reduce permeability.

**Foaming agents**

Foamed concrete or mortar contains up to 80% by volume of void space, with densities as low as 300 kg/m³ and 28 day strengths between 0.2 and 20 MPa. It is typically produced by blending cement, fine sand or fly ash and water into a preformed foam or by mechanically foaming the appropriate mix using a foaming surfactant. For the low densities below 600 kg/m³, no fillers are included, but for densities up to 1600 kg/m³, limestone dust and concreting sand may be incorporated. Foamed concrete is free flowing, can be pumped and requires no compaction. When set the material offers good frost resistance and thermal insulation. It is therefore used for trench reinstatement, filling cellars or providing insulation under floors or in flat roofs.

**Pumping agents**

Not all concrete mixes are suitable for pumping. Mixes low in cement or with some lightweight aggregates tend to segregate and require thickening with a pumping agent. Conversely, high-cement content mixes require plasticising to make them pumpable. A range of pumping agents is therefore produced to suit the requirements of various concrete mixes. Lightweight aggregate concrete is often pumped into place for floor slabs.

**Pigments**

A wide range of coloured pigments is available for incorporation into concrete and mortars (BS EN 12878: 2005). Titanium oxide can be added to enhance the whiteness of white cement. Carbon black is used with grey Portland cement, although the black loses intensity with weathering. The most common colours are the browns, reds and yellows produced with synthetic iron, chromium and manganese oxides, and also with complexes of cobalt, aluminium, nickel and antimony. Additionally, ultramarine and phthalocyanine extend the range of the blues and greens. The depth and shade of colour depends on the dose rate (between 1% and 10%), also the colour of the sand and any other aggregates. To produce pastel shades, pigments can be added to white Portland cement.

**CALCIUM ALUMINATE CEMENT**

Calcium aluminate cement, also known as high alumina cement (HAC), is manufactured from limestone and bauxite (aluminium oxide). The ores, in roughly equal proportions, are charged together into a vertical furnace, which is heated to approximately 1600°C (Fig. 3.4). The mixture melts and is continuously run
off into trays, where it cools to produce the clinker, which is then milled, producing calcium aluminate cement to BS EN 14647: 2005. The dark grey cement composition differs from that of Portland cement as it is based on calcium aluminates rather than calcium silicates. Although calcium aluminate cement can be produced over a wide range of compositions, the standard product has a 40% alumina content.

Calcium aluminate cement should not be used for foundations or structural purposes, but only for specific heat-resisting applications and where deterioration rates can be predicted. However, it is useful where rapid strength gain is required, allowing the fast removal of formwork within 6 to 24 hours. The fast evolution of heat allows concreting to take place at low temperatures. The material also has good heat-resistant properties, so it may be used to produce refractory concrete. When mixed with Portland cement it produces a rapid-setting concrete, suitable for non-structural repairs and sealing leaks. Good-quality calcium aluminate cement is generally resistant to chemical attack by dilute acids, chlorides and oils, but not alkalis.

Some structural failures associated with calcium aluminate cement have been caused by conversion of the concrete, in which changes in the crystal structure, accelerated by high temperatures and humidity, have caused serious loss of strength, increased porosity and subsequent chemical attack. Depending on the degree of conversion, calcium aluminate cement becomes friable and a deeper brown in colour; the exact degree of conversion can only be determined by chemical analysis of a core sample. It is now recognised that such failures can be prevented by using a minimum cement content of 400 kg/m³, limiting the water/cement ratio to a maximum of 0.4, and ensuring controlled curing during the 6- to 24-hour initial hardening stage. The concrete should be covered or sprayed to prevent excessive water loss, particularly where substantial increases in temperature may occur.

Additionally, in order to prevent alkaline hydrolysis of the concrete, aggregates containing soluble alkalis should not be used; hard limestone is generally considered to be the best aggregate. Coloured calcium aluminate cement concrete has the advantage that it is free from calcium hydroxide which causes efflorescence in Portland cements. The BRE Special Digest SD3 (2002) gives methods for assessing existing calcium aluminate cement concrete [high alumina cement concrete (HACC)] constructions and suggests appropriate remedial actions. In some cases where the depth conversion of HACC structural members is significant, with time there is an increasing risk of reinforcement corrosion.

LOW ENVIRONMENTAL IMPACT CEMENTS

Current carbon dioxide emissions from the manufacture of Portland cement in the UK are estimated at 0.82 tonne of CO₂ per tonne of CEM 1 cement. At the current rate of world production this relates to approximately 7.5% of global CO₂ emissions. While the Portland cement industry has significantly increased production efficiency by using the dry manufacturing process and increasing the blended content, ultimately limestone has to be calcined at high temperatures to drive off carbon dioxide.

The overall carbon footprint of Portland cement can be reduced by the addition of ground granulated blast furnace slag (GGBS), natural pozzolanas or fly ash (PFA), and blends containing much higher proportions of pulverised fuel ash than are currently permitted for Portland fly ash cement CEM II are under review.

Belite cement

The main components of Portland cement are alite (tricalcium silicate) and belite (dicalcium silicate). These are formed at 1450°C and 1200°C, respectively.

Belite cements can be manufactured from finely ground raw materials at kiln temperatures of 1350°C, and as the CaO content of belite is less than that of alite, less CO₂ per tonne is driven off during the calcination process. The total CO₂ emissions for making belite cement are estimated at 0.69 tonne/tonne compared to 0.88 tonne/tonne for alite cement. This equates roughly to a 16% energy saving for belite cement compared to Portland cement. Belite cement has good long-term properties but develops its strength very slowly. Therefore, its potential is enhanced by blending with a more active component.

Calcium sulphaaluminate cement

Calcium sulphaaluminate (CSA) is produced from a mixture of bauxite (aluminium oxide), anhydrite (calcium sulphate) and limestone at kiln temperatures between 1000°C and 1300°C.

$$3CaCO_3 + 3Al_2O_3 + CaSO_4 \rightarrow 4CaO \cdot 3Al_2O_3 \cdot SO_3 + 3CO_2$$

calcium sulphaaluminate
Considerably less CO₂ is produced per tonne of calcium sulfoaluminate cement, compared to the production of alite or belite cement. Calcium sulfoaluminate cements develop early strength, so the blending of belite and calcium sulfoaluminate to produce belite-sulfoaluminate cement offers some potential as an alternative to Portland cement. The addition of cheap Fe₂O₃ (iron oxide) in the raw meal leads to the formation of ferrite (calcium ferroaluminate, C₄AF), by the partial substitution of the Al₂O₃ (alumina) component in calcium sulfoaluminate. Belite-sulfoaluminate-ferrite cement therefore has the potential for reduced manufacturing costs. Typical compositions of belite-sulfoaluminate-ferrite cement currently used in China are 35–70% CSA, belite <30% and ferrite 10–30%. Compared to manufacturing Portland cement CEMI, the reduction in CO₂ emissions is estimated at around 20%.

Magnesium oxide cements

Two types of magnesium oxide-based cements are under development, based on magnesium carbonate (magnesite) and magnesium silicate, respectively. Magnesium oxide is produced from magnesium carbonate by heating to 650°C. After the bound carbon dioxide has been evolved, the magnesium oxide is hydrated to magnesium hydroxide (brucite), which acts as the cement binder. This material is rapidly carbonated in use by reabsorbing most of the CO₂ liberated in its manufacture. Magnesium oxide can also be produced from magnesium silicates by heating the mineral to between 650°C and 700°C. In this case no carbon dioxide is evolved in the manufacture, but CO₂ is subsequently absorbed in its use as a cement. Manufacturing a tonne of this cement would produce 0.4 tonnes of CO₂, but its use would absorb 1.1 tonnes leaving a reduction of 0.7 tonnes from the atmosphere. This product therefore has the potential to be ‘zero carbon’ or better. Magnesium silicate minerals are abundant worldwide, but not in large quantities within the UK. This product is the subject of a major research and development programme.

Concrete

Concrete is a mixture of cement, aggregates and water together with any other admixtures which may be added to modify the placing and curing processes or the ultimate physical properties. Initially, when mixed concrete is a plastic material, which takes the shape of the mould or formwork. When hardened it may be a dense, load-bearing material or a lightweight, thermally insulating material, depending largely on the aggregates used. It may be reinforced or prestressed by the incorporation of steel.

Most concrete is crushed and recycled at the end of its useful life, frequently as hard core for new construction work. However, a growth in the use of recycled aggregates for new concrete can be anticipated, as this will have a significant environmental gain in reducing the demand on new aggregate extraction.

AGGREGATES FOR CONCRETE

Aggregates form a major component of concretes, typically approximately 80% by weight in cured mass concrete. Aggregate properties, including crushing strength, size, grading and shape, have significant effects on the physical properties of the concrete mixes and hardened concrete. Additionally, the appearance of visual concrete can be influenced by aggregate colour and surface treatments. The standard BS EN 12620: 2002 specifies the appropriate properties including materials, size, grading and shape. Aggregates for concrete are normally classified as lightweight, dense or high-density. Standard dense aggregates are classified by size as fine (i.e. sand) or coarse (i.e. gravel). Additionally, steel or polypropylene fibres or gas bubbles may be incorporated into the mix for specialist purposes.

Dense aggregates

Source and shape

Dense aggregates are quarried from pits and from the seabed. In the south-east of England, most land-based sources are gravels, typically flint, whereas further north and west, both gravels and a variety of crushed quarried rocks are available. Marine aggregates which account for 18% of production in England and Wales may require washing to remove deleterious matter such as salts, silt and organic debris. The total chloride content should be monitored to ensure that it is within the limits to BS 8500: 2006 for reinforced or unreinforced concrete as appropriate. This may be achieved by using well-drained unwashed marine sand in conjunction with land-based coarse aggregates.
The shape of aggregates can significantly affect the properties of the mix and cured concrete. Generally, rounded aggregates require a lower water content to achieve a given mix workability, compared to the equivalent mix using angular aggregates. However, cement paste ultimately bonds more strongly to angular aggregates with rough surfaces than to the smoother gravels, so a higher crushing strength can be achieved with crushed rocks as aggregate. Excessive proportions of long and flaky coarse aggregate should be avoided, as they can reduce the durability of concrete.

**Aggregate size**

For most purposes the maximum size of aggregate should be as large as possible consistent with ease of placement within formwork and around any steel reinforcement. Typically, 20 mm aggregate is used for most construction work, although 40 mm aggregate is appropriate for mass concrete, and a maximum of 10 mm for thin sections. The use of the largest possible aggregate reduces the quantity of sand and therefore cement required in the mix, thus controlling shrinkage and minimising cost. Large aggregates have a low surface area/volume ratio, and therefore produce mixes with greater workability for a given water/cement ratio, or allow water/cement ratios to be reduced for the same workability, thus producing a higher crushing-strength concrete.

**Grading**

To obtain consistent quality in concrete production, it is necessary to ensure that both coarse and fine aggregates are well graded. A typical continuously graded coarse aggregate will contain a good distribution of sizes, such that the voids between the largest stones are filled by successively smaller particles down to the size of the sand. Similarly, a well-graded sand will have a range of particle sizes, but with a limit on the proportion of fine clay or silt, because too high a content of fines (of size less than 0.063 mm) would increase the water and cement requirement for the mix. Usually a maximum of 3% fines is considered non-harmful. This overall grading of aggregates ensures that all void spaces are filled with the minimum proportion of fine material and expensive cement powder. In certain circumstances, coarse aggregate may be graded as single-sized or gap graded. The former is used for controlled blending in designed mixes whilst the latter is used particularly for exposed aggregate finishes on visual concrete. Sands are classified into three categories according to the proportion passing through a 0.500 mm sieve: coarse C (5–45%), medium M (30–70%) and fine F (55–100%). Only the coarse and medium categories of sands should be used for heavy-duty concrete floor finishes.

**Sampling and sieve analysis**

To determine the grading of a sample of coarse or fine aggregate, a representative sample has to be subjected to a sieve analysis. Normally, at least ten samples would be taken from various parts of the stock pile, and these would be reduced down to a representative sample using a riffle box, which successively divides the sample by two until the required test volume is obtained (Fig. 3.5). Aggregate gradings are determined by passing the representative sample through a set of standard sieves (BS EN 12620 + A1 : 2008). Aggregate size is specified by the lower \(d\) and upper \(D\) sieve sizes. For coarse aggregates the sieve sizes are 63, 31.5, 16, 8, 4, 2 and 1 mm and for fine aggregates 4, 2, 1, 0.250 and 0.063 mm. Coarse aggregates are usually defined as having a minimum size \(d\) of 2 mm, while fine aggregates often have a maximum size \(D\) of 4 mm. The sieve analysis is determined by assessing the cumulative percentage passing through each sieve size. This is plotted against the sieve size and compared to the limits as illustrated for a typical coarse aggregate (Fig. 3.6).

Aggregates for concreting are normally batched from stockpiles of 20 mm coarse aggregate and concreting sand in the required proportions to ensure consistency, although all-in aggregate, which contains
both fine and coarse aggregates, is also available as a less well controlled, cheaper alternative, where a lower grade of concrete is acceptable. Where exceptionally high control on the mix is required, single-size aggregates may be batched to the customer’s specification. The batching of aggregates should normally be done by weight, as free surface moisture, particularly in sand, can cause bulking, which is an increase in volume by up to 40% (Fig. 3.7). Accurate batching must take into account the water content in the aggregates in the calculations of both the required weight of aggregates and the quantity of water to be added to the mix.

**Impurities within aggregates**

Where a high-quality exposed concrete finish is required, the aggregate should be free of iron pyrites, which causes spalling and rust staining of the surface. Alkali–silica reaction (ASR) can occur when active silica, present in certain aggregates, reacts with the alkalis within Portland cement causing cracking.

**Recycled aggregates**

The proportion and nature of constituent materials in recycled aggregate for concrete must be determined according to pr EN 933-11 (pending) and declared according to BS EN 12620: 2002 Amendment A1: 2008 as percentage limits by mass.

**Constituent categories of recycled coarse aggregate:**

- **Rc** Concrete, concrete products, mortar, concrete masonry units
- **Ru** Unbound aggregate, natural stone, hydraulically bound aggregate
- **Rb** Clay masonry units (bricks and tiles), calcium silicate bricks, aerated non-floating concrete
- **Ra** Bituminous materials
- **Rg** Glass
- **FL** Floating material in volume
- **X** Other material (clay, soil, gypsum plaster, metals, wood, plastic, rubber)

Quantities of deleterious material within recycled aggregates must be declared and carefully controlled, to prevent adverse effects on the quality of the concrete.

Current research is evaluating concrete made from a mixture of recycled aggregates, china clay aggregate waste from Cornwall (‘tip sand’ and ‘stent’ rock), together with a high proportion of pulverised fuel ash (PFA) replacing the Portland cement clinker. The mixture has approximately 30% less embodied energy than standard concrete.

**High-density aggregates**

Where radiation shielding is required, high-density aggregates such as barytes (barium sulphate), magnetite (iron ore), lead or steel shot are used. Hardened concrete densities between 3000 and 5000 kg/m$^3$, double that for normal concrete, can be achieved.


**Lightweight aggregates**

Natural stone aggregate concretes typically have densities within the range 2200–2500 kg/m$^3$, but where densities below 2000 kg/m$^3$ are required, then an appropriate lightweight concrete must be used.

Lightweight concretes in construction exhibit the following properties in comparison with dense concrete:

- they have enhanced thermal insulation but reduced compressive strength;
- they have increased high-frequency sound absorption but reduced sound insulation;
- they have enhanced fire resistance over most dense aggregate concretes (e.g. granite spalls);
- they are easier to cut, chase, nail, plaster and render than dense concrete;
- the reduced self-weight of the structure offers economies of construction;
- the lower formwork pressures enable the casting of higher lifts.

The three general categories of lightweight concrete are lightweight aggregate concrete, aerated concrete and no-fines concrete (Fig. 3.8).

Many of the lightweight aggregate materials are produced from by-products of other industrial processes or directly from naturally occurring minerals. The key exception is expanded polystyrene, which has the highest insulation properties, but is expensive due to its manufacture from petrochemical products.

**Pulverised fuel ash**
Pulverised fuel ash, or fly ash, is the residue from coal-fired electricity-generating stations. The fine fly ash powder is moistened, pelleted and sintered to produce a uniform lightweight PFA aggregate, which can be used in load-bearing applications.

**Foamed blastfurnace slag**
Blastfurnace slag is a by-product from the steel industry. Molten slag is subjected to jets of water, steam and compressed air to produce a pumice-like material. The foamed slag is crushed and graded to produce aggregate, which can be used in load-bearing applications. Where rounded pelletised expanded slag is required the material is further processed within a rotating drum.

**Expanded clay and shale**
Certain naturally occurring clay materials are pelletised, then heated in a furnace. This causes the evolution of gases which expand and aerate the interior, leaving a hardened surface crust. These lightweight aggregates may be used for load-bearing applications.

**Expanded perlite**
Perlite is a naturally occurring glassy volcanic rock, which, when heated almost to its melting point, evolves steam to produce a cellular material of low density. Concrete made with expanded perlite has good thermal insulation properties but low compressive strength and high drying shrinkage.

**Exfoliated vermiculite**
Vermiculite is a naturally occurring mineral, composed of thin layers like mica. When heated rapidly the layers separate, expanding the material by up to 30 times, producing a very lightweight aggregate. Exfoliated vermiculite concrete has excellent thermal insulation properties but low compressive strength and very high drying shrinkage.

**Expanded polystyrene**
Expanded polystyrene beads offer the highest level thermal insulation, but with little compressive strength. Polystyrene bead aggregate cement (PBAC) is frequently used as the core insulating material within precast concrete units.

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Fig. 3.8 Lightweight concretes
Aerated concrete

Aerated concrete (aircrete) is manufactured using foaming agents or aluminium powder as previously outlined in the section on foaming agents. Densities in the range 400–1600 kg/m² give compressive strengths ranging from 0.5 to 20 MPa. Drying shrinkages for the lowest-density materials are high (0.3%), but thermal conductivity can be as low as 0.1 W/m K offering excellent thermal insulation properties. Factory-autoclaved aerated concrete blocks have greatly reduced drying shrinkages and enhanced compressive strength over site-cured concrete. Aerated concrete is generally frost resistant but should be rendered externally to prevent excessive water absorption. The material is easily worked on site as it can be cut and nailed.

No-fines concrete

No-fines concrete is manufactured from single-sized aggregate (usually between 10 and 20 mm) and cement paste. Either dense or lightweight aggregates may be used, but care has to be taken in placing the mix to ensure that the aggregate remains coated with the cement paste. The material should not be vibrated. Drying shrinkage is low, as essentially the aggregate is stacked up within the formwork, leaving void spaces; these increase the thermal-insulation properties of the material in comparison with the equivalent dense concrete. The rough surface of the cured concrete forms an excellent key for rendering or plastering which is necessary to prevent rain, air or sound penetration. Dense aggregate no-fines concrete may be used for load-bearing applications.

Fibres

Either steel or polypropylene fibres may be incorporated into concrete, as an alternative to secondary reinforcement, particularly in heavily trafficked floor slabs. The fibres reduce the shrinkage and potential cracking that may occur during the initial setting and give good abrasion and spalling resistance to the cured concrete. The low-modulus polypropylene fibres, which do not pose a corrosion risk after carbonation of the concrete, enhance the energy-absorbing characteristics of the concrete giving better impact resistance. Steel fibres increase flexural strength as well as impact resistance but are more expensive. Alternatively, stainless steel fibres may be used where rust spots on the surface would be unacceptable. Typically, polypropylene fibres are added at the rate of 0.2% by weight (0.5% by volume) and steel at the rate of 3–4% by weight. Both polypropylene and steel fibre concretes can be pumped. Steel fibres to BS EN 14889-1: 2006 may be straight or deformed cold-drawn wire, alternatively sheet fibres. Polymer fibres to BS EN 14889-2: 2006 may be thick or thin monofilaments or fibrillated. (Glass-fibre reinforced cement is described in Chapter 11.)

Ultra-high-performance concrete

Ultra-high-performance concrete (UHPC) has six to eight times the compressive strength of traditional concrete. It is produced from a mixture of Portland cement, crushed quartz, sand, silica fume, superplasticiser, fibres and water with no aggregates larger than a few millimetres. Wollastonite (calcium silicate) filler may also be included in the mix. The fibres most frequently used are either high-strength steel for maximum strength or polypropylene (PP) of approximately 12 mm in length for lower load applications. The concrete can be cast into traditional moulds by gravity or pumped or even injection cast under pressure. When cast into traditional moulds, the material is self-leveling, so only slight external vibration of the formwork may be required to ensure complete filling. The material is designed for use without steel reinforcement bars.

Structural components in ultra-high-performance concrete may, after setting, be subjected to steam treatment for 48 hours at 90°C. This enhances durability and mechanical properties, eliminates shrinkage and reduces creep. The material does not spall under fire test conditions.

The enhanced compressive and flexural strengths of ductile fibre-reinforced ultra-high-performance concrete enable lighter and thinner sections to be used for structural components such as shell roofs and bridges, creating an enhanced sleek aesthetic. A high-quality durable surface is produced from appropriate moulds (e.g. steel) coated with proprietary release agent.

TRANSLUCENT CONCRETE

By embedding parallel fibre-optic threads into fine concrete, the material is made translucent without any appreciable loss of compressive strength. Translucent concrete can be manufactured as blocks or panels provided that the fibres run transversely from face to face. If one face is illuminated, any shadow cast
onto the bright side is clearly visible on the other face, whilst the colour of the transmitted light is unchanged. The material has many potential applications including walls, floor surfaces and illuminated pavements. Recent claims suggest that up to 80% light transmission is possible.

**INSULATING CONCRETE FORMWORK (ICF)**

**Polystyrene**

Large, hollow, interlocking polystyrene system blocks fit together to create permanent insulating formwork, which is then filled with in situ concrete to produce a monolithic concrete structure. A range of units is available giving a central core of 140–300 mm concrete and total insulation thicknesses between 100 and 300 mm according to the structural and thermal requirements. The two faces of the insulation are connected by a matrix of polystyrene links which become embedded into the concrete. The units, typically 250 mm high, are tongued and grooved to ensure correct location, and horizontal steel reinforcement may be incorporated if required for additional structural strength. Special blocks are available for lintels, wall ends, curved walls and fire walls. A pumpable grade of concrete (high slump) will fill the void space by gravity flow without the need for mechanical vibration. Some temporary support for the formwork is required during construction to ensure accurate alignment. Internal and external finishes may be applied directly to the polystyrene which is keyed for plaster or lightweight render. Alternatively masonry, timber or other claddings may be used externally and dry linings (e.g. plasterboard) may be attached to the inner leaf with appropriate adhesives.

**Lightweight concrete**

Concrete shuttering blocks are manufactured in normal, lightweight or wood-chip concrete to Standards BS EN 15435: 2008 and BS EN 15498: 2008, respectively. Systems are available with and without additional thermal insulation for use as internal, external and partition walls when filled with concrete. Some systems have lateral interlocking (e.g. by tongue and groove) and may be laid with or without mortar according to manufacturers’ specifications. Hollow insulated blocks, 300 mm wide, manufactured from 80% woodchip, when constructed and filled with concrete, can give a wall $U$-value of 0.27 W/m² K.

**POLYMER CONCRETE**

The incorporation of pre-polymers into concrete mixes, the pre-polymers then polymerising as the concrete sets and hardens, can reduce the penetration of water and carbon dioxide into cured concrete. Typical polymers include styrene–butadiene rubber and polyester–styrene. Epoxy resin and acrylic-latex modified mortars are used for repairing damaged and spalled concrete because of their enhanced adhesive properties. Similarly, polymer-modified mortars are used for the cosmetic filling of blowholes and blemishes in visual concrete. Resin-bound concrete construction products include street furniture, decorative elements and window sills. The materials are covered by the standard BS EN 15564: 2008.

**WATER FOR CONCRETE**

The general rule is that if water is of a quality suitable for drinking, then it is satisfactory for making concrete. The standards BS EN 1008: 2002 and pr BS ISO 12439: 2009 give the limits on impurities including sulphates.

**CONCRETE MIXES**

Concrete mixes are designed to produce concrete with the specified properties at the most economical price. The most important properties are usually strength and durability, although thermal and acoustic insulation, the effect of fire and appearance in visual concrete may also be critical.

In determining the composition of a concrete mix, consideration is given to the workability or ease of placement and compaction of the fluid mix and to the properties required in the hardened concrete. The key factor which affects both these properties is the free-water content of the mix after any water is absorbed into the aggregates. This quantity is defined by the water/cement ratio.

**Water/cement ratio**

$$\text{Water/cement ratio} = \frac{\text{weight of free water}}{\text{weight of cement}}$$

The free water in a mix is the quantity remaining after the aggregates have absorbed water to the saturated surface-dry condition. The free water is used to hydrate the cement and make the mix workable. With low water/cement ratios below 0.4, some of the
cement is not fully hydrated. At a water/cement ratio of 0.4, the hydrated cement just fills the space previously occupied by the water, giving a dense concrete. As the water/cement ratio is increased above 0.4, the mix becomes increasingly workable but the resulting cured concrete is more porous owing to the evaporation of the excess water leaving void spaces. Figure 3.9 shows the typical relationship between water/cement ratio and concrete crushing strength.

Workability

Workability describes the ability of the concrete mix to be placed within the formwork, around any reinforcement, and to be successfully compacted by hand or mechanical means to remove trapped air pockets. Mixes should be cohesive, so that they do not segregate during transportation or placing. Workability is affected by not only the water/cement ratio but also the aggregate content, size, grading and shape, and the addition of admixtures. It is measured on site with the slump test (Fig. 3.10). Table 3.12 shows the relationship between water/cement ratio and workability for

Fig. 3.9 Typical variation of crushing strengths about the published norm for the range of water/cement ratios

Fig. 3.10 Slump test (after Everett, 1994: Mitchell’s Materials. 5th ed. Longman Scientific and Technical.)
crushed and uncrushed aggregates at different cement contents.

**Free water**
The workability of concrete is highly dependent on the free water within the mix. An increase in free-water content causes a significant increase in workability, which would result in a greater slump measured in a slump test.

**Aggregate shape**
Rounded aggregates make a mix more workable than if crushed angular aggregates are used with the same water/cement ratio. However, because the bonding between cured cement and crushed aggregate is stronger than that to rounded aggregates, when other parameters are comparable, crushed aggregates produce a stronger concrete.

**Aggregate size**
The size of aggregate also affects the workability of the mix. The maximum practical size of coarse aggregate, compatible with placement around reinforcement and within the concrete section size, should be used to minimise the water content necessary for adequate workability. With fine aggregates, excessive quantities of the fine material (passing through a 0.063 mm test sieve) would increase considerably the water requirement of a particular mix to maintain workability. This is because the smaller particles have a larger surface area/volume ratio and therefore require more water to wet their surfaces. As additional water in the mix will decrease the cured concrete strength, for good-quality dense concrete, well-graded coarser sands are preferable.

**Aggregate/cement ratio**
For a particular water/cement ratio, decreasing the aggregate/cement ratio, which therefore increases proportionally both the cement and water content, increases workability. However, as cement is the most expensive component in concrete, cement-rich mixes are more costly than the lean mixes.

**Air-entraining**
Workability may be increased by air-entraining, although 1% voids in the cured concrete produces a decrease in compressive strength of approximately 6%. Thus in air-entraining, there is a balance between the increased workability and resultant improved compaction, versus the void space produced with its associated reduced crushing strength.

**Slump test**
The slump test is used for determining the workability of a mix on site. It gives a good indication of consistency from one batch to the next, but it is not effective for very dry or very wet mixes. The slump test is carried out as shown in Fig. 3.10. The base plate is placed on level ground and the cone filled with the concrete mix in three equal layers, each layer being tamped down 25 times with the 16 mm diameter tamping rod. The final excess of the third layer is struck off and the cone lifted off from the plate to allow the concrete to slump. The drop in level (mm) is the recorded slump, which may be a true slump, a shear slump or a collapse slump. In the case of a shear slump the material is retested. In the case of a collapse slump the mixture is too wet for most purposes. Typical slump values would be 0 to 25 mm for very dry mixes, frequently used
in road making; 10–40 mm (low workability) for use in foundations with light reinforcement; 50–90 mm (medium workability) for normal reinforced concrete placed with vibration and over 100 mm for high-workability concrete. Typically, slump values between 10 and 175 mm may be measured, although accuracy and repeatability are reduced at both extremes of the workability range. The slump test is not appropriate for aerated, no-fines or gap-graded concretes. The European Standard EN 206-1: 2000 classifies consistency classes of concrete mixes by results from the standard tests of slump (Table 3.13), Vebe consistency (a form of mechanised slump test), compaction and flow.

**Compaction**

After placing within the formwork, concrete requires compaction to remove air voids trapped in the mix before it begins to stiffen. Air voids weaken the concrete, increase its permeability and therefore reduce durability. In reinforced concrete, lack of compaction reduces the bond to the steel, and on exposed visual concrete, blemishes such as blowholes and honeycombing on the surface are aesthetically unacceptable and difficult to make good successfully. Vibration, to assist compaction, may be manual by rodding or tamping for small works, but normally poker vibrators and beam vibrators are used for mass and slab concrete, respectively. Vibrators which clamp on to the formwork are sometimes used when the reinforcement is too congested to allow access for poker vibrators.

The degree of compaction achieved by a standard quantity of work may be measured by the compacting factor test. In this test a fresh concrete sample is allowed to fall from one hopper into another. The weight of concrete contained in the lower hopper, when struck off flush, compared with a fully compacted sample gives the compacting factor. The compacting factor for a medium-workability concrete is usually about 0.9.

**Concrete cube and cylinder tests**

To maintain quality control of concrete, representative test samples should be taken, cured under controlled conditions and tested for compressive strength after the appropriate 3, 7 or 28 day period. Steel cylinder and cube moulds (Fig. 3.11) are filled in layers with either

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**Table 3.13 Slump test classes to European Standard EN 206-1: 2000**

<table>
<thead>
<tr>
<th>Slump class</th>
<th>Slump (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>S1</td>
<td>10–40</td>
</tr>
<tr>
<td>S2</td>
<td>50–90</td>
</tr>
<tr>
<td>S3</td>
<td>100–150</td>
</tr>
<tr>
<td>S4</td>
<td>160–210</td>
</tr>
<tr>
<td>S5</td>
<td>≥220</td>
</tr>
</tbody>
</table>

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**Fig. 3.11 Cylinder and cube test**
Table 3.14  Compressive strength classes for dense and lightweight concrete

<table>
<thead>
<tr>
<th>Compressive strength classes for dense concrete</th>
</tr>
</thead>
<tbody>
<tr>
<td>C8/10</td>
</tr>
<tr>
<td>C45/55</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Compressive strength classes for lightweight concrete</th>
</tr>
</thead>
<tbody>
<tr>
<td>LC8/9</td>
</tr>
<tr>
<td>LC45/50</td>
</tr>
</tbody>
</table>

Notes:
Within each compressive strength class the numbers indicate the 28-day crushing strength in MPa (N/mm²) as determined by the 150 mm diameter by 300 mm cylinder and 150 mm cube test, respectively.

hand or mechanical vibration. For hand tamping, a 100 mm cube would be filled in two equal layers, each tamped 25 times with a 25 mm square-end standard compacting bar; mechanical vibration would normally be with a vibrating table or pneumatic vibrator. The mix is then trowelled off level with the mould. Cubes and cylinders are cured under controlled moisture and temperature conditions for 24 hours, then stripped and cured under water at 18–20°C until required for testing.

The European standard concrete tests use cylinders 150 mm in diameter and 300 mm high rather than cubes, as they tend to give more uniform results for nominally similar concrete specimens. For a particular concrete, the characteristic compressive strength as determined by the cylinder test is lower than that obtained from the equivalent cube test. The compressive strength classes (Table 3.14), therefore, have a two-number notation (e.g. C 20/25). The first number, which is used in the European structural design codes, refers to the characteristic cylinder compressive strength, and the second number is the characteristic 150 mm cube compressive strength.

**DURABILITY OF CONCRETE**

While good-quality, well-compacted concrete with an adequate cement content and a low water/cement ratio is generally durable, concrete may be subjected to external agencies which cause deterioration, or, in certain circumstances, such as alkali–silica reaction, it may be subject to internal degradation. The standard BS EN 206-1: 2000 specifies requirements for the specification, constituents, composition, production and properties of concrete.

**Sulphate attack**

Sulphates are frequently present in soils, but the rate of sulphate attack on concrete is dependent on the soluble sulphate content of the groundwater. Thus, the presence of sodium or magnesium sulphate in solution, is more critical than that of calcium sulphate, which is relatively insoluble. Soluble sulphates react with the tricalcium aluminate (C₃A) component of the hardened cement paste, producing calcium sulphaolunate (ettringite). This material occupies a greater volume than the original tricalcium aluminate; therefore, expansion causes cracking, loss of strength and increased vulnerability to further sulphate attack. The continuing attack by sulphates depends on the movement of sulphate-bearing groundwater, and in some cases, delayed ettringite formation may not be apparent for 20 years. Delayed ettringite formation is sometimes observed in precast concrete which had been steam-cured, or when the temperature within the in situ mass concrete had risen excessively during the curing process. With magnesium sulphates, deterioration may be more serious as the calcium silicates within the cured concrete are also attacked. The use of sulphate-resisting Portland cement or combinations of Portland cement and fly ash [pulverised-fuel ash (PFA)] or granulated blastfurnace slag (GGBS) reduces the risk of sulphate attack in well-compacted concrete. In the presence of high soluble sulphate concentrations, concrete requires surface protection. The criteria which increase the resistance of the cement matrix to sulphate attack are described in the document CEN/TR 15697: 2008.

The BRE Special Digest 1: 2005 also describes provision for combating sulphate deterioration, including the more rapid form of attack in which the mineral thaumasite is formed. Thaumasite sulphate attack has
seriously affected concrete foundations and substructures including some bridges on the UK M5 motorway. This type of sulphate attack is most active at temperatures below 15°C.

**Frost resistance**

Weak, permeable concrete is particularly vulnerable to the absorption of water into capillary pores and cracks. On freezing, the ice formed will expand causing frost damage. The use of air-entraining agents, which produce discontinuous pores within concrete, reduces the risk of surface frost damage. Concrete is particularly vulnerable to frost damage during the first two days of early hardening. Where new concrete is at risk, frost precautions are necessary to ensure that the mix temperature does not fall below 5°C until a strength of 2 MPa is achieved. Eurocode 2 (BS EN 1992-1-1: 2004) refers to four levels of exposure class (XF1 to XF4) with respect to freeze/thaw deterioration (Table 3.16 page 86).

**Fire resistance**

Up to 250°C, concrete shows no significant loss of strength, but by 450°C, depending on the duration of heating, the strength may be reduced to half and by 600°C little strength remains. However, as concrete is a good insulator, it may take 4 hours within a building fire for the temperature 50 mm below the surface of the concrete to rise to 650°C (Fig. 3.12).

The effect of heat on the concrete causes colour changes to pink at 300°C, grey at 600°C and buff by 1000°C. The aggregates used within concrete have a significant effect on fire resistance. For fire protection, limestone aggregates may perform slightly better than granites and other crushed rocks, which spall owing to differential expansion. Where the concrete cover over reinforced steel is greater than 40 mm, secondary reinforcement with expanded metal gives added protection to the structural reinforcement. Lightweight-aggregate concretes, owing to their enhanced thermal properties, perform significantly better in fires with respect to both insulation and spalling.

Concrete manufactured without organic materials is Class A1 with respect to reaction to fire. If more than 1% of organic materials are incorporated into the mix, then the material will require testing to the standard (BS EN 13501-1: 2007).

**Chemical attack and aggressive ground**

The resistance of cured concrete to acid attack is largely dependent on the quality of the concrete, although the addition of granulated blastfurnace slag (GGBS) or fly ash (pulverised fuel ash (PFA)) increases the resistance to acids. Limestone-aggregate concrete is more vulnerable to acid attack than concretes with other aggregates. The resistance of cured concrete to chemical attack is defined by the design chemical class number, ranging from DC1 (low resistance) to DC4 (high resistance). The required design chemical class (DC Class) of the concrete is calculated by combining the effects of the sulphate content of the ground, the nature of the groundwater and the anticipated working life of the construction (BRE Special Digest 1: 2005).

Determining the design chemical class required for concrete in a particular ground environment is a three-stage process. The first stage is to determine the design sulphate class (DS) of the site. This is a five-level classification based primarily on the sulphate content of the soil and/or groundwater. It takes into account the concentrations of calcium sulphate and also the more soluble magnesium and sodium sulphates and the presence of chlorides and nitrates if the pH is less than 5.5 (acid).
Design sulphate class | Limits of sulphate (mg/l)
---|---
DS1 | <500
DS2 | 500–1500
DS3 | 1600–3000
DS4 | 3100–6000
DS5 | >6000

The next stage is to determine the aggressive chemical environment for concrete (ACEC) classification. Adverse ground conditions such as acidity (low pH), often found in brownfield sites, and/or mobile groundwater lead to a more severe ACEC classification. Static water is more benign and leads to a less severe ACEC classification. The aggressive chemical environment for concrete classes range from AC1 (the least aggressive) to AC5 (the most aggressive), and are based on a combination of the design sulphate class, groundwater mobility and pH.

The design chemical class (DC1 to DC4) defines the qualities of the concrete required to resist chemical attack. It is determined from the ACEC class of the ground together with factors relating to the concrete, such as section size and intended working life (e.g. 100 years). As there are only four design chemical classes against five ACEC classes, for the severest grade of ACEC (i.e. AC5) there are additional protective measures (APMs) which can be specified to combat the more adverse conditions. Usually APM3 (surface protection to the concrete) is appropriate for AC5 environments, but for increasing the intended working life from 50 to 100 years under the less aggressive AC3 or AC4 conditions, any one APM may be applied.

Additional protective measures (APMs) for buried concrete:
- APM1: enhance the concrete quality;
- APM2: use controlled permeability formwork;
- APM3: provide surface protection to the concrete;
- APM4: increase the thickness of the concrete as a sacrificial layer;
- APM5: reduce groundwater by drainage of the site.

Careful consideration of all these additional factors is required to ensure that a suitably durable concrete, appropriate to the job, is delivered on site for use in aggressive ground and chemical environments (BRE Special Digest 1: 2005).

**Crystallisation of salts**

The crystallisation of salts, particularly from sea water, within the pores of porous concrete can cause sufficient internal pressure to disrupt the concrete.

**Alkali–silica reaction**

Alkali–silica reaction (ASR) may occur between cements containing sodium or potassium alkalis and any active silica within the aggregate. In severe cases, expansion of the gel produced by the chemical reaction causes map cracking of the concrete, which is characterised by a random network of very fine cracks bounded by a few larger ones. Aggregates are defined as having low, normal or high reactivity. The risk of alkali–silica reaction when using normal reactivity aggregates can be controlled by restricting the alkali content of the Portland cement to a maximum of 0.6% (low alkali cement) or the soluble alkali content of the concrete to 3 kg/m³. Additions of controlled quantities of silica fume, ground granulated blastfurnace slag (GGBS) or pulverised-fuel ash (PFA) may be used with low- or normal-reactivity aggregates to reduce the risk of alkali–silica reaction. Alternative methods of minimising the risk of alkali–silica reaction include the addition of lithium salts or metakaolin to the concrete mix.

**Carbonation**

Carbon dioxide from the atmosphere is slowly absorbed into moist concrete and reacts with the calcium hydroxide content to form calcium carbonate. The process occurs mainly at the surface and only penetrates very slowly into the bulk material. The rate of penetration is dependent on the porosity of the concrete, temperature and humidity; generally it becomes problematic only when the concrete surrounding steel reinforcement is affected. Carbonation turns strongly alkaline hydrated cement (pH 12.5) into an almost neutral medium (pH 8.3) in which steel reinforcement will corrode rapidly if subjected to moisture.

\[
\text{Ca(OH)}_2 + \text{CO}_2 \rightarrow \text{CaCO}_3
\]

Good-quality dense concrete may only show carbonation to a depth of 5–10 mm after 50 years, whereas a low-strength permeable concrete may carbonate to a depth of 25 mm within 10 years. If reinforcement is not correctly located with sufficient cover, it corrodes causing expansion, spalling and rust staining. The depth of carbonation can be determined by testing a core sample for alkalinity using phenolphthalein chemical indicator, which turns pink in contact with the uncarbonated alkaline concrete. Where steel reinforcement has become exposed due to carbonation and...
rusting, it may be coated with a rust-inhibiting cement and the cover restored with polymer-modified mortar which may contain fibre reinforcement. Additional protection against further attack can be achieved by the final application of an anti-carbonation coating which acts as a barrier to carbon dioxide. Thermosetting polymers such as polyurethane and chlorinated rubber, also certain acrylic-based polymers, give some protection against carbonation.

**PHYSICAL PROPERTIES OF CONCRETE**

**Thermal movement**

The coefficient of thermal expansion of concrete varies between \((7 \text{ and } 14) \times 10^{-6} \degree \text{C}\), according to the type of aggregate used, mix proportions and curing conditions.

**Moisture movement**

During the curing process, concrete exhibits some irreversible shrinkage which must be accommodated within the construction joints. The extent of the shrinkage is dependent on the restraining effect of the aggregate and is generally larger when smaller or lightweight aggregates are used. High-aggregate content mixes with low workability tend to have small drying shrinkages.

The reversible moisture movement for cured concrete is typically \((2–6) \times 10^{-4} \degree \text{C}\), depending on the aggregate.

**Creep**

Creep is the long-term deformation of concrete under sustained loads (Fig. 3.13). The extent of creep is largely dependent on the modulus of elasticity of the aggregate. Thus an aggregate with a high modulus of elasticity offers a high restraint to creep. The extent of creep may be several times that of the initial elastic deformation of the concrete under the same applied load. Where rigid cladding is applied to a concrete-frame building, compression joints at each storey must be sufficiently wide to take up any deformation due to creep in addition to normal cyclical movements.

**CONCRETE STRENGTH CLASSES**

Concrete should be specified, placed and cured according to BS EN 206-1: 2000. The preferred strength classes of concrete are shown in Table 3.14, in which the numbers refer to the test sample crushing strengths of a \(150 \times 300 \text{ mm cylinder}\) and a \(150 \text{ mm cube}\), respectively.

**SPECIFICATION OF CONCRETE MIXES**

There are five methods for specifying concrete described in BS 8500-1: 2006. All should conform to the standards BS 8500-1: 2006 and BS EN 206-1: 2000.

The five methods are:

- designated concrete
- designed concrete
- prescribed concrete
- standardised prescribed concrete
- proprietary concrete

If the application can be considered to be routine, then designated concrete is usually appropriate. If, however, the purchaser requires specific performance criteria and accepts the higher level of responsibility in the specification, then designed or prescribed concretes may be used. For housing and similar applications, standardised prescribed mixes should give the required performance, provided there is sufficient control over the production and quality of materials used.

**Designated concrete**

Designated concretes are appropriate for most concrete construction including general-purpose work, foundations, reinforced concrete and air-entrained pavement concrete. The purchaser is responsible for correctly specifying the proposed use and the concrete...
mix designation. In addition, the purchaser must specify whether the concrete is to be reinforced, the exposure (or soil) conditions, the nominal aggregate size if other than 20 mm and the consistency class (slump). The producer must ensure that the mix fulfills all the performance criteria. Thus, normally for foundations in design chemical class soil conditions DC3, the designated mix FND 3 would be required. This mix may be supplied with sulphate-resisting Portland cement at 340 kg/m³ and a maximum water/cement ratio of 0.5, or as Portland cement with 25% fly ash or 75% granulated blast furnace slag. Any of these mixes will perform to the required criteria for the specified purpose. For routine work, designated mixes produced by quality-assured plants offer the specifier the least risk of wrong specification. Table 3.15 illustrates typical housing applications for designated mixes.

### Designed concrete

The producer is responsible for selecting a designed concrete which will meet the performance criteria listed by the specifier. The specifier must clearly indicate the required use, curing conditions, exposure conditions, surface finish, maximum aggregate size and any excluded materials. In addition, the compressive strength class, the maximum water/cement ratio, the minimum cement content, the consistency (slump) and permitted cement types should be quoted. Within these constraints, the producer is responsible for producing a concrete which conforms to the required properties and any additional stated characteristics. Designed concrete would be used when the user specifications are outside those covered by designated concrete. Specialist requirements include low heat of hydration, exposure to chlorides or lightweight concrete.

#### Table 3.15  Designated and standardised prescribed concrete for housing and other applications (BS 8500-1: 2006)

<table>
<thead>
<tr>
<th>Typical application</th>
<th>Designated concrete</th>
<th>Standardised prescribed concrete</th>
<th>Consistency class</th>
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</thead>
<tbody>
<tr>
<td>Foundations (Design Chemical Class 1):</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Blinding and mass concrete fill</td>
<td>GEN 1</td>
<td>ST2</td>
<td>S3</td>
</tr>
<tr>
<td>Strip footings</td>
<td>GEN 1</td>
<td>ST2</td>
<td>S3</td>
</tr>
<tr>
<td>Mass concrete foundations</td>
<td>GEN 1</td>
<td>ST2</td>
<td>S3</td>
</tr>
<tr>
<td>Trench fill foundations</td>
<td>GEN 1</td>
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<td>S3</td>
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<tr>
<td>Drainage works—immediate support</td>
<td>GEN 1</td>
<td>ST2</td>
<td>S1</td>
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<td>Other drainage works</td>
<td>GEN 1</td>
<td>ST2</td>
<td>S3</td>
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<tr>
<td>Oversite below suspended slabs</td>
<td>GEN 1</td>
<td>ST2</td>
<td>S3</td>
</tr>
<tr>
<td>Foundations (Design Chemical Classes 2–4):</td>
<td></td>
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<td>Kerb bedding and backing</td>
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</tr>
<tr>
<td>House floors with no embedded metal for screeding</td>
<td>GEN 1</td>
<td>ST2</td>
<td>S2</td>
</tr>
<tr>
<td>House floors with no embedded metal—no finish</td>
<td>GEN 2</td>
<td>ST3</td>
<td>S2</td>
</tr>
<tr>
<td>Garage floors with no embedded metal</td>
<td>GEN 3</td>
<td>ST4</td>
<td>S2</td>
</tr>
<tr>
<td>Wearing surface—light foot and trolley traffic</td>
<td>RC 25/30</td>
<td>ST4</td>
<td>S2</td>
</tr>
<tr>
<td>Wearing surface—general industrial</td>
<td>RC 32/40</td>
<td></td>
<td>S2</td>
</tr>
<tr>
<td>Wearing surface—heavy industrial</td>
<td>RC 40/50</td>
<td></td>
<td>S2</td>
</tr>
<tr>
<td>Paving:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>House drives, domestic parking and external paving</td>
<td>PAV 1</td>
<td></td>
<td>S2</td>
</tr>
<tr>
<td>Heavy duty external paving for rubber tyre vehicles</td>
<td>PAV 2</td>
<td></td>
<td>S2</td>
</tr>
</tbody>
</table>

Note:
m refers to resistance to the higher magnesium levels in the various sulphate classes.
**Prescribed concrete**

The purchaser fully specifies all materials by weight (kg/m³), including admixtures but not the concrete strength. The purchaser is therefore responsible for the performance characteristics of the concrete. Prescribed concretes are used particularly for specialist finishes such as exposed aggregate visual concrete.

**Standardised prescribed concrete**

Standardised prescribed concretes are a set of five standard mixes, which may be mixed on site or delivered by a non-third-party certified producer. Standard mixes ST1 to ST5 may be made to S1, S2, S3 or S4 slump classes, giving low, medium, high or very high workability. The specification must record a maximum aggregate size and whether the concrete is to be reinforced or not. Table 3.15 illustrates typical housing applications for standardised prescribed concrete.

**Proprietary concrete**

Proprietary concrete must conform to the standards BS 8500-2: 2006 and BS EN 206-1: 2000 and be properly identified. This category allows for a concrete supplier to produce a concrete mix with an appropriate performance but without indicating its composition.

**IN SITU CONCRETE TESTING**

The compressive strength of hardened concrete may be estimated in situ by mechanical or ultrasonic measurements. The Schmidt hammer or sclerometer measures the surface hardness of concrete by determining the rebound of a steel plunger fired at the surface. In the pull-out test, the force required to extract a previously cast-in standard steel cone gives a measure of concrete strength. Ultrasonic devices determine the velocity of ultrasound pulses through concrete. Since pulse velocity increases with concrete density, the technique can be used to determine variations within similar concretes. The test gives a broad classification of the quality of concrete, but not absolute data for concretes of different materials in unknown proportions.

**Reinforced concrete**

Concrete is strong in compression, with crushing strengths typically in the range 20–40 MPa, and up to 100 MPa for high-strength concretes. However, the tensile strength of concrete is usually only 10% of the compressive strength. Steel is the universally accepted reinforcing material as it is strong in tension, forms a good bond and has a similar coefficient of thermal expansion to concrete. The location of the steel within reinforced concrete is critical, as shown in Fig. 3.14, to ensure that the tensile and shear forces are transferred to the steel. The longitudinal bars carry the tensile forces while the links or stirrups combat the shear forces and also locate the steel during the casting of the concrete. Links are therefore more concentrated around locations of high shear, although inclined bars may also be used to resist the shear forces. Fewer or thinner steel bars may be incorporated into reinforced concrete to take a proportion of the compressive loads in order to minimise the beam dimensions.

Steel reinforcement for concrete is manufactured, largely from recycled scrap, into round, ribbed, indented or ribbed and twisted bars (Fig. 3.15). Mild steel is frequently used for the plain bars to form bent links. Hot-rolled, high-yield steel is used for ribbed and indented bars. British Standard BS 4482: 2005 refers to 250 MPa yield strength steel for plain bars and to the higher grade 500 MPa steel for plain, ribbed and indented reinforcement of diameters between 2.5 and 12 mm. British Standard BS 4449: 2005 specifies high-yield steel (grade 500 MPa) with three levels of ductility A, B and C (highest) for ribbed bars from 6 to 50 mm diameters. Welded steel mesh reinforcement to BS 4483: 2005 is used for slabs, roads and within sprayed concrete.

Austenitic stainless steels may be used for concrete reinforcement where failure due to corrosion is a potential risk. Grade 1.4301 (18% chromium, 10% nickel) stainless steel is used for most applications, but the higher grade 1.4436 (17% chromium, 12% nickel, 2.5% molybdenum) is used in more corrosive environments. Where long-term performance is required in highly corrosive environments, the duplex grades of stainless steel may be used. The initial cost of stainless steel reinforcement is approximately eight times that of standard steel reinforcement, but in situations where maintenance costs could be high, for example due to chloride attack from sea water or road salts, the overall life cycle costs may be reduced by its use. Additionally, stainless steels have higher strengths than the standard carbon steels. Suitable stainless steels for the reinforcement of concrete are specified in BS 6744: 2001 + A2: 2009.
For reinforced concrete to act efficiently as a composite material the bond between the concrete and steel must be secure. This ensures that any tensile forces within the concrete are transferred to the steel reinforcement. The shape and surface condition of the steel and the quality of the concrete all affect the bond strength.

To obtain the most efficient mechanical bond with concrete, the surface of the steel should be free of flaky rust, loose scale and grease, but the thin layer of rust, typically produced by short-term storage on site, should not be removed before use. The use of hooked ends in round bars reduces the risk of the steel being pulled out under load, but high bond strength is achieved with ribbed or indented bars which ensure a good bond along the full length of the steel. Steel rebars are usually either supplied in stock lengths, or cut and bent ready for making up into cages. Sometimes the reinforcement may be supplied as prefabricated cages, which may be welded rather than fixed with iron wire as on site. Steel reinforcement, although weldable, is rarely welded on site. Rebar joints can easily be made with proprietary fixings, such as steel sleeves fastened by shear bolts. Spacers are used to ensure the correct separation between reinforcement and formwork.

Good-quality dense concrete gives the strongest bond to the steel. Concrete should be well compacted...
Corrosion of steel within reinforced concrete

Steel is protected from corrosion provided that it has adequate cover of a good-quality, well-compacted and cured concrete. The strongly alkaline environment of the hydrated cement renders the steel passive. However, insufficient cover caused by the incorrect fixing of the steel reinforcement or the formwork can allow the steel to corrode. Rust expansion causes surface spalling; then exposure of the steel allows corrosion, followed by rust staining of the concrete surface (Fig. 3.16). Calcium chloride accelerators should not normally be used in reinforced concrete as the residual chlorides cause accelerated corrosion of the steel reinforcement.

Additional protection from corrosion can be achieved by the use of galvanised, epoxy-coated or stainless steel reinforcement. The protective alkalinity of the concrete is reduced at the surface by carbonation. The depth of carbonation depends on the permeability of the concrete, moisture content and any surface cracking. The nominal cover for concrete reinforcement is therefore calculated from the anticipated degree of exposure (Table 3.16) and the concrete strength class as in Table 3.17. The recommended cover specified relates to all reinforcement, including any

<table>
<thead>
<tr>
<th>Exposure classes</th>
<th>Typical environmental conditions</th>
</tr>
</thead>
<tbody>
<tr>
<td>No risk of reinforcement corrosion or attack on concrete</td>
<td></td>
</tr>
<tr>
<td>X0</td>
<td>Concrete with no reinforcement. Dry concrete</td>
</tr>
<tr>
<td>Corrosion induced by carbonation</td>
<td></td>
</tr>
<tr>
<td>XC1</td>
<td>Dry or permanently wet</td>
</tr>
<tr>
<td>XC2</td>
<td>Wet and rarely dry</td>
</tr>
<tr>
<td>XC3</td>
<td>Moderate humidity</td>
</tr>
<tr>
<td>XC4</td>
<td>Cyclic wet and dry</td>
</tr>
<tr>
<td>Corrosion induced by chlorides</td>
<td></td>
</tr>
<tr>
<td>XD1</td>
<td>Humid environment</td>
</tr>
<tr>
<td>XD2</td>
<td>Wet and rarely dry</td>
</tr>
<tr>
<td>XD3</td>
<td>Cyclic wet and dry</td>
</tr>
<tr>
<td>Corrosion induced by seawater</td>
<td></td>
</tr>
<tr>
<td>XS1</td>
<td>Exposure to sea air</td>
</tr>
<tr>
<td>XS2</td>
<td>Submerged under sea water</td>
</tr>
<tr>
<td>XS3</td>
<td>Tidal and sea spray zone</td>
</tr>
<tr>
<td>Freeze/thaw deterioration</td>
<td></td>
</tr>
<tr>
<td>XF1</td>
<td>Moderate saturation</td>
</tr>
<tr>
<td>XF2</td>
<td>Moderate saturation with de-icing agent</td>
</tr>
<tr>
<td>XF3</td>
<td>High saturation</td>
</tr>
<tr>
<td>XF4</td>
<td>High saturation with de-icing agent</td>
</tr>
<tr>
<td>Chemical attack</td>
<td></td>
</tr>
<tr>
<td>XA1</td>
<td>Slightly aggressive agencies</td>
</tr>
<tr>
<td>XA2</td>
<td>Moderately aggressive agencies</td>
</tr>
<tr>
<td>XA3</td>
<td>Highly aggressive agencies</td>
</tr>
</tbody>
</table>
Table 3.17 Minimum cover required to ensure durability of steel reinforcement in structural concrete for exposure classes to Eurocode 2 (BS EN 1992-1-1: 2004)

<table>
<thead>
<tr>
<th>Exposure class</th>
<th>X0</th>
<th>XC1</th>
<th>XC2/XC3</th>
<th>XC4</th>
<th>XD1/YS1</th>
<th>XD2/YS2</th>
<th>XD3/YS3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Recommended cover (mm)</td>
<td>10</td>
<td>15</td>
<td>25</td>
<td>30</td>
<td>35</td>
<td>40</td>
<td>45</td>
</tr>
<tr>
<td>Minimum cover (mm)</td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>15</td>
<td>20</td>
<td>25</td>
<td>30</td>
</tr>
<tr>
<td>Strength class</td>
<td>≥C30/37</td>
<td>≥C30/37</td>
<td>≥C35/45</td>
<td>≥C40/50</td>
<td>≥C40/50</td>
<td>≥C40/50</td>
<td>≥C45/55</td>
</tr>
</tbody>
</table>

Notes:
The recommended cover relates to standard production with a design working life of 50 years.
Increased cover is required for a design working life of 100 years.
The minimum cover relates to very specific conditions combining high quality control for positioning of the reinforcement and the concrete production, additionally the use of 4% (minimum) air entrainment.
The Standard BS 8500-1: 2006 gives a more detailed set of recommendations relating strength classes to nominal cover including the option to add additional cover (Δc) for workmanship deviation.

Table 3.18 Indicative strength classes for durability of concrete to Eurocode 2 (BS EN 1992-1-1: 2004)

<table>
<thead>
<tr>
<th>Corrosion risk</th>
<th>XC1</th>
<th>XC2</th>
<th>XC3 and XC4</th>
<th>XD1 and XD2</th>
<th>XD3</th>
<th>XS1</th>
<th>XS2 and XS3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Indicative strength class</td>
<td>C20/25</td>
<td>C25/30</td>
<td>C30/37</td>
<td>C30/37</td>
<td>C35/45</td>
<td>C30/37</td>
<td>C35/45</td>
</tr>
<tr>
<td>Damage to concrete</td>
<td>X0</td>
<td>XF1</td>
<td>XF2</td>
<td>XF3</td>
<td>XA1</td>
<td>XA2</td>
<td>XA3</td>
</tr>
<tr>
<td>Indicative strength class</td>
<td>C12/15</td>
<td>C30/37</td>
<td>C25/30</td>
<td>C30/37</td>
<td>C30/37</td>
<td>C30/37</td>
<td>C35/45</td>
</tr>
</tbody>
</table>

Notes:
The Standard BS 8500-1: 2006 details a more comprehensive relationship between minimum strength class and exposure class for frost resistance in relation to different maximum aggregate sizes and minimum cement contents.

Wire ties and secondary reinforcement. Some reduction in carbonation rate can be achieved by protective coatings to the concrete surface. It should be noted that the choice of an adequately durable concrete for the protection of the concrete itself against attack and for the prevention of reinforcement corrosion may result in a higher compressive strength concrete being required than is necessary for the structural design (Table 3.18).

Where the depth of concrete cover over reinforcement is in doubt it can be measured with a covermeter. If reinforcement is corroding, cathodic protection by application of a continuous direct current to the steel reinforcement may prevent further deterioration and lead to realkalisation of the carbonated concrete.

Fibre-composite reinforced concrete

In most situations steel is used for reinforcing or prestressing concrete. However, for structures in highly aggressive environments high-modulus continuous fibres embedded in resin offer an alternative. The fibres, either glass, carbon or aramid, are encased in a thermosetting resin and drawn through a die by pultrusion to produce the required cross-section. The extruded material is then overwound with further fibres to improve its bond with concrete. The fibre-composite rods are used as reinforcement or as prestressing tendons within standard concrete construction.

Bendy concrete

Fibre-reinforced concrete of an appropriate mix may be continuously extruded into various sections to produce sheets, cylinders or tubes. The product is more flexible and has a higher impact strength than ordinary concrete. Bendy concrete may be drilled, cut and nailed without damage. It is lighter than ordinary concrete and with its good fire resistance may be used as an alternative to other wall boards.

Fibre-reinforced aerated concrete

Polypropylene fibre-reinforced aerated concrete is used for making lightweight blocks, floor, wall and roofing panels, offering a combination of strength and insulation properties. The material, like standard aerated concrete, can be cut and worked with standard hand tools. Where additional strength is required, steel fibre-reinforced aerated concrete may be used for cast in situ or factory-produced units. The fibre-reinforced material has a greater resilience than standard aerated concrete. Roofing membranes and battens for tiling...
**Table 3.19** Typical cover to concrete reinforcement for fire resistance to Eurocode 2 (BS EN 1992-1-2: 2004)

<table>
<thead>
<tr>
<th>Fire resistance (minutes)</th>
<th>Beams Width (mm)</th>
<th>Simply supported</th>
<th>Continuous beams</th>
</tr>
</thead>
<tbody>
<tr>
<td>R3</td>
<td>0</td>
<td>25</td>
<td>15</td>
</tr>
<tr>
<td>R60</td>
<td>120</td>
<td>40</td>
<td>25</td>
</tr>
<tr>
<td>R90</td>
<td>150</td>
<td>55</td>
<td>35</td>
</tr>
<tr>
<td>R120</td>
<td>200</td>
<td>65</td>
<td>45</td>
</tr>
<tr>
<td>R180</td>
<td>240</td>
<td>80</td>
<td>60</td>
</tr>
<tr>
<td>R240</td>
<td>280</td>
<td>90</td>
<td>75</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Columns Minimum dimensions (mm)</th>
<th>One face exposed</th>
</tr>
</thead>
<tbody>
<tr>
<td>R3</td>
<td>155</td>
</tr>
<tr>
<td>R60</td>
<td>155</td>
</tr>
<tr>
<td>R90</td>
<td>155</td>
</tr>
<tr>
<td>R120</td>
<td>175</td>
</tr>
<tr>
<td>R180</td>
<td>230</td>
</tr>
<tr>
<td>R240</td>
<td>295</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Walls Minimum dimensions (mm)</th>
<th>One face exposed</th>
</tr>
</thead>
<tbody>
<tr>
<td>REI 30</td>
<td>100</td>
</tr>
<tr>
<td>REI 60</td>
<td>110</td>
</tr>
<tr>
<td>REI 90</td>
<td>120</td>
</tr>
<tr>
<td>REI 120</td>
<td>150</td>
</tr>
<tr>
<td>REI 180</td>
<td>180</td>
</tr>
<tr>
<td>REI 240</td>
<td>230</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Slabs Slab thickness (mm)</th>
<th>One-way slabs</th>
<th>Two-way slabs</th>
</tr>
</thead>
<tbody>
<tr>
<td>REI 30</td>
<td>60</td>
<td>10</td>
</tr>
<tr>
<td>REI 60</td>
<td>80</td>
<td>20</td>
</tr>
<tr>
<td>REI 90</td>
<td>100</td>
<td>30</td>
</tr>
<tr>
<td>REI 120</td>
<td>120</td>
<td>40</td>
</tr>
<tr>
<td>REI 180</td>
<td>150</td>
<td>55</td>
</tr>
<tr>
<td>REI 240</td>
<td>175</td>
<td>65</td>
</tr>
</tbody>
</table>

Notes:

Fire resistance class:

R, load-bearing criterion; E, integrity criterion and I, insulation criterion in standard fire exposure.

All reinforcement cover requirements are also dependent on the dimensions and geometry of the concrete components and the degree of fire exposure (BS EN 1992-1-2: 2004).

Where low cover thicknesses are required for fire protection, a higher depth of cover may be required for corrosion protection (BS EN 1992-1-1: 2004).

Concrete manufactured without organic materials is Class A1 with respect to reaction to fire. If more than 1% of organic materials are incorporated into the mix, then the material will require testing to the standard (BS EN 13501-1: 2007).

The depth of concrete cover over the steel reinforcement, to ensure various periods of fire resistance, is listed in Table 3.19. Where cover exceeds 40 mm, additional reinforcement will be required to prevent surface spalling of the concrete. The cover should prevent the temperature of the steel reinforcement from exceeding 550°C (or 450°C for prestressing steel).

**Prestressed concrete**

Concrete has a high compressive strength but is weak in tension. Prestressing with steel wires or tendons ensures that the concrete component of the composite
LIME, CEMENT AND CONCRETE

material always remains in compression when subjected to flexing up to the maximum working load. The tensile forces within the steel tendons act on the concrete putting it into compression, such that only under excessive loads would the concrete go into tension and crack. Two distinct systems are employed; in pre-tensioning, the tendons are tensioned before the concrete is cured, and in post-tensioning, the tendons are tensioned after the concrete is hardened (Fig. 3.17).

Pre-tensioning

Large numbers of precast concrete units, including flooring systems, are manufactured by the pre-tensioning process. Tendons are fed through a series of beam moulds and the appropriate tension is applied. The concrete is placed, vibrated and cured. The tendons are cut at the ends of the beams, putting the concrete into compression. As with precast reinforced concrete it is vital that prestressed beams are installed the correct way up according to the anticipated loads.

Post-tensioning

In the post-tensioning system the tendons are located in the formwork within sheaths or ducts. The concrete is placed, and when sufficiently strong, the tendons are stressed against the concrete and locked off with special anchor grips incorporated into the ends of the concrete. Usually reinforcement is incorporated into post-tensioned concrete, especially near the anchorages, which are subject to very high localised forces. In the bonded system, after tensioning the free space within the ducts is grouted up, which then limits the reliance on the anchorage fixing; however, in the unbonded system the tendons remain free to move independently of the concrete. Tendon ducts are typically manufactured from galvanised steel strip or high-density polythene.

Post-tensioning has the advantage over pre-tensioning that the tendons can be curved to follow the most efficient prestress lines. In turn this enables long spans of minimum thickness to be constructed. During demolition or structural alteration work, unbonded post-tensioned structures should be de-tensioned, although experience has shown that if demolished under tension, structures do not fail explosively. In alteration work, remaining severed tendons may subsequently require re-tensioning and re-anchoring to recover the structural performance. However, the use of post-tensioning does not preclude subsequent structural modifications.

Visual concrete

The production of visual concrete, whether precast or in situ, requires not only a high standard of quality control in manufacture, but also careful consideration to the correct specification and detailing of the material to ensure a quality finish which weathers appropriately. The exposed concrete at St John’s College, Oxford (Fig. 3.18) illustrates the visual qualities of the material when designed, detailed and executed under optimum conditions.
The appearance of visual concrete is affected by four key factors:

- the composition of the concrete mix;
- the formwork used;
- any surface treatment after casting;
- the quality of workmanship.

**DESIGN CONSIDERATIONS**

The satisfactory production of large areas of smooth concrete is difficult due to variations in colour and the inevitability of some surface blemishes, which can be improved, but not eradicated, by remedial work. Externally smooth concrete weathers unevenly due to the build-up of dirt deposits and the flow of rainwater. Therefore, if concrete is to be used externally as a visual material, early design considerations must be given to the use of textured or profiled surfaces to control the flow of rainwater. Generally, the range of finishes and quality control offered by precasting techniques are wider than those available for in situ work, but frequently construction may involve both techniques. The use of external renderings offers an alternative range of finishes for concrete and other substrates. Figure 3.19 illustrates the range of processes available in the production of visual concrete.

**PRECAST CONCRETE**

Precast-concrete units may be cast vertically or horizontally, although most factory operations use the latter, either face-up or face-down, as better quality control can be achieved by this method. Moulds are usually manufactured from plywood or steel. Whilst steel moulds are more durable for repeated use, plywood moulds are used for the more complex forms; they can also be more readily modified for non-standard units. Moulds are designed to be dismantled for the removal of the cast unit and must be manufactured to tight tolerances to ensure quality control on the finished product. As high costs are involved in the initial production of the moulds, economies of construction can be achieved by limiting the number of variations. This can have significant effects on the overall building aesthetic. Fixing and lifting systems
for transportation must be incorporated into precast units, usually in conjunction with the steel reinforcement. In addition to visual concrete panels, units faced with natural stone, brickwork or tiles extend the range of precast architectural claddings (Figs. 3.20 and 3.21). The document PD CEN/TR 15739: 2008 categorises the range of precast concrete finishes according to flatness (P), texture (T) and colour (C).

**Precast aircrete panels**

Precast aircrete panels, 100 mm thick and to standard storey heights, are suitable for the inner leaf of standard cavity construction and internal walls. Tolerances are close, similar to those required by the equivalent
thin-joint masonry system. Maximum dimensions are $600 \times 3000$ mm with a standard thermal conductivity of 0.11 W/m K. The thin mortar jointing system with 2 mm joints is used for fixing the panels. Larger units, 200 mm thick, are manufactured for commercial projects.

**IN SITU CONCRETE**

The quality of in situ visual concrete is heavily dependent on the formwork as any defects will be mirrored in the concrete surface. The formwork must be strong enough to withstand, without distortion, the pressure of the fresh concrete, and the joints must be tight enough to prevent leakage, which can cause honeycombing of the surface. A wide range of timber products, metals and plastics are used as formwork, depending on the surface finish required.

The Millau Viaduct in France (Fig. 3.22), completed in 2004, is an elegant cable-stayed bridge supported by seven slender piers 270 m over the Tarn Valley. The main columns of 1.5 m thick concrete were cast in situ with self-climbing steel formwork externally and by crane hoist internally. The concrete piers are surmounted by 90 m steel pylons, which support the steel box section road platform. Each pier is supported by four reinforced concrete piles splayed at the foot to spread the loading. During construction work of the 2.46 km viaduct, rapid-hardening concrete was being placed at the rate of 80 m$^3$ per hour and ultimately a total mass of 205,000 tonnes was used.

**CONCRETE FINISHES**

**Smooth finishes**

In direct as-cast concrete, the surface texture and water absorbancy of the formwork or any formwork lining directly determine the final exposed fairfaced finish. A high level of quality control is therefore required to ensure a visually acceptable finish. Hard, shiny, non-absorbent formwork materials, such as steel, glass-fibre reinforced polyester (GRP) or plastic-coated plywood, can give surfaces which suffer from map crazing due to differential shrinkage between the surface and underlying bulk material. Additionally, blow-holes caused by air bubbles trapped against the form face may spoil the surface if the concrete has not been sufficiently vibrated. Where the absorbency of the formwork varies, because of the mixing of new and reused formwork, or variations within the softwood timbers, or because of differing application of release agent to the formwork, permanent colour variations may be visible on the concrete surface. Release agents prevent bonding between the concrete and the formwork, which might cause damage to the concrete on striking the formwork. Cream emulsions and oils with surfactant are typically used as release agents for timber and steel, respectively. Formwork linings with controlled porosity can improve the quality of off-the-form finishes, by substantially reducing the number of blow-holes. The linings allow the escape of air and excess moisture but not cement, during vibration. A good-quality direct-cast concrete should exhibit only a few small blow-holes and modest colour variation.

*Fig. 3.22* Concrete columns — Millau Viaduct above the Tarn Valley, France. *Architect: Foster + Partners. Photograph: Arthur Lyons*
The application of paint to off-the-form concrete will emphasise the surface blemishes such as blow-holes. These become particularly noticeable if a light-coloured gloss paint is used. Surface defects must therefore be made good with filler before priming and subsequent painting of the concrete.

Textured finishes

A variety of textured finishes can be achieved by the use of rough-sawn boards as formwork. The grain effect can be enhanced by abrasive blasting, and a three-dimensional effect can be achieved by using variations in board thickness. Plastic materials, such as glass-fibre reinforced polyester (GRP), vacuum-formed thermoplastic sheeting, neoprene rubber and polystyrene, can be used as formwork linings to give different pattern effects. Colour variations are reduced by the use of matt finishes, which retain the mould release agent during compaction of the concrete. The number of blow-holes is reduced by the use of the slightly absorbent materials such as timber and polystyrene. Concrete panels cast face-up can be textured by rolling or tamping the concrete whilst it is still plastic.

Ribbed and profiled finishes

Ribbed concrete is typically cast in situ against vertical timber battens fixed to a plywood backing. In order to remove the formwork, without damage to the cured concrete, the battens must be splayed and smooth. A softer ribbed appearance is achieved by hammering off the projecting concrete to a striated riven finish. Profiled steel formwork and rope on plywood produce alternative finishes. Where deep profiles are required, expanded polystyrene and polyurethane foam can be carved out to produce highly sculptural designs.

Abraded, acid etched and polished finishes

Light abrasion with sand paper may be applied to in situ or precast concrete. Acid etching is normally limited to precast concrete due to the hazards associated with the use of acids on site. Both techniques remove the surface laitance (cement-rich surface layer) to create a more stone-like finish with some exposure of the aggregate. Polishing with carborundum abrasives produces a hard shiny finish, imparting full colour brightness to the aggregate. It is, however, a slow and therefore expensive process.

Exposed aggregate finishes

The exposure of the coarse aggregate in concrete, by removal of the smooth surface layer formed in contact with the formwork, produces a concrete with a more durable finish and better weathering characteristics, which is frequently aesthetically more pleasing. Smooth, profiled and deeply moulded concrete can all be treated, with the visual effects being largely dependent on the form and colour of the coarse aggregates used. While gap-graded coarse aggregates can be used in both precast and in situ exposed aggregate finishes, precasting gives additional opportunities for the uniform placement of the aggregate. In face-down casting, flat stones can be laid on the lower face of the mould, which can be pretreated with retardant to slow the hardening of the surface cement. In face-up casting, individual stones can be pressed into the surface either randomly or to prescribed patterns without the use of retardants. Alternatively, a special facing mix can be used on the fairfaced side of the panel, with the bulk material made up with a cheaper standard mix. The aggregate has to be exposed by washing and brushing when the concrete has cured sufficiently to be self-supporting. The use of a retarder applied to the formwork face enables the timing of this process to be less critical. The surface should be removed to a depth no more than one third of the thickness of the aggregate to eliminate the risk of it becoming detached. An alternative method of exposing the aggregate in both precast and in situ concrete involves the use of abrasive blasting. Depending on the size of grit used and the hardness of the concrete, a range of finishes including sculptural designs can be obtained.

Tooled concrete finishes

A range of textures can be obtained by tooling hardened concrete either by hand or mechanically. Generally, a high-quality surface must be tooled as blemishes can be accentuated rather than eliminated by tooling. Only deep tooling removes minor imperfections such as blow-holes and the effects of slight formwork misalignment. Hand tooling is suitable for a light finish on plain concrete and club hammering can be used on a ribbed finish. Where deep tooling is anticipated, allowance must be made for the loss of cover to the steel reinforcement. The exposed aggregate colour in tooled concrete is less intense than that produced by wash-and-brush exposure due to the effect of the hammering on the aggregate. Standard mechanical
tools are the needle-gun, the bush hammer and the point-tool (Fig. 3.23). A range of visual concrete finishes is illustrated in Fig. 3.24.

Weathering of concrete finishes

The weathering of exposed visual concrete is affected by the local microclimate, the concrete finish itself and the detailing used to control the flow of rainwater over the surface. It is virtually impossible to ensure that all sides of a building are equally exposed, as inevitably there will be a prevailing wind and rain direction which determines the weathering pattern. It is therefore likely that weathering effects will differ on the various elevations of any building. Some elevations will be washed regularly, whilst others may suffer from an accumulation of dirt which is rarely washed. However, this broad effect is less likely to cause unsightly weathering than the pattern streaking on individual facades.

The choice of concrete finish can have a significant effect on the weathering characteristics. Good-quality dense uniform concrete is essential if patchy weathering is to be avoided, and generally a rougher finish is likely to perform better than a smooth as-cast finish. Profiling and the use of exposed aggregates have the advantage of dictating the flow of rainwater, rather than letting it run in a random manner, but dirt becomes embedded in the hollows. Dark aggregates and bold modelling minimise the change in appearance on weathering, but generally, exposed non-absorbent aggregates are likely to give the best weathering performance. Horizontal surfaces may be subject to organic growths and this effect is increased by greater surface permeability.

Careful detailing is necessary to ensure a dispersed and controlled flow of water over the washed areas. The water should then be collected or shed clear by bold details to prevent pattern staining below. Water collected onto horizontal surfaces should not be allowed to run down facades below, so copings, sills and string courses all should be provided with drips to throw the water off the building face; alternatively, water should be removed by gutters. Multistorey facades should be articulated with horizontal features to throw the water off, at least at each storey height. Only on seriously exposed facades where strong winds are likely to cause rain to be driven upwards, should small horizontal drip projections be avoided. Where concrete is modelled, due consideration should be given to the direction of flow and the quantity of rainwater anticipated.

EXTERNAL RENDERING

 Renders are used to provide a durable and visually acceptable skin to sound but unattractive construction. Renders can reduce rain penetration and maintain the thermal insulation of walls. The finishes illustrated in Fig. 3.25 are all appropriate for external use. In each case it is essential to ensure good adhesion to the background. Where a good mechanical key, such as raked-out brickwork joints, is not present, an initial stipple coat of sand, cement, water and appropriate
Fig. 3.24  Selection of visual concrete finishes
bonding agent (e.g. styrene-butadiene-rubber) is required to create a key. Bonding is also affected by the suction or absorbency of the background; where suction is very high, walls may be lightly wetted before the rendering is applied. Metal lathing may be used over timber, steel or friable masonry to give a sound background. Two or three coats of rendering are normally applied; in either case the successive coats are weaker by a reduction in thickness or strength of the mix. Smooth renders require careful workmanship for external work, as they may craze if finished off with a steel rather than a wooden float.

Generally, permeable renders are more durable than dense impermeable renders as the latter may suffer cracking and subsequent localised water penetration. Sands for external renderings should be sharp rather than soft. The design detailing of rendering is important to ensure durability. The top edges of rendering should be protected from the ingress of water by flashings, copings or eaves details. Rendering should stop above damp-proof course level and be formed into a drip with an appropriate edging bead. Rainwater run-off from sills and opening heads should be shed away from the rendering to prevent excessive water absorption at these points, which would lead to deterioration and detachment of the rendering. Figure 3.26 illustrates the striking visual effect of the rendered blockwork student halls of residence at the University of East London adjacent to the Royal Albert Dock.

**Roughcast render**

Roughcast consists of a wet mix of cement (1 part), lime (0.5 part), sand (3 parts) and 5–15 mm shingle or crushed stone (1.5 parts), which is applied to walls by throwing from a hand scoop.

**Dry-dash render**

A 10 mm coat of cement (1 part), lime (1 part) and sand (5 parts) is applied to the wall, and whilst it is still wet, calcined flint, spar or shingle is thrown onto the surface and tamped in with a wooden float.

**Scraped finish**

A final coat of cement (1 part), lime (2 parts) and sand (9 parts) is applied and allowed to set for a few hours, prior to scraping with a rough edge (e.g. saw blade) to remove the surface material. After it has been scraped, the surface is lightly brushed over to remove loose material.

**Textured finishes**

A variety of finishes can be obtained by working the final rendering coat with a float, brush, comb or other tool to produce a range of standard textured patterns. Pargeting, in which more sophisticated patterns are produced, has its cultural roots in Suffolk and Essex.

**Tyrolean finish**

For a Tyrolean finish, cement mortar is spattered onto the wall surface from a hand-operated machine. Coloured mixes may be used.
Painted rendered finishes

Most renderings do not necessarily need painting; however, smooth renderings are frequently painted with masonry paint to reduce moisture absorption and give colour. Once painted, walls will need repainting at regular intervals.

Concrete components

In addition to the use of concrete for the production of large in situ and precast units, concrete bricks (Chapter 1) and concrete blocks (Chapter 2), the material is widely used in the manufacture of small components, particularly concrete tiles, slates and paving slabs.

CONCRETE ROOFING TILES AND SLATES

Concrete plain and interlocking slates and tiles form a group of highly competitive pitched roofing materials, with concrete interlocking tiles remaining the cheapest visually acceptable unit pitched-roof product. Plain and feature double-lap and interlocking tiles are manufactured to a range of designs, many of which emulate the traditional clay tile forms (Fig. 3.27). Concrete plain tiles may be used on pitches down to 35°, whilst the ornamental tiles are appropriate for vertical hanging and pitches down to 70°. The ranges of colours usually include both granular and through colour finishes. Standard ranges of concrete interlocking tiles and slates can be used in certain cases down to roof pitches of 17.5°, and for some shallow-pitched roofs the concrete tiles are laid to broken bond. One interlocking product emulates the appearance of plain tiles, but can be used down to a minimum pitch of 22.5°. Colours include brown, red, rustic and grey in granular and smooth finish. A limited range of polymer-surfaced concrete interlocking tiles may be used at roof rafter pitches as low as 12.5°, provided that all tiles are clipped to prevent wind lift.

Concrete interlocking slates are manufactured with either a deep flat profile, giving a stone/slate
Concrete roofing tiles and slates appearance, or with a thin square or chamfered leading edge to simulate natural slate. Surfaces can be simulated riven or smooth in a range of colours including grey, blue, brown, buff and red. Matching accessories for either traditional mortar bedding or dry-fixing for ridges, hips and verges are available, together with appropriate ventilation units.

CONCRETE PAVING SLABS AND TILES

Grey concrete paving slabs are manufactured from Portland cement mixes with pigments added to produce the standard buff, pink and red colours. Standard sizes include 900 × 600 mm, 750 × 600 mm and 600 × 600 mm × 50 mm, but a wide range of smaller and thinner units is available for the home improvement market including 600 × 600 mm, 600 × 450 mm, 450 × 450 mm and 400 × 400 mm by 30–40 mm. Thicker units (65 and 70 mm) are manufactured to withstand light traffic. Plain pressed slabs may have slightly textured surfaces, whilst cast slabs are available with smooth, simulated riven stone, terrazzo or textured finishes. Toolled textured-finished slabs and associated products are available for use in visually sensitive locations. In addition to the standard square and rectangular units, a wide range of decorative designs including hexagonal, simulated bricks and edging units is generally available.

Tile units for roof terraces, balconies and external pedestrian areas in frost-resistant Portland cement concrete are manufactured to square and hexagonal designs in a range of standard red, brown and buff colours. They are suitable for laying on asphalt, bitumen membrane roofing, inverted roofs and sand/cement screed. Typical sizes are 305 × 305 mm and 457 × 457 mm with thicknesses ranging from 25 to 50 mm.

NITROGEN OXIDE ABSORBING CEMENT, PAVERS AND TILES

Titanium oxide-coated paving stones and roofing tiles absorb nitrogen oxides produced by road traffic and convert them by a photochemical reaction into nitrogen and oxygen, thus reducing the harmful pollution within trafficked zones. Nitrogen oxide levels can be reduced in urban areas by between 10% and 20%. Furthermore, the titanium-coated paving slabs are easier to clean than standard concrete pavers. Cement incorporating nano-crystalline titanium oxide is commercially available for construction use where pollution control is particularly required.

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Introduction

Timber, arguably the original building material, retains its prime importance within the construction industry because of its versatility, diversity and aesthetic properties. About 20% of the earth’s land mass is covered by forests, divided roughly two thirds as hardwoods in temperate and tropical climates and one third as softwoods within temperate and colder regions. Approximately a third of the annual worldwide timber harvest is used in construction, and the rest is consumed for paper production, as a fuel, or wasted during the logging process.

Environmental issues, raised by the need to meet the current and future demands for timber, can only be resolved by sustainable forest developments. In temperate climate forests, clear cutting, in which an area is totally stripped, followed by replanting, is the most economical, but the shelterwood method, involving a staged harvest over several years, ensures that replacement by young trees becomes established as the mature ones are felled. The managed forests of North America and Scandinavia are beginning to increase in area due to additional planting for future use. The deforestation of certain tropical regions has allowed wind and rain to erode the thin topsoil, leaving inhospitable or desert conditions; furthermore, the overall reduction in world rain forest areas is contributing significantly to the greenhouse effect by reducing the rate of extraction of carbon dioxide from the atmosphere.

Compared to the other major construction materials, timber as a renewable resource is environmentally acceptable. As illustrated in Fig. 4.1, brick, steel, plastics and particularly aluminium, all use more energy in their production, thus contributing considerably to carbon dioxide emissions. Trees require little energy for their conversion into usable timber, and young replacement trees are particularly efficient at absorbing carbon dioxide and releasing oxygen into the atmosphere. Temperate and tropical hardwoods, when suitably managed, can be brought to maturity within a human lifespan, whereas softwoods take half that period to mature. Timber products manufactured from reconstituted and waste wood add to the efficient use of forestry. Timber frame, one of the modern methods of construction (MMC), now accounts for approximately 22% of new-build housing within the UK.

Great emphasis is now placed on timber certification schemes, which track the material as a chain of custody from source to user, to ensure the accuracy of environmental claims being made. In the UK, the two international schemes, the Forest Stewardship Council (FSC) and the Programme for the Endorsement of Forest Certification (PEFC), ensure that timber with their labels has been harvested from properly managed sustainable sources. Products purchased under the labels of the Canadian Standards Association (CSA) or the American Sustainable Forestry Initiative (SFI) are also certified by PEFC. In addition TRADA, as an accreditation body, undertakes to certify other chains of custody, which are subsequently routinely verified. One scheme accredited by PEFC for tropical timber is the Malaysian Timber Certification Scheme (MTCS). Materials available under these certified schemes include structural timber, joinery timber and the timber products, plywood, particleboard, flooring, MDF and OSB.
Timber

The Study Centre at Darwin College, Cambridge (Fig. 4.2), which occupies a narrow site overlooking the River Cam, is designed to accommodate both books and computers. It is a load-bearing masonry and timber building which features the extensive use of English oak, including massive paired columns to the first-floor reading room which is partly cantilevered over the river. The columns in green oak have characteristic shakes and splits giving an impression of great age, and these contrast with the refined oak and oak veneer of the floors, windows frames and furniture. Joints in the green oak are held by stainless steel fixings, which can be tightened as the timber dries and shrinks. The use of oak throughout gives unity to the building, which sits comfortably within its highly sensitive location.

Metabolism of the Tree

The tree, a complex living organism, can be considered in three main sections: the branches with their leaves, the trunk (or bole) and the roots (Fig. 4.3). The roots anchor the tree to the ground and absorb water with dissolved minerals from the soil. The leaves absorb carbon dioxide from the air and in the presence of sunlight, together with chlorophyll as a catalyst, combine carbon dioxide with water to produce sugars. Oxygen, a by-product of the process, diffuses out of the leaves. The sugars in aqueous solution are transported down the branches and trunk to be subsequently converted, where required for growth, into the cellulose of the tree. The trunk gives structural strength to the tree and acts as a store for minerals and food such as starch and also as a two-way transport medium.

The tree is protected from extremes of temperature and mechanical damage by the bark, inside which is the bast layer which transports the sugars synthesised in the leaves downwards. Radial rays then move the food into the sapwood cells for storage. Inside the bast is the thin and delicate cambium, which is the growing layer for the bark and sapwood. Growth only takes place when the cambium layer is active, which in temperate climates is during the spring and summer seasons.

A transverse section through the bole shows the growth rings. These are sometimes referred to as annual rings, but unusual growth patterns can lead to multiple rings within one year, and in tropical climates, where seasonal changes are less pronounced, growth rings may be indistinct and not annual. The growth rings are apparent because the early wood produced at the start of the growing season tends to be made from larger cells of thinner walls and is thus softer and more porous than the late wood produced towards the end of the growing season. Each year as the tree matures with the production of an additional growth ring, the cells of an inner ring are strengthened by a process of secondary thickening. This is followed by lignification in which the cell dies. These cells are no longer able to act as food stores, but now give increased structural strength to the tree. The physical changes are often associated with a darkening of the timber due to the incorporation into the cell walls of so-called extractives, such as resins in softwoods or tannins in oak. These are natural wood preservatives which make heartwood more durable than sapwood.
CONSTITUENTS OF TIMBER

The main constituents of timber are cellulose, hemi-cellulose and lignin, which are natural polymers. Cellulose, the main constituent of the cell walls, is a polymer made from glucose, a direct product of photosynthesis within the leaves of the tree. Glucose molecules join together to form cellulose chains containing typically 10,000 sugar units (Fig. 4.4). Alternate cellulose chains, running in opposite directions to each other, form a predominantly well-ordered crystalline material. It is this crystalline chain structure which
Fig. 4.3 Metabolism of the tree (after Everett, A. 1994: *Mitchell’s Materials*. 5th edition. Longman Scientific & Technical)

![Metabolism of the tree](image)

**Photosynthesis**

\[ 6\text{CO}_2 + 6\text{H}_2\text{O} \xrightarrow{\text{sunlight/chlorophyll}} \text{C}_6\text{H}_{12}\text{O}_6 + 6\text{O}_2 \]

carbon dioxide \hspace{1cm} water \hspace{1cm} glucose \hspace{1cm} oxygen

![Structure of cellulose](image)

Fig. 4.4 Structure of cellulose
gives cellulose its fibrous properties, and accounts for approximately 45% of the dry weight of the wood.

Hemicelluloses, which account for approximately 25% of the weight of wood, have more complex partially crystalline structures composed of a variety of other sugars. The molecular chains are shorter than those in cellulose producing a more gelatinous material. Lignin (approximately 25% by weight of the timber) is an insoluble non-crystalline polymeric material. Its main constituents are derivatives of benzene combined to form a complex branched-chain structure.

The three major components are combined to form microfibrils which are in turn the building blocks for the cell walls. Crystalline cellulose chains are surrounded by semi-crystalline hemicellulose, and then a layer of non-crystalline cellulose and are finally cemented together with lignin (Fig. 4.5). Millions of these microfibrils are built up in layers to form the individual cell walls. It is this composite structure which gives timber its physical strength, with the cellulose contributing mainly to the tensile properties and the hemicellulose and lignin to the compressive strength and elasticity.

In addition to the three major constituents and significant quantities of water, timbers contain many minor constituents; some, such as resins, gums and tannins, are associated with the conversion of sapwood to heartwood. Starch present in sapwood is attractive to fungi, and inorganic granules such as silica make working certain tropical hardwoods, such as teak, difficult. The various colours present in different timbers arise from these minor constituents, as the various celluloses and lignin are virtually colourless. Some colours are fixed to the polymeric chains, but others are light-sensitive natural dyes which fade on prolonged exposure to sunlight, unless the timber is coated with an ultraviolet-absorbing finish.

HARDWOODS AND SOFTWOODS

Commercial timbers are defined as hardwoods or softwoods according to their botanical classification rather than their physical strength. Hardwoods (angiosperms) are from broad-leaved trees, which in temperate climates are deciduous, losing their leaves in autumn, although in tropical climates, when there is little seasonal variation, old leaves are constantly being replaced by new. Softwoods (gymnosperms) are from conifers, characteristically with needle-shaped leaves, and growing predominantly in the northern temperate zone. Mostly they are evergreen, with the notable exception of the European larch (Larix decidua) and the Californian redwood (Sequoia sempervirens), the world’s largest tree with a height of over 100 metres.

Although the terms hardwood and softwood arose from the physical strength of the timbers, paradoxically, balsa (Ochroma lagopus) used for model making is botanically a hardwood, whilst yew (Taxus baccata), a strong and durable material, is defined botanically as a softwood. Under microscopic investigation, softwoods show only one type of cell which varies in size between the rapid growth of spring and early summer (early wood) and the slow growth of the late summer and autumn (late wood). These cells, or tracheids, perform the food- and water-conducting functions and give strength to the tree. Hardwoods, however, have a more complex cell structure with large cells or vessels for the conducting functions and smaller cells or wood fibres providing the mechanical support. According to the size and distribution of the vessels, hardwoods are divided into two distinct groups. Diffuse-porous hardwoods, which include beech (Fagus sylvatica), birch (Betula pendula) and most tropical hardwoods,
have vessels of a similar diameter distributed approximately evenly throughout the timber. Ring-porous hardwoods, however, including oak (*Quercus robur*), ash (*Fraxinus excelsior*) and elm (*Ulmus procera*), have large vessels concentrated in the earlywood, with only small vessels in the latewood (Fig. 4.6). The Jerwood library of Trinity Hall, Cambridge (Fig. 4.7) illustrates the visual quality of limed oak as an architectural feature within the context of a sensitive built environment.

**TIMBER SPECIES**

Any specific timber can be defined through the correct use of its classification into family, genus and species. Thus oak, and beech are members of the Fagaceae family; beech is one genus (*Fagus*) and oak (*Quercus*) another. The oak genus is subdivided into several species, including the most common, the pedunculate oak (*Quercus robur*) and the similar but less common sessile oak (*Quercus petraea*). Such exact timber nomenclature is, however, considerably confused by the use of lax terminology within the building
industry; for example, both Malaysian meranti and Philippine lauan are frequently referred to as Philippine mahogany, and yet they are from a quite different family and genus to the true mahogany (Swietenia) from the West Indies or Central America. This imprecision can cause the erroneous specification or supply of timber, with serious consequences. Where there is the risk of confusion, users should specify the correct genus and species.

The standard BS EN 13556: 2003 lists both hardwoods (dicotyledons) and softwoods (gymnosperms) used within Europe, with a four-letter code. The first two letters are a distinctive combination referring to the genus (e.g. oak – Quercus – QC). The third and fourth letters refer to the particular species; thus, European oak – Quercus petraea is QC and American red oak – Quercus rubra is QCXR. Typical softwoods are western red cedar – Thuja plicata – THPL and Scots pine – Pinus sylvestris – PNSY.

Softwood accounts for approximately 80% of the timber used in the UK construction industry. Pine (European redwood) and spruce (European white-wood) are imported from Northern and Central Europe, whilst western hemlock, spruce, pine, and fir are imported in quantity from North America. Forest management in these areas ensures that supplies will continue to be available. Smaller quantities of western red cedar, as a durable lightweight cladding material, are imported from North America, together with American redwood from California, pitch pine from Central America and para pine from Brazil. Increasingly, New Zealand, South Africa and Chile are becoming significant exporters of renewable timber. The UK production of pine and spruce provides only about 10% of the national requirements while Ireland plans to be self-sufficient early in the next century.

Over 100 different hardwoods are used in the UK, although together beech, oak, sweet chestnut, meranti, lauan, elm, American mahogany and ramin account for over half of the requirements. Approximately half of the hardwoods used in the UK come from temperate forests in North America and Europe including Britain, but the remainder, including the durable timbers such as iroko, mahogany, sapele and teak, are imported from the tropical rain forests. The Great Oak Hall at Westonbirt Arboretum, Gloucestershire (Fig. 4.8), illustrates the use of ‘medieval’ construction systems within a modern building by using ‘green’ oak fixed with dowels and wedges.

Since 1965, 6.5% of the Amazon forest has been lost, but much of this deforestation has been for agricultural purposes, with more than three quarters of the timber felled used as a local fuel rather than exported as timber. With the growing understanding of the environmental effects of widespread deforestation, some producer governments are now applying stricter controls to prevent clear felling and encourage sustainable harvesting through controlled logging. Other imported naturally durable hardwoods, available in long lengths, include eki, greenheart and opepe, whilst UK-produced sweet chestnut is durable and an appropriate structural timber. Some timbers not previously used within the UK, such as jatoba (Hymenaea courbaril), are now being imported from South America.

**CONVERSION**

Conversion is the process of cutting boles or logs into sections prior to seasoning. Subsequent further cutting into usable sizes is called manufacture. Finishing operations involving planing and sanding produce a visually smooth surface but reduce the absorption of penetrating wood stains. Timber for solid sections is sawn, whereas thin layers for plywood are peeled and veneers are usually sliced across the face of the log to maximise the visual effect of colour and figure, which is the pattern effect seen on the longitudinal surface of cut wood.

**Types of cut**

The two main types of cut, plain sawn and quarter sawn, refer to the angle between the timber face and the growth rings. This is best observed from the end of the timber, as in Fig. 4.9. If the cut is such that the growth rings meet the surface at less than 45° then the timber is plain sawn. Timber with this type of cut tends to have a more decorative appearance but a greater tendency to distort by cupping. Timber cut with the growth rings meeting the surface at not less than 45° is quarter sawn. Such timber is harder wearing, weather-resistant and less likely to flake. If a log is cut through and through, which is most economical, then a mixture of plain and quarter-sawn timber is produced. Quarter sawing is more expensive as the log requires resetting for each cut and more waste is produced; however, the larger sections will be more dimensionally stable. The centre of the tree, the pith, is frequently soft and may be weakened by splits or shakes. In this case, the centre is removed as a boxed heart.
Fig. 4.8 Traditional oak construction—Great Oak Hall, Westonbirt Arboretum, Gloucestershire. Architects: Roderick James Architects. Photograph: Arthur Lyons

Fig. 4.9 Conversion of timber

**Sizes**

BS EN 1313-1: 1997 defines the standard sizes of sawn softwood timbers at 20% moisture content (Table 4.1). Widths over 225 mm and lengths over 5 m are scarce and expensive, but finger jointing (BS EN 385: 2001), which can be as strong as the continuous timber, does allow longer lengths to be specified. Regularising, which ensures uniformity of width of a sawn timber, reduces the nominal section by 3 mm (5 mm over 150 mm) and planing on all faces or ‘processed all round’ (PAR) reduces, for example, a 47 × 100 mm section to 44 × 97 mm (Table 4.2). Hardwood sizes are more variable due to the diversity of hardwood species, but preferred sizes to BS EN 1313-2: 1999 are specified in Table 4.1. Hardwoods are usually imported in random widths and lengths; certain structural hardwoods such as iroko (*Chlorophora excelsa*) are available in long lengths (6–8 m) and large sections. Tolerances for acceptable deviations from target sizes for softwood are given in BS EN 1313-1: 1997 and BS EN 336: 2003 (Table 4.3). The latter defines two tolerance levels for
Table 4.1  Standard sizes of softwoods and hardwoods

Standard sizes of sawn softwood (20% moisture content) to BS EN 1313-1: 1997.

<table>
<thead>
<tr>
<th>Thickness (mm)</th>
<th>Width (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>75 100 115 125 138 150 200 225 250 275 300</td>
</tr>
<tr>
<td>16</td>
<td>V V V V V V</td>
</tr>
<tr>
<td>19</td>
<td>V V V V V V</td>
</tr>
<tr>
<td>22</td>
<td>V V V V V V V V</td>
</tr>
<tr>
<td>25</td>
<td>V V V V V V V V V V V</td>
</tr>
<tr>
<td>32</td>
<td>V V V V V V V V V V V</td>
</tr>
<tr>
<td>38</td>
<td>x √ x √ x √ x x x x x x</td>
</tr>
<tr>
<td>47</td>
<td>x x x x x x x x x x</td>
</tr>
<tr>
<td>50</td>
<td>x √ √ √ √ √ √ x x</td>
</tr>
<tr>
<td>63</td>
<td>√ √ √ √ x x</td>
</tr>
<tr>
<td>75</td>
<td>x x √ √ √ √ x x x</td>
</tr>
<tr>
<td>100</td>
<td>x x x x x x</td>
</tr>
<tr>
<td>150</td>
<td>x x x x</td>
</tr>
<tr>
<td>250</td>
<td>x x x x x x</td>
</tr>
<tr>
<td>300</td>
<td>x x x x x x</td>
</tr>
</tbody>
</table>

Sizes marked with a tick indicate preferred EU sizes.
Sizes marked with a cross are the complementary UK preferred sizes.
Sizes marked with a V are the additional UK customary sizes.

Customary lengths of timber to BS EN1313-1: 1997

<table>
<thead>
<tr>
<th>Length m</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.80</td>
</tr>
<tr>
<td>2.40</td>
</tr>
<tr>
<td>2.70</td>
</tr>
<tr>
<td>3.90</td>
</tr>
</tbody>
</table>

Lengths over 5.70 m may not be readily available without finger jointing.

Standard sizes of sawn hardwood (20% moisture content) to BS EN1313-2: 1999

<table>
<thead>
<tr>
<th>Preferred thicknesses</th>
</tr>
</thead>
<tbody>
<tr>
<td>EU 20 27 32 40 50 60 65 70 80 100 mm</td>
</tr>
<tr>
<td>Complementary thicknesses</td>
</tr>
<tr>
<td>UK 19 26 38 52 63 75 mm</td>
</tr>
</tbody>
</table>

Preferred widths

<table>
<thead>
<tr>
<th>Preferred lengths</th>
</tr>
</thead>
<tbody>
<tr>
<td>EU 10 mm intervals for widths between 50 mm and 90 mm, 20 mm intervals for widths of 100 mm or more.</td>
</tr>
<tr>
<td>Preferred lengths</td>
</tr>
<tr>
<td>EU 100 mm intervals for lengths between 2.0 m and 6.0 m, 50 mm intervals for lengths less than 1.0 m.</td>
</tr>
</tbody>
</table>

sawn surface dimensions (tolerance class 1, T1 and tolerance class 2, T2) with T2 specifying the smaller tolerance limits, also appropriate to planed timber. Customary lengths for structural softwood timber and hardwood are given in Table 4.1.

MOISTURE CONTENT AND SEASONING

As a tree is a living organism, the weight of water within it is frequently greater than the dry weight of wood itself. The water content of a tree is equal in winter and in summer, but one advantage of
Table 4.2  Maximum permitted reduction from target sawn sizes of softwoods and hardwoods by planing two opposed faces

Maximum reductions from sawn softwood sizes by planing two opposed faces (BS EN1313-1: 1997)

<table>
<thead>
<tr>
<th>Typical application</th>
<th>Reduction from basic size (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>15–35</td>
</tr>
<tr>
<td>Constructional timber</td>
<td>3</td>
</tr>
<tr>
<td>Matching and interlocking boards (not flooring)</td>
<td>4</td>
</tr>
<tr>
<td>Wood trim</td>
<td>5</td>
</tr>
<tr>
<td>Joinery and cabinet work</td>
<td>7</td>
</tr>
</tbody>
</table>

Maximum reductions from sawn hardwood sizes by planing two opposed faces (BS EN1313-2: 1999)

<table>
<thead>
<tr>
<th>Typical application</th>
<th>Reduction from basic size (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>15–25</td>
</tr>
<tr>
<td>Flooring, matchings, interlocked boarding and planed all round</td>
<td>5</td>
</tr>
<tr>
<td>Trim</td>
<td>6</td>
</tr>
<tr>
<td>Joinery and cabinet work</td>
<td>7</td>
</tr>
</tbody>
</table>

The primary aim of seasoning is to stabilise the timber to a moisture content that is compatible with the equilibrium conditions under which it is to be used, so that subsequent movement will be negligible. At the same time, the reduction in water content to below 20% will arrest any incipient fungal decay, which can only commence above this critical level. Drying occurs with evaporation of water from the surface, followed by movement of moisture from the centre of the timber outwards due to the creation of a vapour–pressure gradient. The art of successful seasoning is to control the moisture loss to an appropriate rate. If the moisture loss is too rapid then the outer layers shrink while the centre is still wet and the surface sets in a distended state (case hardening) or opens up in a series of cracks or checks. In extreme cases as the centre subsequently dries out and shrinks it may honeycomb.

Air seasoning

Timber, protected both from the ground and from rain, is stacked in layers separated by strips of wood called stickers which, depending on their thickness, control the passage of air (Fig. 4.10). The air, warmed by the sun and circulated by the wind, removes moisture from the surface of the timbers. The timber ends are protected by waterproof coatings (bituminous paint) to prevent rapid moisture loss, which would cause splitting. Within the UK a moisture content of between 17% and 23% may be achieved within a few months for softwoods, or over a period of years for hardwoods.
Kiln drying

Kiln drying or seasoning is effected by heating within a closed chamber, which can be programmed to a precise schedule of temperature and humidity. Thus, drying to any desired moisture content can be achieved without significant degradation of the timber, although some early examples of kiln-dried timber showed serious damage through the use of inappropriate drying schedules. For economic reasons, timber is frequently air seasoned to fibre saturation point, followed by kiln drying to the required moisture content. This roughly halves the necessary kiln time and fuel costs. A typical softwood load would be dried from fibre saturation point within a few days and hardwood within two to three weeks.

Seasoned timber, if exposed to rain on site, will reabsorb moisture. Good site management is therefore necessary to protect timber from both physical damage and wetting prior to its use. The heating up of new buildings by central heating systems can cause rapid changes in the moisture content of joinery timber and lead to shrinkage, cracking and splitting.

MOISTURE MOVEMENT

Wood is an anisotropic material, with differing moisture movements along the three principal axes; tangential, radial and longitudinal (Fig. 4.11). The highest moisture movement is tangential to the grain, next being radial with the least along the grain. Typical figures are given in Table 4.4. The larger the ratio between tangential and radial movement, the greater the distortion. Moisture movements are conventionally quoted for a change in relative humidity from 90% to 60% at 25°C. The BRE classifies woods into three categories according to the sum of radial and tangential movements effected by this standard change in relative humidity.

<table>
<thead>
<tr>
<th>Tree</th>
<th>Moisture movement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Afzelia</td>
<td>Medium moisture movement</td>
</tr>
<tr>
<td>Agba</td>
<td>Large moisture movement</td>
</tr>
<tr>
<td>Iroko</td>
<td>Hardwoods</td>
</tr>
<tr>
<td>Jelutong</td>
<td>Medium moisture movement</td>
</tr>
<tr>
<td>Lauan</td>
<td>Small moisture movement (less than 3.0%)</td>
</tr>
<tr>
<td>Mahogany, African</td>
<td>Medium moisture movement (3.0–4.5%)</td>
</tr>
<tr>
<td>Mahogany, American</td>
<td>Large moisture movement (over 4.5%)</td>
</tr>
<tr>
<td>Meranti</td>
<td>Hardwoods</td>
</tr>
<tr>
<td>Merbau</td>
<td>Hardwoods</td>
</tr>
<tr>
<td>Obeche</td>
<td>Hardwoods</td>
</tr>
<tr>
<td>Teak</td>
<td>Hardwoods</td>
</tr>
</tbody>
</table>

Softwoods

<table>
<thead>
<tr>
<th>Tree</th>
<th>Moisture movement</th>
</tr>
</thead>
<tbody>
<tr>
<td>Corsican pine</td>
<td>European redwood</td>
</tr>
<tr>
<td>Douglas fir</td>
<td>European whitewood</td>
</tr>
<tr>
<td>Sitka spruce</td>
<td>Parana pine</td>
</tr>
<tr>
<td>Western hemlock</td>
<td>Radiata pine</td>
</tr>
<tr>
<td>Western red cedar</td>
<td>Scots pine</td>
</tr>
</tbody>
</table>

Moisture movement is assessed on the sum of the radial and tangential movements for a change in environmental conditions from 60 to 90% relative humidity.
humidity. Small movement is defined as less than 3%, medium between 3% and 4.5% and large over 4.5%. Large movement timbers are not recommended for use as cladding.

**TIMBER DEFECTS**

Timber, as a natural product, is rarely free from blemishes or defects, although in some instances, such as knotty pine, waney-edge fencing timber or burr veneers, the presence of the imperfections enhances the visual quality of the material. Timber imperfections can be divided into three main categories: natural, conversion and seasoning defects, according to whether they were present in the living tree, or arose during subsequent processing. Additionally, timber may be subject to deterioration by weathering, fungal and insect attack and fire. These latter effects are discussed later in the chapter.

**Natural defects**

**Knots**

Knots are formed where branches of the tree join the trunk (Fig. 4.12). Where the wood fibres of the branch are continuous with the trunk, then a live knot is produced. If, however, the branch is dead, or bark becomes incorporated into the trunk, a dead knot is produced. This is liable to be loose, leads to incipient decay and causes structural weakness.

Knots are described as face, edge, splay, margin or arris, depending on how they appear on the faces of converted timber. Additionally, knots may appear as clusters, and range in size from insignificant to many millimetres across. Frequently they are hard to work, and in softwoods contain quantities of resin, which will continue to seep out unless the wood is sealed before painting.

**Natural inclusions**

Many minor defects occur to varying degrees in different varieties of timber. Bark pockets occur where pieces of bark have been enclosed within the timber as a result of earlier damage to the cambium or growth layer. Pitch pockets and resin streaks, containing fluid resin, are frequently seen along the grain of softwoods; their extent in usable timber is limited by BS EN 942: 2007.

**Compression and tension wood**

Trees leaning owing to sloping ground, or subject to strong prevailing winds, produce reaction wood to counteract these forces. In softwoods, compression wood is produced which is darker in colour due to an increased lignin content. In hardwoods, tension wood is produced, which is lighter in colour owing to the presence of an extra cellulose layer in the cell walls. Both types of reaction wood have an abnormally high longitudinal shrinkage, causing distortion on seasoning; furthermore, tension wood tends to produce a rough surface when it is machined.

**Abnormal growth rings**

The width of the growth rings is an indicator of the growth rate and timber strength, with the optimum ranged around five rings per centimetre for softwoods and three rings per centimetre for hardwoods depending on the species. Excessively fast or slow growth rates give rise to weaker timber owing to a reduction in the proportion of the stronger late wood or its production with thinner walled fibres.

**Conversion defects**

**Sloping grain**

For maximum strength, timber should be approximately straight grained, as with increasing slope of the grain (Fig. 4.13), there is a proportionate reduction in bending strength, ranging typically from 4% at 1 in 25 to 19% at 1 in 10. The British Standard (BS 5756: 2007) limits the slope of grain in visual strength graded structural tropical hardwood (HS) to 1 in 11. BS 4978:
Fig. 4.13 Sloping grain

2007 limits the slope of grain in visual strength graded softwoods to 1 in 6 for the general structural grade (GS) and 1 in 10 for the special structural grade (SS). The slope of grain in timber for internal and external windows, doors and door frames is limited to 1 in 10.

Wane
Wane is the loss of the square edge of the cut timber due to the incorporation of the bark or the curved surface of the trunk. A degree of wane is acceptable in structural and floor timbers (BS 4978: 2007 and BS 1297: 1987), and is a special feature in waney-edge fencing.

Seasoning defects
Some of the commonest defects in timber are associated with the effects of seasoning. During the seasoning process, the contraction of the timber is different in the three major directions; furthermore, as described in Section ‘Moisture movement’ (page 115), the outside of the timber tends to dry out more rapidly than the interior. These combined effects cause distortion of the timber including warping and the risk of rupture of the timber to produce surface checks and splits (Fig. 4.14).

Shakes
Major splits within timber are termed shakes, and these may result from the release of internal stresses within the living tree on felling and seasoning; however, some fissures may be present within the growing timber. Commonly, shakes are radial from the exterior of the trunk, but star shakes which originate at the centre or pith may be associated with incipient decay. Ring shakes follow round a particular growth ring and are frequently caused by the freezing of the sap in severe winters.

Fig. 4.14 Warping splits and checks

Appearance grading
The standard BS EN 975 Part 1: 2009 and Part 2: 2004 set the criteria for appearance grading of hardwoods, specifically for oak and beech and European poplars, respectively. Grades relate the number and size of the natural, seasoning and conversion defects visible on the timber face. Generally, four quality categories are listed for different forms such as boules, individual boards, edged boards and finished timber. The standard BS EN 1611-1: 2000 specifies five grades of appearance grading for the softwoods; European spruces, firs, pines, Douglas fir and larches according to the number and size of observable defects.

SPECIFICATION OF TIMBER
The building industry uses timber for a wide range of purposes from rough-sawn structural members to claddings, trim and highly machined joinery. The specification of timber for each use may involve defining the particular hardwood or softwood, where particular visual properties are required. However, for the majority of general purposes, where strength and
durability are the key factors, timber is specified by either a strength class, or a combination of timber species and strength grade.

In addition to strength class or grade, the specification of structural timber should include lengths and cross-section sizes, surface finish or tolerance class, moisture content and any preservative or special treatments (BRE Digest 416: 1996).

**Strength grading**

Strength grading is the measurement or estimation of the strength of individual timbers, which allows each piece to be used to its maximum efficiency. It may be done visually, a slow and skilled process, or within a grading machine which tests flexural rigidity. The European standard BS EN 14081-1: 2005 broadly outlines the requirements for visual and machine grading of timber, including the requirements for marking. The standards for the visual grading of hardwood and softwood within the UK are BS 5756: 2007 and BS 4978: 2007, respectively.

**Visual strength grading**

Each piece of timber is inspected for distortions, growth ring size and slope of grain, then checked against the set permissible limits for the number and severity of the natural defects, such as knots, waney-edge, and fissures. The timber is then assigned to a grade and stamped accordingly. Softwood timber is assessed as special structural grade (SS), general structural grade (GS) or reject. Hardwoods are graded to THA or THB (heavy structural temperate hardwood), TH1 or TH2 (general structural temperate hardwood) or reject as appropriate. (The higher grades THA and TH1 in each category have fewer natural defects such as knots or sloping grain.) The one grade for tropical hardwood is HS (structural tropical hardwood).

**Machine strength grading**

Each piece of timber is quickly inspected for any distortions which may cause it to be rejected manually, or any serious defects within 500 mm of either end, at which point machine testing is ineffective. It is then tested, usually by one of two systems of contact or bending-type grading machines. In both techniques, the timber is moved through a series of rollers and the machine measures either the load required to produce a fixed deflection or the deflection produced by a standard load. Either technique is measuring stiffness which is then related to timber strength and therefore a grading standard.

The three grading machines used within the UK are shown in Fig. 4.15. In the constant deflection machine, the timber is then moved through a series of rollers which press it firmly against a curved metal plate. The force required to bend the timber to this standard deflection is determined by a series of transducers and from this data the timber strength is computed. However, a second pass through this machine is required to eliminate the effects of bow. The constant load system applies a defined lateral load, depending on the sample thickness, and the resulting deflection, with automatic adjustment for bow, indicates the timber grade. A more sophisticated system measures the forces necessary to bend the timber into an S-shape with two fixed deflections, thus neutralising the effects of any natural bow in the material.

As the timber leaves the machine it is stamped with the standard number, the strength class, M (machine graded), together with information on its species, wet/dry graded state, the producer and the certification body. Timber single species are coded with four letters to BS EN 13556: 2003 (e.g. Scots pine – *Pinus sylvestris* PNSY). Species combinations have four letters commencing with W (e.g. British spruce – *Picea sitchensis* and *Picea abies* – WPCS). Timber for trussed rafters may be colour-coded according to its strength class. The machine classes and associated dye colour codes are given in Table 4.5.

Recent non-contact techniques for strength grading include X-ray and stress wave systems. X-ray machines assess the density of the timber, which is then related to strength. Stress wave techniques measure either the speed through the timber, or the natural frequency, of a stress wave produced by a small impact, and relate this to strength. Both techniques offer the potential for faster throughput than conventional contact strength grading systems, although they are sensitive to timber moisture content. Some grading machines combine physical bending techniques with the use of X-ray or microwave systems for the detection of natural defects such as knots or sloping grain, respectively.

### Table 4.5 | Strength classes and the associated machine-graded colour codings

<table>
<thead>
<tr>
<th>Strength class</th>
<th>Colour code</th>
</tr>
</thead>
<tbody>
<tr>
<td>C27</td>
<td>red</td>
</tr>
<tr>
<td>C24</td>
<td>purple</td>
</tr>
<tr>
<td>C22</td>
<td>blue</td>
</tr>
<tr>
<td>C16</td>
<td>green</td>
</tr>
</tbody>
</table>
Fig. 4.15 Strength-grading machines and timber certification mark. (Grading machine diagram reproduced from Digest 476 by permission of BRE.)
Strength classes

Strength classes to BS EN 338: 2003 (Table 4.6) are defined as C14 to C50 and D18 to D70, where the prefix C refers to softwoods (coniferous) and D to hardwoods (deciduous). The number refers to the characteristic bending strength in MPa. The full specification of the strength classes gives characteristic values for density and a wide range of strength and stiffness properties, all based on sample test values. The data do not take into account any safety factors to be included in the design process. For trussed rafters, the grades TR20 and TR26 defined in BRE Digest 445: 2000 are applicable.

Table 4.7 shows softwood species and strength grade combinations in relation to strength classes for visual and machine-graded softwood timber.

Table 4.6  Relationship between strength classes and physical properties

| Strength classes to BS EN 338: 2003 — Characteristic values — softwood species |
|---------------------------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|
| C14                            | C16            | C18            | C20            | C22            | C24            | C27            | C30            | C35            | C40            | C45            | C50            |
| Strength properties MPa        |                |                |                |                |                |                |                |                |                |                |                |                |                |
| Bending                        | 14             | 16             | 18             | 20             | 22             | 24             | 27             | 30             | 35             | 40             | 45             | 50             |
| Stiffness properties GPa       |                |                |                |                |                |                |                |                |                |                |                |                |                |
| Mean modulus of elasticity     | 7              | 8              | 9              | 9.5            | 10             | 11             | 11.5           | 12             | 13             | 14             | 15             | 16             |
| parallel to the grain          |                |                |                |                |                |                |                |                |                |                |                |                |                |
| Average density kg/m³          | 350            | 370            | 380            | 390            | 410            | 420            | 450            | 460            | 480            | 500            | 520            | 550            |

| Strength classes to BS EN 338: 2003 — Characteristic values — hardwood species |
|---------------------------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|
| D18                            | D24            | D30            | D35            | D40            | D50            | D60            | D70            |
| Strength properties MPa        |                |                |                |                |                |                |                |                |                |                |                |                |                |
| Bending                        | 18             | 24             | 30             | 35             | 40             | 50             | 60             | 70             |
| Stiffness properties GPa       |                |                |                |                |                |                |                |                |                |                |                |                |                |
| Mean modulus of elasticity     | 10             | 11             | 12             | 12             | 13             | 14             | 17             | 20             |
| parallel to the grain          |                |                |                |                |                |                |                |                |                |                |                |                |                |
| Average density kg/m³          | 610            | 630            | 640            | 650            | 660            | 750            | 840            | 1080           |

| Strength classes to BS 5268-2: 2002 — Grade stresses for permissible stress design code |
|---------------------------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|----------------|
| C14                            | C16            | C18            | C22            | C24            | C27            | C30            | C35            | C40            | D30            | D35            | D40            | D45            | D50            | D60            | D70            |
| Strength properties MPa        |                |                |                |                |                |                |                |                |                |                |                |                |                |                |                |                |
| Bending parallel to the grain  | 4.1            | 5.3            | 5.8            | 6.8            | 7.5            | 9.5            | 11             | 12             | 13             | 9              | 11             | 12.5           | 16             | 18             | 23             |
| Stiffness properties GPa       |                |                |                |                |                |                |                |                |                |                |                |                |                |                |                |                |
| Minimum modulus of elasticity  | 4.6            | 5.8            | 6.0            | 6.5            | 7.2            | 8.2            | 8.2            | 9.0            | 10.0           | 6.0            | 6.5            | 7.5            | 12.6           | 15.6           | 18.0           |
| parallel to the grain          |                |                |                |                |                |                |                |                |                |                |                |                |                |                |                |                |

Notes:
Additional hardwood species strength classes D18 and D24 have been added within pr EN 338: 2008.
Timber to classes C45 and C50 may not be readily available.
C refers to coniferous softwoods and D refers to deciduous hardwoods.

Service class

The service class defines the conditions in which the timber will be used and thus the anticipated moisture content. There are three categories defined within Eurocode 5: Design of timber structures: Part 1-1: (BS EN 1995-1-1: 2004). Timber to be used in service classes 1 and 2 must be adequately protected from the weather when on site.

Service classes of wood in use:

Service class 1  Timber with a moisture content corresponding to an ambient temperature of 20°C and a relative humidity of the surrounding air only exceeding 65% for a few weeks each year.
Table 4.7  Softwood species/grade combinations which satisfy the requirements of BS EN 338: 2003 and BS 5268 Part 2: 2002 strength classes as listed in Eurocode 5 Part 1:1

<table>
<thead>
<tr>
<th>Species</th>
<th>Origin</th>
<th>European Standard BS EN 338: 2003 Strength Classes</th>
<th>Grading standard</th>
<th>C14</th>
<th>C16</th>
<th>C18</th>
<th>C22</th>
<th>C24</th>
<th>C27</th>
<th>C30</th>
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<td><strong>Grading standard</strong></td>
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<td>Radiata pine</td>
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<td>S. African pine</td>
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<td>Western red cedar</td>
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</tbody>
</table>

Notes:
Grading: BS EN 519 refers to machine grading; BS 4978 refers to visual grading.
√ indicates available machine graded to the class indicated.
GS and SS are General Structural and Special Structural visual grades, respectively.
US/Can refers to the US and Canadian visual grading standards for lumber.
The Canadian and US grades 1, 2, 3, and Sel refer to No.1, No.2, No.3 and Select, respectively.
BS EN 519: 1995 has now been superseded by BS EN 14081 – Parts 1 - 3: 2005.
Average moisture content not exceeding 12% (e.g. internal walls, internal floors except ground floor, warm roofs).

**Service class 2**

Timber with a moisture content corresponding to an ambient temperature of 20°C and a relative humidity of the surrounding air only exceeding 85% for a few weeks each year.

Average moisture content not exceeding 20% (e.g. ground floors, inner leaf of cavity walls, single leaf walls with external cladding).

**Service class 3**

Timber exposed to conditions leading to higher moisture contents than in service class 2.

Average moisture content 20% and above (e.g. exposed parts of buildings and marine structures).

### Limit state design

Eurocode 5 represents a significant change for designers in timber and timber products as it is based on limit state design, rather than permissible stress. This brings timber into line with steel and concrete for which this approach has already been taken. There are generally two limit states to be considered; first, the ultimate limit state beyond which parts of the structure may fail or collapse, and secondly the serviceability limit state beyond which excessive deformation, deflection or vibration would render the structure unfit for its purpose. The ultimate limit states are determined from the characteristic values of the loads or actions and the material properties, to which partial safety factors are applied. Generally, the characteristic values of the material properties are higher in BS EN 338: 2003 than in the BS 5268-2: 2002 grade stresses, as they are derived from laboratory tests without reductions for long-term loading or safety factors which become the responsibility of the designer. The characteristic values are used in limit state design (Eurocode 5), and the grade stresses apply to the BS 5268-2: 2002 permissible stress design code.

### MODIFIED TIMBER

The physical properties of timber can be significantly changed by either thermal treatment to produce thermal modified timber (TMT), or reagent treatment producing chemical modifications.

**Thermally modified timber**

Thermal modification involves heating the timber, usually pine or spruce but also hardwoods such as beech, maple, birch, oak and ash, to between 180°C and 240°C within an inert atmosphere to prevent combustion. Industrial processes variously use vacuum, an atmosphere of steam or nitrogen or a bath of heated oil, with process times around 36 hours. The Finnish process at 190°C produces, from pine or spruce, Thermowood S to durability Class 3, or at 212°C Thermowood D to durability Class 2 (equivalent to European oak). The product is significantly more durable and more resistant to termite attack than untreated timber. It is less hygroscopic producing less moisture movement, has improved thermal insulation and the resins are removed from the knots. The timber is darker in colour and has a reduced strength in relation to splitting. In an alternative commercial process using temperatures between 150°C and 180°C, with aqueous, drying and heating cycles over several days, highly durable but brown softwoods and hardwoods are produced. Thermal modified timber is described in the British Standards document DD CEN/TS 15679: 2007. Where thermal modified timber is to be used for structural purposes, allowances must be made for the reduced strength of the material. Frake or limba (*Terminalia superba*), a non-durable West African timber, is currently being imported and thermally treated to produce a durable hardwood. The Beehive Centre, Cambridge (Fig. 4.16), is clad with thermally modified timber.

**Thermally modified timber classes:**

<table>
<thead>
<tr>
<th>Classes</th>
<th>Typical uses</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Softwoods</strong></td>
<td></td>
</tr>
<tr>
<td>Class D (212°C)</td>
<td>Garden structures, external cladding, exterior doors, window frames, decking, internal floors and internal decoration</td>
</tr>
<tr>
<td>Class S (190°C)</td>
<td>Construction materials, structural components, garden furniture, interior floors, exterior doors and window frames</td>
</tr>
<tr>
<td><strong>Hardwoods</strong></td>
<td></td>
</tr>
<tr>
<td>Class D (212°C)</td>
<td>Garden furniture, patio floors, interior floors and interior decoration</td>
</tr>
<tr>
<td>Class S (190°C)</td>
<td>Constructional materials, mouldings, furniture, interior floors and interior decoration</td>
</tr>
</tbody>
</table>

**Chemical modified timber**

Chemical modification of timber involves acetylation of the timber by adding acetyl groups to the free
hydroxyl groups within the cellulose macromolecules. The process involves treatment with acetic anhydride, which penetrates the full thickness of the material. The modified timber (Accoya) with a reduced number of free hydroxyl groups has significantly reduced water absorption and thus greater dimensional stability. For a timber such as radiata pine, a maximum moisture content of approximately 6% is achieved with no colour change, producing a timber durable to Class 1. The acetylation process does not affect the physical properties of the timber, but the significantly increased durability to biological deterioration makes it less susceptible to termite attack.

**TIMBER FRAME CONSTRUCTION**

The two standard forms of timber frame construction are balloon frame and platform frame. The latter modern method of construction (MMC) is more common within the UK. For two-storey housing the balloon frame is constructed with double-storey height panels, with the intermediate floor supported from the framework. In platform frame, the panels are single-storey height, which are easier to manoeuvre, and the intermediate floor is supported directly on the lower-storey panels. Panels are normally constructed of 100 × 50 mm softwood, sheathed in plywood or particleboard, and filled with insulation between the studs and with an internal finish of plasterboard. The subsequent external weatherproofing may be of brickwork or blockwork and rendering as required.

**GRIDSHELL CONSTRUCTION**

The Weald and Downland Open Air Museum in Sussex (Fig. 4.17) illustrates a new approach to timber construction combining modern computing technologies with traditional craftsmanship. The large barn-like construction (50 m long × 12 m wide × 10 m high) is formed from a double-skin grid of 35 × 50 mm × 40 m long green oak laths at 1 m centres, which generate an undulating envelope of curved walls blending into three domes. Continuous curvature of the walls and roof are necessary for structural integrity.

Freshly sawn green oak was used as it is supple and easily formed. Initially, it was bolted into a flat grid with stainless steel bolts. The supporting scaffolding was then gradually removed, allowing the construction to settle into its design form which was finally fixed around the perimeter. Once the correct form is established, the geometry is locked to ensure stability against wind and snow loading. The construction requires no interior supports, which would have inhibited the free use of the internal space. The structure was glazed with
polycarbonate clerestory panels and clad in western red cedar vertical boarding. Gridshell construction has been used previously in Germany and Japan, but the Weald and Downland Museum by Edward Cullinan Architects is the first of its type within the UK. A more recent gridshell building is the Savill Building at Windsor Great Park by Glenn Howells Architects (Fig. 4.18), constructed from locally sourced larch and green oak cladding.

**TIMBER PILES AND FOUNDATIONS**

Timber piled foundations have been used for many centuries and have a good record of durability. The city of Venice is largely built on timber piles and their use as an alternative construction system is current within North America for the foundations of bridges and other significant structures. The use of timber pile foundations, in appropriate ground conditions, offers an economical alternative to concrete, with the environmental advantage of creating carbon dumps to reduce global warming.

Historically, a range of softwoods and hardwoods has been used for timber piles, but in the UK Douglas fir, up to 500 mm square and 12–15 m long, is a standard material (BS 8004: 1986). Other suitable timbers are treated: Scots pine or larch, oak in non-salt water soils, elm, beech and sycamore. Untreated timber below the water table is virtually immune to decay, but it is at risk from biological degradation above this level. It is therefore appropriate to treat timber with preservatives if it is to be used above the water table. Preservative-treated timber piles, cut off below ground level and capped with concrete, should have a service life of 100 years.

**ROUND TIMBER**

Forest thinnings, which are too small to be converted into rectangular standard sections for construction, have the potential to be used directly for certain low-technology forms of building. Currently, much of this material is used for paper and particleboard production or burnt as firewood.

The advantage of this material is that it is a renewable resource with a relatively short production cycle and rapid carbon dioxide sequestration. Timber poles are naturally tapered, but the effects on mechanical properties of defects such as knots and sloping grain, which are significant in converted timber, are virtually eliminated. Also, as little machining is required energy and labour costs are low.
Forest thinnings up to 200 mm in diameter are generally available, but typical small-scale construction usually requires poles in the 50–150 mm range, with lengths between 3 and 15 m. Figure 4.19 illustrates an experimental building at Hooke Park using Norway spruce timber pole construction.

**Dimensional classes for round timber**

The draft standard pr EN 1315-1: 2008 describes nine dimensional classes (D0–D8) for both hardwood and softwood round timber, measured at the mid-diameter and ranging from <10 to ≥80 cm. The current standards BS EN 1315 Part 1: 1997 and Part 2: 1997 are specific to softwood and hardwood round timber, respectively.

**Qualitative classification of round timber**

Round timbers may be classified within four quality classes which should be listed with their species. Classification depends on the extent of natural defects and any deterioration by rot or insect attack. The criteria are listed BS EN 1927 Parts 1–3: 2008 for spruces and...
The qualitative classifications for oak and beech, poplar and ash, maple and sycamore are in BS EN 1316 Parts 1 and 2: 1997 and Part 3: 1998, respectively. For example, QA is the highest grade of oak (*Quercus*) and FrB is the normal quality of Ash (*Fraxinus*).

**Quality class A**  First quality timber with only minor defects.

**Quality class B**  Average to first quality with knots typical of the species.

**Quality class C**  Average to low quality but without serious loss of the characteristics of the wood.

**Quality class D**  Timber which may be sawn into usable wood.

**SOFTWOOD CLADDING**

Western red cedar has long been the preferred timber for external timber cladding because of its durability and warm colour. However, recently the popularity of larch and Douglas fir as softwood cladding has increased as greater emphasis is placed on the use of renewable resources from sustainable forests. Large quantities of these materials, which are classified as moderately durable to decay, will become available as plantation-grown stocks reach maturity. Both larch and Douglas fir are more resistant to impact damage than western red cedar, and for cladding purposes they should not need additional preservative treatment. The timbers have been used successfully for school and health centre buildings as both vertical and horizontal cladding. As the timbers are acidic, all fixings must be in corrosion-resistant material; also, some resin bleed can be expected which will penetrate any applied surfaces finishes.

**HARDWOOD FLOORING**

Hardwood flooring has a proven track record for durability and aesthetic impact. Both solid timber and plywood laminates with a 4 mm hardwood wearing layer are commercially available. The standard timbers are the European oak, beech, birch, ash, chestnut, walnut and maple, but additionally some imported hardwoods with darker grain colours are available and interesting effects are produced with bamboo. The timbers are frequently offered with minimal knots and uniform graining or as *rustic* with knots and a larger variation of colour. Laminites are usually prefinished but solid timber may be sealed with oil or lacquer after installation on site.

**JOINERY TIMBER**

The term joinery applies to the assembly of worked timber and timber panel products, using timber which has been planed to a smooth finish. By contrast, carpentry refers to the assembly of the structural carcase of a building usually with rough sawn timbers. Joinery work, including the production of windows, doors, staircases, fitted furniture, panelling and mouldings, requires timber that is dimensionally stable, appropriately durable with acceptable gluing properties, and can be machined well to a good finish. Joinery grade timber is categorised into seven quality classes (*Table 4.8*) according to the number and size of natural defects, particularly knots. These classes are sub-divided into two surface categories, visible and concealed, according to whether the timber is to be visible in use. (BRE Digest 407: 1995 and BS EN 942: 2007 list some softwoods and hardwoods suitable for joinery.) Softwood flooring, cladding and profiled boards should not be specified as joinery.

**DETERIORATION OF TIMBER**

The major agencies causing the deterioration of timber in construction are weathering, fungi, insects and fire. The natural durability of timber is defined into five categories in relation to the resistance of the heartwood to wood-decaying fungi (BS EN 350–1: 1994).

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**Table 4.8 Classes of timber for joinery use and maximum knot sizes (BS EN 942: 2007)**

<table>
<thead>
<tr>
<th>Class</th>
<th>Visible faces</th>
<th>Concealed faces</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>J2</td>
<td>J5</td>
</tr>
<tr>
<td>Maximum knot size (mm)</td>
<td>2</td>
<td>5</td>
</tr>
<tr>
<td>Maximum percentage of width of finished piece (%)</td>
<td>10</td>
<td>20</td>
</tr>
</tbody>
</table>

Note:
The standard also refers to spiral and slope of grain, fissures, shakes, resin pockets, bark, discoloured sapwood, exposed pith and Ambrosia beetle damage.
Natural durability of timber:
Class 1 Very durable
Class 2 Durable
Class 3 Moderately durable
Class 4 Slightly durable
Class 5 Not durable (perishable)

Weathering

On prolonged exposure to sunlight, wind and rain, external timbers gradually lose their natural colours and turn grey. Sunlight and oxygen break down some of the cellulose and lignin into water-soluble materials which are then leached out of the surface leaving it grey and denatured. Moisture movements, associated with repeated wetting and drying cycles, raise the surface grain, open up surface checks and cracks and increase the risk of subsequent fungal decay. Provided the weathering is superficial, the original appearance of the timber can be recovered by removing the denatured surface.

Fungal attack

Fungi are simple plants, which unlike green plants, cannot synthesise chlorophyll, and therefore must obtain their nutrients by metabolising organic material, breaking it down into soluble forms for absorption into their own system. For growth they need oxygen and a supply of food and water, a minimum moisture content of 20% being necessary for growth in timber. The optimum temperature for growth is different for the various species of fungi, but usually within the range 20–30°C. Little growth takes place below 5°C and fungi will be killed by prolonged heating to 40°C. Some timbers, particularly the heartwoods of certain hardwoods, are resistant to attack because their minor constituents or extractives are poisonous to fungi.

All fungi have a similar life cycle (Fig. 4.20), commencing with the microscopic spores which are always present in quantity in the air. Under favourable conditions, spores within the surface cracks of timber will germinate and produce fine filaments or hyphae, which feed on the cellulose of the timber. The hyphae branch and grow through the timber cells feeding on both the walls and their contents. With increasing colonisation of the timber, the fine hyphae combine to produce a white matrix or mycelium, which is then visible to the eye. After a period of growth, the mycelium at the surface produces fruiting bodies which generate many thousands of spores to continue the life cycle. The spores, which are less than 10 μm in size, are readily distributed by air movement.

Fig. 4.20  Fungus life cycle
**Moulds and stains**

Moulds and stains are fungi that metabolise only the starch and sugar food reserves stored within the timber cells; therefore, sapwoods are generally more vulnerable than heartwoods, since during the conversion of sapwood to heartwood the stored food is removed. Generally, there is little loss of strength associated by such an attack, although one variety, **blue-stain**, aesthetically degrades large quantities of timber and its presence may indicate conditions for incipient wood-rotting fungal attack. It is best prevented by kiln drying to quickly reduce the surface moisture content unless infection has already occurred within the forest. Generally, softwoods are more susceptible to attack than hardwoods. However, the light-coloured hardwoods ramin, obeche and jelutong are sometimes affected.

**Wet and dry rots**

The name dry rot, attributed to one variety of fungus, is a misnomer, as all fungal growths require damp conditions before they become active. Destructive fungi can be categorised as soft, brown, or white rots.

Soft rots, which belong to a group of micro-fungi, are restricted to very wet conditions such as timbers buried in the ground and are therefore not experienced within normal construction. They are usually found only within the timber surface, which becomes softened when wet and powdery when dry.

The brown rots preferentially consume the cellulose within timber, leaving more of the lignin, tannin and other coloured extractives; thus, the timber becomes progressively darker. In contrast, the white rots consume all the constituents of the cells, so the timber becomes lighter in colour as the attack proceeds.

A major cause of deterioration of timber within buildings is **Serpula lacrymans**, the so-called dry rot. Under damp conditions, above 20% moisture content, the mycelium forms cotton-wool like masses over the surface of the timber, which becomes wet and slimy. The mycelium strands, up to 20 mm in diameter, can grow through brickwork and past inert materials to infect otherwise dry timber. Under drier conditions the mycelium forms a grey-white layer over the timber, with patches of bright yellow and occasionally lilac. The fruiting bodies, or fructifications, are plate-like forms, which disperse the rust-red spores. In some circumstances the fruit bodies may be the first signs of attack by dry rot. After an attack by dry rot the timber breaks up both along and across the grain into cube-shaped pieces, becoming dry and friable, hence the name dry rot.

Wet rot or cellar fungus (**Coniophora puteana**) is the most common cause of timber decay within buildings in the UK. It requires a higher moisture content than dry rot (40–50%), and is therefore frequently associated with water ingress due to leaks or condensation. The decayed timber is darkened and tends to crack mainly along the grain. The thin individual strands or hyphae are brown or black, and the fruit bodies, rarely seen, are olive green in colour. Frequently, the decay is internal without significantly affecting the exposed faces of the timber.

**Phellinus contiguus** (**Poria contigua**) causes decay to external softwood joinery, particularly window frames, causing the timber to decompose into fibrous lengths. Another variety, **Phellinus megaloporus**, is known to attack oak timbers ultimately leaving a white mass.

**Figure 4.21** illustrates the relative vulnerability of sapwood compared to the naturally more resistant darker heartwood, which has been partially protected from rot by secondary thickening and the inclusion of extractives.

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**Fig. 4.21** Deterioration of sapwood timber illustrating the relative durability of heartwood over sapwood
Insect attack

Insect attack on timber within the UK is limited to a small number of species, and tends to be less serious than fungal attack. This is the reverse of the situation in hotter climates where termites and other insects can cause catastrophic damage, although the recent unintentional importation and subsequent establishment of subterranean termites in North Devon shows that this species may pose a future threat to UK buildings.

The main damage by insects within the UK comes from beetles, which during their larval stage bore through the timber, mainly within the sapwood, causing loss of mechanical strength. For other species, such as the pinhole borers (*Platypus cylindricus*), the adult beetle bores into the timber to introduce a fungus on which the larvae live. The elm bark beetle (*Scolytus scolytus*) was responsible for the spread of Dutch elm disease in the 1970s. The larvae tunnelled under the bark, within the bast and cambium layers, preventing growth and spreading the destructive fungus which eventually killed large numbers of trees across the UK.

The typical life cycle (Fig. 4.22) commences with eggs laid by the adult beetle in cracks or crevices of timber. The eggs develop into the larvae which tunnel through the timber leaving behind their powdery waste or frass. Depending on the species, the tunnelling process can continue for up to several years before the development of a pupa close to the surface of the timber, prior to the emergence of the fully developed adult beetle, which eats its way out leaving the characteristic flight hole. The insects which attack well-seasoned timber within the UK are the common furniture beetle, death watch beetle, house longhorn beetle and powder post beetle. Wood-boring weevils only attack timber that has been previously affected by fungal decay (Fig. 4.23).

**Common furniture beetle**

The common furniture beetle (*Anobium punctatum*) attacks mainly the sapwood of both hardwoods and softwoods. It can be responsible for structural damage in the cases of severe attack, and is thought to be present in up to 20% of all buildings within the UK. The brown beetle is 3–5 mm long and leaves flight holes of approximately 2 mm in diameter. Both waterborne and organic-solvent insecticides offer effective treatments.

**Death watch beetle**

The death watch beetle (*Xestobium rufovillosum*) characteristically attacks old hardwoods, particularly oak, and is therefore responsible for considerable damage to historic buildings. Attack is normally on the sapwood, but heartwood softened by moisture and fungal decay will attract infestation; adjacent softwood may also be affected. The brown beetle is approximately 8 mm long and leaves a flight hole of 3 mm diameter. Remedial measures should include the eradication of damp and the application of organic-solvent insecticides.

**House longhorn beetle**

The house longhorn beetle (*Hylotrupes bajulus*) is a serious pest in some parts of southern England, particularly in Surrey, and it is referenced in the Building Regulations 2000 — Approved Document to support regulation 7: 1999 (amended 2000). House longhorn beetle can infest and cause serious structural damage to the sapwood of seasoned softwood roof timbers. With an average life cycle of six years and a larva that is up to 35 mm long, this beetle can cause serious damage before evidence of the infestation is observed. The affected timbers bulge where tunnelling occurs just below the surface, and the eventual flight holes of the black beetle are oval and up to 10 mm across. Where sufficient serviceable timber remains, remedial treatment with organic-solvent or paste formulations is appropriate.

**Powder post beetle**

The powder post beetle (*Lyctus brunneus*) attacks the sapwood of certain hardwoods, particularly oak and...
Wood-boring beetles common within the UK

**Common furniture beetle (Anobium punctatum)**
Actual size approx. 3 to 5 mm long
Flight holes
Bore dust

**Death watch beetle (Xestobium rufovillosum)**
Actual size approx. 6 mm long
Typical damage
Bore dust

**House longhorn beetle (Hylotrupes bajulus)**
Actual size approx. 25 mm long
Typical damage to rafters
Bore dust

**Powder post beetle (Lyctus brunneus)**
Actual size approx. 5 to 6 mm long
Severe internal damage and apparently superficial external damage
Bore dust

**Wood-boring weevil (Pentarthrum huttoni)**
Actual size approx. 3 to 5 mm long
Typical internal damage
Bore dust
ash. The sapwood of large-pored tropical hardwoods, such as ramin and obeche, may also be affected. Only timbers with sufficient starch content within the sapwood are vulnerable to attack as the larvae feed on starch rather than on the cell walls. The eggs are laid by the adult female beetle into the vessels which are the characteristically large cells within hardwoods. Timbers with low starch content or fine vessels are immune, and the extended soaking of vulnerable timbers in water can reduce the risk of attack, but owing to the long time scale involved, this is not commercially viable. The 4 mm reddish brown beetle leaves a flight hole of approximately 1.5 mm diameter. Timbers are attacked only until all the sapwood is consumed, so in older buildings damage is usually extinct. In new buildings, coatings of paint or varnish make treatment impractical, so replacement is the usual option.

**Ambrosia beetle**

A large number of ambrosia beetle species attack freshly felled hardwood and softwood logs, in both temperate and tropical regions. A high moisture content of over 35% is necessary for this beetle attack, which is therefore eliminated on seasoning. The circular pinholes range from 0.5 to 3 mm depending on the particular beetle species and the tunnelling is across the grain of the timber.

**Wood-boring weevils**

Wood-boring weevils attack only timber previously softened by fungal decay. The most common weevil (*Pentarthrum huttoni*) produces damage similar in appearance to the common furniture beetle, but removal of decayed timber will eliminate the secondary infestation.

**Termites**

Termites are social insects, similar in size to ants (4–5 mm), which live in colonies containing millions of individuals. Most species of termites are beneficial to nature in breaking down organic matter, but a few varieties can cause catastrophic damage to buildings. The subterranean termite family *Rhinotermitidae* is the major cause of building damage, with the genus *Reticulitermes* being a significant threat in the UK. The species *R. santonensis* is already widely distributed in Europe and the species *R. lucifugus* was established at Saunton, North Devon. However, treatment with hexaflumuron, an insect growth regulator, within bait, appears to have eliminated the infestation, as no further evidence has been observed since 2000. The primary source of food for termites is the cellulose in wood, particularly from structural softwood timbers near to ground level or in partially decayed timber. Timber may be only slightly affected or it may be heavily excavated leaving only the surface and any protective coatings. In areas at risk of termites, termite protection may be appropriate, but where attack has already occurred, specialist advice from the Forestry Commission is essential. Colony elimination by physical, chemical or biological techniques can be a slow process taking months or even years to complete.

**Preservation of timber**

Wood preservatives contain pesticides in the form of insecticides and fungicides. Their use is therefore strictly controlled to limit unnecessary or accidental environmental damage. Preservative treatments should involve only materials in current approval by the Control of Pesticides Regulations (1986 as amended 1997) and they should be used in accordance with the Control of Substances Hazardous to Health (COSHH) regulations, the manufacturers’ instructions and by operatives wearing appropriate protective clothing. Timber treatments may be divided into the application of preservatives to new timber, and remedial treatments used to eradicate or reduce an existing problem.

**Preservative treatments for new timber**

A wide range of effective timber preservatives is commercially available for use under controlled industrial conditions. However, within the current climate of increasing health and environmental awareness, the drive towards more ecologically friendly products has led to considerable changes within industrial timber preservation processes. The use of the two traditional principal timber preservatives for external use, creosote and chromated copper arsenate (CCA), is now restricted by EU legislation. Creosote (BS 144: 1997) is only available for industrial use and may be used only for piles and bridges. Similarly for CCA, current legislation now prohibits the use of arsenic- and chromium-containing products. New products are predominantly organic biocides.

The industrial timber preservation processes involve the use of vacuum and either high- or low-pressure impregnation, alternatively dip diffusion. Chemicals are either water-borne, solvent based or micro-emulsions.

The double-vacuum process, using organic solvent-borne preservatives, is suitable for low- and
medium-risk timber such as external joinery. The timber, at less than 28% moisture content, is loaded into a low-pressure vessel which is evacuated to extract the air from within the timber. The vessel is flooded with preservative and a low positive pressure applied for several minutes to one hour depending on the permeability of the timber. The vessel is then drained and evacuated to remove excess preservative from the timber surface. Formulations consist of either fungicides or insecticides or both dissolved in volatile organic solvents. The solvents penetrate well into the timber but have a strong odour and are highly flammable. A water repellent may also be incorporated into the preservative formulation. Organic-solvent preservatives will eventually be limited in use to timbers where it is critical that the dimensions are not affected by the preservative treatment. To reduce the environmental effects of volatile organic compound (VOC) emissions, organic solvents are being replaced by micro-emulsions of biodegradable biocide formulations, with significantly reduced organic-solvent content. The treated timbers generally meet the requirements of Use Classes 1, 2 and 3.1 to BS EN 335-1: 2006.

The pressure/vacuum process is similar to the double-vacuum process, but uses waterborne preservatives and the application of low or high pressure within a pressure vessel to ensure the appropriate level of penetration. The standard preservatives are based on copper compounds and azole biocides. Some treatments also include boron derivatives. Timber treated with proprietary products such as Tanalith is coloured slightly green but can be directly painted. Low-pressure treatments are generally to Use Classes 1, 2 and 3.1. High-pressure treatment can give additional protection to Use Class 4.

Dip-diffusion treatments involve the immersion or spraying of the freshly sawn green timber using boron derivatives (disodium octaborate tetrahydrate). Two or three immersions are used to ensure complete coverage of all faces of the timber, and larger sections require a second treatment by spraying or immersion. After treatment the timber should be then stored for an appropriate period to allow the diffusion of the preservative into the timber to the required depth of penetration. However, the product is water soluble and therefore not suitable for pressure impregnation of timber for external structures.

Timbers to be built in high-risk situations, such as industrial roofs, frames and floors, timbers embedded in masonry, sole plates, sarking boards, tile battens, etc., should be treated with appropriate timber preservative.

**Remedial treatment for timber**

Remedial treatments to existing buildings should be limited to those strictly necessary to deal with the fungal or insect attack. The use of combined fungicides and insecticides is not advised when the attack is by one agent only. Within the UK, much timber decay is caused by building failures. As fungal decay can only occur in damp conditions, the first remedial measure must be to restore dry conditions. This should remove the necessity for frequent chemical applications.

The orthodox approach to the eradication of fungal and wood-boring beetle attack involves the removal of severely decayed or affected timber, followed by appropriate preservative treatment to the remaining timber. For fungal attack, 300 mm of apparently sound timber should be removed beyond the last visible sign of decay, and the adjoining timbers treated with fungicide. For wood-boring beetle, unless the infestation is widespread, preservative treatment should be applied only up to 300 mm beyond the visible holes. Organic-solvent fungicides and insecticides applied by brush or spray offer some protection from further attack, but applications of pastes or rods which deliver higher quantities of the active ingredients are usually more effective. Insecticidal smoke treatments need to be repeated annually as they are only effective against emerging beetles, but they may be useful in situations where brushing or spraying is impracticable.

The environmental approach to the eradication of fungal decay relies heavily on the removal of the causes of damp. On the basis that fungal attack will cease when timber is at less than 20% moisture content, increased ventilation and the rectification of building defects should prevent further attack. Only seriously affected timbers need to be replaced, and affected masonry sterilised; however, continual monitoring is required, as dormant fungal decay will become active if the timber moisture content rises again above 20%. *Rothounds* (specially trained sniffer dogs), fibre optics and chemical detection systems offer non-destructive methods for locating active dry rot before it becomes visible.

Pesticides used professionally for remedial treatment include disodium octaborate or azoles as fungicides and fluoroxy (an insect growth regulator) as insecticide. These pesticides are currently considered acceptable for treatment in areas inhabited by bats, which are a protected species under the Wildlife and Countryside Act 1981.
Guidance on timber treatments

Under the European Standards EN 351-1: 2007 and BS EN 335-1: 2006, timber preservative treatments against wood-destroying organisms are categorised by performance standards and not to the individual chemical preservative treatments. The standards define wood preservatives according to their effectiveness in a range of environmental use conditions.

Use classes for wood and wood-based products against biological attack (BS EN 335-1: 2006):

<table>
<thead>
<tr>
<th>Use Class</th>
<th>Interior or under cover and dry.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Use Class 1</td>
<td>3.1 Exterior, above ground, protected. Occasionally wet.</td>
</tr>
<tr>
<td>Use Class 3</td>
<td>3.2 Exterior, above ground, unprotected. Frequently wet.</td>
</tr>
<tr>
<td>Use Class 4</td>
<td>4.1 Exterior in ground contact. Predominantly or permanently wet. (fresh water)</td>
</tr>
<tr>
<td>Use Class 5</td>
<td>In salt water, permanently wet.</td>
</tr>
</tbody>
</table>

Key: 0 natural durability sufficient, A preservative treatment may be advisable for certain end uses, B preservative treatment may be necessary for certain end uses, C preservative treatment is normally advised, X preservative treatment necessary.

Note: Use Classes were previously described as Hazard Classes and this terminology remains in BS EN 460: 1994 and BS EN 599-1: 1997.

The standard BS EN 335 Part 1: 2006 gives general guidance on the likely occurrence of biological agents under the various class conditions. BS EN 335 Part 2: 2006 refers specifically to the particular biological agents which attack solid wood under the various class conditions and describes a decision-making sequence for the selection of the correct solid wood appropriate to the use class. Additional guidance on preservation is given in BS EN 460: 1994 (Table 4.9) which relates the timber durability class to the use class, indicating where additional treatment may be necessary.

The level of treatment required to give the necessary performance is classified according to the depth of penetration into the timber and by retention or loading within the defined zone of the timber. The depth of penetration is defined in BS EN 351-1: 2007, by six classes (NP1–NP6) of increasing zones of preservative retention. As in not all cases full preservative treatment is appropriate, a range of service factors (A–D) define the level of safety and economic considerations appropriate to preservative treatment. These are listed in the British Standard BS 8417: 2003 (Table 4.10).

The British Standard (BS 8417: 2003) also gives guidance on the types of preservative treatment appropriate for timbers to be used in more severe situations; for example, occasionally wet roof timbers in a house longhorn beetle affected location:

<table>
<thead>
<tr>
<th>Use class</th>
<th>Durability class</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0 0 0 0 0</td>
</tr>
<tr>
<td>2</td>
<td>0 0 0 A A</td>
</tr>
<tr>
<td>3</td>
<td>0 0 A B B</td>
</tr>
<tr>
<td>4</td>
<td>0 A C X X</td>
</tr>
<tr>
<td>5</td>
<td>0 C C X X</td>
</tr>
</tbody>
</table>

Key: 0 natural durability sufficient, A preservative treatment may be advisable for certain end uses, B preservative treatment may be necessary for certain end uses, C preservative treatment is normally advised, X preservative treatment necessary.

Table 4.9  Relationship between environmental use class and timber durability class and requirement for preservative treatment against wood-destroying fungi (BS EN 460: 1994)

The British Standard (BS 8417: 2003) also gives guidance on the types of preservative treatment appropriate for timbers to be used in more severe situations; for example, occasionally wet roof timbers in a house longhorn beetle affected location:

Building component (e.g. dry roof timbers in a non-Hylotrupes area)

Use class of wood against biological attack (e.g. 1 for dry roof timbers)

Service factor - safety and economic considerations (e.g. B for dry roof timbers)

Desired Service life - 15, 30 or 60 years (e.g. 60 years for dry roof timbers)

Durability class 5 - non-durable timber is appropriate.

Building component - roof timbers (with risk of wetting in an Hylotrupes area)

Timber species - e.g. European whitewood

Treatment - required for insect hazard and desirable for fungal hazard

Service life required - e.g. 60 years

Preservative type - organic solvent containing active fungicide and insecticide
Table 4.10  Service factors for preservative treatment of timber to BS 8417: 2003

<table>
<thead>
<tr>
<th>Service factors</th>
<th>Need for treatment</th>
<th>Safety and economic considerations</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Unnecessary</td>
<td>Negligible</td>
</tr>
<tr>
<td>B</td>
<td>Optional</td>
<td>Remedial action is easy</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Preservation is an insurance</td>
</tr>
<tr>
<td></td>
<td></td>
<td>against cost of repairs</td>
</tr>
<tr>
<td>C</td>
<td>Desirable</td>
<td>Remedial action is expensive</td>
</tr>
<tr>
<td></td>
<td></td>
<td>and difficult</td>
</tr>
<tr>
<td>D</td>
<td>Essential</td>
<td>Consequences of structural</td>
</tr>
<tr>
<td></td>
<td></td>
<td>collapse would be serious</td>
</tr>
</tbody>
</table>

(The British Standard (BS 8417: 2003) gives guidance on the required depth of penetration and retention of the preservative.)

Fire

Timber is an organic material and therefore combustible. As timber is heated it initially evolves any absorbed water as vapour. By about 230–250°C, decomposition takes place with the production of charcoal and combustible gases such as carbon monoxide and methane are evolved, which cause the flaming. Finally, the charcoal smoulders to carbon dioxide and ash.

However, despite its combustibility, timber particularly in larger sections performs better in a fire than the equivalent sections of exposed steel or aluminium. Timber has a low thermal conductivity, which, combined with the protection afforded by the charred surface material, insulates the interior from rapid rises in temperature and loss of strength. The rate of charring of timber under the standard Fire Resistance Test ranges between 30 and 50 mm per hour per surface exposed, according to the timber density (Table 4.11). It is therefore possible to predict the fire resistance of any timber component using the British Standards (BS 5268-4.1: 1978 and 5268-4.2: 1990). Additionally, as all timbers have a low coefficient of expansion, timber beams will not push over masonry walls as sometimes occurs with steel beams and trusses during fires. Solid timber with a minimum density of 350 kg/m³ and a thickness equal to or greater than 10 mm may be assigned to Euroclass D with respect to fire without testing.

Flame retardants

Within a fire, volatile combustible components are evolved from the surface of the timber and these cause the flaming. The two alternative types of treatment which may be used on timber to reduce this effect are impregnation or the application of surface coatings. Treatments need to be appropriate to the dry, humid or external conditions to which the timber will be exposed.

Impregnation involves forcing into the timber under high pressure and then vacuum, inorganic chemicals which on heating evolve non-combustible gases. Timbers should be machined to their final dimensions before treatment. For interior use, typical compositions include water-borne inorganic salts, such as ammonium sulphate or phosphate with sodium borate or zinc chloride. As these materials are hygroscopic, the timber should not be used in areas of high humidity. For exterior use, a leach-resistant flame-retardant material based on an organo-phosphate is used, as this is heat fixed by polymerisation within the timber.

Surface treatments, which cause the evolution of non-combustible gases in fire, include antimony trioxide flame-retardant paints which are suitable for both interior and exterior use. Intumescent coatings, which swell up and char in fire, are suitable for most environments if overcoating is applied. However, the protection afforded by surface treatments may be negated by unsuitable covering or removal by redecoration.

Untreated timber, which is normally Class 3 Spread of Flame to BS 476-7: 1997, can be improved to Class 1 by surface treatments. Class 0 can only be achieved by heavy impregnation, a combination of impregnation and surface coating or certain very specific surface coating treatments.

European fire classification of construction materials

The European fire classification of construction products and building elements is defined in BS EN 13501-1: 2007. All construction products except floorings may be classified to one of the following seven
classes: A1, A2, B, C, D, E or F based on performance. Class A1 represents products which do not contribute to the fire load even within a fully developed fire. Class A2 products do not significantly contribute to the fire load and growth, while the other classes reflect decreasingly stringent fire performance criteria down to Class E products which can resist a small flame for a short period of time without substantial flame spread. Class F products are outside the other classes or have no determined performance. In addition to fire load factors, the standard includes classifications relating to smoke production (s1, s2 and s3, where s1 is the most stringent criterion) and flaming droplets (d0, d1 and d2, where d0 indicates no flaming droplets or particles). For floorings the seven classes are A1FL, A2FL, BFL, CFL, DFL, EFL and FFL with sub-classifications for smoke production (s1, s2 and s3). Correlation between the UK and European classes is not exact and materials require testing to the European Standard before assignment to a particular class to BS EN 13501-1: 2007.

The Euroclass fire performance rating under the conditions specified in BS EN 13986: 2004 for 12 mm untreated solid wood panels of minimum density 400 kg/m³ is Class D-s2, d0 for non-floor use and Class DFL-s1 for floorings. The Euroclass fire performance of structural timber with a minimum density of 350 kg/m³ and a minimum thickness of 22 mm is Class D-s2, d0 to BS EN 14081-1: 2005.

The durability of fire retardant treatments in relation to their interior or exterior end use applications is described in pr EN 15912: 2009.

Durability of reaction to fire performance (DRF):

<table>
<thead>
<tr>
<th>DRF Class</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>ST</td>
<td>Short term use (e.g. less than one year)</td>
</tr>
<tr>
<td>INT1</td>
<td>Permanent interior use (service class 1)</td>
</tr>
<tr>
<td>INT2</td>
<td>Permanent interior use (service class 2)</td>
</tr>
<tr>
<td>EXT</td>
<td>Permanent exterior use (service class 3)</td>
</tr>
</tbody>
</table>

Note:
Service classes to BS EN 1995-1-1: 2004

TIMBER CONNECTORS

A range of steel timber connectors is commercially available. Trussed rafters, which account for a large market, are usually constructed with galvanised-steel nail plates (Fig. 4.24). Plates are hydraulically pressed into both sides of the timbers to be connected at the butt joints. Other types of connector include single- and double-sided circular toothed plates fixed with a central bolt, also joist hangers and roof truss clips.

Metal web floor joists (Fig. 4.25) are manufactured from strength graded timber flanges with steel web units nail-plated to sides of the softwood flanges. They have the advantage over solid timber joists of lightness and the open web void spaces for services.
Timber products

A wide range of products is manufactured from wood material, ranging in size from small timber sections and thin laminates through chips and shavings down to wood fibres. The physical properties of the materials produced reflect a combination of the sub-division of the wood, the addition of any bonding material and the manufacturing process. The physical properties then determine the products’ appropriate uses within the building industry. Many of the products are manufactured from small timber sections or timber by-products from the conversion of solid timber which otherwise would be wasted. Compressed straw slabs and thatch are additionally included in this section.

The product range includes:
- laminated timber;
- cross-laminated timber;
- structural insulated panels;
- laminated veneer lumber;
- plywood;
- blockboard and laminboard;
- particleboard;
- fibreboard;
- wood wool slabs;
- compressed straw slabs;
- thatch;
- shingles;
- ‘Steko’ blocks;
- flexible veneers.

Within the European Union, whenever wood-based panels are used in construction, compliance with the Construction Products Directive must be demonstrated. This may be achieved by adherence to European Harmonised Standard for wood-based panels EN 13986 (BS EN 13986: 2004 in the UK). This requires that products used in construction comply with its specifications and also to the additional performance-based criteria within the various EN standards listed for each specific material. Most European countries now use the € mark on boards and panels to show compliance with this harmonised standard.

### LAMINATED TIMBER

**Manufacture**

Large solid timber sections are limited by the availability of appropriate lumber; in addition, their calculated strength must be based on the weakest part of the variable material. Laminated-timber sections overcome both of these difficulties and offer additional opportunities to the designer. Laminated timber is manufactured by curing within a jig, layers of accurately cut smaller timber sections which are continuously glued together with a resin adhesive. Laminates may be vertically or horizontally orientated. The use of strength-graded timber and the staggering of individual scarf or finger joints ensures uniformity of strength within the product; although, under BS 5268-2: 2002 and BS EN 387: 2001, large finger joints through the whole section of a glulam member are permissible. The manufacturing process ensures greater dimensional stability and fewer visual defects than in comparable solid timber sections. Laminated timber may be homogeneous, with all laminates of the same strength class of timber, or combined, in which lower strength class laminates are used for the centre of the units. Table 4.12 gives the European strength classes to BS EN 1194: 1999 for the two alternatives. Laminated timber manufactured from spruce or pine and phenolic or urea/melamine formaldehyde resins would normally achieve a Euroclass fire performance rating, under the conditions specified in BS EN 13986: 2004 of Class D, subject to testing.

**Forms**

Sections can be manufactured up to any transportable size, typically 30 m, although spans over 50 m are possible. Standard-size straight sections (315 × 65 and 90 mm; 405 × 90 and 115 mm; 495 × 115 mm) are stock items, but common sizes range from 180 × 65 to 1035 × 215 mm. Sections may be manufactured to order, to any uniform or non-uniform linear or curved form. The majority of laminated timber structures are manufactured from softwoods such as European redwood or whitewood, although the rib members within the roof structure of the Thames Flood Barrier

<table>
<thead>
<tr>
<th>Table 4.12</th>
<th>Strength classes to BS EN 1194: 1999 for homogeneous and combined glulam</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glulam strength classes</td>
<td>GL24</td>
</tr>
<tr>
<td>Homogeneous glulam:</td>
<td></td>
</tr>
<tr>
<td>Strength class of laminates</td>
<td>C24</td>
</tr>
<tr>
<td>Combined glulam:</td>
<td></td>
</tr>
<tr>
<td>Strength class of outer laminates</td>
<td>C24</td>
</tr>
<tr>
<td>Strength class of inner laminates</td>
<td>C18</td>
</tr>
</tbody>
</table>
(Fig. 4.26) were manufactured from the West African hardwood, iroko. Figure 4.27 illustrates typical laminated timber arches, columns and portal frames as generators of structural forms. The aesthetic properties of laminated timber can be enhanced by the application of suitable interior or exterior timber finishes. Steel fixing devices and joints may be visually expressed (Fig. 4.27) or almost unseen (Fig. 4.28) by the use of concealed bolted steel plates. Laminated timber performs predictably under fire conditions with a charring rate of 40 mm per hour as defined within BS 5268:4-1: 1978. Preservative treatments are necessary when the material is to be used under conditions in which the moisture content is likely to exceed 20%. The three service classes of glulam structures relate the environmental conditions.

Service classes for glulam:

<table>
<thead>
<tr>
<th>Service Class</th>
<th>Condition Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Service Class 1</td>
<td>Internal conditions with heating and protection from damp. (typical moisture content &lt;12%)</td>
</tr>
<tr>
<td>Service Class 2</td>
<td>Protected, but unheated conditions. (typical moisture content &lt;20%)</td>
</tr>
<tr>
<td>Service Class 3</td>
<td>Exposed to the weather. (typical moisture content &gt;20%)</td>
</tr>
</tbody>
</table>

**CROSS-LAMINATED TIMBER**

Cross-laminated timber (CL) is similar to conventional plywood, except that the laminates are thicker, and the panel thicknesses are generally between 50 and 300 mm, although 500 mm can be produced. The maximum panel size is governed by transportation limits which are normally $13.5 \times 3$ m, but the production limit is $20 \times 4.8$ m. The panels manufactured from spruce, larch or pine are classified by surface quality which is planed or sanded.

**Surface quality classification of cross-laminated timber:**

- Class C Standard grade (non-visible)
- Class AB Interior grade (residential visible)
- Class BC Interior grade (industrial visible)

The structural panels, which may be used to form walls, roofs and floors, are easily clad with brickwork, tiling or rendering as appropriate.

The thermal properties of CL panels depend on the timber used. Density is usually within the range 470–590 kg/m³, and an average of 500 kg/m³ has a thermal conductivity $\lambda$ value 0.13 W/m K. To achieve a target $U$-value of 0.35 W/m² K a 100 mm CL panel, with internally a 25 mm service gap and 12 mm...
plasterboard, and externally standard brickwork cladding and a 50 mm cavity, would require 75 mm of mineral wool insulation or the thermal equivalent. A well-ventilated cavity and breathable insulation is required.

**STRUCTURAL INSULATED PANELS**

Structural insulated panels (SIPs) are prefabricated lightweight building components, used for load-bearing internal and external walls and roofs. Unlike cladding sandwich panels, structural insulated panels can support considerable vertical and horizontal loads without internal studding. They are manufactured from two high-density face layers separated by a lightweight insulating core. The three layers are strongly bonded together to ensure that the composite acts as a single structural unit. The outer layers of oriented strand board (OSB), cement-bonded particleboard or gypsum-based products are typically
8–15 mm thick. The core is composed of a rigid cellular foam, such as polyurethane (PUR), polyisocyanurate (PIR), phenolic foam (PF), expanded (EPS) or extruded (XPS) polystyrene giving an overall unit thickness between 70 and 250 mm. The structural performance is predominantly influenced by the thickness and physical properties of the outer layers, and the thermal performance is largely determined by the width and insulating characteristics of the core material. Large panels (e.g. 5.9 m × 2.4 m × 162 mm, $U$-value 0.15 W/m² K) may be used for two-storey construction to Code for Sustainable Homes Level 3.

Structural insulated panels (typically 1.2 m wide by 2.4 m high) offer a thermally efficient and air-tight form of construction, which is rapidly erected on site. Joining between panels is usually some form of tongue and groove system. Sound reduction for separating walls is typically 58 dB depending on construction details. External cladding may be brickwork, wooden panelling or rendering with plasterboard as the standard internal finish. Table 4.13 gives typical thermal performance data for structural insulated panels. The use of SIPs is one of the Modern Methods of Construction (MMC) promoted by the UK Government to reduce energy waste in new building. Currently, SIPs account for approximately 10% of new building methods construction.

### LAMINATED VENEER LUMBER

Laminated veneer lumber (LVL) (Fig. 4.29), also known as microlam, is more economical than laminated timber as there is little waste in the production process. It is manufactured to three grades by laminating timber strands with polyurethane resin under heat and pressure. In one process, logs are cut into flat timber strands 300 mm long; these are then treated with resin, aligned and hot-pressed into billets of reconstituted wood. In the other processes, 3 mm thick timber strands or sheets of veneer are coated with waterproof adhesive and bundled together with the grain parallel. The strands or veneers are pressed together and microwave cured to produce structural timber billets or sheets up to 26 m long. The versatile material is suitable for use in columns, beams, purlins and trusses and can be machined as solid timber (Fig. 4.30). I-section joists, with LVL flanges and OSB webs, are suitable for flat and pitched roofs and floor construction. Metal-web timber joists combine LVL flanges with metal strutting webs. Untreated LVL has a Class 3 surface spread of flame classification (BS 476-7: 1997). Three grades of laminated veneer lumber are classified by BS

---

**Table 4.13** Thermal performance data for structural insulated panels (SIPs)

<table>
<thead>
<tr>
<th>Core material</th>
<th>Face material</th>
<th>Panel thickness (mm)</th>
<th>Thermal performance (W/m² K)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Polyurethane foam</td>
<td>cement bonded</td>
<td>86</td>
<td>0.28</td>
</tr>
<tr>
<td>Polyurethane foam</td>
<td>particleboard</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Polyisocyanurate foam</td>
<td>oriented strand</td>
<td>100</td>
<td>0.23</td>
</tr>
<tr>
<td></td>
<td>board</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Polystyrene foam</td>
<td>oriented strand</td>
<td>140</td>
<td>0.22</td>
</tr>
<tr>
<td></td>
<td>board</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Notes:
The thermal performance data (U-values) are typical for the listed SIPs when constructed with a brick outer leaf and 50 mm clear vented cavity.
EN 14279: 2004 according to their serviceability in dry and exposed conditions.

Grades of laminated veneer lumber:

<table>
<thead>
<tr>
<th>Purpose/loading</th>
<th>Environmental conditions</th>
<th>Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>Load-bearing</td>
<td>Dry (service class 1)</td>
<td>LVL/1</td>
</tr>
<tr>
<td>Load-bearing</td>
<td>Humid (service class 2)</td>
<td>LVL/2</td>
</tr>
<tr>
<td>Load-bearing — exterior conditions</td>
<td>Exterior (service class 3)</td>
<td>LVL/3</td>
</tr>
</tbody>
</table>

Notes:
Service classes are according to BS EN 1995-1-1: 2004.
Laminated veneer lumber for structural applications should be marked ‘S’ for structural application and ‘G’ for general application to pr BS ISO 18776.

Monocoque structures

Interesting and innovative built forms can be created using LVL (and other timber products) to create flat or curved form monocoque structures. These work on the well-established principles from the motor industry, in which the hard body skin acts in concert with any stiffeners to form the structure. Using this technology, structurally efficient and elegant forms, which may be slender, tapered, flat or curved can be produced. LVL is quickly becoming a significant material to complement the more established products such as plywood, OSB and glulam, particularly because of its availability in very large sections.

PLYWOOD

Manufacture

Plywood is manufactured by laminating a series of thin timber layers, or plies, to the required thickness. The timber log is softened by water or steam treatment and rotated against a full-length knife to peel off a veneer or ply of constant thickness (Fig. 4.31). The ply is then cut to size, dried and coated with adhesive prior to laying up to the required number of layers. Not all the plies are of the same thickness; often, thicker plies of lower-grade material are used in the core. However, the sheets must be balanced about the centre to prevent distortions caused by differential movement. Plies are normally built up with adjacent grain directions at right angles to each other to give uniform strength and reduce overall moisture movement, although with even plywoods, the central pair of plies has parallel grains. The laminate of plies and glue is cured in a hot press, sanded and trimmed to standard dimensions for packaging. Decorative veneers of hardwood or plastic laminate may be applied to one or both faces. Most plywood imported into the UK is
made from softwood (largely pine and spruce), from North America and Scandinavia. Smaller quantities of plywood produced from temperate hardwoods are imported from Finland (birch) and Germany (beech), while tropical hardwood products are imported from Indonesia, Malaysia, South America and Africa. Bamboo plywood is made from a core of tightly compressed fibres with bamboo veneers on either face. It is coated with a lacquer but should not be used externally without preservative pre-treatment as bamboo is inherently not durable.

The standard sheet size is 2440 × 1220 mm, with some manufacturers producing sheet sizes of up to 3050 × 1525 mm or slightly larger. Sheet thicknesses range from 4 to 25 mm for normal construction use, although thinner sheets down to 1.5 mm are available for specialist purposes.

Under the European fire classification of construction materials (BS EN 13501-1: 2007), an untreated plywood panel would normally achieve a class Ds2, d0 rating, excluding its use as flooring when the rating is class DFL-s1 (depending on a minimum thickness of 9 mm, a minimum density of 400 kg/m³ and fixing to a non-combustible substrate [class A1 or A2] without an air gap. The secondary classifications ‘s’ and ‘d’ relate to smoke production and flaming droplets.)

**Grades**

Plywood is classified according to its general appearance and physical properties (BS EN 313-1: 1996). The key characteristics are the form of construction, durability, and nature of the surface. The durability of plywood is largely determined by the bonding class of the adhesive used. This ranges from Class 1, to the most durable Class 3 (BS EN 314-2: 1993), which can be used externally without delamination, provided that the timber itself is durable or suitably protected against deterioration.

**Bonding classes for plywood:**

Class 1 Dry conditions (suitable for interior use).

Class 2 Humid conditions (protected external
applications, e.g. behind cladding or under roof coverings).

Class 3 Exterior conditions (exposed to weather over sustained periods).

Phenol formaldehyde resins are most frequently used for the most durable plywoods. Marine plywood (BS 1088-1: 2003) is a combination of a moderately durable timber bonded with phenolic or melamine-formaldehyde resin. The standard class of marine plywood is suitable for regular wetting or permanent exposure to salt or fresh water. The lower grades of plywood are bonded with melamine-urea formaldehyde or urea-formaldehyde resins. The relationship between the natural durability of the timber against wood-destroying fungi and the Use Class to which the plywood can be assigned is described in DD CEN/TS 1099: 2007.

The quality of plywood is also affected by the number of plies for a particular thickness and the surface condition of the outer plies which range from near perfect, through showing repaired blemishes, to imperfect. Factory-applied treatments to improve timber durability and fire resistance are normally available.

The standard BS EN 635: 1995 describes five classes of allowable defects (E, and I to IV) according to decreasing quality of surface appearance; class E is practically without surface defects. These are related to hardwood and softwood surfaced plywoods in BS EN 635: 1995 Parts 2 and 3, respectively.

The performance specifications for plywood to be used in dry, humid or exterior conditions against the criteria of bonding strength and durability with respect to biological decay are described in the standard BS EN 636: 2003.

Biological use class conditions (formerly hazard class) for the use of plywood:

- **Use Class 1** Dry conditions (average moisture content < 10%).
- **Use Class 2** Humid conditions (average moisture content < 18%).
- **Use Class 3** Exterior conditions (average moisture content > 18%).

These biological use classes correspond to the service classes in BS EN 1995-1-1: 2004.

The standard BS EN 636: 2003 also has a classification system based on the strength and stiffness of plywood based on bending tests. Plywood is assigned to a four part code specifying bending strength and modulus in both the length and width directions. Plywood sheets should be identified according to their intended application with ‘S’ for structural and ‘G’ for general use.

### Uses

Considerable quantities of plywood are used by the construction industry because of its strength, versatility and visual properties. The strength of plywood in shear is used in the manufacture of plywood box and I-section beams in which the plywood forms the web. Increased stiffness can be generated by forming the plywood into a sinusoidal web. Plywood box beams can be manufactured to create pitched and arched roof forms as illustrated in Fig. 4.32. Stiffened and stressed skin panels, in which plywood and softwood timbers are continuously bonded to act as T or I-beams, will span greater distances as floor structures than the same depths of traditional softwood joists with nailed boarding. Such structural units can also be used to form pitched roofs or to form folded plate roof structures or barrel vaulting (Fig. 4.32). Plywood of 8–10 mm thickness is frequently used as the sheeting material in timber frame construction and for complex roof forms such as domes. The lower-grade material is extensively used as formwork for in situ concrete.

### CORE PLYWOOD

The standard core-plywood products are blockboard and laminboard. Both are manufactured with a core of usually softwood strips sandwiched between one or two plies (Fig. 4.33). In blockboard the core strips are between 7 and 30 mm wide, but in laminboard, the more expensive product, they are below 7 mm in width and continuously glued throughout. As with plywood, the grain directions are perpendicular from layer to layer. Most core plywoods are bonded with urea formaldehyde adhesives appropriate to interior applications only. The standard sheet size is 2440 × 1220 mm with a thickness range from 12 to 25 mm, although larger sheets up to 45 mm thick are available. Blockboard may be finished with a wide range of decorative wood, paper or plastic veneers for use in fitted furniture. Variants on the standard products include plywood with phenolic foam, polystyrene or a particleboard core.
PARTICLEBOARDS

Particleboards are defined as panel materials produced under pressure and heat from particles of wood, flax, hemp or other similar lignocellulosic materials. The wood particles may be in the form of flakes, chips, shavings, sawdust, wafers or strands (BS EN 309: 2005). Boards may be uniform through their thickness or of a multilayer structure. Wood particleboard and cement-bonded particleboard are made from wood chips with resin and cement binder, respectively. Oriented strand board is manufactured from large wood flakes and is classified in BS EN 300: 2006.

Wood particleboard (chipboard)

Manufacture

Wood particleboard (chipboard) is manufactured from wood waste or forest thinnings which are converted into wood chips, dried and graded according to size. The chips are coated with adhesive to approximately 8% by weight and then formed into boards (Fig. 4.34). The woods chips are either formed randomly into boards giving a uniform cross-section or distributed with the coarse material in the centre and the finer chips at the surface to produce a smoother product. The boards are then compressed and cured between the plates of a platen press at 200°C. Boards
are finally trimmed, sanded and packed. In the Mende process a continuous ribbon of 3–6 mm particleboard is produced by calendering the mix around heated rollers.

The standard sizes are 2440 × 1220 mm, 2750 × 1220 mm, 3050 × 1220 mm and 3660 × 1220 mm, with the most common thicknesses ranging from 12 to 38 mm, although much larger sheet sizes and thicknesses from 2.5 mm are available.

Extruded particleboard (BS EN 14755: 2005) is manufactured by extruding the mixture of wood chip and resin through a die into a continuous board; however, in this method the wood chips are predominantly orientated at right angles to the board face, thus generating a weaker material. Extruded particleboard is specified within four grades according to its density and whether it is solid or has tube voids.

Types

The durability of particleboards is dependent on the resin adhesive. Much UK production uses urea formaldehyde resin, although the moisture-resistant grades are manufactured with melamine–urea formaldehyde or phenol formaldehyde resins. Wood chipboards are categorised into seven colour-coded types to BS EN 312: 2003 according to the anticipated loading and environmental conditions. The standard specifies requirements for mechanical and swelling properties and also formaldehyde emissions. The first
colour code defines the loading and the second colour the moisture conditions.

**Grades of wood particleboard:**

<table>
<thead>
<tr>
<th>Purpose/loading</th>
<th>Environmental conditions</th>
<th>Colour codes</th>
<th>Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>General purpose</td>
<td>Dry</td>
<td>White, white blue</td>
<td>P1</td>
</tr>
<tr>
<td>Interior fitments</td>
<td>Dry</td>
<td>White blue</td>
<td>P2</td>
</tr>
<tr>
<td>Non-load-bearing</td>
<td>Humid</td>
<td>White green</td>
<td>P3</td>
</tr>
<tr>
<td>Load-bearing</td>
<td>Dry</td>
<td>Yellow, yellow blue</td>
<td>P4</td>
</tr>
<tr>
<td>Load-bearing</td>
<td>Humid</td>
<td>Yellow, yellow green</td>
<td>P5</td>
</tr>
<tr>
<td>Heavy duty, load-bearing</td>
<td>Dry</td>
<td>Yellow blue</td>
<td>P6</td>
</tr>
<tr>
<td>Heavy duty, load-bearing</td>
<td>Humid</td>
<td>Yellow green</td>
<td>P7</td>
</tr>
</tbody>
</table>

**Grades of extruded particleboard:**

ES Extruded Solid: Board with a minimum density of 550 kg/m³.

ET Extruded Tubes: Board with a minimum solid density of 550 kg/m³.

ESL Extruded Solid Light: Board with a density of less than 550 kg/m³.

ESL Extruded Tubes Light: Board with a solid density of less than 550 kg/m³.

(Grade ET must have at least 5 mm of material over the void spaces.)

Standard particleboards are hygroscopic and respond to changes in humidity. A 10% change in humidity will typically increase the sheet length and breadth by 0.13% and the thickness by 3.5%. Dry grades should not be exposed to moisture even during construction. Humid-tolerant grades are resistant to occasional wetting and relative humidities over 85%. However, no particleboards should be exposed to prolonged wetting, as they are all susceptible to wet rot fungal attack.

All untreated wood particleboards have Class 3 spread of flame (BS 476-7: 1997). However, they can be treated to the requirements of Class 1 by chemical addition in manufacture, impregnation or the use of intumescent paints. Class 0 may also be achieved. For untreated particleboard with a minimum density of 600 kg/m³ and a minimum thickness of 9 mm, the Euroclass fire performance rating under the conditions specified in BS EN 13986: 2004 is Class D-s2, d0 for non-floor use and Class D_{FL}-s1 for floorings.

A wide range of wood veneer, primed/painted, paper and plastic (PVC, phenolic film or frequently melamine) finishes is available as standard products. Pre-cut sizes are available edged to match. Domestic flooring grade particleboard, usually 18 or 22 mm, may be square-edged or tongued and grooved. The industrial flooring grades are typically from 38 mm upwards in thickness.

**Uses**

Significant quantities of wood particleboard (chipboard) are used in the furniture industry. Much flat-pack DIY furniture is manufactured from painted or veneered particleboard. Particleboard can be effectively jointed by use of double-threaded particleboard wood screws and various specialist fittings. Where high humidity is anticipated the moisture-resistant grades should be used. The domestic housing market uses large quantities of flooring-grade particleboard as it is competitively priced compared to traditional tongued and grooved softwood. Joist centres should be at 450 and 610 mm centres maximum for 18/19 and 22 mm particleboard, respectively. Edges should be tongued and grooved or fully supported and the standard panel size is 2400 × 600 mm. For heavy-duty flooring, flat-roof decking and structural work, the moisture-resistant structural grade must be used. Phenolic film-coated particleboard offers a suitable alternative to plywood as formwork to concrete.

**Cement-bonded particleboard**

**Manufacture**

Cement-bonded particleboard is manufactured from a mixture of wood particles or filaments (usually softwood) and cement. The boards which are light grey in colour have a uniform cementitious surface. The material has up to 75% cement by weight, with the cement filling all the void spaces, producing a material with a density of 1000–1250 kg/m³ (cf. 650–690 kg/m³ for standard grade particleboard).

**Types and uses**

The material based on Portland cement has good resistance to fire, water, fungal attack and frost. The standard (BS EN 634-2: 2007) specifies only one grade, which is suitable for use both internally and externally. It should be colour-coded white, white (non-load-bearing) and brown (suitable for dry, humid and exterior conditions) with 25 mm vertical stripes near one corner of the board. Within the one grade there are two Technical Classes 1 and 2 which relate only to modulus of elasticity in bending. The standard BS EN 633: 1994 refers to both Portland and magnesium-based cements. Magnesite-bonded particleboard is used as a lining board but it is not frost-resistant and is unsuitable for use in humid conditions.

Boards frequently have a core of coarse wood chips, sandwiched between finer material, producing a good finish, which may be further treated by sanding
and priming. Because of its density, cement-bonded particleboard has good sound-insulation properties. Typically, 18 mm board will give sound reduction of 31–33 dB. The material is frequently used for soffits, external sheathing and roofing on both modular and timber-frame buildings, particularly where racking resistance is required. The heavier grades, generally tongued and grooved, are suitable for flooring, due to their resistance to moisture, fire, impact and airborne sound.

The material has a Class 0 Surface Spread of Flame to Building Regulations (Class 1 to BS 476: Part 7: 1997). The Euroclass fire performance rating under the conditions specified in BS EN 13986: 2004 for 10 mm cement-bonded particleboard is Class B-s1, d0 for non-floor use and Class BFL-s1 for floorings.

Board sizes are typically 1200 × 2440, 2600 or 3050 mm, with standard thicknesses of 12 and 18 mm, although sheets up to 40 mm in thickness are made. However, because of the density of the material, a 1200 × 2440 × 12 mm board weighs approximately 45 kg and should not be lifted by one operative alone.

Gypsum-bonded particleboard
Gypsum-bonded particleboard, available in sheets of 6 mm thickness upwards, is an alternative multipurpose building board. It is not included in the scope of BS EN 633: 1994.

Oriented strand board

Manufacture
Oriented strand board (OSB) is manufactured from 0.5 mm thick timber flakes tangentially cut and measuring approximately 75 × 35 mm. These are dried and coated with wax and 2.5% of either phenol formaldehyde or melamine–urea formaldehyde resin. The mix is laid up in three (or occasionally five) layers with the strands running parallel to the sheet on the outer faces and across or randomly within the middle layer. The boards are then cured under heat and pressure, sanded and packaged (Fig. 4.35). Standard panel sizes are 2440 or 3660 × 1220 mm with densities usually in the range 600–680 kg/m³.

Grades and uses
Oriented strand board is graded according to the anticipated loading and environmental conditions (BS EN 300: 2006). Large quantities are used as sheathing in timber frame housing. The moisture-resistant grade is suitable for roof sarking, whilst the higher specification grade with enhanced strength properties is suitable for flat roof decking. Thicker panels are used for heavy-duty flooring, and OSB is often used as the web material in timber I-beams. Oriented strand board is manufactured to a thickness range of 6–40 mm. In Europe it is manufactured from Scots pine and spruce, but in North America from aspen and Southern pine. Within the UK and Ireland, OSB is manufactured from managed forest thinnings certified by the Forest Stewardship Council.

<table>
<thead>
<tr>
<th>Grade</th>
<th>Purpose/loading</th>
<th>Environmental conditions</th>
<th>Colour codes</th>
</tr>
</thead>
<tbody>
<tr>
<td>OSB 1</td>
<td>General purpose, interior fitments</td>
<td>Dry conditions (use class 1)</td>
<td>White blue</td>
</tr>
<tr>
<td>OSB 2</td>
<td>Load-bearing</td>
<td>Dry conditions (use class 1)</td>
<td>Yellow, yellow blue</td>
</tr>
<tr>
<td>OSB 3</td>
<td>Load-bearing</td>
<td>Humid conditions (use class 2)</td>
<td>Yellow, yellow green</td>
</tr>
<tr>
<td>OSB 4</td>
<td>Heavy duty, load-bearing</td>
<td>Humid conditions (use class 2)</td>
<td>Yellow green</td>
</tr>
</tbody>
</table>

Note:
Use class environmental conditions were previously called hazard class.

The Euroclass fire performance rating under the conditions specified in BS EN 13986: 2004 for 9 mm untreated oriented strand board with a minimum density of 600 kg/m³ is Class D-s2, d0 for non-floor use and Class DFL-s1 for floorings.

Flaxboard

Manufacture
Flaxboard is manufactured from a mixture of at least 70% flax shives (thin slices) and adhesive. Other raw materials such as wood flakes, chips and saw dust may be incorporated. The boards are formed under heat and pressure.

Grades and uses
The standard BS EN 15197: 2007 describes four types of flaxboard, according to their potential use and environmental conditions.

<table>
<thead>
<tr>
<th>Grade</th>
<th>Purpose/loading</th>
<th>Environmental conditions</th>
</tr>
</thead>
<tbody>
<tr>
<td>FB1</td>
<td>General purpose/filling</td>
<td>Dry</td>
</tr>
<tr>
<td>FB2</td>
<td>Non-load-bearing/veneering</td>
<td>Dry</td>
</tr>
<tr>
<td>FB3</td>
<td>Non-load-bearing/interior fitment and furniture</td>
<td>Dry</td>
</tr>
<tr>
<td>FB4</td>
<td>Non-load-bearing</td>
<td>Humid</td>
</tr>
</tbody>
</table>
Fig. 4.35  Oriented strand board, wood wool and compressed straw slabs
Scrimber

Scrimber is manufactured from small forest thinnings, typically radiata pine, aspen, birch and larch. The small section timber, in the range 70–120 mm diameter, is debarked and crushed to a web of fibrous material, largely maintaining the orientation of the timber fibres. The material is dried and reconstituted with adhesive under heat and pressure into a solid product. The scrimber produced has many of the good properties of natural timber, including ease of nailing, and can be manufactured to any required dimensions, restricted only by the size of the press. Scrimber has also been manufactured from crushed bamboo.

FIBREBOARDS

Fibreboards are manufactured from wood or other plant fibres by the application of heat and/or pressure. They are bonded by the inherent adhesive properties and felting of the fibres or by the addition of a synthetic binder. In the wet process used for the manufacture of hardboard, medium board and softboard, no adhesive is added to the wood fibres. In the case of medium-density fibreboard (MDF), a resin-bonding agent is incorporated during the production process.

Manufacture

**Wet process**

Forest thinnings and wood waste are chipped and then softened by steam heating. The chips are ground down into wood fibres and made into a slurry with water. The slurry is fed onto a moving wire-mesh conveyor, where the excess water is removed by suction and light rolling which causes the fibres to felt together. The wet lap is then cut to lengths and transferred to a wire mesh for further pressing and heat treatment to remove the remaining water and complete the bonding process. Boards are then conditioned to the correct moisture content and packaged. The range of products primarily arise from the differing degrees of compression applied during the manufacturing process (Fig. 4.36).

**Dry process – medium-density fibreboard**

The manufacture of medium-density fibreboard (MDF) involves the addition of adhesive, usually urea formaldehyde, to the dry wood fibres, which are laid up to an appropriate thickness, slightly compressed to a density of at least 450 kg/m³ and cut to board lengths. The boards are cured under heat and pressure in a press, trimmed to size and sanded. MDF has the advantage of a high-quality, machinable finish, and is now used for the production of various mouldings as well as boards. Decorative profiled sheets can

Fig. 4.36 Manufacture of fibreboards
be manufactured by laser cutting of MDF panels to individual client designs. Because of the uniformity of the material, solid sections can be routed to any form. It is therefore widely used for furniture panels as well as internal load-bearing applications.

Nine grades are described in BS EN 622-5: 2006 relating to their anticipated loading and environmental conditions. Most MDF is based on urea formaldehyde resin, but to ensure safety, the quantities of formaldehyde used are strictly controlled by appropriate standards (BS EN 622-1: 2003). Where improved moisture resistance is required, a melamine–urea formaldehyde resin is used, but this material is not suitable for exterior applications. MDF sheets and mouldings can be finished with a range of coatings including paints, lacquers, stains, plastic laminates, wood veneers and foils.

Moisture-resistant dense MDF (690–800 kg/m³) with all-through colour is available in a range of colours and thicknesses from 8 to 30 mm. The material is manufactured from organic-dyed, fade-resistant wood fibres and melamine resin with a low residual formaldehyde content. The material can be machined to decorative forms and patterns with a high-quality surface finish which requires only sealing to enhance the colours.

Grades of medium-density fibreboard (MDF):

<table>
<thead>
<tr>
<th>Grade</th>
<th>Purpose/loading</th>
<th>Environmental conditions</th>
<th>Colour codes</th>
</tr>
</thead>
<tbody>
<tr>
<td>MDF</td>
<td>General purpose</td>
<td>Dry</td>
<td>White, white blue</td>
</tr>
<tr>
<td>MDF.H</td>
<td>General purpose</td>
<td>Humid</td>
<td>White, white green</td>
</tr>
<tr>
<td>MDF.LA</td>
<td>Load-bearing</td>
<td>Dry</td>
<td>Yellow, yellow blue</td>
</tr>
<tr>
<td>MDF.HLS</td>
<td>Load-bearing</td>
<td>Humid</td>
<td>Yellow, yellow green</td>
</tr>
<tr>
<td>L-MDF</td>
<td>Non-load-bearing/ general purpose</td>
<td>Dry</td>
<td>Yellow, yellow green</td>
</tr>
<tr>
<td>L-MDF.H</td>
<td>Non-load-bearing/ general purpose</td>
<td>Humid</td>
<td>Yellow, yellow green</td>
</tr>
<tr>
<td>UL1-MDF</td>
<td>Non-load-bearing/ general purpose</td>
<td>Dry</td>
<td>White, white blue</td>
</tr>
<tr>
<td>UL2-MDF</td>
<td>Non-load-bearing/ general purpose</td>
<td>Dry</td>
<td>White, white blue</td>
</tr>
<tr>
<td>MDF.RWH</td>
<td>Rigid underlays in roofs and walls</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Notes:
L-MDF refers to light MDF and UL-MDF to ultra light MDF, respectively.
Grades to BS EN 622-5: 2006.
MDF for exterior use is colour coded grey.
MDF treated with flame retardant is colour coded red.

The Euroclass fire performance rating under the conditions specified in BS EN 13986: 2004 for 9 mm untreated MDF with a minimum density of 600 kg/m³ is Class D-s2, d0 for non-floor use and Class DFL-s1 for floorings.

**Hardboard**

Hardboards are the densest fibreboards, with a minimum density of 900 kg/m³. The boards range in colour from light to dark brown, usually with one smooth surface and a mesh-textured surface on the underside, although *duo-faced* hardboard—smooth on two faces—is available. Standard sheet sizes are 1220 × 2440 to 3600 mm and 1700 × 4880 mm; there are also door sizes. Standard thicknesses range from 3.2 to 6.4 mm, although a wider range is available.

The standard BS EN 622-2: 2004 specifies six grades of hardboard according to load-bearing properties and environmental conditions.

Grades of hardboard:

<table>
<thead>
<tr>
<th>Grade</th>
<th>Purpose/loading</th>
<th>Environmental conditions</th>
<th>Colour codes</th>
</tr>
</thead>
<tbody>
<tr>
<td>HB</td>
<td>General purpose</td>
<td>Dry</td>
<td>White, white blue</td>
</tr>
<tr>
<td>HB.H</td>
<td>General purpose</td>
<td>Humid</td>
<td>White, white green</td>
</tr>
<tr>
<td>HB.E</td>
<td>General purpose</td>
<td>Exterior</td>
<td>White, white brown</td>
</tr>
<tr>
<td>HB.LA</td>
<td>Load-bearing</td>
<td>Dry</td>
<td>Yellow, yellow blue</td>
</tr>
<tr>
<td>HB.LA1</td>
<td>Load-bearing</td>
<td>Humid</td>
<td>Yellow, yellow green</td>
</tr>
<tr>
<td>HB.LA2</td>
<td>Heavy duty, load-bearing</td>
<td>Humid</td>
<td>Yellow, yellow green</td>
</tr>
</tbody>
</table>

Standard hardboard is suitable for internal use, typically panelling, wall and ceiling linings, floor underlays and furniture. A range of perforated, embossed and textured surfaces is available. Applied coatings include primed or painted and various printed woodgrain, PVC or melamine foils.

The Euroclass fire performance rating under the conditions specified in BS EN 13986: 2004 for 6 mm untreated hardboard with a minimum density of 900 kg/m³ is Class D-s2, d0 for non-floor use and Class DFL-s1 for floorings.

**Tempered hardboard**

Tempered hardboards, impregnated with oils during manufacture, are denser and stronger than the standard hardboards, with enhanced water- and abrasion-resistance. Tempered hardboards are dark brown to black in colour and have a density usually exceeding 960 kg/m³. Tempered hardboards are
suitable for structural and exterior applications. The high shear strength of the material is used within hardboard-web structural box and I beams. Typical exterior applications include claddings, fascias and soffits, where weather resistance is important. The moisture resistance of tempered hardboard makes it suitable for lining concrete formwork.

Mediumboard and softboard

Mediumboard and softboard are manufactured by the wet process. Mediumboard (high-density and low-density) and softboard exhibit a range of physical properties which reflects the degree of compression applied during the manufacturing process. High-density mediumboard (density 560–900 kg/m³) has a dark brown shiny surface like hardboard. Low-density mediumboard (density 400–560 kg/m³) has a light brown soft finish. Softboard (density 210–400 kg/m³) is light in colour with a fibrous, slightly textured finish. Softboard impregnated with bitumen offers an increased moisture resistance over the untreated material.

The general-purpose exterior grades (E) may be used for exterior cladding. The higher-density grades (H) are used for wall linings, sheathing, partitioning, ceilings and floor underlays. Low-density mediumboard (L) is used for wall linings, panelling, ceilings and notice boards. Softboard is used for its acoustic and thermal insulating properties. Bitumen impregnated softboard is suitable for use as a floor underlay to chipboard on concrete.

The standard BS EN 622-3: 2004 specifies ten grades of low (L) and high (H) density mediumboard according to load-bearing requirements and environmental conditions.

Grades of mediumboard:

<table>
<thead>
<tr>
<th>Grade</th>
<th>Purpose/loading</th>
<th>Environmental conditions</th>
<th>Colour codes</th>
</tr>
</thead>
<tbody>
<tr>
<td>MB.L</td>
<td>General purpose</td>
<td>Dry</td>
<td>White, white blue</td>
</tr>
<tr>
<td>MB.H</td>
<td>General purpose</td>
<td>Dry</td>
<td>White, white blue</td>
</tr>
<tr>
<td>MBL.H</td>
<td>General purpose</td>
<td>Humid</td>
<td>White, white green</td>
</tr>
<tr>
<td>MBH.H</td>
<td>General purpose</td>
<td>Humid</td>
<td>White, white green</td>
</tr>
<tr>
<td>MBLE</td>
<td>General purpose</td>
<td>Exterior</td>
<td>White, white brown</td>
</tr>
<tr>
<td>MBH.E</td>
<td>General purpose</td>
<td>Exterior</td>
<td>White, white brown</td>
</tr>
<tr>
<td>MBH.LA1</td>
<td>Load-bearing</td>
<td>Dry</td>
<td>Yellow, yellow blue</td>
</tr>
<tr>
<td>MBH.LA2</td>
<td>Heavy duty, load-bearing</td>
<td>Dry</td>
<td>Yellow, yellow blue</td>
</tr>
<tr>
<td>MBH.HLS1</td>
<td>Load-bearing</td>
<td>Humid</td>
<td>Yellow, yellow green</td>
</tr>
<tr>
<td>MBH.HLS2</td>
<td>Heavy duty, load-bearing</td>
<td>Humid</td>
<td>Yellow, yellow green</td>
</tr>
</tbody>
</table>

The Euroclass fire performance rating under the conditions specified in BS EN 13986: 2004 for 9 mm untreated high-density mediumboard of 600 kg/m³ is Class D-s2, d0 for non-floor use and Class DFL-s1 for floorings. For untreated low-density mediumboard of 400 kg/m³ the equivalent rating is Class E, pass for non-floor use and Class EFL for floorings.

The standard BS EN 622-4: 1997 specifies five grades of softboard according to load-bearing properties and environmental conditions.

Grades of softboard:

<table>
<thead>
<tr>
<th>Grade</th>
<th>Purpose/loading</th>
<th>Environmental conditions</th>
<th>Colour codes</th>
</tr>
</thead>
<tbody>
<tr>
<td>SB</td>
<td>General purpose</td>
<td>Dry</td>
<td>White, white blue</td>
</tr>
<tr>
<td>SB.H</td>
<td>General purpose</td>
<td>Humid</td>
<td>White, white green</td>
</tr>
<tr>
<td>SB.E</td>
<td>General purpose</td>
<td>Exterior</td>
<td>White, white brown</td>
</tr>
<tr>
<td>SB.LS</td>
<td>Load-bearing</td>
<td>Dry</td>
<td>Yellow, yellow blue</td>
</tr>
<tr>
<td>SB.HLS</td>
<td>Load-bearing</td>
<td>Humid</td>
<td>Yellow, yellow green</td>
</tr>
</tbody>
</table>

The Euroclass fire performance rating under the conditions specified in BS EN 13986: 2004 for 9 mm untreated softboard of 250 kg/m³ is Class E, pass for non-floor use and Class EFL-s1 for floorings.

WOOD WOOL SLABS

Manufacture

Wood wool slabs are manufactured by compressing long strands of chemically stabilised wood fibres coated in Portland cement (Fig. 4.35, page 147). The grey product has an open texture which may be left exposed, spray painted or used as an effective substrate for plastering. It is also a suitable material for permanent shuttering for concrete. Slabs are available in a range of thicknesses from 25 to 150 mm, typically 500, 600 or 625 mm wide and up to 3 m in length. Standard sizes are listed in BS EN 13168: 2001.

Types and uses

Wood wool slabs are available either plain-edged or with interlocking galvanised-steel channels to the longitudinal edges. Thicknesses in the range 15–50 mm are suitable for ceilings, partitions, wall linings and permanent concrete shuttering. The thicker grades from 50 to 150 mm may be used for roof decking, with spans up to 3 m according to the anticipated loading. Some products incorporate an additional insulation layer of
mineral wool or cellular plastics to enhance thermal properties.

The material is rated as Class 0 with respect to Building Regulations and Class I (BS 476-7: 1997) in terms of surface spread of flame. Classification to the European Standard (BS EN 13501-1: 2007) is subject to manufacturer’s testing. The material is resistant to fungal attack and is unaffected by wetting. Wood wool slabs offer good sound-absorption properties due to their open-textured surface. This is largely unaffected by the application of sprayed emulsion paint. The material is therefore appropriate for partitions, internal walls and ceilings where sound absorption is critical. Acoustic insulation for a pre-screeded 50 mm slab is typically 30 dB. The relatively high proportion of void space affords the material good thermal insulating properties with a typical thermal conductivity of 0.077 W/m K at 8% moisture content.

The material is workable, being easily cut and nailed. Where a gypsum plaster or cement/lime/sand external rendering is to be applied all joints should be reinforced with scrim. Wood wool slabs form a suitable substrate for flat roofs finished with reinforced-bitumen membranes, mastic asphalt or metals.

COMPRESSED STRAW SLABS

Compressed straw slabs are manufactured by forming straw under heat and pressure, followed by encapsulation in a fibreglass mesh and plastering grade paper (Fig. 4.35, page 147). Typically used for internal partitioning, the panels are mounted onto a timber sole plate and butt-jointed with adhesive or dry-jointed with proprietary sherardised clip fixings. All joints are jute scrimmed and the whole partitioning finished with a 3 mm skim of board plaster. The slabs are 58 mm thick by 1200 mm wide in a range of standard lengths from 2270 to 2400 mm. Service holes may be incorporated into the panels at 300 mm centres for vertical electrical wiring. While normal domestic fixtures can be fitted directly to the panels, heavier loads require coach-bolt fixings through the panels. The product should not be used where it will be subjected to moisture. Compressed straw slabs, when skim plastered, have a 30 minute fire resistance rating, a Class 0 spread of flame, and a sound reduction of typically 35 dB over the range 100–3200 Hz. The standard BS 4046: 1991 describes four types of compressed straw slabs depending on boron-based insecticide treatment and the provision of continuous longitudinal voids for services. Alternative finishes are plain paper for direct decoration or showerproof paper.

Types of compressed straw slabs:

- Type A Untreated Solid core
- Type B Insecticide treated Solid core
- Type C Untreated Continuous voids in core
- Type D Insecticide treated Continuous voids in core

THATCH

Thatch was the roof covering for most buildings until the end of the Middle Ages, and remained the norm in rural areas until the mid-nineteenth century. For most of the twentieth century thatch was only used in conservation work; however, with the new resurgence of interest in the material, partially associated with the reconstruction of the Globe Theatre in London (Fig. 4.37), thatch once again has become a current construction material.

Materials

The three standard materials for thatching within the UK are water reed (*Phragmites australis*), long straw (usually wheat) and combed wheat reed (also known as Devon reed). Water reed is associated with the Norfolk broads, the fens, south Hampshire and the Tay estuary, but much is imported from Turkey, Poland, Romania and China. Long straw is the standard thatch in the Midlands and Home Counties, while combed wheat reed is more common in Devon and Cornwall. Water reed is the most durable, lasting typically 50–60 years, but long straw and combed wheat reed last approximately 20 and 30 years, respectively, depending on location and roof pitch. All thatched roofs will need reridding at 10–15 year intervals; in the case of water reed this is often done with saw sedge (*Cladium mariscus*) which is more flexible than the reed itself.

Both long straw and combed wheat reed are often grown and harvested specifically as thatching materials to ensure long, undamaged stems. Long straw is threshed winter wheat, whereas combed wheat straw is wheat with any leaves and the grain head removed. An alternative to combed wheat reed is triticale (*Triticale hexaploide*), which is a cross between wheat and rye. Triticale produces a more reliable harvest than other forms of wheat straw and it is indistinguishable from combed wheat reed when used as a thatching material. Water reed for thatching is usually between 915 and
1830 mm in length. Typical lengths for long straw and combed wheat reed are 760 and 915 mm, respectively.

**Appearance**

Long straw roofs show the lengths of the individual straws down the roof surface and are also characterised by the use of split hazel rodding around the eaves and gables to secure the thatch. To prevent attack by birds they are frequently covered in netting. Combed wheat reed and water reed both have a closely packed finish with the straw ends forming the roof surface. A pitch of about $50^\circ$ is usual for thatch with a minimum of $45^\circ$, the steeper pitches being more durable. The ridge, which may be a decorative feature, is produced by either wrapping wheat straw over the apex or butting up reeds from both sides of the roof. Traditionally, hazel twigs are used for fixings, although these can be replaced with stainless steel wires. The durability of thatch is significantly affected by the climate. All materials tend to have shorter service lives in warmer locations with high humidity, which encourage the development of fungi. Chemical treatment, consisting of an organic heavy metal compound, may be used, preferably on new thatch, to delay the biological decomposition. Thatch is usually laid to a thickness of between 220 and 400 mm.

**Properties**

**Fire**

The fire hazards associated with thatched roofs are evident; however, fire retardants can be used, although these may denature the material. In the case of the Globe Theatre in London, a sparge water-spray system has been installed. In other new installations, the granting of planning consent has been facilitated by the location of permanent water drenching systems near the ridge and by the use of fire-resisting board and foil under the thatch to prevent internal fire spread. Electrical wiring and open fire chimneys are the most common causes of thatch fires, although maintenance work on thatched roofs is also a risk if not carefully managed.

**Insulation**

Thatch offers good insulation, keeping buildings cool in summer and warm in winter, a typical $300 \text{ mm}$ of water reed achieving a $U$-value of $0.35 \text{ W/m}^2 \text{ K}$. 
Western red cedar (Thuja plicata), as a naturally durable material, is frequently used as shingles or shakes for roofing or cladding. Shingles are cut to shape, whereas shakes are split to the required thickness, usually between 10 and 13 mm. Both shakes and shingles may be tapered or straight. Shingles, typically 400, 450 or 600 mm long and between 75 and 355 mm wide, may be treated with copper-based wood preservatives to improve their durability. Additionally, they may be treated with flame retardant to satisfy the AA fire rating of BS 476-3: 1975. Shingles should be fixed with corrosion-resistant nails, leaving a spacing of 5–6 mm between adjacent shingles. A minimum pitch of 14° is necessary and three layers are normally required. Whilst the standard laying pattern is straight coursing, staggered patterns and the use of profiled shingles on steeper pitches can create decorative effects. Shingles should normally be laid over a waterproof breather membrane. The standard BS 5534: 2003 suggests a minimum side lap of 40 mm for shakes and shingles.

Figure 4.38 illustrates the typical detailing of cedar shingles as a lightweight cladding material. The natural colour of the material gives an overall warmth to the exterior envelope of the building. In exposed locations the red-brown cedar wood surface gradually weather to a silver-grey, while in very sheltered locations the shingles become green with lichen. Decorative shape shingles include fish-scale, arrow, hexagonal, half cove, diagonal and round.

**STEKO BLOCKS**

Steke is an innovative wall system, which uses very accurately engineered large hollow timber blocks which simply slot together. No fixings or glue is needed to form load-bearing wall construction. The blocks are manufactured from small off-cuts of spruce timber, which can be readily obtained from renewable sources. The blocks (640 × 320 × 160 mm wide) weighing 6.5 kg, are manufactured from two 20 mm thick panels glued to horizontal battens and separated by vertical studs (Fig. 4.39). Units fit snugly together with their tongued and grooved profiles and vertical dowels. Special units include quarter, half and three-quarter blocks, also components for corners, lintels, wall closers, base plates, wall plates and solid blocks for point loads. Walls can be up to 20 m long without additional bracing, and 3 m high, but up to four or five storeys high, if the necessary horizontal bracing is provided by the intermediate floors and roof. Where two-storey unrestrained walls are required they can be post-tensioned with threaded steel rods. The blocks are untreated except when borax protection against house longhorn beetle is required. Electrical
services can run in the voids, which are finally dry-injected with cellulose insulation. Steko construction must work from a 300 mm upstand to avoid contact with surface water, and should be protected externally with a breather paper to prevent rain damage during construction. Internal finishes may be left exposed or finished with 15 mm gypsum plasterboard, which gives 30 minutes’ fire resistance. Additional insulation is required externally to achieve the current Building Regulation requirements. Typically, 100 mm of mineral wool and a 20 mm render externally on cellulose filled and internally plastered blocks will achieve a $U$-value of 0.20 W/m K. The first UK house built with the Steko system, which originated in Switzerland, has been designed and built on a cliff-top site in Downderry, Cornwall.

**FLEXIBLE VENEERS**

Flexible veneers are sheets of wood veneer which have been backed with paper or fibre reinforcement to allow the material to be handled without splitting. Flexible veneers can be moulded onto profiled components of MDF or plywood by a rolling process. Typical standard components are cornice and pelmet trims for kitchen furniture. The material can be rolled up for storage without damage, unlike traditional veneer. The veneer is fixed to the substrate with polyvinyl acetate (PVA) or urea formaldehyde (UF) adhesive under pressure or with contact adhesive. The material is increasingly being specified by designers for creating high-quality polished wood finishes to complex curved forms such as reception desks and wall panelling.

**DECORATIVE TIMBERS**

Certain organic materials, such as coconut shell and palm wood are being developed as decorative materials for surface finishes. Palm wood is also being used to manufacture palm plywood, it is illustrated in Fig. 4.40. Bamboo is used for the interior of Madrid Barajas International Airport (Fig. 4.41) to create a relaxing atmosphere, whilst using a renewable resource. Potentially bamboo can be used as a structural material as in South America, but durability and jointing systems need resolution.

**Recycling timber**

According to Timber Research and Development Association (TRADA), the UK generates approximately 11 million tonnes of timber waste annually of which 80% enters landfill sites. Not only is this costly, but it also generates the greenhouse gas, methane, as it degrades with time. About half the total of timber waste arises from building construction and demolition. Of the 20% of timber waste which is currently recovered, only 16% is recycled and 4% used for energy production. Furthermore, poor handling and bad storage of new timber products on site leads to an average 10% wastage, and this can rise to 20%, according to TRADA, in extreme cases.
Options for reducing the waste include better construction site management and direct reuse as in timber flooring. However, the main market is for reprocessing by chipping the material for the manufacture of particleboard and cement bonded permanent concrete shuttering or for its use as mulch or compost. Energy generation from waste timber and biomass crops requires further development, but has good potential, and the distillation of timber can produce ethanol as a biofuel.

Timber products such as particleboard with high quantities of polymer resin, and timber treated with some of the older hazardous wood preservatives such as creosote and chromated copper arsenate (CCA), cannot be recycled into new products.

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BUILDING RESEARCH ESTABLISHMENT PUBLICATIONS
BRE Special digests

BRE SD7: 2008 Insulation of timber-frame construction. U-values and regulations for the UK, Republic of Ireland and the Isle of Man.

BRE Digests
BRE Digest 423: 1997 The structural use of wood-based panels.
BRE Digest 431: 1998 Hardwoods for joinery and construction (Parts 1, 2 and 3).
BRE Digest 443: 1999 Termite and UK buildings (Parts 1 and 2).
BRE Digest 470: 2002 Life cycle impacts of timber.
BRE Digest 477 Part 5: 2004 Wood-based panels: Medium density fibreboard (MDF).
BRE Digest 479: 2003 Timber piles and foundations.
BRE Digest 492: 2005 Timber grading and scanning.
BRE Digest 496: 2005 Timber frame building.
BRE Good building guides
BRE GBG 32: 1999 Ventilating thatched roofs.

BRE Information papers
BRE IP 2/96 Assessment of exterior medium density fibreboard.
BRE IP 8/96 Moisture resistance of laminated veneer lumber (LVL).
BRE IP 9/96 Preservative-treated timber for exterior joinery - European standards.
BRE IP 4/97 Preservative-treated timber for exterior joinery - Applying European standards.
BRE IP 2/01 Evaluating joinery preservatives.
BRE IP 13/01 Preservative-treated timber - The UK’s code of best practice.
BRE IP 14/01 Durability of timber in ground contact.
BRE IP 1/03 European Standards for wood preservatives and treated wood.
BRE IP 9/03 Best practice of timber waste management.
BRE IP 13/04 An introduction to building with structural insulated panels.
BRE IP 10/05 Green gluing of timber: a feasibility study.
BRE IP 13/05 Incising UK-grown Sitka spruce.
BRE IP 3/07 Modern methods of construction (MMC) in housing (Parts 1 – 4).

BRE Reports
BR 276: 1995 Long-term field trials on preserved timber in ground contact.

TRADA PUBLICATIONS
Wood information sheets
WIS 0-3: 2006 Introduction to timber framed construction.
WIS 4-7: 2006  Timber strength grading and strength classes.
WIS 4-12: 2008  Care of timber and wood-based products on building sites.
WIS 4-14: 2006  Moisture in timber.
WIS 4-16: 2002  Timber in joinery.
WIS 4-29: 2002  Dry-graded structural softwood.
WIS 4-31: 2008  Life cycle costing.
WIS 4-33: 2009  Life cycle assessment.

ADVISORY ORGANISATIONS

British Woodworking Federation, 55 Tufton Street, London SW13QL, UK (0870 458 6939).

Coed Cymru, The Old Sawmill, Tregynon, Newtown, Powys SY16 3PL, UK (01686 650777).
Glued Laminated Timber Association, Chiltern House, Stocking Lane, Hughenden Valley, High Wycombe, Bucks HP14 4ND, UK (01494 565180).
Thatching Advisory Services Ltd., The Old Stables, Redenham Park Farm, Redenham, Andover, Hampshire SP11 9AQ, UK (01264 773820).
Timber Research and Development Association, Stocking Lane, Hughenden Valley, High Wycombe, Buckinghamshire HP14 4ND, UK (01494 569600).
Trussed Rafter Association, P.O. Box 571, Chesterfield, Sheffield S40 9DH, UK (01246 230036).
UK Timber Frame Association Ltd., The e-Centre, Cooperage Way, Business Village, Alloa FK10 3LP, UK (01259 272140).
Wood for Good Ltd., 211 High Road, London N28AN, UK (020 8365 2700).
Wood Panel Industries Federation, 28 Market Place, Grantham, Lincolnshire NG31 6LR, UK (01467 563707).
FERROUS AND NON-FERROUS METALS

Introduction

A wide range of ferrous and non-ferrous metals and their alloys are used within the construction industry, but iron, steel, aluminium, copper, lead and zinc predominate. Over the past decade titanium has featured significantly in construction, having previously been used mainly in the chemical process industry and for military purposes. Recent trends have been towards the development of more durable alloys and the use of coatings to both protect and give visual diversity to the product ranges. Generally, the metals require a large energy input for their production from raw materials; however, this high embodied energy is partially offset by the long life and recycling of most metals. The recovery rate of steel from demolition sites in the UK approaches 99%, with 10% being reused and the remainder recycled. Approximately 60% of current steel production is from scrap, and steel can be recycled any number of times without any degradation of the material.

Ferrous metals

Ferrous metals are defined as the metals in which the element iron predominates. The earliest use of the metal was for the manufacture of implements and weapons in the Iron Age commencing in Europe circa 1200 BC. Significant developments were the use by Wren in 1675 of a wrought iron chain in tension to restrain the outward thrust from the dome of St Paul’s Cathedral, the use of cast iron in compression for the Ironbridge at Coalbrookdale in 1779, and by Paxton in the prefabricated sections of the Crystal Palace in 1851.

Steel is a relatively recent material, only being available in quantity after the development of the Bessemer converter in the late nineteenth century. The first steel-frame high-rise building of 10 storeys was built in 1885 in Chicago by William le Baron Jenney.

The platform level of the Waterloo train terminal in London (Fig. 5.1) is covered by curved and tapered 3-pin steel arches, which are designed to accommodate the flexing inevitably caused by the movement of trains at this level. The steel arches, each consisting of two prismatic bow-string trusses connected by a knuckle joint, are asymmetrical to allow for the tight curvature of the site. The tops of the longer trusses are covered with toughened glass providing views towards old London, with profiled stainless steel spanning between. The area spanned by the shorter trusses is fully glazed. The structure is designed for a minimum lifetime of 100 years.

MANUFACTURE OF STEEL

The production of steel involves a sequence of operations which are closely interrelated in order to ensure maximum efficiency of a highly energy-intensive process. The key stages in the production process are the making of pig iron, its conversion into steel, the casting of the molten steel and its formation into sections or strip. Finally, coils of steel strip are cold-rolled into thin sections and profiled sheet.

Manufacture of pig iron

The raw materials for the production of iron are iron ore, coke and limestone. Most iron ore is imported from America, Australia and Scandinavia, where the
Fig. 5.1 Structural steelwork — Waterloo train terminal, London. Architects: Grimshaw Architects. Photographs: Courtesy of Jo Reid & John Peck, Peter Strobel
iron content of the ore is high. Coke is produced from coking coal, mainly imported from Europe, in batteries of coking ovens. Some of this coke is then sintered with iron ore prior to the iron-making process.

Iron ore, coke, sinter and limestone are charged into the top of the blastfurnace (Fig. 5.2). A hot air blast, sometimes enriched with oxygen, is fed through the tuyères into the base of the furnace. This heats the furnace to white heat, converting the coke into carbon monoxide, which then reduces the iron oxide to iron.

The molten metal collects at the bottom of the furnace. The limestone forms a liquid slag, floating on the surface of the molten iron. Purification occurs as impurities within the molten iron are preferentially absorbed into the slag layer.

\[
2\text{C (coke)} + \text{O}_2 \rightarrow 2\text{CO (carbon monoxide)}
\]

\[
\text{Fe}_2\text{O}_3 + 3\text{CO} \rightarrow 2\text{Fe} + 3\text{CO}_2
\]
The whole process is continuous, as relining the blastfurnace with the special refractory bricks is expensive and time-consuming. From time to time, as the molten slag level rises, excess is tapped off for subsequent disposal as a by-product of the steel-making industry. When hot metal is required for the subsequent steel-making process it is tapped off into huge ladles for direct transportation to the steel converter. At this stage the iron is only 90–95% pure with sulphur, phosphorus, manganese and silicon as impurities and a carbon content of 4–5%. Waste gases from the blastfurnace are cleaned and recycled as fuel within the plant. A blastfurnace will typically operate non-stop for 10 years, producing 40,000 tonnes per week.

Steel-making

There are two standard processes used within the UK for making steel. The basic oxygen process is used for the manufacture of bulk quantities of standard-grade steels, and the electric arc furnace process is used for the production of high-quality special steels and particularly stainless steel. The Manchester Stadium (Fig. 5.3) built for the Commonwealth Games in 2002 and Manchester City Football Club used approximately 2000 tonnes of structural steel.

Basic oxygen process

Bulk quantities of steel are produced by the basic oxygen process in a refractory lined steel furnace which can be tilted for charging and tapping. A typical furnace (Fig. 5.4) will take a charge of 350 tonnes and convert it into steel within 30 minutes. Initially scrap metal, accounting for one quarter of the charge, is loaded into the tilted furnace, followed by the remainder of the charge as hot metal direct from the blastfurnace. A water-cooled lance is then lowered to blow
Fig. 5.5 Electric arc furnace

high-pressure oxygen into the converter. This burns off impurities and reduces the excess carbon content while raising the temperature. Argon and a small quantity of nitrogen are introduced at the bottom of the furnace. Lime is added to form a floating slag to remove further impurities and alloying components are added to adjust the steel composition, prior to tapping. Finally, the furnace is inverted to run out any remaining slag, prior to the next cycle.

Electric arc process
The electric arc furnace (Fig. 5.5) consists of a refractory-lined hearth, covered by a removable roof, through which graphite electrodes can be raised and lowered. With the roof swung open, scrap metal is charged into the furnace, the roof is closed and the electrodes are lowered to near the surface of the metal. A powerful electric arc is struck between the electrodes and the metal, which heats it up to melting point. Lime and fluorspar are added to form a slag, and oxygen is blown into the furnace to complete the purification process. When the temperature and chemical analysis are correct, the furnace is tilted to tap off the metal, to which appropriate alloying components may then be added. A typical furnace will produce 150 tonnes of high-grade or stainless steel within 90 minutes.

Casting
Traditionally, the molten steel was cast into ingots, prior to hot rolling into slabs and then sheet. However, most steel is now directly poured, or teemed, and cast into continuous billets or slabs, which are then cut to appropriate lengths for subsequent processing. Continuous casting (Fig. 5.6), which saves on reheating, is not only more energy efficient than processing through the ingot stage but also produces a better surface finish to the steel.

Hot-rolled steel
Sheet steel is produced by passing 25 tonne hot slabs at approximately 1250°C through a series of computer-controlled rollers which reduce the thickness to typically between 1.5 and 20 mm prior to water cooling and coiling. A 25 tonne slab would produce 1 km coil of 2 mm sheet. Steel sections such as universal beams and columns, channels and angle (Fig. 5.7) are rolled from hot billets through a series of stands to the appropriate section.

Cold-rolled steel
Sheet steel may be further reduced by cold rolling, which gives a good surface finish and increases its tensile strength. Light round sections may be processed into steel for concrete reinforcement, whilst coiled sheet may be converted into profiled sheet or light steel sections (Fig. 5.7). Cold-reduced steel for construction is frequently factory finished with zinc, alloys including terne (lead and tin) or plastic coating. Cold-reduced sheet steel of structural quality is covered by BS ISO 4997: 2007.
CARBON CONTENT OF FERROUS METALS

The quantity of carbon alloyed with iron has a profound influence on the physical properties of the metal due to its significant effect on the microscopic crystal structure (Fig. 5.8). At ambient temperature a series of crystal forms (ferrite, pearlite and cementite) associated with different proportions of iron and carbon are stable. However, on increasing the temperature, crystal forms that were stable under ambient conditions become unstable and are recrystallised into the high-temperature form (austenite). This latter crystal structure can be trapped at room temperature by the rapid quenching of red-hot steel, thus partially or completely preventing the natural recrystallisation processes which otherwise would occur on slow cooling. These effects are exploited within the various heat treatments that are applied to steels in order to widen the available range of physical properties.

Wrought iron

Wrought iron contains only about 0.02% of carbon. It was traditionally made by remelting and oxidising pig iron in a reverberatory furnace. The process was continued until virtually all the high carbon content of the pig iron had been burnt off to produce a pasty wrought iron, which was withdrawn from the furnace and then hammered out. Wrought iron is fibrous in character due to the incidental incorporation within the metal of slag residues and impurities such as magnesium sulphide, which are formed into long veins by the hammering process. Wrought iron has a high melting point, approaching 1540°C, depending on its purity. It was traditionally used for components in tension due to its tensile strength of about 350 MPa. It is ductile and easily worked or forged when red hot, and thus eminently suitable for crafting into ornamental ironwork, an appropriate use because of its greater resistance to corrosion than steel. Because of its high melting point, wrought iron cannot be welded or cast. Production ceased in the UK in 1974 and modern wrought iron is either recycled old material or, more frequently, low-carbon steel, with its attendant corrosion problems.

Cast iron

Cast iron contains in excess of 2% carbon in iron. It is manufactured by the carbonising of pig iron and scrap with coke in a furnace. The low melting point of around 1130°C and its high fluidity when molten, give rise to its excellent casting properties but, unlike wrought iron, it cannot be hot worked and is generally a brittle material. The corrosion resistance of cast iron has been exploited in its use for boiler castings, street furniture and traditional rainwater goods. Modern foundries manufacture castings to new designs and as reproduction Victorian and Edwardian components.

Differing grades of cast iron are associated with different microscopic crystal structures. The common grey cast iron contains flakes of graphite, which cause the characteristic brittleness and impart the grey
Fig. 5.7 Hot-rolled and cold-rolled sections (after Trebilcock 1994)

colour to fractured surfaces. White cast iron contains the carbon as crystals of cementite (iron carbide, $\text{Fe}_3\text{C}$) formed by rapid cooling of the melt. This material may be annealed to reduce its brittle character. A more ductile cast iron (spheroidal cast iron) is produced by the addition of magnesium and ferrosilicon and annealing which causes the carbon to crystallise into graphite nodules. This material has an increased tensile strength and significantly greater impact resistance. All cast irons are strong in compression. The designation system for cast irons is described in BS EN 1560: 1997.

Road iron goods, such as manhole covers, made from largely recycled grey cast iron are heavy but brittle. Where increased impact resistance is required for public roads, lighter and stronger ductile iron components are used. Traditional sand-cast rainwater goods are usually manufactured from grey cast iron, while cast iron drainage systems are manufactured from both grey and ductile iron. Unlike steel, cast iron does not soften prematurely in a fire, but may crack if cooled too quickly with water from a fire hose. Cast iron drainage systems in both grey and spheroidal cast iron are covered by the standard BS EN 877: 1999. Cast iron drainage systems are particularly appropriate in heritage and conservation areas.

Steels

A wide range of steels are commercially available reflecting the differing properties associated with carbon content, the various heat treatments and the addition of alloying components.

Carbon contents of steels range typically between 0.05% and 1.7% and this alone is reflected in a wide spectrum of physical properties. The low-carbon (0.05–0.15%) and mild steels (0.15–0.25%) are...
relatively soft and can be subjected to extensive cold working. Medium-carbon steels (0.25–0.5%), which are often heat treated, are hard wearing. High-carbon steels (0.5–0.9%) and carbon tool steels (0.9–1.7%) exhibit increasing strength and wear resistance with increasing carbon content. In addition, extra-low-carbon steel (<0.02%) and ultra-low-carbon steel (<0.01%), with carbon contents similar to that of traditional wrought iron, are used for high formability and drawing applications.

HEAT TREATMENT OF STEELS

The physical properties of steels can be modified by various heat treatments which involve heating to a particular temperature followed by cooling under controlled conditions. The full range of heat treatments is described in BS EN 10052: 1994. Construction steels for the manufacture of component parts requiring quenching and tempering (+QT) or normalising (+N) are covered by BS EN 10343: 2009.

Hardening

Rapidly quenched steel, cooled quickly from a high temperature in oil or water, thus retaining the high-temperature crystalline form, is hard and brittle. This effect becomes more pronounced for the higher carbon content steels, which are mostly unsuitable for engineering purposes in this state. Quenching is often followed by tempering to reduce excessive hardness and brittleness.

Annealing and normalising

These processes involve the softening of the hard steel, by recrystallisation, which relieves internal stresses within the material and produces a more uniform grain structure. For annealing, the steel is reheated and soaked at a temperature of approximately 650°C, then cooled slowly at a controlled rate within a furnace or cooling pit. This produces the softest steel for a given composition. With normalising, the steel is reheated to 830–930°C for a shorter period and then allowed to cool more rapidly in air. This facilitates subsequent cold working and machining processes.

Tempering

Reheating the steel to a moderate temperature (400–600°C), followed by cooling in air, reduces the brittleness, allowing some recrystallisation of the metal. The magnitude of the effect is directly related to the tempering temperature with ductility increasing, and tensile strength reducing, for the higher process temperatures.

Carburising

Components may be case hardened to produce a higher carbon content on the outer surface, whilst leaving the core relatively soft, thus giving a hard wearing surface without embrittlement and loss of impact resistance to the centre. Usually, this process involves heating the components surrounded by charcoal or other carbon-based material to approximately 900°C for several hours. The components are then heat treated to develop fully the surface hardness.

SPECIFICATION OF STEELS

Steels within the European Union are designated by a series of European Standards, BS EN 10025: 2004.

Hot rolled structural steels:

- BS EN 10025-1: 2004 General technical data
- BS EN 10025-2: 2004 Non-alloy structural steels
- BS EN 10025-3: 2004 Weldable fine-grained structural steels
- BS EN 10025-4: 2004 Rolled weldable fine-grained structural steels
- BS EN 10025-5: 2004 Steels with improved atmospheric corrosion resistance
- BS EN 10025-6: 2004 High yield strength structural steels

In addition, BS EN 10210-1: 2006 and BS EN 10219-1: 2006 relate to hot- and cold-formed structural hollow sections, respectively. The standard grades and their associated characteristic strengths are illustrated in Tables 5.1–5.5. In the standards, S refers to structural steel and the subsequent coding numbers relate to the minimum yield strength. The sub-grade letters refer to impact resistance and other production conditions and compositions, such as W for weather-resistant steel. Steel numbers for each grade of steel are defined by BS EN 10027-2: 1992.

The following example illustrates the two coding systems for one standard grade of steel:
Table 5.1  Steel designations for standard grades to BS EN 10025-2: 2004 (hot rolled products of non-alloy structural steels)

<table>
<thead>
<tr>
<th>Grade</th>
<th>Number</th>
<th>Ultimate tensile strength (MPa)</th>
<th>Minimum yield strength (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>S185</td>
<td>1.0035</td>
<td>290–510</td>
<td>185</td>
</tr>
<tr>
<td>S235JR</td>
<td>1.0038</td>
<td>360–510</td>
<td>235</td>
</tr>
<tr>
<td>S235JO</td>
<td>1.0114</td>
<td>360–510</td>
<td>235</td>
</tr>
<tr>
<td>S235J2</td>
<td>1.0117</td>
<td>360–510</td>
<td>235</td>
</tr>
<tr>
<td>S275JR</td>
<td>1.0044</td>
<td>410–560</td>
<td>275</td>
</tr>
<tr>
<td>S275JO</td>
<td>1.0143</td>
<td>410–560</td>
<td>275</td>
</tr>
<tr>
<td>S275J2</td>
<td>1.0145</td>
<td>410–560</td>
<td>275</td>
</tr>
<tr>
<td>S355JR</td>
<td>1.0045</td>
<td>470–630</td>
<td>355</td>
</tr>
<tr>
<td>S355JO</td>
<td>1.0553</td>
<td>470–630</td>
<td>355</td>
</tr>
<tr>
<td>S355J2</td>
<td>1.0577</td>
<td>470–630</td>
<td>355</td>
</tr>
<tr>
<td>S355K2</td>
<td>1.0596</td>
<td>470–630</td>
<td>355</td>
</tr>
</tbody>
</table>

Notes:
Sub-grades JR, JO, J2 and K2 indicate increasing impact resistance as measured by the Charpy V-notch test.
K has a higher impact energy than J, the symbols R, O and 2 refer to the impact test at room temperature, 0 and −20°C, respectively.
Data is for thicknesses of 16 mm or less.

Table 5.2  Steel designations for higher grade structural steels to BS EN 10025-3: 2004 (hot-rolled products in weldable fine grain structural steels)

<table>
<thead>
<tr>
<th>Grade</th>
<th>Number</th>
<th>Ultimate tensile strength (MPa)</th>
<th>Minimum yield strength (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>S275N</td>
<td>1.0490</td>
<td>370–510</td>
<td>275</td>
</tr>
<tr>
<td>S275NL</td>
<td>1.0491</td>
<td>370–510</td>
<td>275</td>
</tr>
<tr>
<td>S355N</td>
<td>1.0545</td>
<td>470–630</td>
<td>355</td>
</tr>
<tr>
<td>S355NL</td>
<td>1.0546</td>
<td>470–630</td>
<td>355</td>
</tr>
<tr>
<td>S420N</td>
<td>1.8902</td>
<td>520–680</td>
<td>420</td>
</tr>
<tr>
<td>S420NL</td>
<td>1.8912</td>
<td>520–680</td>
<td>420</td>
</tr>
<tr>
<td>S460N</td>
<td>1.8901</td>
<td>550–720</td>
<td>460</td>
</tr>
<tr>
<td>S460NL</td>
<td>1.8903</td>
<td>550–720</td>
<td>460</td>
</tr>
</tbody>
</table>

Notes:
Sub-grade N (normalised or normalised rolled) relates to the physical state of the steel and L (low temperature impact) to high impact resistance.
Data is for thicknesses of 16 mm or less.

S275JR (BS EN 10027-1: 2005)
1.0044 (BS EN 10027-2: 1992)

S275JR  S refers to structural steel.
The yield strength is 275 MPa.
J is the lower impact strength at room temperature R.
### Table 5.3  Steel designations for higher grades to BS EN 10025-4: 2004 (hot rolled products in thermomechanical-rolled weldable fine-grain structural steels)

<table>
<thead>
<tr>
<th>Designation</th>
<th>Number</th>
<th>Ultimate tensile strength (MPa)</th>
<th>Minimum yield strength (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>S275M</td>
<td>1.8818</td>
<td>370–530</td>
<td>275</td>
</tr>
<tr>
<td>S275ML</td>
<td>1.8819</td>
<td>370–530</td>
<td>275</td>
</tr>
<tr>
<td>S355M</td>
<td>1.8823</td>
<td>470–630</td>
<td>355</td>
</tr>
<tr>
<td>S355ML</td>
<td>1.8834</td>
<td>470–630</td>
<td>355</td>
</tr>
<tr>
<td>S420M</td>
<td>1.8825</td>
<td>520–680</td>
<td>420</td>
</tr>
<tr>
<td>S420ML</td>
<td>1.8836</td>
<td>520–680</td>
<td>420</td>
</tr>
<tr>
<td>S460M</td>
<td>1.8827</td>
<td>540–720</td>
<td>460</td>
</tr>
<tr>
<td>S460ML</td>
<td>1.8838</td>
<td>540–720</td>
<td>460</td>
</tr>
</tbody>
</table>

Notes:
Sub-grade M (thermomechanical rolled) relates to the physical state of the steel and L (low temperature impact) to high impact resistance. Data is for thicknesses of 16 mm or less.

### Table 5.4  Steel designations for weather-resistant grades to BS EN 10025-5: 2004

<table>
<thead>
<tr>
<th>Designation</th>
<th>Number</th>
<th>Ultimate tensile strength (MPa)</th>
<th>Minimum yield strength (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>S235JOW</td>
<td>1.8958</td>
<td>360–510</td>
<td>235</td>
</tr>
<tr>
<td>S235J2W</td>
<td>1.8961</td>
<td>360–510</td>
<td>235</td>
</tr>
<tr>
<td>S355JOWP</td>
<td>1.8945</td>
<td>470–630</td>
<td>355</td>
</tr>
<tr>
<td>S355J2WP</td>
<td>1.8946</td>
<td>470–630</td>
<td>355</td>
</tr>
<tr>
<td>S355JOW</td>
<td>1.8959</td>
<td>470–630</td>
<td>355</td>
</tr>
<tr>
<td>S355J2W</td>
<td>1.8965</td>
<td>470–630</td>
<td>355</td>
</tr>
<tr>
<td>S355K2W</td>
<td>1.8966</td>
<td>470–630</td>
<td>355</td>
</tr>
</tbody>
</table>

Notes:
Sub-grades JO, J2 and K2 indicate increasing impact resistance, respectively. Sub-grade W refers to weather-resistant steel. P indicates a high phosphorus grade. Data is for thicknesses of 16 mm or less.

### Table 5.5  Steel designations for high yield strength quenched and tempered steels to BS EN 10025-6: 2004

<table>
<thead>
<tr>
<th>Designation</th>
<th>Number</th>
<th>Ultimate tensile strength (MPa)</th>
<th>Minimum yield strength (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>S460Q</td>
<td>1.8908</td>
<td>550–720</td>
<td>460</td>
</tr>
<tr>
<td>S500Q</td>
<td>1.8924</td>
<td>590–770</td>
<td>500</td>
</tr>
<tr>
<td>S550Q</td>
<td>1.8904</td>
<td>640–820</td>
<td>550</td>
</tr>
<tr>
<td>S620Q</td>
<td>1.8914</td>
<td>700–890</td>
<td>620</td>
</tr>
<tr>
<td>S690Q</td>
<td>1.8931</td>
<td>770–940</td>
<td>690</td>
</tr>
<tr>
<td>S890Q</td>
<td>1.8940</td>
<td>940–1100</td>
<td>890</td>
</tr>
<tr>
<td>S960Q</td>
<td>1.8941</td>
<td>980–1150</td>
<td>960</td>
</tr>
</tbody>
</table>

Notes:
Q indicates quenched steel. Data is for thicknesses between 3 and 50 mm.
Weldable structural steels, as used in the Wembley Stadium, London (Fig. 5.9), have a carbon content within the range 0.16–0.25%. Structural steels are usually normalised by natural cooling in air after hot rolling. The considerable size effect, which causes the larger sections to cool more slowly than the thinner sections, gives rise to significant differences in physical properties; thus, an 80 mm section can typically have a 10% lower yield strength compared to a 16 mm section of the same steel. Whilst grade S275 had previously been considered to be the standard-grade structural steel and is still used for most small beams, flats and angles, the higher grade S355 is increasingly being used for larger beams, columns and hollow sections.

### Hollow sections

Circular, oval, square and rectangular hollow sections are usually made from flat sections which are progressively bent until almost round. They are then passed through a high-frequency induction coil to raise the edges to fusion temperature, when they are forced together to complete the tube. Excess metal is removed from the surface. The whole tube may then be reheated to normalising temperature (850–950°C), and hot rolled into circular, oval, rectangular or square sections. For smaller sizes, tube is heated to 950–1050°C, and stretch reduced to appropriate dimensions. The standard steel grades to BS EN 10210-1: 2006 are S275J2H and S355J2H (Table 5.6). Cold-formed hollow sections differ in material characteristics from the hot-finished sections and conform to BS EN 10219: 2006. The lowest grade S235, with minimum yield strength of 235 MPa is imported, but the standard non-alloy grades are S275 and S355. Grades S420 and S460 are designated as alloy special steels (Table 5.7).

For larger hollow sections, the rotary forge process is used to produce seamless tubes. A hot tapered ingot is pierced by a hydraulic ram, and the central void is opened up by the action of rollers and a rotating mandrel. The steel subsequently passes through a series of eccentric rollers which elongate the tube reducing the section to the required dimensions.

### Bending of structural sections

Castellated beams, rolled, hollow and other sections can be bent into curved forms by specialist metal bending companies. The minimum radius achievable depends on the metallurgical properties, thickness and cross-section. Generally, smaller sections can be curved to smaller radii than the larger sections, although for a given cross-section size the heavier-gauge sections can be bent to smaller radii than the thinner-gauge sections. Normally, universal sections can be bent to tighter radii than hollow sections of the same dimensions. Elegant structures, such as Merchants Bridge, Manchester (Fig. 5.10), can be produced with curved standard sections and also curved tapered beams. The cold bending process work hardens the steel, but without significant loss of performance within the elastic range appropriate to structural steelwork. Tolerances on units can be as low as ±2 mm with multiple bends, reverse curvatures and bends into three dimensions all possible. Increasingly, cold bending is replacing...
Table 5.6  Steel designations for hot finished structural hollow sections to BS EN 10210: 2006 (hot finished structural sections of non-alloy and fine grain structural steels)

<table>
<thead>
<tr>
<th>Designation</th>
<th>Properties</th>
</tr>
</thead>
<tbody>
<tr>
<td>BS EN 10027-1: 2005 and BS EN 10027-2: 1992</td>
<td>BS EN 10210-1: 2006 limits</td>
</tr>
<tr>
<td>Grade</td>
<td>Number</td>
</tr>
<tr>
<td>S235JRH</td>
<td>1.0039</td>
</tr>
<tr>
<td>S275JOH</td>
<td>1.0149</td>
</tr>
<tr>
<td>S275J2H</td>
<td>1.0138</td>
</tr>
<tr>
<td>S355JOH</td>
<td>1.0547</td>
</tr>
<tr>
<td>S355J2H</td>
<td>1.0576</td>
</tr>
<tr>
<td>S355K2H</td>
<td>1.0512</td>
</tr>
<tr>
<td>S275NH</td>
<td>1.0493</td>
</tr>
<tr>
<td>S275NLH</td>
<td>1.0497</td>
</tr>
<tr>
<td>S355NH</td>
<td>1.0539</td>
</tr>
<tr>
<td>S355NLH</td>
<td>1.0549</td>
</tr>
<tr>
<td>S420NH</td>
<td>1.8750</td>
</tr>
<tr>
<td>S420NLH</td>
<td>1.8751</td>
</tr>
<tr>
<td>S460NH</td>
<td>1.8953</td>
</tr>
<tr>
<td>S460NLH</td>
<td>1.8956</td>
</tr>
</tbody>
</table>

Notes:
H refers to hollow sections.
Sub-grades JR, JO and J2 indicate impact resistance at room temperature, 0 and $-20^\circ C$, respectively.
K2 refers to higher impact energy than J2.
Sub-grade N (normalised or normalised rolled) relates to the physical state of the steel and L (low temperature impact) to high impact resistance.
The standard UK production grades are the S275J2H and S355J2H designations.
Data is for thicknesses between 3 and 16 mm.

induction or hot bending which requires subsequent heat treatment to regain the initial steel properties.

**Cast sections**

Components such as the nodes for rectangular and circular hollow-section constructions and large pin-joint units are manufactured directly as individual castings. They may then be welded to the standard milled steel sections to give continuity of structure. Structural steels appropriate for casting are listed in the standard BS EN 10340: 2007. Alloy steels for casting may be subjected to quenching and tempering, normalising or annealing according to the specific component requirements.

**Steel frame construction**

The use of light steel frame construction is an alternative to timber frame technology. Most light framing is manufactured from cold-formed galvanised steel channel sections, factory assembled, ready for subsequent bolting together on site. Panel units are typically 600 mm wide, storey-height modules, lined with a vapour check layer and plasterboard. This modern method of construction (MMC) considerably reduces the time on site compared to building traditional load-bearing masonry.

**BI-STEEL**

*Bi-steel* panels consist of two steel plates held apart by an array of welded steel bar connectors (Fig. 5.11). The panels are usually assembled into larger modules for delivery to site, where they are erected and the void space filled with concrete. The combination of permanent steel formwork and concrete fill acts as reinforced concrete, with the steel providing resistance to in-plane and bending forces and the concrete offering resistance to compression and shear. Units are manufactured up to 2 m wide and 18 m long in S275 or S355 steel to thicknesses between 200 and 700 mm and may be flat or curved. Adjacent panels may be bolted or fixed with proprietary connectors, giving fast erection times on site.
Table 5.7  Steel designations for cold-formed structural hollow sections to BS EN 10219-1: 2006 (cold-formed welded structural hollow sections of non-alloy and fine grain steels)

<table>
<thead>
<tr>
<th>Designation</th>
<th>Grade</th>
<th>Number</th>
<th>Ultimate tensile strength (MPa)</th>
<th>Minimum yield strength (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>S275JOH</td>
<td>1.0149</td>
<td>410–560</td>
<td>275</td>
</tr>
<tr>
<td></td>
<td>S275J2H</td>
<td>1.0138</td>
<td>410–560</td>
<td>275</td>
</tr>
<tr>
<td></td>
<td>S355JOH</td>
<td>1.0547</td>
<td>470–630</td>
<td>355</td>
</tr>
<tr>
<td></td>
<td>S355J2H</td>
<td>1.0576</td>
<td>470–630</td>
<td>355</td>
</tr>
<tr>
<td></td>
<td>S355K2H</td>
<td>1.0512</td>
<td>470–630</td>
<td>355</td>
</tr>
<tr>
<td></td>
<td>S275NH</td>
<td>1.0493</td>
<td>370–510</td>
<td>275</td>
</tr>
<tr>
<td></td>
<td>S275NLH</td>
<td>1.0497</td>
<td>370–510</td>
<td>275</td>
</tr>
<tr>
<td></td>
<td>S355NH</td>
<td>1.0539</td>
<td>470–630</td>
<td>355</td>
</tr>
<tr>
<td></td>
<td>S355NLH</td>
<td>1.0549</td>
<td>470–630</td>
<td>355</td>
</tr>
<tr>
<td></td>
<td>S460NH</td>
<td>1.8953</td>
<td>540–720</td>
<td>460</td>
</tr>
<tr>
<td></td>
<td>S460NLH</td>
<td>1.8956</td>
<td>540–720</td>
<td>460</td>
</tr>
<tr>
<td></td>
<td>S275MH</td>
<td>1.8843</td>
<td>360–510</td>
<td>275</td>
</tr>
<tr>
<td></td>
<td>S275MLH</td>
<td>1.8844</td>
<td>360–510</td>
<td>275</td>
</tr>
<tr>
<td></td>
<td>S355MH</td>
<td>1.8845</td>
<td>450–610</td>
<td>355</td>
</tr>
<tr>
<td></td>
<td>S355MLH</td>
<td>1.8846</td>
<td>450–610</td>
<td>355</td>
</tr>
<tr>
<td></td>
<td>S420MH</td>
<td>1.8847</td>
<td>500–660</td>
<td>420</td>
</tr>
<tr>
<td></td>
<td>S420MLH</td>
<td>1.8848</td>
<td>500–660</td>
<td>420</td>
</tr>
<tr>
<td></td>
<td>S460MH</td>
<td>1.8849</td>
<td>530–720</td>
<td>460</td>
</tr>
<tr>
<td></td>
<td>S460MLH</td>
<td>1.8850</td>
<td>530–720</td>
<td>460</td>
</tr>
</tbody>
</table>

Notes:
H refers to hollow sections.
Sub-grades JR, JO and J2 indicate impact resistance at room temperature, 0 and –20°C, respectively.
K2 refers to higher impact energy than J2.
Sub-grades M (thermomechanical rolled) and N (normalised or normalised rolled) relate to the physical state of the steel and L (low temperature impact) to high impact resistance.
The standard UK production grades are the S275J2H and S355J2H designations.
Data is for thicknesses between 3 mm and 16 mm.

FIRE PROTECTION OF STRUCTURAL STEEL

The two approaches to the design of fire precautions in buildings rely on periods of fire resistance as described within the Building Regulations Approved Document B and in BS 9999: 2008, or according to a risk profile based on occupancy, fire growth rate, ventilation conditions and building geometry also described in BS 9999: 2008. This standard includes consideration of more complex spaces such as auditoria and atria as well as less sophisticated building configurations.

The fire protection of structural steel may be approached either by the traditional method involving the application of insulation materials with standard fire resistance periods (Fig. 5.12) or by a structural fire engineering method, which predicts the potential rate of rise of temperature of exposed steel members in each situation, based on the calculated fire load and particular exposure of the steel.

Applied protection to structural steel

Intumescent coatings
Thin film intumescent coatings, which do not seriously affect the aesthetic of exposed structural steelwork, offer up to 120 minutes' fire protection. A full colour range for application by spray, brush or roller can be used on steel and also for remedial work on old cast
iron or wrought iron structures. The three standards of finish usually specified are basic, normal decorative finish and high decorative finish. The normal finish has a slight orange peel effect, whereas the high decorative finish is smooth and even.

*Sprayed coatings*
Sprayed coatings based on either vermiculite cement or mineral-fibre cement may be applied directly to steel to give up to 240 minutes’ fire protection. The process is particularly appropriate for structural steel in ceiling voids, where the over-spray onto other materials is less critical. The finish, which can be adjusted to the required thickness, is heavily textured, and the products are relatively cheap.

*Boarded systems*
Lightweight boards boxed around steel sections offer between 30 and 240 minutes’ fire protection according to their thicknesses. Products are generally based on vermiculite or mineral fibres within cement, calcium silicate or gypsum binders. Boarded systems are screwed directly to the structural steel, to light-gauge steel fixings or to a box configuration. Precoated
products are available, or the standard systems may be subsequently decorated.

Preformed casings
Preformed sheet-steel casings, which encase lightweight vermiculite plaster, give a high-quality appearance and up to 240 minutes’ fire resistance. The calculated fire resistance is based solely on the thickness of insulation and does not take into account any additional protection afforded by the sheet steel.

Masonry and concrete
Structural steel may be fully encased with masonry or suitably reinforced lightweight concrete in which non-spalling aggregates should be used. Hollow steel columns may be filled with plain, fibre-reinforced or bar-reinforced concrete to give up to 120 minutes’ fire resistance. For plain or fibre-reinforced concrete a minimum section of $140 \times 140$ mm or $100 \times 200$ mm is required and $200 \times 200$ mm or $150 \times 250$ mm for bar-reinforced concrete filling.

Water-filled systems
Interconnecting hollow steel sections can be given fire protection by filling with water as part of a gravity feed or pumped system. Water loss is automatically replaced from a tank, where corrosion inhibitor and anti-freeze agents are added to the system as appropriate.

Fire engineering
The heating rate of a structural steel section within a fire depends upon the severity of the fire and the degree of exposure of the steel. Where a steel section has a low surface/cross-sectional area (Hp/A) ratio (Fig. 5.13), its temperature will rise at a slower rate than a section with a high Hp/A ratio. Fire-engineered solutions calculate the severity of a potential fire based on the enclosure fire loads, ventilation rates and thermal characteristics, and then predict temperature rises within the structural steel based on exposure. The stability of the structural member can therefore be predicted,

Fig. 5.11 Bi-steel unit. Photograph: Courtesy of Corus

Fig. 5.12 Structural steelwork, typical fire-protection systems
Fig. 5.13 Hp/A ratios and rates of heating in fire

taking into consideration its steel grade, loading and any structural restraint. From these calculations it can be determined whether additional fire protection is required and at what level to give the required fire resistance period.

Depending on the particular circumstances, a fully loaded unprotected column with a section factor (Hp/A) of less than $50 \text{ m}^{-1}$ may offer 30 minutes' structural fire resistance; similarly, lighter columns with lightweight concrete blocks in the web can achieve 30 minutes' fire resistance. Shelf angle floors of suitable section, and in which a high proportion of the steel is encased by the concrete floor construction, can achieve 60 minutes' fire resistance (Fig. 5.14).

**PROFILED STEEL SHEETING**

The majority of profiled sheet steel is produced by shaping the precoated strip through a set of rollers, which gradually produce the desired section without damage to the applied coating. The continuous profiled sheet is then cut and packaged to customer requirements. The standard sections have a regular trapezoidal profile, with the depth of the section dependent on the loading and required span (Fig. 5.15). In cases where there is the risk of buckling, stiffeners are incorporated into the profile. Curved profiled sheets for eaves and soffits are manufactured by brake pressing from the same coated strip. Trapezoidal profiles may be crimped in this process, although sinusoidal sheets and shallow trapezoidal sections can be curved without this effect. The rigidity of curved sections reduces their flexibility and thus the tolerances of these components. Proprietary spring-clip fixings may be used when concealed fixings are required for

Fig. 5.14 Fire resistance of structural-steel systems
certain profiled sheet sections. The standard BS EN 14782: 2006 requires a minimum thickness of 0.4 mm (excluding any coating) for self-supporting profiled sheet steel in construction.

**STEEL CABLES**

Steel cables are manufactured by drawing annealed thin steel rod through a series of lubricated and tapered tungsten carbide dies, producing up to a 10-fold elongation. The drawing process increases the strength and reduces the ductility of the steel; thus, the higher carbon steels, required for the production of high tensile wires, need special heat treatment before they are sufficiently ductile for the sequence of drawing processes.

In order to manufacture steel cables for suspended structures or prestressed concrete, a set of individual wires are twisted into a strand, then a series of strands are woven around a central core of steel or fibre strand to produce a rope. A series of ropes are then woven to produce cable to the required specifications. Typical configurations use 7 or 19 strands or ropes to form heavy duty cables, frequently in grade 1.4401 stainless steel for exposed structures.

**PERFORATED STEEL SHEETS, STEEL MESHES AND NETS**

Perforated steel sheets are manufactured in mild steel, galvanised steel and stainless steel for use in architectural features, sunshades, balustrades, and also wall and ceiling panels. Perforated sheets, also available in aluminium, copper, brass and bronze, may have round, square or slotted holes within a wide range of sizes and spacings to produce the desired aesthetic effect. Metal sheets are either punched or plasma profile cut.

Stainless steel meshes are available as flexible or rigid self-supporting weaves, each in a wide variety of patterns offering choice for use as external facades and sun-screening, also internal space dividers, balustrades, wall coverings and suspended ceilings. Patterns range from traditional weave and expanded metal to chain mail, with a wide variation in texture and transparency. Some patterns are available in mild steel and also in non-ferrous metals. Stainless steel netting can be used creatively to form open tent and canopy structures.

**Ferrous alloys**

**WEATHERING STEEL**

Weathering steels are structural steels which have been alloyed with small proportions of copper, usually between 0.25% and 0.55%, together with silicon, manganese, chromium and either vanadium or phosphorus as minor constituents (BS 7668: 2004 and BS EN 10025-5: 2004). The alloying has the effect of making the naturally formed brown rust coating adhere tenaciously to the surface, thus preventing further loss by spalling as in Fig. 5.16. The use of weathering steels is not appropriate within marine environments, and all weathering steel must be carefully detailed to ensure that the rainwater run-off does not impinge on other materials, particularly concrete or glass where it will cause severe staining during the first few years of exposure to the elements. Corten is the commercial name for weathering steels. Table 5.4 (page 173) gives the steel specification and steel number to the European Standards. Weathering steels are used for structural
and cladding applications also for sculptural works of art.

**STAINLESS STEELS**

Stainless steels are a range of alloys containing at least 10.5% chromium. The corrosion resistance of the material is due to the natural passive film of chromium oxide which immediately forms over the material in the presence of oxygen; thus, if the surface is subsequently scratched or damaged the protective film naturally reforms. The corrosion resistance is increased by the inclusion of nickel and molybdenum as additional alloying components. The standard (austenitic) grades used within construction are 18% chromium, 10% nickel (1.4301) and 17% chromium, 12% nickel, 2.5% molybdenum (1.4401). The 18/10 alloy is suitable for use in rural and lightly polluted urban sites, while the 17/12/2.5 higher specification alloy is more appropriate for use within normal urban, marine and industrial environments. For certain aggressive environments, the high-alloy (duplex) stainless steel (number 1.4462) should be used. Ferritic stainless steel (1.4016) containing only chromium, with a reduced corrosion resistance, is appropriate for internal building use where corrosion is a less critical factor. Standard grades to BS EN 10088-1: 2005 for stainless steels are given in Table 5.8. Extended listings of grades for construction are detailed in BS EN 10088-4: 2009 for flat products and in BS EN 10088-5: 2009 for sections, bars and rods.

Stainless steel is manufactured by a three-stage process. Scrap is melted in an electric arc furnace, then refined in an argon–oxygen decarburiser and alloyed to the required composition in a ladle furnace by the addition of the minor constituents. Most molten metal is continuously cast into billets or slabs for subsequent forming. Stainless steel is hot rolled into plate, bar and sheet, while thin sections may be cold rolled. Heavy universal sections are made up from plates. Stainless steel may be cast or welded and is readily formed into small components such as fixings and architectural ironmongery. Polished, brushed, matt, patterned and profiled finishes are available; additionally, the natural oxide film may be permanently coloured by chemical and cathodic treatment to bronze, blue, gold, red, purple or green according to its final thickness. Approximately 50% of stainless steel derives from recycled scrap steel.

Stainless steel is available in square, rectangular, oval and circular hollow sections as well as the standard sections for structural work. Its durability is illustrated by the Lloyd’s Building (Fig. 5.17), which maintains its high-quality finish within the urban context of the City of London. Stainless steel is widely used for roofing, cladding, interior and exterior trim owing to its combined strength, low maintenance and visual impact (Fig. 5.18). Stainless steels for self-supporting profile sheet and tile roofing systems are listed in BS EN 508-3: 2008. A minimum thickness of 0.4 mm is specified

### Table 5.8  Stainless steel compositions and grades to BS EN 10088-1: 2005 for different environmental conditions

<table>
<thead>
<tr>
<th>Designation</th>
<th>Name (indicating composition of alloying components)</th>
<th>Suitable environments</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Type</strong></td>
<td></td>
<td><strong>Number</strong></td>
</tr>
<tr>
<td>Austenitic</td>
<td>X5CrNi18-10</td>
<td>1.4301</td>
</tr>
<tr>
<td></td>
<td>X5CrNiMo17-12-2</td>
<td>1.4401</td>
</tr>
<tr>
<td>Ferritic</td>
<td>X6G17</td>
<td>1.4016</td>
</tr>
<tr>
<td>Duplex</td>
<td>X2CrNiMoN22-5-3</td>
<td>1.4462</td>
</tr>
</tbody>
</table>

Notes:
- Cr, Ni, Mo and N refer to chromium, nickel, molybdenum and nitrogen, respectively.
- X2, X5 and X6 refer to the carbon contents of 0.02, 0.05 and 0.06%, respectively.
in BS EN 14782: 2006. The standard also lists optional organic finishes including polyester, silicone-modified polyester, polyurethane, polyvinylidene fluoride and PVC plastisol. The corrosion resistance of stainless steel also makes it eminently suitable for masonry fixings, such as corbels, anchor bolts, cavity wall ties and for concrete reinforcement. Austenitic stainless steels are used for the manufacture of pipework, catering and drainage products where durability and corrosion resistance are critical. Exposed exterior stainless steel should be washed regularly to retain its surface characteristics. Pitting corrosion causing surface pin-point attack, crevice corrosion under tight-fitting washers and stress corrosion cracking, where the material is under high tensile load, may occur where inappropriate grades are used in aggressive environments.

High-performance (superduplex) stainless steel (1.4507) wire ropes, which have 50% more tensile strength than standard (austenitic) stainless steel (1.4401), are appropriate for architectural support and restraint systems. These alloys containing around 1.6% copper have a lower elastic stretch than standard stainless steels and a higher fatigue resistance, which makes them appropriate for architectural tensile elements including those within marine and swimming pool environments.

In addition to organic coatings, stainless steel may be coated with terne (see terne-coated steel) or electroplated with ≥10 g/m² tin to BS EN 508-3: 2008.

**HEAT-TREATED STEELS**

The size effect, which causes a reduction in yield strength in large sections due to their slower cooling rates than the equivalent thin sections, can be ameliorated by the addition of small quantities of alloying elements such as chromium, manganese, molybdenum and nickel.

**Coated steels**

To inhibit corrosion, steel may be coated with metallic or organic finishes. Metallic finishes are typically zinc (Z), zinc–iron alloy (ZF), zinc–aluminium alloy (ZA), aluminium–zinc alloy (AZ), aluminium–silicon–alloy (AS) and aluminium (A) all of which may be applied by hot-dipping of steel into the molten metal (Table 5.9). These metallic coatings, except for pure aluminium, are covered by the standard BS EN 10346: 2009. The standard BS EN ISO 14713: 1999 indicates an
Table 5.9  Typical examples for zinc- and aluminium-based coatings on steel

<table>
<thead>
<tr>
<th>Coating</th>
<th>Designation</th>
<th>Coating mass (g/m²) (total mass)</th>
<th>Nominal thickness (μm) (on each face)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zinc</td>
<td>Z275</td>
<td>275</td>
<td>20</td>
</tr>
<tr>
<td>Zinc–iron</td>
<td>ZF120</td>
<td>120</td>
<td>8</td>
</tr>
<tr>
<td>Zinc–aluminium</td>
<td>ZA255</td>
<td>255</td>
<td>20</td>
</tr>
<tr>
<td>Aluminium–zinc</td>
<td>AZ185</td>
<td>185</td>
<td>25</td>
</tr>
<tr>
<td>Aluminium–silicon</td>
<td>AS100</td>
<td>100</td>
<td>17</td>
</tr>
</tbody>
</table>

Note:
The standard BS EN 10346: 2009 gives a range of coating thicknesses for each of the coatings.

anticipated annual corrosion loss of between 0.7 and 2 μm of zinc per year from galvanised steel in most UK locations. Organic coatings (BS EN 10169: 2003) may be divided into liquid paints, powder coatings and films. Certain products are suitable only for interior applications.

ZINC-COATED STEEL

The zinc coating of steel has for many years been a standard method for its protection against corrosion. The zinc coating may be applied by hot-dipping or spraying with the molten metal, sherardising in heated zinc powder (BS EN 13811: 2003) or electrolytically (ZE) to BS EN 10152: 2009. In hot-dip galvanising the steel is cleaned by pickling in acid followed by immersion in molten zinc at 450°C. The zinc coating (Z) protects the steel by acting as a physical barrier between the steel and its environment, and also by sacrificially protecting the steel where it is exposed by cutting or surface damage. If annealed, the pure zinc coating becomes alloyed from the substrate steel producing an alloy of 8–12% iron (ZF) which gives a better surface for painting or welding.

The durability of the coated steel is dependent on the thickness of the coating and the environment. Coating designations to BS EN 10346: 2009 (Table 5.9) indicate the total mass and nominal thickness of applied metallic coatings.

Coastal situations and industrial environments with high concentrations of salt and sulphur dioxide, respectively, may cause rapid deterioration. The alkalies in wet cement, mortar and plaster etch zinc coatings, but once dry, corrosion is slow; however, calcium chloride used as an accelerator in plaster is aggressive and should only be used sparingly. Fixings for zinc-coated sheet should be carefully chosen to avoid the formation of bimetallic couples, which can cause accelerated corrosion. In particular, no copper or brass should make contact with either zinc or iron–zinc alloy coated steel. Other metals, such as lead, aluminium and stainless steel, have less serious effects in clean atmospheres, but generally all fixings should be sealed and insulated by rubber-faced washers. Where zinc-coated steel is to be fixed to unseasoned timber or timber impregnated with copper-based preservatives, the wood should be coated with bitumen paint. Where damaged in cutting, fixing or welding, the zinc coating should be repaired with the application of zinc-rich paint.

Zinc-coated steel may be painted for decoration or improved corrosion resistance. However, the normal spangle zinc finish will show through paint and the minimised spangle or iron–zinc alloy uniform matt grey finish is more appropriate for subsequent painting.

ALUMINIUM–ZINC ALLOY COATED STEEL

The two distinct zinc/aluminium alloys used for coating steel contain 5% (ZA) and 55% (AZ) aluminium. The ZA coating to BS ISO 14788: 2005 is usually a substrate for an organic finish. For profile sheet metal roofing to BS EN 508-1: 2008, the minimum coating mass totals are 255 and 185 g/m², respectively, compared to 350 g/m² for pure zinc. Steel coated with the AZ alloy of aluminium (55%), zinc (43.4%) and silicon (1.6%) is more durable than that coated with an equivalent thickness of pure zinc, and may be used without further protection in non-aggressive environments. It is also used as the substrate for certain organic coatings. The finish is a metallic lustre due to crystal formation.

ALUMINIUM-COATED STEEL

Hot dip aluminium on steel for profiled roofing sheet or tile to BS EN 508-1: 2008 may be applied to three grades with total mass levels of 195, 230 or 305 g/m². However, the 195 g/m² product is only suitable for subsequent organic coating. Usually, an alloy containing 8–11% silicon is used, which produces a durable alloy finish of aluminium, iron and silicon.

TERNE-COATED AND LEAD-CLAD STEEL

Lead and terne, an alloy of lead (80–90%) and tin (20–10%), are used as finishes to steel and stainless steel for cladding and roofing units. Terne can be
applied to sheet stainless steel as a 20 μm layer by immersion in the molten alloy. Terne-coated stainless steel does not suffer from bimetallic corrosion and can normally be used in contact with lead, copper, aluminium or zinc. Thermal movement is similar to stainless steel, allowing for units up to 9 m in length to be used for roofing and cladding. The composite material, lead-clad steel, is produced by cold-roll bonding 0.75 mm lead to 1.0 mm terne-plated steel or to 0.8 mm terne-plated stainless steel. Lead-clad steel is suitable for cladding and roofing systems and has the appearance and corrosion resistance of milled lead. Because of the support afforded by the steel substrate, lead-clad steel or stainless steel can be used for self-supported fascias, soffits, gutters and curved sections. Joints can be lead burned and soldered as traditional lead. Cut ends should be protected by soldering in the case of lead-clad steel, although the stainless steel version requires no protective treatment. Unlike traditional lead, the material is virtually theft-proof and does not suffer significantly from creep. Patination oil should be applied to the lead surface after installation to prevent staining effects. Details of the various steel and coating grades are listed in BS ISO 4999: 2005.

ORGANIC COATED STEEL

Since the 1960s, a range of heat-bonded organic coatings for steel has been developed, including PVC plastisol (Colourcoat), polyvinylidene fluoride (PVDF), polyesters and PVC film (Stelvetite). Within this product range the PVC plastisol currently has the largest market share for cladding and roofing within the UK.

PVC plastisol coating

PVC plastisol is applied to zinc or aluminium/zinc coated steel to a thickness of 0.2 mm. It has a tough leather grain finish and is available in a wide range of colours, although the pastel shades are recommended for roofing applications. The reverse side is usually coated with a grey corrosion-resistant primer and polyester finish, although PVC plastisol may be specified for unusually aggressive internal environments. Careful site storage and handling is required to prevent physical damage to the surface. For non-marine environments the most durable colours will give a period to first maintenance of greater than 20 years. Very deep colours, and the non-pastel shades in coastal locations, will have reduced periods to first repainting.

Polyvinylidene fluoride coating

Polyvinylidene fluoride (PVDF), an inert fluoro-carbon, when applied as a 0.027 mm coating to zinc-coated steel, has good colour stability at temperatures up to 120°C, making it suitable for world-wide use and for buildings which are likely to be extended at a later date. The finish is smooth and self-cleaning, although considerable care is required on site to prevent handling damage. A period to first maintenance of 15 years is typical within the UK for non-coastal locations. The wide colour range includes metallic silver. Polyvinylidene fluoride finished zinc/tin (terne) coated steel or stainless steel is also available in a range of colour finishes which include copper, copper patina and stainless steel. The material can be used for cladding and roofing; the solar reflective paint finish reduces excessive solar gain.

Polyester coating

Polyester and silicone polyester coated galvanised steels are economic products, but offer only medium-term life in non-aggressive environments. Externally, the period to first maintenance will be typically 10 years in unpolluted inland locations, but they are suitable for internal use. Silicone polyester should not be used in marine or hot, humid environments. Polyester and silicone polyester coatings are smooth and typically 0.025 mm in thickness.

Polyurethane coating

Polyurethane is applied as a 0.05 mm coating to zinc or zinc/aluminium alloy coated steel. It is available in a range of solid and metallic colours with a 25 year period to first maintenance on zinc/aluminium-coated steel for roofs. A textured finish on zinc-coated steel with a 20 year period to first maintenance is available for exterior walls in non-marine environments.

Enamel coating

Organic enamel-coated steels offering good light reflectance are suitable for internal use as wall and roof linings. Coatings, usually 0.022 mm thick, are typically applied to hot-dip zinc/aluminium alloy-coated steel and are easily cleaned. The standard colour is brilliant white, but a range of light colours is also available.
PVC film coating

PVC film (0.02 mm) in a range of colours, decorative patterns and textured finishes is calendered to zinc-coated steel strip. The product is suitable only for internal applications.

Paints

The wide range of paints used to protect steel from corrosion fall into several categories listed in BS EN ISO 12944-5: 2007. Paints including solvent-borne chlorinated rubber, acrylic polymers and vinyl chloride copolymers dry by solvent evaporation which can be reversed by solvent action. The majority of other products dry irreversibly by a loss of solvent or coalescence from the water emulsion. Alkyds, urethane alkyds and epoxy esters dry by a combination of a loss of solvent and aerial oxidation. Water-borne acrylics, vinyls and urethanes coalesce as the water evaporates. Two-component paints such as epoxys and polyurethanes set through chemical processes activated on mixing.

Paint is normally applied in three coats. The primer adheres to the cleaned surface and gives corrosion protection. The second coat builds up thickness and the finishing coat provides protection from the environment and the required aesthetic finish. The standard BS EN ISO 12944: 2007 gives guidelines on the suitability of a range of paint systems in relation to environmental conditions and durability.

Steel tiles and slates

Lightweight steel tile and slate units, manufactured from galvanised or aluminium–zinc alloy hot-dipped steel, coated with acrylic or polyester resin and a granular finish, give the appearance of traditional slate or pantile roofs. The products have the advantage, particularly for refurbishment work, of lightness in comparison to the traditional materials. Units can typically be used for roof pitches between 12° and 90°. A span of 1200 mm allows for wider spacing of roof trussed rafters. Units in a range of traditional material colours are available with appropriate edge and ventilation accessories. The standard BS EN 14782: 2005 requires a minimum thickness of 0.4 mm (excluding any coating) for self-supporting sheet steel in construction, but a typical 0.7 mm polyester coated galvanised sheet steel system will span 1 m.

Aluminium

Aluminium has only been available as a construction material for about a hundred years. Possibly the most well-known early use of the metal was for the cast statue of Eros, which has stood in Piccadilly Circus, London, since 1893. Because of its durability, it is used widely in construction, particularly for secondary components as illustrated in the permanent shading devices on the Faculty of Divinity building of the University of Cambridge (Fig. 5.19). Aerofoil-shaped shading systems manufactured from extruded aluminium, either fixed or linked to an active building management system, are standard features in many new buildings.

MANUFACTURE

Aluminium, the most common metallic element in the earth’s crust, is extracted from the ore bauxite, an impure form of aluminium oxide or alumina. Large reserves of bauxite are available in West Africa, Australia and the West Indies. The bauxite is dissolved in caustic soda, filtered, reprecipitated to remove impurities and dried. The pure alumina is then dissolved in fused cryolite (sodium aluminium fluoride) within a carbon-lined electrolytic cell. Electrolysis of the aluminium oxide produces oxygen and the pure aluminium, which is tapped off periodically and cast. The process is highly energy intensive, and typically the production of 1 tonne of aluminium requires 15,000 kWh of electrical energy. Globally, 50% of this energy is from renewable hydroelectric power. Currently, 63% of new aluminium used in the UK is from recycled sources, and recycling requires only 5% of the energy input compared to primary production. Recycling from building construction in Europe is approximately 95%. Cast ingots or slabs are hot rolled at 500°C into 5 mm coiled sheet, which subsequently can be cold rolled into thinner sheet or foil. Due to the ductility of aluminium, the metal can be extruded into complex shapes or drawn into wire. Forming and machining processes are generally easier than with steel. Aluminium components may also be formed by casting.

PROPERTIES

Aluminium is one of the lightest metals with a density of 2700 kg/m³ compared to steel 7900 kg/m³.
Standard-grade aluminium (99% pure) has a tensile strength between 70 and 140 MPa, depending on temper; however, certain structural aluminium alloys (e.g. alloy 5083) achieve 345 MPa comparable to the 410–560 MPa for S275 steel. This compares favourably on a strength-to-weight basis, but the modulus of elasticity for aluminium is only one third that of steel, so deflections will be greater unless deeper sections are used. For an aluminium section to have the same stiffness as an equivalent steel member, the aluminium section must be enlarged to approximately half the weight of the steel section.

**DURABILITY**

The durability of aluminium as a construction material is due to the protection afforded by natural oxide film, which is always present on the surface of the metal. The aluminium oxide film, which is immediately produced when the surface of the metal is cut or scratched, is naturally only 0.01 μm thick, but may be thickened by the process of anodisation.

**FIRE**

The strength of aluminium is halved from its ambient value at a temperature of 200°C, and for many of the alloys is minimal by 300°C.

**CONTACT WITH OTHER BUILDING MATERIALS**

Whilst dry cement-based materials do not attack aluminium, the alkalinity of wet cement, concrete and mortar causes rapid corrosion. Thus, where these materials make contact during the construction process, the metal should be protected by a coating of bitumen paint. Furthermore, anodised and particularly coloured sections, such as glazing units, can be permanently damaged by droplets of wet cement.
products, and should be protected on site by a removable lacquer or plastic film. Under dry conditions aluminium is unaffected by contact with timber; however, certain timber preservatives, particularly those containing copper compounds, may cause corrosion under conditions of high humidity. Where this risk is present the metal should be protected with a coating of bitumen.

Although aluminium is highly resistant to corrosion in isolation, it can be seriously affected by corrosion when in contact with other metals. The most serious effects occur with copper and copper-based alloys, and rainwater must not flow from a copper roof or copper pipes into contact with aluminium. Except in marine and industrial environments it is safe to use stainless steel fixings or lead with aluminium, although zinc and zinc-coated steel fixings are more durable. Unprotected mild steel should not be in electrical contact with aluminium.

**ALUMINIUM ALLOYS**

Aluminium alloys fall into two major categories, either cast or wrought. Additionally, the wrought alloys may be subjected to heat treatment. The majority of aluminium used in the construction industry is wrought, the content and degree of alloying components being directly related to the physical properties required, with the pure metal being the most malleable. BS EN 573-1: 2004 designates aluminium alloys into categories according to their major alloying components (Table 5.10).

The following example illustrates the coding system for aluminium and its alloys:

**Table 5.10**  Broad classification of aluminium alloys to BS EN 573-1: 2004

<table>
<thead>
<tr>
<th>Alloy series</th>
<th>Major alloying components</th>
</tr>
</thead>
<tbody>
<tr>
<td>1000</td>
<td>Greater than 99% aluminium</td>
</tr>
<tr>
<td>2000</td>
<td>Copper alloys</td>
</tr>
<tr>
<td>3000</td>
<td>Manganese alloys</td>
</tr>
<tr>
<td>4000</td>
<td>Silicon alloys</td>
</tr>
<tr>
<td>5000</td>
<td>Magnesium alloys</td>
</tr>
<tr>
<td>6000</td>
<td>Magnesium and silicon alloys</td>
</tr>
<tr>
<td>7000</td>
<td>Zinc alloys</td>
</tr>
<tr>
<td>8000</td>
<td>Other elements</td>
</tr>
</tbody>
</table>

Note:
In many cases minor alloying components are also present.

Structural aluminium is alloy EN AW-6082 where EN refers to the European Norm, A is for aluminium and W is for wrought products. 6082 is within the 6000 series of magnesium and silicon alloys, and the final three digits refer to the exact chemical composition as listed in BS EN 573-3: 2009.

For flashings where on-site work is necessary, 99.8% pure aluminium (alloy EN AW-1080A) or 99.5% (EN AW-1050A) offer the greatest malleability, although the standard commercial grade 99% pure aluminium (alloy EN AW-1200) is suitable for insulating foils and for continuously supported sheet roofing. The minimum nominal thickness for fully supported aluminium sheet roofing and cladding is 0.6 mm to BS EN 14783: 2006.

Profiled aluminium for roofing and cladding, requiring additional strength and durability is alloyed with 1.25% manganese (alloy EN AW-3103). It is produced from the sheet by roll forming, and can be manufactured into curved sections to increase design flexibility. Preformed rigid flashings to match the profile sheet are manufactured from the same alloy and finish. The alloy with 2% magnesium (alloy EN AW-5251) is more resistant to marine environments. The standard BS EN 508-2: 2008 details the full range of alloys for self-supporting roofing systems and illustrates examples of profiles applicable to aluminium sheets and tiles.

Aluminium rainscreen cladding panels up to $2.8 \times 1.5$ m in size, may also be shaped using the superplastic forming (SPF) process, which relies on the high extensibility of the alloy EN AW-5083SPF. Sheet alloy, typically 2 mm in thickness, is heated to $380–500^\circ$C, and forced by air pressure into the three-dimensional form of the mould. Horizontal or vertical ribs are frequently manufactured to give enhanced rigidity, but cladding panels may be formed to individual designs including curvature in two directions. Coloured finishes are usually polyester powder or polyvinylidene fluoride coatings.

Extruded sections for curtain walling, doors and windows require the additional strength imparted by alloying the aluminium with magnesium and silicon (alloy EN AW-6063). Thermal insulation within such extruded sections is achieved by a hidden thermal break or by an internal plastic or timber insulating cladding.

Structural aluminium for load-bearing sections and space frames typically contains magnesium, silicon and manganese (alloy EN AW-6082). Tempering
increases the tensile strength to the range 270–310 MPa, which is more comparable to the standard grade of structural steel S275 (minimum tensile strength 410 MPa).

**FINISHES FOR ALUMINIUM**

**Anodising**

The process of anodising thickens the natural aluminium oxide film to typically 10–25\(\mu\)m. The component is immersed in sulphuric acid and electrolytically made anodic, which converts the surface metal into a porous aluminium oxide film, which is then sealed by boiling in water. The anodising process increases durability and can be used for trapping dyes within the surface to produce a wide range of coloured products. Some dyes fade with exposure to sunlight, the most durable colours being gold, blue, red and black. Exact colour matching for replacement or extensions to existing buildings may be difficult, and manufacturers will normally produce components within an agreed band of colour variation. If inorganic salts of tin are incorporated into the surface during the anodising process then colour-fast bronzes are produced. Depending on the period of exposure to the electrolytic anodisation process, a range of colours from pale bronze to black may be produced. Different aluminium alloys respond differently to the anodising treatment. Pure aluminium produces a silver mirror finish, whereas the aluminium–silicon alloys (e.g. alloy EN AW-6063) produce a grey finish.

**Surface textures**

A range of surface textures is achieved by mechanical and chemical processing. Finishes include bright polished, matt, etched and pattern-rolled according to the pretreatments applied, usually before anodising, and also the particular alloy used. The aluminium discs on the facade of the Selfridges building in Birmingham give an innovative decorative finish to the steel-frame store, forming an elegant contrast to the blue painted rendered surface (Fig. 5.20).

**Coatings**

**Metallic**

Zinc-coated aluminium is a foldable cladding material combining the durability of aluminium with the appearance of pre-weathered zinc. Matching rainwater goods including half-round gutters and down pipes are available. An equivalent pre-patinated or pre-oxidised titanium-coated aluminium with the appearance of bright steel is under development.

**Organic**

Polyester coatings, predominantly white, but with a wide range of colour options, are used for double glazing systems, cladding panels and rainwater goods. Electrostatically applied polyester powder is heat-cured to a smooth self-cleaning finish. PVC and PVDF simulated wood-grain and other pattern film finishes may also be applied to aluminium extrusions and curtain-wall systems. The range of factory-applied organic coatings, including polyester, acrylic, polyvinylidene fluoride (PVDF), alkyd and polyurethane, suitable for self-supporting roofing systems including tiles is described in BS EN 508-2: 2008.

**Paint**

Where aluminium is painted for decorative purposes it is important that the appropriate primer is used. The aluminium should be abraded or etched to give a good key to the paint system, although cast aluminium normally has a sufficiently rough surface. Oxide primers are appropriate but red lead should be avoided.

**Maintenance of finished aluminium**

For long-term durability, all external aluminium finishes should be washed regularly, at intervals not normally exceeding three months, with a mild detergent solution. Damaged paint coatings may be touched up on site, but remedial work does not have the durability of the factory-applied finishes.

**ALUMINIUM IN BUILDING**

Typical applications for aluminium and its alloys in building include; roofing and cladding, curtain-wall and structural glazing systems, flashings, rainwater goods, vapour barriers, and internal ceilings, panelling, luminaires, ducting, architectural hardware and walkways. Minimum sheet thicknesses for self-supporting sheet aluminium roofing and other applications to BS EN 14782: 2006 are 0.6 and 0.4 mm, respectively. For fully supported aluminium roofing and cladding to BS EN 14783: 2006, the nominal minimum sheet thickness is 0.6 mm.
Fig. 5.20 Aluminium discs — Selfridges store, Birmingham. Architects: Foster + Partners. Photographs: Arthur Lyons
Monocoque construction

The Lord’s Cricket Ground Media Centre (Fig. 5.21) was the world’s first semi-monocoque building in aluminium. The media centre is a streamlined pod raised 14 m off the ground on two concrete support towers, giving journalists and commentators an uninterrupted view over the cricket ground. The structure consists of a curved 6 and 12 mm aluminium-plate skin welded to a series of ribs. Thus acting together, the skin and the ribs provide both the shape and the structural stability, a system typically used in the boat building and aircraft industries. The building was made in 26 sections and transported to the site for assembly.

Thermal breaks in aluminium

In order to overcome thermal bridging effects, where aluminium extrusions are used for double-glazing systems, thermal breaks are inserted between the aluminium in contact with the interior and exterior spaces. These may be manufactured from preformed polyamide strips, or alternatively the appropriate extrusions are filled with uncured polymer, then the bridging aluminium is milled out after the plastic has set.

Jointing methods

Aluminium components may be joined mechanically with aluminium bolts or rivets; non-magnetic stainless steel bolts are also appropriate. If aluminium is to be electric arc welded, the use of an inert-gas shield, usually argon, is necessary to prevent oxidation of the metal surface. A filler rod, compatible with the alloy to be welded, supplies the additional material to make up the joint. Strong adhesive bonding of aluminium components is possible, provided that the surfaces are suitably prepared.

Copper

Copper was probably one of the first metals used by man, and evidence of early workings suggests that the
metal was smelted as early as 7000 BC. Later it was discovered that the addition of tin to copper improved the strength of the material and by 3000 BC the Bronze Age had arrived. The Romans made extensive use of copper and bronze for weapons, utensils and ornaments. Brass from the alloying of copper and zinc emerged from Egypt during the first century BC. By the mid-eighteenth century South Wales was producing 90% of the world’s output of copper, with the ore from Cornwall, but now the main sources are the United States, Chile, Peru and Australia. The traditional visual effect of copper is illustrated in a modern context by the millennium project, Swan Bells in Perth, Australia (Fig. 5.22). The copper was initially clear-coated to prevent gradual oxidation and patination within the marine environment of the harbour.

**MANUFACTURE**

The principal copper ores are the sulphides (e.g. chalcopyrite), and sulphides in association with iron (e.g. chalcoite). Ores typically contain no more than 1% copper and therefore require concentration by flotation techniques before the copper is extracted through a series of furnace processes. The ores are roasted then smelted to reduce the sulphur content and produce matte, which contains the copper and a controlled proportion of iron sulphide. The molten matte is refined in a converter by a stream of oxygen. This initially oxidises the iron which concentrates into the slag and is discarded; sulphur is then burnt off to sulphur dioxide, leaving 99% pure metal, which on casting evolves the remaining dissolved gases and solidifies to blister copper. The blister copper is further refined in a furnace to remove remaining sulphur with air and then oxygen with methane or propane. Finally, electrolytic purification produces 99.9% pure metal. Approximately 40% of copper and the majority of brass and bronze used within the UK are recycled from scrap. Recycling requires approximately 25% of the energy used in the primary production of copper depending on the level of impurities present.

**GRADES OF COPPER**

Grades of copper and copper alloys are designated by both a symbol and a number system to the current standards BS EN 1172: 1997 and BS EN 1412: 1996. Thus, for example, phosphorus deoxidised non-arsenical copper, typically used for roofing and cladding, would be defined as:

<table>
<thead>
<tr>
<th>Table 5.11</th>
<th>Broad classification of copper alloys</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number series</td>
<td>Letters</td>
</tr>
<tr>
<td>000–099</td>
<td>A or B</td>
</tr>
<tr>
<td>100–199</td>
<td>C or D</td>
</tr>
<tr>
<td>200–299</td>
<td>E or F</td>
</tr>
<tr>
<td>300–349</td>
<td>G</td>
</tr>
<tr>
<td>350–399</td>
<td>H</td>
</tr>
<tr>
<td>400–449</td>
<td>J</td>
</tr>
<tr>
<td>450–499</td>
<td>K</td>
</tr>
<tr>
<td>500–599</td>
<td>L or M</td>
</tr>
<tr>
<td>600–699</td>
<td>N or P</td>
</tr>
<tr>
<td>700–799</td>
<td>R or S</td>
</tr>
</tbody>
</table>

Note: The three-digit number designates each material and the letter indicates copper or the alloy group.

Symbol system:

Cu-DHP
(Deoxidised high-residual phosphorus)

Number system:

CW024A-R240
(C refers to copper, W refers to wrought products, 024A identifies the unique composition (Table 5.11). The subsequent letters and numbers define a specific requirement such as tensile strength or hardness, in this case - a minimum tensile strength of 240 MPa, which is half-hard temper.)

Only four of the numerous grades of copper are commonly used within the construction industry.

**Electrolytic tough pitch high-conductivity copper (Cu-ETP or CW004A)**

Electrolytic tough pitch high-conductivity copper is used mainly for electrical purposes; however, the sheet material is also used for fully supported traditional and long strip copper roofing. It contains approximately 0.05% dissolved oxygen which is evolved as steam if the copper is heated to 400°C in a reducing flame, thus rendering the metal unsuitable for welding or brazing.

**Fire refined tough pitch copper (Cu-FRHC or CW005A)**

Fire refined tough pitch copper has a similar specification to Cu-ETP, but with marginally more impurities.
Fig. 5.22 Copper cladding — Swan Bells Tower, Perth Australia. Architects: Hames Sharley Architects. Photographs: Arthur Lyons
**Tough pitch non-arsenical copper (Cu-FRTP or CW006A)**

Tough pitch non-arsenical copper is used for general building applications. It is suitable for sheet roofing.

**Phosphorus deoxidised non-arsenical copper (Cu-DHP or CW024A)**

Phosphorus deoxidised non-arsenical copper is the standard grade for most building applications including roofing, but not for electrical installations. The addition of 0.05% phosphorus to refined tough pitch copper isolates the oxygen rendering the metal suitable for welding and brazing. It is therefore used for plumbing applications where soldering is inappropriate.

The detailed chemical compositions of these and all other grades of copper are listed in DD CEN/TS 13388:2008.

**COPPER FORMS AND SIZES**

Copper is available as wire, rod, tube, foil, sheet and plate. Typical roofing grades are 0.45, 0.6 and 0.7 mm. (The minimum nominal thickness for self-supporting sheet copper to BS EN 14782: 2005 and for the fully supported material to BS EN 14783: 2006 is 0.5 mm.) The metal is supplied dead soft (fully annealed), one eighth or one quarter hard, half-hard or full-hard. It rapidly work hardens on bending, but this can be recovered by annealing at red heat. Copper can be worked at any temperature, since, unlike zinc, it is not brittle when cold. The standard grade of copper used for roofs, pipes and domestic water-storage cylinders is phosphorus deoxidised non-arsenical copper CW024A, although the other tough-pitch grades CW004A, CW005A and CW006A may also be used for roofs. Copper for pipework is supplied in annealed coils for mini/microbore systems, in 6 m lengths half-hard and hard for general plumbing work. The hard temper pipes cannot be bent. Plastic coated tubes, colour coded to identify the service (e.g. yellow–gas), are available.

**PATINA**

The green patina of basic copper sulphate or carbonate on exposed copper gradually develops according to the environmental conditions. On roofs within a marine or industrial environment the green patina develops within five years; under heavy pollution it may eventually turn dark brown or black. Within a town environment, the patina on roofs will typically develop over a period of ten years. However, vertical copper cladding will normally remain a deep brown, due to the fast rainwater run-off, except in marine environments when the green colour will develop. On-site treatment to accelerate the patinisation process is unreliable, but pre-patinised copper sheet is available if the effect is required immediately. Green pre-patinised copper sheet should not be welded, brazed or soldered as heat treatment causes discolouration of the patina. The factory-generated bright green patina will weather according to the local environmental conditions, often turning quickly to a blue-green. However, if the metallic appearance is to be retained externally, then a copper/aluminium/zinc alloy which resists weathering and retains its bright gold colour may be used. The Urbis Centre in Manchester illustrates well the visual effect of a large feature patinised copper roof (Fig. 5.23).

**CORROSION**

Generally, copper itself is resistant to corrosion; however, rainwater run-off may cause staining on adjacent materials and severe corrosion to other metals. Zinc, galvanised steel and non-anodised aluminium should not be used under copper, although in this respect lead, stainless steel and brass are unaffected. Copper may cause corrosion to steel or anodised aluminium in direct contact, if moisture is present. Specifically, copper should not be installed below exposed bitumen, bitumen paint, or cedarwood shingles where leaching action producing acid solutions can cause localised attack on the metal. Additionally, some corrosion may arise from the acid produced by algae on tiled roofs. The accidental splashing of lime or cement mortar onto copper causes a blue-green discolouration; however, this can readily be removed with a soft brass brush. Some corrosion of copper pipework may be caused by soft water, particularly if high levels of dissolved carbon dioxide are present; hard waters generally produce a protective film of calcium compounds, which inhibits corrosion. Pitting corrosion has been reported in rare cases associated with either hard, deep-well waters or hot, soft waters with a significant manganese content. Additionally, excessive acidic flux residues not removed by flushing the system may cause corrosion. Within heating systems in which oxygen in the primary circulating water is constantly being replenished through malfunction or poor design, bimetallic corrosion will occur between steel radiators and copper pipework. This will result in the
build-up of iron oxide residues at the bottom of the radiators. The use of appropriate inhibitors will reduce this effect.

PROTECTIVE COATINGS FOR COPPER

A range of clear coatings is available for external and internal application to copper where the original colour and surface finish is to be retained. Air-drying acrylic thermosetting resins and soluble fluoropolymers are suitable for exterior use. The acrylic resins, which incorporate benzotriazole to prevent corrosion if the coating is damaged, have a life expectancy in excess of 10 years, whilst the recently developed fluoropolymers should last for 20 years. For internal use polyurethane, vinyls, epoxy and alkyd resins are appropriate. Silicones are necessary for high temperature applications.

COPPER ROOFING SYSTEMS

Traditional and long-strip systems

Copper roofing systems may be categorised as traditional or long strip. The latter has the advantage that bays between 8.1 and 14.6 m, depending on the pitch, may be constructed without the necessary crosswelts on sloping roofs or drips on flat roofs appropriate to the traditional system. This has significant cost benefits in terms of installation costs. The long-strip copper roof system (Fig. 5.24) with bays up to approximately 600 mm wide may be laid on roofs with pitches between 3° and 90° and uses one-quarter or half-hard temper 0.6 or 0.7 mm copper strip. The system requires specified areas of the roof to be fixed with conventional
welted joint clips, and the remainder with expansion clips, which allow for the longitudinal expansion of the bays, but ensure a secure fixing to the substructure. Lateral thermal movement is accommodated by a space at the base of the standing seams. Long-strip copper is laid on a breather membrane which allows free movement between the metal and the structure, whilst isolating the copper from any ferrous fixings in the structure and providing some sound reduction from the effects of wind and rain. All fixings should be made from the same copper as the roof. Nails with minimum 6 mm diameter heads should be copper or brass.

Within the traditional copper roofs (Fig. 5.25), standing seams or batten roll jointing systems are used depending on the pitch and appearance required. For pitches of 5° or less, batten rolls are appropriate, as standing seams are vulnerable to accidental flattening and subsequent failure by capillary action. Cross welts may be continuous across roofs where batten rolls are used, but should be staggered where standing seams are used. Bays should not exceed approximately 1700 mm in length. Either soft or one quarter hard temper copper is normally used. The substructure, breather membrane and fixed clips are as used in long-strip roofing. These differences in articulation within the traditional roofing systems and particularly by contrast to the smooth line of long-strip system offer alternative visual effects to the designer of copper roofs. Copper rainwater systems are available with a range of standard components. Copper shingles offer an additional aesthetic to traditional copper roofing and cladding systems.

**Self-supporting systems**

Typical sections for profiled sheet copper and copper tiles are illustrated in the standard BS EN 506: 2008. Phosphorus deoxidised copper (Cu-DHP) is specified for this purpose. The standard BS EN 14782: 2006 defines a minimum thickness of 0.5 mm for self-supporting copper sheet for roofing and cladding.

**Bonded copper systems**

Proprietary systems offer visual effects similar to those of traditional copper, aluminium, stainless and terne-coated stainless steel roofing systems, by using the metal bonded to either particleboard or roofing sheet. (The latter is referred to within Chapter 6 on Bitumen and Flat Roofing Materials.) Copper bonded to 18 mm high-density moisture-resistant chipboard offers a smoother finish than that achieved by traditional roof- ing and cladding methods, whilst still showing the articulation of standing or flat seams.

**COPPER ALLOYS**

Copper may be alloyed with zinc, tin, aluminium, nickel or silicon to produce a range of brasses and bronzes. The full range of copper alloys is listed in DD CEN/TS 13388: 2008.

**Brass**

Brass is an alloy of copper and zinc, most commonly with zinc content between 10% and 45%. It is used for small components such as architectural ironmongery, door and window furniture, handrails and balustrades. It may be lacquered to prevent deterioration of the polished finish, although externally and in humid environments the lacquer fails, requiring the brass to be cleaned with metal polish to remove the tarnish. Brass plumbing fittings manufactured from 60/40 brass may corrode in soft, high-chloride content waters by dezincification. The process produces insoluble zinc corrosion products and ultimately porous metal fittings which may cause failure of the system. In situations where this problem is likely, dezincification-resistant (DZR) fittings made from alloy CW602N containing 2% lead should be used. Such components are marked with the ‘CR’ dezincification-resistant symbol.

**Bronze**

Bronze is an alloy of copper and tin, used for high-quality door furniture and recently as a woven fabric cladding material for the Theatre Royal rehearsal centre, Plymouth. Bronzes are usually harder and more durable than the equivalent brasses and exhibit a greater resistance to corrosion. Phosphor bronze contains up to 0.5% phosphorus in an 8% tin bronze. Because of its load-bearing properties and durability, it is frequently used as corbel plates and fixings for stone, and precast concrete cladding panels. Aluminium bronze (copper and aluminium), silicon bronze (copper and silicon) and gunmetals (copper–tin–zinc alloys) are also used for masonry fixings and cast components by virtue of their strength and durability. Nickel bronze alloys (copper, nickel and zinc) can be manufactured to highly polished silver finishes, particularly appropriate for interior fittings.
Fig. 5.25 Traditional copper roofing and copper shingles. *Photograph: Courtesy of Copper Development Association*
ANTI-MICROBIAL SURFACES

It is currently recognised that copper and copper alloys are naturally anti-microbial materials and therefore have a vital role to play in the reduction of hospital-acquired infections such as MRSA. It is known that a large proportion of infections are spread by touch, so it is appropriate that touch surfaces, such as door handles, push plates, taps and light switches within hospitals and other healthcare facilities should be manufactured from copper, brass or bronze rather than stainless steel or other inactive materials.

LEAD

The Egyptians used lead in the glazing of their pottery and for making solder by 5000 BC. It was mined in Spain by the Phoenicians around 2000 BC.

MANUFACTURE

Lead occurs naturally as the sulphide ore, galena. The manufacturing process involves the concentration of the ore by grinding and flotation. The sulphide is converted to the oxide by roasting, and then reduced to the metal in a blastfurnace charged with limestone and coke. Further refining removes impurities which otherwise would reduce the softness of the metal. Approximately 75% of lead used within the UK is recycled material.

LEAD SHEET

The majority of lead sheet for roofing, cladding, flashings and gutter linings is produced by milling thick sheet down to the required thickness. Continuous machine cast lead, which accounts for approximately 10% of the UK market, is manufactured by immersing a rotating water-cooled metal drum in a bath of molten lead at constant temperature. The lead solidifies on the surface of the drum and is peeled off as it emerges from the melt. The thickness can be adjusted by altering the speed of rotation of the drum. The sheet produced is without the anisotropic directional grain structure associated with the standard rolling process. The standard BS EN 12588: 2006 defines lead sheet by thickness rather than by the code system formerly used within the UK (Table 5.12). Sand cast lead sheet is still manufactured by the traditional method, which involves pouring molten lead onto a prepared bed of sand. The sheet thickness is controlled by drawing a piece of timber across the molten metal surface to remove the excess material. Sand cast lead is normally only used for conservation work on key historic buildings, when much of the old lead may be recycled in the process. A typical cast lead sheet size is 6 × 1.5 m.

Table 5.12 Lead sheet thickness and colour codes to BS EN 12588: 2006 with typical applications

<table>
<thead>
<tr>
<th>European designation</th>
<th>Green</th>
<th>Yellow</th>
<th>Blue</th>
<th>Red</th>
<th>Black</th>
<th>White</th>
<th>Orange</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nominal thickness (mm)</td>
<td>1.25</td>
<td>1.50</td>
<td>1.75</td>
<td>2.00</td>
<td>2.50</td>
<td>3.00</td>
<td>3.50</td>
</tr>
<tr>
<td>UK designation</td>
<td>3</td>
<td>4</td>
<td>5</td>
<td>6</td>
<td>7</td>
<td>8</td>
<td></td>
</tr>
<tr>
<td>Lead codes</td>
<td>3.12</td>
<td>1.59</td>
<td>1.80</td>
<td>2.24</td>
<td>2.65</td>
<td>3.15</td>
<td>3.55</td>
</tr>
<tr>
<td>Nominal weight (kg/m²)</td>
<td>15.0</td>
<td>18.0</td>
<td>20.4</td>
<td>25.4</td>
<td>30.1</td>
<td>35.7</td>
<td>40.3</td>
</tr>
</tbody>
</table>

Typical application:
- Flat roofing
- Pitched roofing
- Vertical cladding
- Soakers
- Hip and ridge flashings
- Parapets, box/tapered valley gutters
- Pitched valley gutters
- Weatherings to parapets
- Apron and cover flashings
- Chimney flashings
The minimum nominal thickness for fully supported lead sheet roofing and cladding is 1.25 mm to BS EN 14783: 2006. The handling of lead requires appropriate on-site care, particularly due to weight of the material and the toxic fumes produced during lead burning.

**CORROSION**

Freshly cut lead has a bright finish, but it rapidly tarnishes in the air with the formation of a blue-grey film of lead carbonate and lead sulphate. In damp conditions a white deposit of lead carbonate is produced, and in cladding this can both be aesthetically unacceptable and cause some staining of the adjacent materials. The effect can be prevented by the application of patination oil after the lead has been fixed. Lead is generally resistant to corrosion due to the protection afforded by the insoluble film; however, it is corroded by organic acids. Acidic rainwater run-off from mosses and lichens may cause corrosion and contact with damp timbers, particularly oak, teak and western red cedar, should be avoided by the use of building paper or bitumen paint. Trapped condensation under sheet lead may cause significant corrosion, so consideration must be given to the provision of adequate ventilation underneath the decking which supports the lead. Dew points must be checked to ensure that condensation will not occur and be trapped under the lead sheet in either new work or renovation. Generally, lead is stable in most soils; however, it is attacked by the acids within peat and ash residues. Electrolytic corrosion rarely occurs when lead is in contact with other metals, although within marine environments aluminium should not be used in association with lead. Corrosion does occur between wet Portland cement or lime products and lead during the curing process; thus, in circumstances where the drying out will be slow, lead should be isolated from the concrete with a coat of bitumen paint.

**FATIGUE AND CREEP**

In order to prevent fatigue failure due to thermal cycling or creep, that is, the extension of the metal under its own weight over extended periods of time, it is necessary to ensure that sheet sizes, thicknesses and fixings are in accordance with the advice given by the Lead Sheet Association in their technical manuals. The metal must be relatively free to move with temperature changes, so that alternating stresses are not focused in small areas leading to eventual fatigue fracture. A geotextile separating underlay may be used. The addition of 0.06% copper to 99.9% pure lead refines the crystal structure giving increased fatigue resistance without significant loss of malleability. The composition of lead sheet is strictly controlled by BS EN 12588: 2006.

**LEAD ROOFING**

Lead roofing requires a smooth, continuous substrate. Generally, the bay sizes depend on the roof geometry and the thickness of lead to be used (Fig. 5.26). For flat roofs (from 1 in 80 [approximately 1°] to 10°), joints are generally wood-cored rolls down the fall and drips across. For pitched roofs (10–80°), joints in the direction of the fall may be wood-cored or hollow rolls, with laps across the fall, unless for aesthetic reasons the bays are to be divided by drips. For steep pitches welts are used and over 80°, standing seams are appropriate (Fig. 5.27). Fixings are copper or stainless steel nails and clips within the rolls, welts or standing seams. Lead as a highly malleable material can be formed or bossed into shape with the specialist tools including the bossing stick and bossing mallet. Welding or leadburning involves the joining of lead to lead using additional material to make the joint thicker by one third than the adjacent material.

The David Mellor Cutlery Factory, Hathersage, Derbyshire (Fig. 9.14, Chapter 9, page 291), illustrates a traditionally detailed lead roof. The wood-cored roll-jointed lead is supported on a stepped deck manufactured from prefabricated stressed-skin insulated plywood boxes, tapered to fit the radial design. These units are supported on a series of lightweight steel trusses, tied at the perimeter by a steel tension ring and at the centre lantern by a ring-truss. Around the perimeter, the lead is burnt to ensure a vertical seal. A steeper roof without the need for drips is illustrated in Fig. 5.27.

**LEAD SHEET CLADDING**

For cladding the thickness of lead to be used dictates the maximum spacing between vertical joints and distance between laps. Vertical joints may be wood-cored rolls or welts and occasionally standing seams or hollow rolls, where the risk of physical damage from ladders is negligible. The lead is hung by nailing at the head, with allowance for up to 6 mm thermal movement to occur within the lap joints.

An alternative form of lead cladding is the use of preformed lead-faced cladding panels, which are...
then fixed to the building facade (Fig. 5.28). Typically, 25 mm exterior grade plywood covered with 1.80 or 2.24 mm (Code 4 or Code 5) lead is used. The panels are set against a lead-faced timber structural support leaving 25 mm joints for thermal movement. Standard details are illustrated in the relevant Code of Practice BS 6915: 2001.

**LEAD TILES**

An innovative use of lead as a roofing material is illustrated by the Haberdashers’ Hall in London (Fig. 5.29). The two-storey building, constructed around a courtyard, features a roof clad with diamond-shaped lead tiles. The individual units are formed from $1 \times 1.5$ m
marine plywood diamonds each dressed with lead sheet, and incorporating a flashing on two edges to seal under the panels above.

**FLASHINGS**

Lead, because of its malleability and durability, is an ideal material from which to form gutters and gutter linings, ridge and hip rolls, and the full range of standard and specialist flashings, including ornamental work to enhance design features. For most flashing applications, lead sheet of 1.32, 1.80 or 2.24 mm (Codes 3–5) is used, fixed with copper or stainless steel and occasionally lead itself.

**ACRYLIC-COATED LEAD SHEET**

Acrylic-coated lead sheet is produced to order and in the standard colours, white, slate grey, terracotta and dark brown for use in colour-coordinated flashings. The colour-coated 1.80 mm (Code 4) milled lead is produced in widths of 250, 300, 450 and 600 mm. The material is moulded by bossing as for standard lead sheet, and where welded, the exposed grey metal may be touched up if necessary.

**Zinc**

Zinc was known to the Romans as the alloy, brass, but it was not produced industrially until the mid-eighteenth century, and was not in common use on buildings until the nineteenth century. The cut surface tarnishes quickly to a light grey due to the formation of a patina of basic zinc carbonate. The metal is hard at ambient temperatures and brittle when cold. It should therefore not be worked at metal temperatures below 10°C without prior warming and heavy impacts should not be used within the forming processes of bending and folding.

**MANUFACTURE**

Zinc occurs naturally as the sulphide ore, zinc blende. The ore is first concentrated and then roasted to produce zinc oxide. The addition of coal reduces zinc oxide to the metal, which is evolved as the vapour and then condensed. High-grade zinc is produced by the electrolysis of a purified zinc sulphate solution. Zinc is classified according to its purity as specified in the standard BS EN 1179: 2003 (Table 5.13).

Approximately 3610 kW hours of energy is consumed in the primary production of one tonne of sheet zinc; however, a large proportion of the metal is recycled.

<table>
<thead>
<tr>
<th>Table 5.13</th>
<th>Colour codes for grades of zinc</th>
</tr>
</thead>
<tbody>
<tr>
<td>Grade classification</td>
<td>Z1</td>
</tr>
<tr>
<td>Zinc content (%)</td>
<td>99.995</td>
</tr>
<tr>
<td>Colour code</td>
<td>White</td>
</tr>
</tbody>
</table>
ZINC SHEET

Zinc sheet is manufactured by continuous casting and rolling in a range of thicknesses (Table 5.14) to a maximum coil width of 1000 mm. The two standard products are the pure metal (99.995% zinc) and its alloy with small additions of titanium and copper (e.g. 0.06% and 0.08% minima, respectively). The rolling process modifies the grain structure, particularly in the pure metal; however, this does not affect
the working of the sheets. The alloy has improved performance with respect to strength and creep resistance but also a reduced coefficient of thermal expansion which enables the construction of roof bays up to 10 m, or in certain cases up to 16 m in length, depending on design considerations including bay width. The zinc–copper–titanium alloy (BS EN 988: 1997) may be folded or curved to produce interlocking cladding panels for vertical, horizontal or diagonal installation. Both the pure metal and the titanium alloy can be worked by hand at room temperature and do not work-harden. The minimum nominal thickness for self-supporting zinc alloy to BS EN 14782: 2006 and for fully supporting zinc roofing and cladding to BS EN 14783: 2006 is 0.6 mm. The zinc–copper–titanium alloy is used to add a unity to the suite of elegant buildings forming the University of Cambridge, Centre for Mathematical Sciences, by Edward Cullinan Architects (Fig. 5.30).

**Table 5.14** Zinc–copper–titanium sheet thicknesses and weights

<table>
<thead>
<tr>
<th>Nominal thickness (mm)</th>
<th>0.6</th>
<th>0.65</th>
<th>0.7</th>
<th>0.8</th>
<th>1.0</th>
<th>1.20</th>
<th>1.50</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nominal weight (kg/m²)</td>
<td>4.3</td>
<td>4.7</td>
<td>5.0</td>
<td>5.8</td>
<td>7.2</td>
<td>8.6</td>
<td>10.8</td>
</tr>
</tbody>
</table>

**PATINA**

Bright zinc tarnishes in the air with the production of a thin oxide film, which is rapidly converted into basic zinc carbonate by the action of water and carbon dioxide. The patina then prevents further degradation of the surface. Ordinary zinc has a lighter blue-grey patina than the alloyed sheet, so the two materials should not be mixed within the same construction. Pre-weathered alloys are available if light or slate grey patinated surfaces are required immediately. The lifetime of zinc depends directly on the thickness. A 0.8 mm roof should last for 40 years in urban conditions, whereas the same sheet as cladding, washed clean by rain, could last for 60 years. The titanium alloy with considerably improved durability has a predicted life of up to 100 years in a rural environment depending on the pitch of the application.

**LACQUERED ZINC SHEET**

A factory-applied 25 μm heat-treated polyester lacquer finish to zinc gives a range of colour options through white, brown, terracotta, red, green grey and blue. Alternative organic coatings include acrylic, silicone-polyester, polyvinylidene fluoride and PVC plastisol to BS EN 506: 2008.
CORROSION

Zinc should not be used in contact with copper or where rainwater draining from copper or copper alloys would discharge onto zinc. It may, however, be used in association with aluminium. In contact with steel or stainless steel, the zinc must be the major component to prevent significant corrosion effects. unprotected cut edges of galvanised steel located above zinc can cause unsightly rust stains and should be avoided. If the underside of zinc sheet remains damp due to condensation for extended periods of time then pitting corrosion will occur, causing eventual failure. It is therefore necessary to ensure that the substructure is designed appropriately with vapour barrier, insulation and ventilation to prevent interstitial condensation. Sulphur dioxide within polluted atmosphere prevents the formation of the protective carbonate film and causes corrosion.

Zinc is not affected by Portland cement mortars or concrete, although it should be coated with acrylic resin paint where it will be in contact with soluble salts from masonry or cement additives. Zinc may be laid directly onto seasoned softwoods, unless impregnated with copper-salt preservatives, which have a slight corrosion-promoting effect. However, zinc should not be used on acidic timbers such as oak, chestnut and western red cedar. Furthermore, zinc should not be used in association with western red cedar shingles, which generate an acidic discharge. The acidic products from the effect of ultraviolet radiation on bitumen can cause corrosion in zinc. If the bitumen is not
protected from direct sunlight by reflective chippings, then any zinc must be separated from the bitumen with an impermeable material.

**FIXINGS**

Fixings for zinc should be of galvanised or stainless steel. Clips are made of zinc, cut along the rolled direction of the sheet and folded across the grain. Watertight joints may be made by soldering using tin/lead solder in conjunction with zinc chloride flux.

**ROOFING AND CLADDING**

Both the roll cap and standing seam systems (Fig. 5.31) are appropriate for fully supported zinc and zinc alloy roofing (BS EN 501: 1994). W elted joints are standard practice across the bays at pitches steeper than 15°; below 15° drips are necessary. A minimum fall of 3° is recommended, although a pitch in excess of 7° will ensure self-cleaning, preventing the accumulation of dirt, which reduces service life. Where the bay length is greater than 3 m, a section 1.3 m in length is fixed rigidly, whilst the remaining area is secured to the substructure with sliding clips which accommodate the thermal movement. Timber roof boarding, oriented strand board or plywood form the ideal substructures for zinc, but chipboard is inappropriate except cement-bonded particleboard for cladding. Where concrete is used it must be sealed against trapped moisture. For cladding, the vertical joints may be welted, standing seam or roll cap with the horizontal joints welted. The maximum bay length for cladding is 6 m, although 3 m is more practical on site. Titanium zinc rainwater systems are available with an appropriate range of standard components.

Zinc may also be used as the surface material for interlocking facade panel systems, with recessed horizontal or vertical joints, fixed to a sub-frame of timber, aluminium or stainless steel. A range of colours is available with lacquered finish.

Titanium zinc interlocking square and diamond tiles are appropriate for vertical hanging and roof pitches down to 25°. They are fixed with soldered and sliding clips to timber battens. A range of sizes is available to give a choice of scale. Tiles are either preweathered or bright and manufactured in 0.7 or 0.8 mm alloy. To muffle the sound of rain, a full-surface substructure is advisable.
RECYCLING OF ZINC

Approximately 30% of newly formed zinc is from recycled sources, and within Europe approximately 90% of sheet material and rainwater goods zinc is recycled.

Titanium

Titanium ore is abundant in the earth’s crust, with reserves well exceeding currently anticipated demands. The main producing countries are Russia, USA, Australia and Japan, although the ores rutile (titanium oxide) and ilmenite (iron-titanium oxide), are also found in Europe, China and South America. Originally isolated in 1887, it was developed for use in the aerospace industry in the 1950s and has been used as a building cladding material in Japan for over thirty years. The Glasgow Science Centre (Fig. 5.32) illustrates titanium’s eye-catching appearance as a modern construction material.

MANUFACTURE

The ore is treated with chlorine to produce titanium tetrachloride, which is then purified to remove other unwanted elements. Treatment with metallic magnesium or sodium reduces the titanium tetrachloride to a sponge of titanium metal, which is then melted under vacuum to produce solid ingots. Ingots are then forged into slabs and rolled out into sheet. Where required an embossed finish can be applied during the final rolling process. Other sections and forms can be produced by hot rolling or cold forming as for steel. Titanium has a high embodied energy; however, this is to some extent balanced against its life-cycle costing and ultimate full recycling.

PROPERTIES AND USES

Titanium is an appropriate material for construction due to its corrosion resistance. It is resistant to acids and alkalis, also industrial and marine environments. Titanium has a density of 4510 kg/m³, intermediate between aluminium (2700 kg/m³) and steel (7900 kg/m³), giving it the advantage of a good strength-to-weight ratio. It is less ductile than steel, so hot forming is required for severe bending. The metal has a modulus of elasticity half that of steel. Titanium has a low coefficient of expansion ($8.9 \times 10^{-6}$), half that of stainless steel and copper and one third that for aluminium. This reduces the risk of thermal stress, and enables titanium sheet roofing to be laid

**Fig. 5.32**  Titanium cladding — the Glasgow Science Centre. Architects: BDP, Building Design Partnership. Photograph: Courtesy of Don Clements
in longer lengths than other metals. The use of relatively thin roofing and cladding panels (0.3–0.4 mm) minimises both the dead load and the supporting structural system. Titanium with its very high melting point of 1670°C can withstand fire tests at 1100°C and has been certified as a ‘non-combustible material’ in Japan for roofing and cladding. Further applications include fascias, panelling, protective cladding for piers and columns and three-dimensional artwork.

**Durability**

The corrosion resistance of titanium arises from its self-healing and tenacious protective oxide film. However, rainwater run off from zinc, lead or copper roofs should be avoided. The Guggenheim Museum in Bilbao, clad in 32,000 m² of commercially pure 0.3–0.4 mm titanium sheet panels, shows some staining due to lack of protection during the construction process and also rainwater run-off. Although initially expensive, on a life-cycle basis, due to its low maintenance costs, titanium may prove to be a highly competitive cladding and roofing material. Already one manufacturer is offering a 100-year guarantee against corrosion failure in roofing applications. Titanium can cause the corrosion of contact aluminium, steel or zinc, but austenitic stainless steel (grade 1.4401) is not affected.

**Finishes**

The normal oxide film can be thickened by heat treatment or anodising, giving permanent colours ranging from blue and mauve to cream and straw. Control is necessary to ensure the absence of colour variations within a project. Surface finishes range from reflective bright to soft matte and embossed, as used on the Glasgow Science Centre buildings (Fig. 5.32). In this case, the rolling grain direction was maintained over the building facades to ensure no visible variation of the embossed stipple effect.

**Welding titanium**

Titanium may be arc welded, but this requires the exclusion of air usually by the use of argon gas shielding. Other welding technologies such as plasma arc and laser or electron beam are used for more specialist applications.

**TITANIUM ALLOYS**

Titanium is available as a wide range of alloys classified according to increased corrosion resistance, higher strength or higher temperature resistance. However, their current use is mainly confined to aerospace, industrial and medical applications. The standard architectural cladding material is 99% pure titanium (grade 1 or grade 2).

**Process of metallic corrosion**

Corrosion is an electro-chemical process, which can only occur in the presence of an electrolyte, that is moisture containing some dissolved salts. The process may be understood by considering the action of a simple Daniell cell as shown in Fig. 5.33.

When the cell operates, two key processes occur. At the anode, the zinc gradually dissolves, generating zinc ions in solution and electrons which flow along the wire and light up the lamp as they move through its filament. At the copper cathode, the electrons are received on the surface of the metal, and combine with copper ions in solution to plate out new shiny metal on the inside of the copper container.

\[
\text{Anode: } Zn \rightarrow Zn^{2+} + 2e^{-}
\]

![Fig. 5.33 Daniell cell](image)
An equivalent process takes place in the dry Leclanché cell—the standard torch battery (Fig. 5.34). However, in this case the central carbon rod replaces the copper and the liquid is replaced by an aqueous paste. The anode process is the same as in the Daniell cell with the gradual dissolution of the zinc container.

At the cathode the carbon rod is surrounded by manganese dioxide, which oxidises the hydrogen gas which otherwise would have been produced there by the reaction between water and the electrons.

This sequence is similar to that seen in the corrosion of iron (Fig. 5.35). In this case the presence of both an electrolyte and oxygen is necessary for corrosion to occur.
Fig. 5.37  Bimetallic corrosion between aluminium and steel

Anode
\[
\text{Fe} \rightarrow \text{Fe}^{2+} + 2e^{-}
\]

Cathode
\[
2\text{H}_2\text{O} + \text{O}_2 + 4e^- \rightarrow 4\text{OH}^{-}
\]

Rust formation
\[
\text{Fe}^{2+} + 2\text{OH}^- \rightarrow \text{Fe(OH)}_2 \rightarrow \text{Fe}_2\text{O}_3 \cdot \text{H}_2\text{O}
\]

Overall summary
\[
4\text{Fe} + 3\text{O}_2 + 2\text{H}_2\text{O} \rightarrow 2\text{Fe}_2\text{O}_3 \cdot \text{H}_2\text{O}
\]

FACTORS AFFECTING THE RATE OF CORROSION

The key factors which accelerate the rate of corrosion are the presence of two dissimilar metals in mutual contact and the degree of pollution within any moisture surrounding the metals. If the more base metal is small in quantity compared to the more noble metal, then rapid corrosion of the more base metal will occur. Figure 5.36 shows which pairs of metals commonly used in construction should not generally be allowed into contact. Within a single metal, minor surface variations, such as crystal grain boundaries, the effects of cold working or welding, the presence of impurities or alloying components within the metal, variable cleanliness or access to aerial oxygen, may all cause accelerated corrosion. Figure 5.37 illustrates the effect of corrosion between an extruded aluminium gutter and a steel rainwater pipe. The aluminium corroded producing a white deposit near the point of contact between the two metals; this was followed by rapid corrosion of the steel.

References

FURTHER READING

Copper Development Association. 2004: *Copper and copper alloys, compositions, applications and properties*. Publication 120, Hemel Hempstead: CDA.

Copper Development Association. 2006: *The guide to copper in architecture*. Publication 154, Hemel Hempstead: CDA.


**STANDARDS**

BS 4 Structural steel sections:


BS 416 Discharge and ventilating pipes and fittings sand-cast or spun in cast iron:

BS 417 Galvanised mild steel cisterns and covers, tanks and cylinders:
  Part 2: 1987 Metric units.

BS 437: 2008 Specification for cast iron drain pipes, fittings and their joints.

BS 449 Specification for the use of structural steel in building:
  Part 2: 1969 Metric units.


BS 493: 1995 Airbricks and gratings for wall ventilation.

BS 779: 1989 Cast iron boilers for central heating and indirect water supply (rated output 44 kW and above).

BS 1161: 1977 Specification for aluminium alloy sections for structural purposes.

BS 1189: 1986 Specification for baths made from porcelain enamelled cast iron.

BS 1202 Nails:
  Part 2: 1974 Copper nails.
BS 1210: 1963 Wood screws.
BS 1245: 1975 Metal door frames (steel).
BS 1329: 1974 Metal hand rinse basins.
BS 1390: 1990 Baths made from vitreous enameled sheet steel.
BS 1449 Steel plate, sheet and strip:
BS 1494-1: 1964 Specification for fixing accessories for building purposes. Fixings for sheet, roof and wall coverings.
BS 1566 Copper indirect cylinders for domestic purposes:
BS ISO 3575: 2005 Continuous hot-dip zinc-coated carbon steel sheet of commercial and drawing qualities.
BS 3987: 1991 Anodic oxide coatings on wrought aluminium for external architectural applications.
BS 4604 The use of high-strength friction grip bolts in structural steelwork. Metric series:
BS 4868: 1972 Profiled aluminium sheet for building.
BS 4873: 2009 Aluminium alloy windows and door sets. Specification.
BS 4921: 1988 Specification for sherardised coatings on iron or steel.
BS ISO 4999: 2005 Continuous hot dip terne (lead alloy) coated cold reduced carbon steel sheet of commercial, drawing and structural qualities.
BS 5950 Structural use of steelwork in building:
BS ISO 5954: 2007 Cold reduced carbon steel sheet according to hardness requirements.
BS 5977 Lintels:
BS 6496: 1984 Specification for powder organic coatings for application and stoving to aluminium alloy extrusions, sheet and preformed sections for external architectural purposes.
BS 6510: 2005 Specification for steel windows, sills, window boards and doors.
BS 6582: 2000 Specification for continuously hot dip lead alloy (terne) coated cold reduced carbon steel flat rolled products.
BS 6722: 1986 Recommendations for dimensions of metallic materials.
BS 6744: 2001 Stainless steel bars for the reinforcement of and use in concrete.
BS 6915: 2001 Design and construction of fully supported lead sheet roof and wall coverings.
BS 7364: 1990 Galvanised steel studs and channels for stud and sheet partitions and linings using screw fixed gypsum wallboards.
BS 8118 Structural use of aluminium:
BS 8202 Coatings for fire protection of building elements:
BS ISO 13887: 2004 Cold reduced steel sheet of higher yield strength with improved formability.
BS ISO 14788: 2005 Continuous hot dip zinc 5% aluminium alloy coated steel sheet.
BS ISO 16020: 2005 Steel for the reinforcement and prestressing of concrete. Vocabulary.
BS EN 124: 1994 Gully tops and manholes. Design requirements, type, testing, marking, quality control.
BS EN 485 Aluminium and aluminium alloys. Sheet, strip and plate:
   Part 2: 2008 Mechanical properties.
   Part 4: 1994 Tolerances on shape and dimensions for cold-rolled products.
BS EN 508 Roofing products from metal sheet. Specification for self-supported products of steel, aluminium or stainless steel sheet:
BS EN 545: 2006 Ductile iron pipes, fittings, accessories and their joints for water pipelines.
BS EN 573 Aluminium and aluminium alloys. Chemical composition and form of wrought products:
   Part 2: 1995 Chemical symbol based designation system.
BS EN 586 Aluminium and aluminium alloys. Forgings:
   Part 2: 1994 Mechanical properties.
   Part 3: 2001 Tolerances on dimensions and form.
BS EN 598: 2007 Ductile iron pipes, fittings, accessories and their joints for sewerage applications. Requirements and test methods.
BS EN 754 Aluminium and aluminium alloys:
   Parts 1-8 Cold drawn rod/bar and tube.
BS EN 755 Aluminium and aluminium alloys:
   Parts 1-9 Extruded drawn rod/bar, tube and profiles.
BS EN 845 Specification for ancillary components for masonry:
   Part 1: 2003 Ties, tension straps, hangers and brackets.
BS EN 877: 1999 Cast iron pipes and fittings, their joints and accessories.

BS EN 969: 2009 Specification for ductile iron pipes, fittings, accessories and their joints for gas applications. Requirements and test methods.


BS EN 1057: 2006 Copper and copper alloys. Seamless round copper tubes for water and gas in sanitary and heating applications.

BS EN 1090 Execution of steel structures and aluminium structures:

BS EN 1172: 1997 Copper and copper alloys. Sheet and strip for building purposes.

BS EN 1173: 2008 Copper and copper alloys. Material condition designation.

BS EN 1179: 2003 Zinc and zinc alloys. Primary zinc.

BS EN 1254: 1998 Copper and copper alloys. Plumbing fittings.

BS EN 1412: 1996 Copper and copper alloys. European numbering system.

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  Part 5: 2004 Structural steels with improved atmospheric corrosion resistance.

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  Part 2: 1992 Steel numbers.

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   Part 2: 1996 Delivery conditions for thermomechanically rolled steels.
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   Part 2: 2006 Products for building exterior applications.
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   Part 1: 1999 General requirements.
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ADVISORY ORGANISATIONS

Aluminium Federation Ltd., National Metalforming Centre, 47 Birmingham Road, West Bromwich, West Midlands B70 6PY, UK (0121 601 6363).
Aluminium Rolled Products Manufacturers Association, National Metalforming Centre, 47 Birmingham Road, West Bromwich, West Midlands B70 6PY, UK (0121 601 6361).
British Non-Ferrous Metals Federation, 5 Grovelands Business Centre, Boundary Way, Hemel Hempstead HP2 7TE, UK (01442 275705).
British Stainless Steel Association, Broomgrove, 59 Clarkehouse Road, Sheffield, South Yorkshire S10 2LE, UK (0114 267 1260).
Cast Iron Drainage Development Association, 72 Francis Road, Edgbaston, Birmingham, W. Midlands B16 8SP, UK (0121 693 9909).
Cold Rolled Sections Association, National Metalforming Centre, Birmingham Road, West Bromwich, West Midlands B70 6PY, UK (0121 601 6350).
Copper Development Association, 5 Grovelands Business Centre, Boundary Way, Hemel Hempstead, Hertfordshire HP2 7TE, UK (01442 275705).
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Galvanizers Association, Wren’s Court, 56 Victoria Road, Sutton Coldfield, West Midlands B72 1SY, UK (0121 355 8838).
Lead Sheet Association, Unit 10, Archers Park, Brandon Road, East Peckham, Tonbridge, Kent TN12 5HP, UK (01622 872432).
Metal Cladding and Roofing Manufacturers Association, 18 Mere Farm Road, Prenton, Wirral, Cheshire CH43 9TT, UK (0151 652 3846).
Stainless Steel Advisory Centre, Broomgrove, 59 Clarkehouse Road, Sheffield, South Yorkshire S10 2LE, UK (0114 267 1265).
Steel Construction Institute, Silwood Park, Ascot, Berkshire SL5 7QN, UK (01344 636525).
Zinc Information Centre, 6 Wrens Court, 56 Victoria Road, Sutton Coldfield B72 1SY, UK (0121 362 1201).
Introduction

The flat roofing materials, which form an impermeable water barrier, include reinforced bitumen membrane systems, mastic asphalt, single-ply plastic membranes and liquid coatings. All these require continuous support on an appropriate roof decking system. Green roofs are considered as an extension of the standard roofing systems. Metal roofing systems are described in Chapter 5.

FIRE EXPOSURE OF ROOFS

All materials used as finishes for roofs, both pitched and flat, are classified with respect to external fire exposure. The classification system (BS 476-3: 2004) indicates whether the roof is flat or pitched followed by a two-letter coding on fire performance.

<table>
<thead>
<tr>
<th>Roof system</th>
<th>Fire penetration (first letter)</th>
<th>Fire spread (second letter)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cold-deck, warm-deck, and inverted roofs</td>
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FIRE EXPOSURE OF ROOFS

All materials used as finishes for roofs, both pitched and flat, are classified with respect to external fire exposure. The classification system (BS 476-3: 2004) indicates whether the roof is flat or pitched followed by a two-letter coding on fire performance.

- **Roof system**: EXT. F. (flat) or EXT. S. (sloping)
- **Fire penetration (first letter)**: A, B, C or D
  - A = no penetration in 1 hour
  - B = no penetration in 30 minutes
  - C = penetration within 30 minutes
  - D = penetrated by preliminary flame test
- **Fire spread (second letter)**: A, B, C or D
  - A = no spread of flame
  - B = spread of flame less than 533 mm
  - C = spread of flame more than 533 mm
  - D = specimens that continue to burn after the test flame was removed or which had a spread of flame more than 381 mm in the preliminary test

A suffix ‘X’ is added where the material develops a hole or suffers mechanical failure.

Thus, a flat roof material classified as EXT. F. AA suffers no fire penetration or spread of flame during the standard one hour fire test.

Cold-deck, warm-deck, and inverted roofs

COLD-DECK ROOFS

In cold-deck roof construction, the weatherproof layer is applied directly onto the roof decking, usually particleboard or plywood, and this is directly supported by the roof structure, frequently timber joists (Fig. 6.1). Thermal insulation is laid over the gypsum plasterboard ceiling, leaving cold void spaces between the
structural timbers or steel. In this form of roof construction, there is a significant risk of condensation forming on the underside of the decking, and this may cause deterioration of the structure. Precautions must be taken to ensure adequate ventilation of the cold voids, and also, the underside of the deck must not cool below the dew point when the external temperature is $-5^\circ C$. Any vapour check under the insulation layer is vulnerable to leakage around electrical service cables. In remedial work on cold-deck roofs, if adequate ventilation cannot be achieved, then conversion to a warm-deck or inverted roof system may be advantageous. Cold-deck roof construction is not recommended for new building work in the Code of Practice (BS 8217: 2005).

WARM-DECK ROOFS

In warm-deck roof construction the thermal insulation is laid between the roof deck and the weatherproof covering (Fig. 6.1). This ensures that the roof deck and its supporting structure are insulated from extremes of temperature, thus limiting excessive thermal movement which may cause damage. As the insulating material is directly under the waterproof layer, it must be sufficiently strong to support any foot traffic associated with maintenance of the roof. The waterproof and insulation layers will require mechanical fixing or ballasting to prevent detachment in strong winds. Surface condensation on the underside of a roof deck within warm-deck roof construction would normally indicate insufficient thermal insulation. Warm-deck construction is the preferred method for lightweight roofs.

INVERTED ROOFS

In inverted roof construction, both the structural deck and the weatherproof membrane are protected by externally applied insulation (Fig. 6.1). This ensures that the complete roof system is insulated from extremes of hot and cold, and also from damage by solar radiation and maintenance traffic. The insulation layer is usually either ballasted with gravel or fully protected with paving slabs. Disadvantages of inverted roof construction are the greater dead-weight and the difficulty in locating leaks under the insulation layer. Inverted roof construction is the preferred method for concrete and other heavyweight roof systems.

Reinforced bitumen membranes

Reinforced bitumen membrane (RBM) roofs (formerly built-up roofing) consist of two or more layers of bitumen sheets bonded together with self-adhesive or hot bitumen. The individual bitumen sheets are
Table 6.1  Reinforced bitumen membranes for roofing to BS 8747: 2007

<table>
<thead>
<tr>
<th>Description</th>
<th>Type (previous coding)</th>
<th>Colour code</th>
<th>Tensile class</th>
<th>Resistance to impact—subclass D</th>
<th>Static loading—subclass L</th>
<th>Puncture class P (derived from D and L)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glass-fibre—fine granule surface</td>
<td>3B</td>
<td>Red</td>
<td>S1</td>
<td>D1</td>
<td>L1</td>
<td>D1L1 = P1</td>
</tr>
<tr>
<td>Polyester—fine granule underlayer</td>
<td>5U</td>
<td>Blue</td>
<td>S2</td>
<td>D2</td>
<td>L2</td>
<td>D2L2 = P2</td>
</tr>
<tr>
<td>Polyester—fine granule surface</td>
<td>5B/180</td>
<td>Blue</td>
<td>S3</td>
<td>D2</td>
<td>L3</td>
<td>D2L3 = P3</td>
</tr>
<tr>
<td>Polyester—fine granule surface</td>
<td>5B/250</td>
<td>Blue</td>
<td>S4</td>
<td>D2</td>
<td>L4</td>
<td>D2L4 = P4</td>
</tr>
<tr>
<td>Polyester—fine granule surface</td>
<td>5B/350</td>
<td>Blue</td>
<td>S5</td>
<td>D3</td>
<td>L4</td>
<td>D3L4 = P5</td>
</tr>
</tbody>
</table>

Notes:
These are typical combinations, but specifiers must check the manufacturers’ specifications as different combinations may apply.

There are five levels of tensile strength from S1 (lowest) to S5 (highest).
There are three levels of resistance to impact D1 (lowest) to D3 (highest).
There are four levels of resistance to static loading L1 (lowest) to L4 (highest).
Numbers 180, 250 and 350 refer to the mass (g/m²) of the base layer (not including any granular finish).

Manufactured from a base carrier of either glass or fibre matting which is impregnated and coated with bitumen. Some products also have an applied adhesive backing and/or a surface protective finish.

Bitumen is the residual material produced after the removal by distillation of all volatile products from crude oil. The properties of bitumen are modified by controlled oxidation, which produces a more rubbery material suitable for roofing work. In the manufacture of bitumen sheets, the continuous base layer of organic fibres, glass fibre or bitumen-saturated polyester is passed through molten oxidised bitumen containing inert filler; the material is then rolled to the required thickness. The bitumen sheet is coated with sand to prevent adhesion within the roll or with mineral chippings to produce the required finish. A range of thicknesses is available, and for ease of recognition of the base fibre layer, the rolls are colour-coded along one edge.

TYPES OF REINFORCED BITUMEN MEMBRANES

Reinforced bitumen membranes for roofing systems are classified by BS EN 13707: 2004 according to a range of physical properties but without specific reference to the materials of manufacture. Guidance given in BS 8747: 2007 relates the key physical properties of tensile strength (S class) and puncture resistance (P class) to the types of product. Puncture resistance is derived from a combination of resistance to impact (D subclass) and resistance to static loading (L subclass). Both the S and P classes have five criteria levels of increasing standard from 1 to 5. The classification does not apply to venting and partial bonding base layers or underlay for mastic asphalt. The polyester-based products have greater strength and durability at a higher initial cost than the glass-fibre-based products. Fine finishes are appropriate for the underlayers or where surface protection is applied, and granular finishes may be used for the exposed layer. Table 6.1 shows the typical relationship between generic products and the derived membrane S and P classes.

The requirement for a particular membrane performance (e.g. S2P3) is determined from a risk analysis, including the geometry of the roof (slope and membrane substrate), access (maintenance, pedestrian and vehicle) and degree of protection (self-protection, ballast, pavers and gardens). BS 8747: 2007 includes a matrix as guidance for specification of the appropriate membrane classification required.

Examples:

Flat concrete roof, 0–5° pitch, gravel ballast, light maintenance access only – membrane classification required S2P3.

Plywood substrate, >5° pitch, self-protected membrane, light maintenance access only—membrane classification required S3P2.

The classification applies to both single-layer and multi-layer reinforced bitumen membrane systems. For multi-layer systems, one layer (usually the underlayer) should have a minimum S2P2 classification.

The Annex to BS 8747: 2007 describes Type 3G glass-fibre reinforced bitumen perforated venting layer used as a first layer of built-up RBM roofing systems where partial bonding or venting is required.
Table 6.2  Reinforced bitumen membranes for roofing to Annex of BS 8747: 2007

<table>
<thead>
<tr>
<th>Class</th>
<th>Base material</th>
<th>Type</th>
<th>Use</th>
<th>Colour code</th>
</tr>
</thead>
<tbody>
<tr>
<td>Class 1</td>
<td>Organic fibres and jute hessian base</td>
<td>Type 1F</td>
<td>Pitched roof underslating sheet</td>
<td>White</td>
</tr>
<tr>
<td>Class 3</td>
<td>Glass-fibre base</td>
<td>Type 3G</td>
<td>RBM roof venting base layer</td>
<td>None</td>
</tr>
<tr>
<td>Class 5</td>
<td>Polyester base</td>
<td>Type 5U</td>
<td>Pitched roof underslating sheet</td>
<td>Blue</td>
</tr>
</tbody>
</table>

Underlay materials for discontinuous roofing such as slates and tiles are described in BS EN 13859: 2005. They are characterised by three classes of resistance to water penetration, namely class W1 (highest resistance) to class W3 (least resistance). The Annex to BS 8747: 2007 describes Type 1F fibre-based bitumen underlay reinforced with hessian and possibly incorporating an aluminium foil heat-reflecting layer on the underside. Type 5U polyester reinforced bitumen roofing underlay has greater mechanical strength with a higher resistance to tearing than the Type 1F underlay. Generally, where the risk of condensation is high, sarking membranes, which are waterproof but vapour permeable, should be used. Alternative breather materials for tile underlay include a range of polyolefin laminates and modified PVC products. The colour codes for the bitumen-based membranes are listed in Table 6.2.

A typical reinforced bitumen membrane system is illustrated in Fig. 6.2. All oxidised bitumen sheets, when exposed to ultraviolet light and ozone, gradually age harden and become less resistant to fatigue failure.

**ROOFING SYSTEMS**

Reinforced bitumen roofing systems of substrate, insulation, membrane and finishes are described in the Code of Practice BS 8217: 2005. Roofing sheets may be applied to roofs constructed from precast or in situ reinforced concrete, plywood (exterior grade), timber (19 mm tongued and grooved, with preservative treatment and conditioning), oriented strand board, wood wool slabs or profile metal decking (galvanised steel or aluminium). Certain proprietary composite decking systems, such as units comprising plywood, rigid urethane foam and aluminium foil, are also appropriate except in areas of high humidity. In warm roof applications a vapour check is applied over the decking. Appropriate vapour check materials include lapped and bonded bitumen membranes, lapped polythene sheets or 12 mm mastic asphalt on glass fibre tissue depending on the structural material.

Insulation materials include cork, rigid mineral (MW) or glass wool, perlite (EPB), cellular foamed glass (CG), rigid polyurethane foam (RUP), extruded (XPS) or expanded moulded (EPS) polystyrene, phenolic foam (PF), polyisocyanurate (PIR), bitumen impregnated fibreboard and various proprietary composite systems. Most manufacturers are now able to supply CFC-free insulation products. The heat-sensitive expanded plastics are frequently supplied pre-bonded to a cork, perlite or fibreboard layer to receive the *pour and roll* hot bitumen or *torch-on* sheet systems. Where suitable falls are not incorporated into the roof structure, the insulation can be supplied ready cut-to-falls. A minimum in service fall of 1 in 80 is required to prevent *ponding*; this requires a design fall of 1 in 40 to allow for settlement.

![Fig. 6.2  Typical reinforced bitumen membrane roofing system](image)
On sloping roofs the first layer of roofing sheet is applied down the slope, but on flat roofs (less than 10°) the direction of the first layer need not relate to the falls. The first layer is either partly or fully bonded depending on the substrate. Perforated sheet is frequently laid loose as the first layer, and becomes spot bonded as the hot bitumen for the second layer is applied. On timber the first layer is nailed. Partial bonding permits some thermal movement between the sheet system and the decking, and also allows for the escape of any water vapour trapped in the decking material. The use of proprietary breather vents on large roofs allows the escape of this entrapped air from the roof structure by migration under the partially bonded layer. Side laps of 50 mm and end laps of typically 100 mm should all be staggered between layers. On sloping roofs the first layer should be nailed at the top of each sheet at 50 mm centres and higher melting point 115/15 bitumen should be used for bonding the subsequent layers to prevent slippage. Protection from the effects of ultraviolet light is afforded by either the factory-applied mineral surface finish to the cap sheet, by the application of reflective paint or on flat roofs, typically a 12 mm layer of reflective white spar stone chippings.

POLYMER-MODIFIED BITUMEN MEMBRANES

High-performance bitumen membranes based on polyester base for toughness, and polymer-modified bitumen coatings for increased flexibility, strength and fatigue resistance, offer considerably enhanced durability over the standard oxidised bitumen sheets. The two types are based on styrene–butadiene–styrene (SBS) and atactic polypropylene (APP) modified bitumen. These systems are covered by BS 8747: 2007.

Elastomeric SBS high-performance membranes

Styrene–butadiene–styrene polymer-modified bitumen membranes have greater elasticity than standard oxidised bitumen sheets. They are laid either by the traditional pour and roll technique which is used for standard bitumen sheets or by torching-on (Fig. 6.3). In the pour and roll process, bonding bitumen is heated to between 200 and 250°C and poured in front of the sheet as it is unrolled, giving continuous adhesion between the layers. In the torching-on process as the sheet is unrolled, the backing is heated to the molten state with propane burners. Alternatively, cold adhesive or mechanical fastening may be used.

Plastomeric APP high-performance membranes

Atactic polypropylene polymer-modified bitumen contains typically 25% atactic polymer in bitumen with some inert filler. The product is more durable than oxidised bitumen and has enhanced high-temperature resistance and low-temperature flexibility. Sheets are manufactured with a polyester and/or glass-fibre core. Some additionally have a glass-fibre reinforced weathering surface. The APP polymer-modified bitumen sheets are bonded with cold adhesive or by torching the heat-sensitive backing, as the temperature of hot-poured bitumen is too low to form a satisfactory bond.
METAL-FACED ROOFING SHEETS

Copper- and aluminium-faced high-performance SBS membranes give visual quality and enhanced durability compared with standard mineral surfaced reinforced bitumen membrane roofing systems. The small squared pattern of indentations (Fig. 6.4) allows for thermal movement between the bonded sheet and metal finish. The copper finish weathers similarly to traditional copper roofing systems producing a green patina. The metal foil thickness is typically 0.08 mm.

INVERTED ROOFS

In the inverted roof the reinforced bitumen membrane waterproofing membrane is laid directly onto the roof deck. Non-absorbent insulation, such as extruded polystyrene, is laid onto the membrane, which is then covered with a filter sheet to prevent the ingress of excessive organic material. River-washed ballast or pavings on supports protect the system from mechanical and wind damage. Inverted roofs have the advantage that the waterproof membrane is protected from thermal stress by the insulating layer. This in turn is protected from damage by the paving or ballast finish. High-performance reinforced bitumen membrane systems are suitable for inverted roofs.

Table 6.3 Mastic asphalt grades to BS 6925: 1988

<table>
<thead>
<tr>
<th>Type</th>
<th>Composition</th>
</tr>
</thead>
<tbody>
<tr>
<td>BS 988B</td>
<td>100% bitumen</td>
</tr>
<tr>
<td>BS 988T25</td>
<td>75% bitumen, 25% lake asphalt</td>
</tr>
<tr>
<td>BS 988T50</td>
<td>50% bitumen, 50% lake asphalt</td>
</tr>
<tr>
<td>Specified by manufacturers</td>
<td>polymer-modified grades</td>
</tr>
</tbody>
</table>

Mastic asphalt

Mastic asphalt is a blended bitumen-based product. It is manufactured either from the bitumen produced by the distillation of crude oil, or from lake asphalt, a naturally occurring blend of asphalt containing 36% by weight of finely divided clay, mainly imported from Trinidad. The bitumen is blended with limestone powder and fine limestone aggregate to produce the standard roofing types listed in BS 6925: 1988 (Table 6.3). Most frequently specified are the BS 988T25 and the polymer-modified grades.

The effect of the finely divided clay particles within lake asphalt type BS 988T confers better laying characteristics and enhanced thermal properties; these are advantageous when the material is to be exposed to

Fig. 6.4 Copper-faced bitumen membrane roofing
wide temperature changes, particularly in warm roof construction systems.

Mastic asphalt is usually delivered as blocks for melting on site prior to laying, although hot molten asphalt is occasionally supplied for larger contracts. Laid mastic asphalt is brittle when cold but softens in hot, sunny weather. The hardness is increased by the re-melting process, and also by the addition of further limestone aggregate. Polymer-modified mastic asphalts, usually containing styrene–butadiene–styrene block copolymers, are more durable and have enhanced flexibility and extensibility at low temperatures, allowing for greater building movements and better resistance to thermal shock. Where mastic asphalt roofs are subjected to foot or vehicle access, paving-grade mastic asphalt should be applied as a wearing layer over the standard roofing grade material. Two key grades S and H are available; the softer grade is suitable for footways and rooftop car parks, and the harder grade for heavily stressed areas. For standard flooring, type F1076 mastic asphalt is required, for coloured flooring type F1451, and for tanking and damp-proof courses T1097 is necessary. The flooring types of mastic asphalt are available in four grades (hard, light, medium and heavy duty) according to the required wearing properties.

**ROOFING SYSTEMS**

Mastic asphalt may be laid over a wide variety of flat or pitched roof decking systems to the Code of Practice BS 8218: 1998. As mastic asphalt is a brittle material it requires continuous firm support. Appropriate decks are concrete (in situ or precast), plywood (exterior grade), timber boarding (19 mm minimum thickness) particleboard, wood wool slabs (50 mm minimum thickness) and profiled metal decking. A typical concrete warm roof system is illustrated in Fig. 6.5.

In dense concrete construction, a sand and cement screed is laid to falls over the in situ slab. The falls should be designed such that even with inevitable variations on site, they are never less than 1 in 80, as this is essential to ensure the immediate removal of the surface water and to prevent ponding. Plywood, profiled metal and other decking systems would be similarly laid to falls. A layer of bitumen-bonded Type 3B glass-fibre roofing sheet is applied over the structure to act as a vapour check.

![Fig. 6.5 Typical mastic asphalt roofing system](image)
Insulation

Thermal insulation, to provide the necessary roof $U$-value, is bonded with hot bitumen. A wide variety of insulation boards or blocks, including compressed cork, high-density mineral wool, fibreboard, perlite, cellular foamed glass, high-density extruded polystyrene and polyisocyanurate are suitable, although where insufficient rigidity is afforded by the insulation material, or if it would be affected by heat during application of the hot asphalt, it must be overlaid with firm heat-resistant boards to prevent damage during maintenance or construction. Where falls are not provided by the structure, the insulation may be set appropriately, provided that the thinnest section gives the required thermal properties. A separating layer of loose-laid geotextile material or sheathing membrane is then applied to allow differential thermal movement between the decking system and the mastic asphalt waterproof finish.

Mastic asphalt application

Mastic asphalt is laid to 20 mm in two layers on roofs up to 30° and in three layers to 20 mm on slopes greater than 30°. Upstands of 150 mm are required to masonry, rooflights, pipes, etc. where they penetrate the roof membrane. Where adhesion on vertical surfaces is insufficient, expanded metal lathing should be used to support the mastic asphalt. An apron flashing should protect the top of the upstand. A layer of sand is rubbed into the top of the final layer whilst it is hot to break up the skin of bitumen-rich material which forms at the worked surface.

Surface protection

Mastic asphalt gradually hardens over a period of a few years and should be protected from softening under bright sunlight by the application of surface protection. Reflective paint coatings – for example, titanium oxide in polyurethane resin or aluminium-pigmented bitumen – are effective until they become dirty, but for vertical surfaces they are the only appropriate measure. Reflective coatings are available in a range of colours giving differing levels of solar reflectivity. For horizontal surfaces and pitches up to 10°, a layer of 10–14 mm white stone chippings will give better protection not only from sunlight but also from ultraviolet light, which gradually degrades bitumen products. Additionally, a layer of stone will act to reduce the risk of thermal shock during very cold periods. Where traffic is anticipated, the mastic asphalt should be protected with glass-fibre reinforced cement (GRC) tiles or concrete pavings.

INVERTED ROOFS

Mastic asphalt forms a suitable waterproof membrane for externally insulated or inverted roofs. The application of the insulating layer over the mastic asphalt has the advantage that it protects the waterproof layer from thermal shock, impact damage and degradation by ultraviolet light. The insulation, usually extruded polystyrene boards, is held down by either gravel or precast concrete paving slabs.

Single-ply roofing systems

Single-ply roofing systems consist of a continuous membrane usually between 1 and 3 mm thick, covering any form of flat or pitched roof (Fig. 6.6). As waterproofing is reliant on the single membrane, a high quality of workmanship is required, which is normally provided by the specialist installer. In refurbishment work where the substrate may be rough, a polyester fleece may be used to prevent mechanical damage to the membrane from below. Life expectancies are typically quoted as 25 years. The wide range of membrane materials used may generally be categorised into thermoplastic, elastomeric and modified bitumen products. In many cases the single-layer membrane is itself a laminate, incorporating either glass fibre or polyester to improve strength and fatigue resistance or dimensional stability, respectively. Both thermoplastic and elastomeric products are resistant to ageing under the severe conditions of exposure on roofs. Fixings offered by the proprietary systems include fully bonded, partially bonded, mechanically fixed and loose laid with either ballast or concrete slabs. Joints are lapped and either heat or solvent welded, usually with tetrahydrofuran (THF). A final seal of the plastic in solvent may be applied to joint edge after the lap joint has been checked for leaks. Most manufacturers provide a range of purpose-made accessories such as preformed corners, rainwater outlet sleeves and fixings for lightning protection. The standard BS EN 13956: 2006 lists an extensive range of materials used for single-ply roofing systems across Europe, but the market in the UK for non-bitumen-based products is dominated by plasticised PVC and EPDM.
THERMOPLASTIC SYSTEMS

Thermoplastic systems, made from non-cross-linked plastics, can be joined by solvent or heat welding. They generally exhibit good weathering properties and chemical resistance. The dominant thermoplastic systems are based on plasticised polyvinyl chloride (PVC) which is normally available in a range of colours. Certain PVC products contain up to 35% by weight of plasticisers, which can migrate to adjacent materials, leaving the membrane less flexible and causing incompatibility with extruded polystyrene insulation or bitumen products. The Imperial War Museum hangar at Duxford by Foster and Partners (Fig. 6.7) is finished with a PVC roofing membrane.

Products manufactured from vinyl ethylene terpolymer (VET, a blend of 35% ethyl vinyl acetate and 65% PVC with only 4% plasticiser added as lubricant) are more compatible with bitumen- and polystyrene-based insulation products. Other products include chlorinated polyethylene (CPE) which has enhanced chemical resistant properties, chlorosulphonated polyethylene (CSM) which is highly weather resistant and polyisobutylene (PIB), a relatively soft material which is easily joined. Thermoplastic polyolefin (TPO) membranes, based on
an alloy of ethylene propylene rubber (EPR) and polypropylene, are halogen-free and combine the temperature, chemical resistance and weldability of PVC with the flexibility of the elastomeric single-ply systems. The thermoplastic elastomer (TPE) single-ply roofing membranes based on polyolefins are similar to the well-established TPOs but have enhanced properties of elasticity and flexibility resulting from the particular catalytic process which generates very pure highly elastic polyolefins.

**Elastomeric Systems**

Elastomeric systems are dominated by ethylene propylene diene monomer (EPDM) which is a cross-linked or cured polymer. It is characterised by high elongation and good weathering resistance to ultraviolet light and ozone. The standard material is black or grey in colour but white is also available. Most products are seamed with adhesives or applied tapes as EPDM cannot be softened by solvents or heat; however, EPDM laminated with thermoplastic faces can be heat welded on site. Products may be mechanically fixed, ballasted or adhered to the substrate.

**Modified Bitumen Systems**

Most modified-bitumen systems are based on SBS (styrene–butadiene–styrene), APP (atactic polypropylene), APAO (poly α-olefin), TPO (thermoplastic polyolefin) or rubber-modified bitumen. Some products are now combining the durability of APP-modified bitumens with the enhanced flexibility of SBS-modified bitumen. Systems usually incorporate polyester or glass-fibre reinforcement for increased dimensional stability. Modified bitumen roofing membranes are usually thicker (e.g. 5 mm) than polymer-based single-ply systems.

**Liquid Coatings**

A range of bitumen- and polymer-based materials is used in the production of liquid-roof waterproofing membranes. Whilst some products are installed in new work, many products are used for remedial action on failed existing flat roofs as an economic alternative to re-roofing (Fig. 6.8). They may be appropriate when the exact location of water ingress cannot be located or when re-roofing is not practicable due to the disruption that it would cause. It is essential that the nature of the existing roof is correctly determined so that an appropriate material can be applied; also failures in the substrate must be identified and rectified. The surface of the existing material must be free of loose material and dust to ensure good adhesion with the liquid coating which can be applied by brush, roller or airless spray. While achieving a uniform thickness is difficult, the systems have the advantage of being seamless. Solar reflective white and a range of colours are available. Without further protection, roofs should only be subjected to light maintenance pedestrian traffic. Installation should normally be carried out by specialist roofing contractors. Systems include glass-fibre reinforced polyester and reinforced elastomeric polyurethane.

**Bitumen-Based Systems**

Most bitumen systems require a primer to seal the existing roof membrane and provide a base for the liquid coating. Two or three coats of bitumen solution or emulsion will normally be required for the waterproofing layer, and a solar reflective finish should be applied after the membrane has fully dried. A layer of glass-fibre reinforcement is usually incorporated during application of the waterproofing membrane to give dimensional stability. Two-component systems mixed during the spraying process cure more rapidly, allowing a seamless 4 mm elastomeric coat to be built up in one layer on either flat or pitched roofs. Where the material is ultraviolet light resistant, a solar reflective layer may not be necessary.

**Polymer-Based Systems**

The range of polymers used for liquid roof finishes is extensive, including acrylic resins, polyurethanes,
polyesters, silicones, rubber copolymers and modified bitumens. Some manufacturers offer a range of colours incorporating the necessary solar reflecting properties. Glass-fibre or polyester mat is used as reinforcement within the membrane layer, which is applied in a minimum of two coats. Additives to improve fire resistance, such as antimony trioxide and bromine compounds, may be incorporated into the formulations and good fire ratings in respect of flame penetration and surface spread can be achieved with some products. Solvent-based products have the advantage of rapid drying times, whilst solvent-free products have ‘greener’ credentials and some are odour-free, reducing the disruption caused during the refurbishment of occupied buildings.

Green roofs

Green roofs are flat or low-pitched roofs landscaped over the waterproofing layer. The landscaping may include some hard surfaces and have access for leisure and recreational functions as well as the necessary routine maintenance. Green roofs offer not only increased life expectancy for the waterproofing layer by protecting it from physical damage, ultraviolet light and temperature extremes, but also increased usable space. Environmental advantages include reduced and delayed rainwater run-off as a Sustainable Drainage System (SUDS), also considerable environmental noise, thermal control, air quality and wildlife habitat benefits. Green roofs (Fig. 6.9) may be waterproofed using modified-bitumen high-performance membrane systems, single-ply membrane systems or mastic asphalt. Under planting, T-grade mastic asphalt should be laid to 30 mm in three layers rather than the usual two layers to 20 mm thickness. Green roofs are divided between the intensive and the extensive systems.

EXTENSIVE GREEN ROOFS

Extensive green roofs are designed to be lighter in weight, relatively cheap, not open to recreational use and requiring the minimum of maintenance. Their prime purpose is either ecological or for the environmental masking of buildings. Planting should be of drought-tolerant, wind- and frost-resistant species, such as sedums, herbs and grasses. Instant cover can be created by the installation of pre-cultivated vegetation blankets where the immediate visual effect is required. Alternatively, a mixture of seeds, plant cuttings, mulch and fertiliser is sprayed onto the growing medium, and this will mature into the finished green roof over a period between one and two years. The complete system with planting, soil, filter sheet, drainage, moisture retention layer and root barrier will add between 60 and 200 kg/m² loading to the roof structure, which must be capable of this additional imposed load. Typical soil depth is up to 100 mm. Limited maintenance is required to remove unwanted weeds, fill bare patches and apply organic fertilisers in the spring, and to remove dead plants and weeds in the autumn. An
alternative approach to the fully planted extensive green roof is the biodiverse green roof which has some initial planting and incorporates natural features such as logs and boulders but which is then left to nature to develop with the local flora and fauna. Figure 6.10 illustrates a typical low-maintenance green roof at the Westonbirt Arboretum, Gloucestershire; a similar effect is achieved at the former Earth Centre, Doncaster, which is planted with sedum (Fig. 17.1 on page 397).

INTENSIVE GREEN ROOFS

Intensive green roofs are generally designed to accept recreational activity and include the wider range of vegetation from grass and herbaceous plants to shrubs. Depths of soil are typically between 200 and 300 mm, which, together with the necessary minimum 50 mm of water reservoir and drainage systems, generate an additional imposed load of typically 400 kg/m² on the existing or proposed structural system. Intensive green roofs may incorporate both soft and hard landscaping and slopes up to 20° are practicable. To conform to health and safety requirements edge protection (e.g. hand rail) or a fall arrest system (e.g. harness attachment points) must be incorporated into the design.

A typical intensive green roof system requires the following construction:

- soil, compost and planting—hard and soft landscaped areas;
- filter fleece to prevent soil blocking the drainage system;
- moisture retention material and drainage system;
- protection mat (to prevent damage to root barrier and waterproofing layers);
- polyethylene foil (isolating layer);
- root barrier;
- waterproof layer;
- insulation;
- vapour barrier.

Where trees are required soil depths may need to be increased to 750–1000 mm, with the associated increase in overall weight. Care must be taken to ensure that the roof membrane is not damaged by gardening implements. Intensive green roofs will require careful choice of planting to cope with the exposed conditions and regular long-term maintenance including weeding, trimming and pest control.

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ADVISORY ORGANISATIONS

European Liquid Waterproofing Association, Roofing House, 31 Worship Street, London EC2A 2DX, UK (020 7448 3859).
Flat Roofing Alliance, Roofing House, 31 Worship Street, London EC2A 2DX, UK (020 7448 3857).
Institute of Asphalt Technology, Paper Mews Place, 290 High Street, Dorking, Surrey RH4 1QT, UK (01306 742792).
Livingroofs, 7 Dartmouth Grove, London SE10 8AR, UK (020 8692 2109).
Mastic Asphalt Council, PO Box 77, Hastings, East Sussex TN35 4WL, UK (01424 814400).
Metal Cladding and Roofing Manufacturers Association, 18 Mere Farm Road, Prenton, Wirral, Cheshire CH43 9TT, UK (0151 652 3846).
National Federation of Roofing Contractors Ltd., 24 Weymouth Street, London W1G 7LX, UK (020 7436 0387).
Roofing Industry Alliance, Roofing House, 31 Worship Street, London EC2A 2DX, UK (020 7448 3857).
**Introduction**

The term glass refers to materials, usually blends of metallic oxides, predominantly silica, which do not crystallise when cooled from the liquid to the solid state. It is the non-crystalline or amorphous structure of glass (Fig. 7.1) that gives rise to its transparency.

Glass made from sand, lime and soda ash has been known in Egypt for 5000 years, although it probably originated in Assyria and Phoenicia. The earliest man-made glass was used to glaze stone beads, later to make glass beads (circa 2500 BC), but it was not until about 1500 BC that it was used to make hollow vessels.

For many centuries glass was worked by drawing the molten material from a furnace. The glass was then rolled out or pressed into appropriate moulds and finally fashioned by cutting and grinding. Around 300 BC the technique of glass blowing evolved in Assyria, and the Romans developed this further by blowing glass into moulds. Medieval glass produced in the Rhineland contained potash from the burning of wood rather than soda ash. Together with an increase in lime content this gave rise to a less durable product which has caused the subsequent deterioration of some church glass from that period.

The various colours within glass derived from the addition of metallic compounds to the melt. Blue was obtained by the addition of cobalt, whilst copper produced blue or red and iron or chromium produced green. In the fifteenth century white opaque glass was produced by the addition of tin or arsenic, and by the seventeenth century ruby red glass was made by the addition of gold chloride. Clear glass could only be obtained by using antimony or manganese as a decolouriser to remove the green colouration caused by iron impurities within the sand.

By the late twentieth century, with the advent of fully glazed facades, as illustrated by the Faculty of Law building at the University of Cambridge (Fig. 7.2), the construction industry had become a major consumer of new glass, and a proactive force in the development of new products. The wide range of glass materials used within the building industry is classified in the standard BS 952-1: 1995.

Within the UK, thousands of tonnes of glass are recycled each year, but this is mainly domestic waste, which cannot be used for the production of window glass as this requires pure materials. Even architectural waste glass is of variable composition with contamination from wire, sealants and special glasses, making it not usable as cullet in the manufacturing process without careful sorting. However, clean cullet from glass processing companies is recycled, and new float glass may incorporate up to 30% of recycled material, significantly reducing overall energy and CO₂ emissions. Re-melting clean cullet uses 25% less energy than making the equivalent quantity of new glass from raw materials. In addition, there is saving on extraction of raw materials. Excess recycled glass, not required for remaking bottles, has been used for making decorative paving surfaces and incorporated into the manufacture of bricks.

Recent experimentation with pulverised domestic glass waste has shown that when expanded by CO₂ from added calcium or magnesium carbonate or carbon black, the resulting material can be used as a lightweight aggregate in concrete. Expansion of 15–20 times gives a granular product with a density range 200–900 kg/m³.
Manufacture

COMPOSITION

Soda lime silicate glass

Modern glass is manufactured from sand (silica), soda ash (sodium carbonate) and limestone (calcium carbonate), with small additions of salt-cake (calcium sulphate) and dolomite (magnesian limestone). This gives a final composition of typically 70–74% silica, 12–16% sodium oxide, 5–12% calcium oxide, 2–5% magnesium oxide with small quantities of aluminium, iron and potassium oxides. The addition of 25% broken glass or cullet to the furnace mix accelerates the melting process and recycles the production waste. Most raw materials are available within the UK, although some dolomite is imported. The production process is relatively energy intensive at 15,000 kWh/m$^3$ (c.f. concrete: 625 kWh/m$^3$), but the environmental pay back arises from its appropriate use in energy-conscious design. Soda lime silicate glass may be chemically strengthened by an ion exchange process which replaces small surface ions by larger ones, thus putting the surfaces and edges into compression.

Alkaline earth silicate, borosilicate and ceramic glass

Other products used within construction include the alkaline earth silicate and borosilicate glasses; these have significantly different chemical compositions giving rise to their particular physical properties. The composition of alkaline earth silicate glass is typically 55–70% silica, 5–14% potassium oxide, 3–12% calcium oxide, 0–15% aluminium oxide with quantities of zirconium, strontium and barium oxides. Borosilicate glass is typically 70–87% silica, 0–8% sodium oxide, 0–8% potassium oxide, 7–15% boron oxide, 0–8% aluminium oxide with small quantities of other oxides. A particular characteristic of borosilicate glass is that it has a coefficient of expansion one third that of other glasses.

Fig. 7.1 Structure of glass (after Button and Pye: 1993)

Fig. 7.2 Glazed facade—Faculty of Law, University of Cambridge. Architects: Foster + Partners. Photograph: Arthur Lyons
of standard soda lime silicate glass, making it significantly more resistant to thermal shock in case of fire.

Ceramic glass is characterised by a near zero coefficient of expansion, making it highly resistant to thermal shock. The composition of ceramic glass is complex, comprising typically 50–80% silica, 15–27% alumina, with small quantities of sodium oxide, potassium oxide, barium oxide, calcium oxide, magnesium oxide, titanium oxide, zirconium oxide, zinc oxide, lithium oxide and other minor constituents. The glass is initially produced by a standard float or rolling technique but subsequent heat treatment converts part of the normal glassy phase into a fine-grained crystalline form, giving rise to the particular physical properties. Ceramic glass has a high softening point compared to other glass products used in construction.

FORMING PROCESSES

Early methods

Early crown glass was formed by spinning a 4 kg cylindrical gob of molten glass on the end of a blow pipe. The solid glass was blown, flattened out and then transferred to a solid iron rod or punty. After reheating it was spun until it opened out into a 1.5 m diameter disc. The process involved considerable wastage including the bullion in the centre, which nowadays is the prized piece. An alternative process involved the blowing of a glass cylinder which was then split open and flattened out in a kiln. This process was used for the manufacture of the glass for the Crystal Palace in 1851.

Subsequently in a major development, a circular metal bait was lowered into a pot of molten glass and withdrawn slowly, dragging up a cylindrical ribbon of glass 13 m high, the diameter of the cylinder being maintained with compressed air. The completed cylinder was then detached, opened up and flattened out to produce flat window glass.

It was only by the early twentieth century with the development of the Fourcalt process in Belgium and the Colburn process in America that it became possible to produce flat glass directly. A straight bait was drawn vertically out of the molten glass to produce a ribbon of glass, which was then drawn directly up a tower, or in the Colburn process turned horizontally, through a series of rollers; finally, appropriate lengths were cut off. However, such drawn sheet glass suffered from manufacturing distortions. This problem was overcome by the production of plate glass, which involved horizontal casting and rolling, followed by grinding to remove distortions and polishing to give a clear, transparent but expensive product. The process was ultimately fully automated into a production line in which the glass was simultaneously ground down on both faces. The plate glass manufacturing process is now virtually obsolete having been replaced by the float process, which was invented in 1952 by Pilkington and developed into commercial production by 1959. Drawn glass is only manufactured for conservation work, where gaseous and solid inclusions in the glass are required to emulate the historic material.

Float glass

A furnace produces a continuous supply of molten glass at approximately 1100°C, flowing across the surface of a large shallow bath of molten tin contained within an atmosphere of hydrogen and nitrogen, which prevents oxidation of the surface of the molten metal (Fig. 7.3). The glass ribbon moves across the molten metal, initially at a sufficiently high temperature for the irregularities on both surfaces to become evened out leaving a flat and parallel ribbon of glass. The temperature of the glass is gradually reduced as it moves forward until at the end of the molten tin,
it is sufficiently solid at 600°C not to be distorted when supported on rollers. Thickness is controlled by the speed at which the glass is drawn from the bath. Any residual stresses are removed as the glass passes through the 200 m annealing lehr or furnace, leaving a fire-polished material. The glass is washed and substandard material discarded for recycling. The computer-controlled cutting, firstly across the ribbon and then removal of the edges, is followed by stacking, warehousing and dispatch. A typical float glass plant will manufacture 5000 tonnes of glass per week, operating continuously for several years.

Float glass for the construction industry is made within the thickness range 2–25 mm, although 0.5 mm is available for the electronics industry. Many surface-modified glasses are produced, by incorporating metal ions into the glass within the float process, or under vacuum by magnetically enhanced cathodic sputtering or by spraying the surface with metal oxides or silicon. Body-tinted glass, which is of uniform colour, is manufactured by blending additional metal oxides into the standard melt. The use of all-electric melting processes offers higher quality control and less environmental pollution than previously produced from earlier oil- or gas-fired furnaces.

Non-sheet products

GLASS FIBRES

Continuous filament

Continuous glass fibres are manufactured by constantly feeding molten glass from a furnace into a forehearth fitted with 1600 accurately manufactured holes through which the glass is drawn at several thousand metres per second. The fibres (as small as 9 μm in diameter) pass over a size applicator and are gathered together as a bundle prior to being wound up on a collet. The material may then be used as rovings, chopped strand or woven strand mats for the production of glass-fibre reinforced materials, such as glass-fibre reinforced polyester (GRP), glass-fibre reinforced cement (GRC), or glass-fibre reinforced gypsum (GRG) – see Chapter 11.

Glass wool

Glass wool is made by the Crown process, which is described in Chapter 13.

CAST GLASS

Glass may be cast and pressed into shape for glass blocks and extruded sections.

Profiled sections

Profile trough sections in clear or coloured 6 or 7 mm cast glass are manufactured in sizes ranging from 232 to 498 mm wide, 41 and 60 mm deep and up to 7 m long, with or without stainless steel longitudinal wires (Fig. 7.4) (BS EN 572-7: 2004). The system can be used horizontally or vertically, as single or double glazing, and as a roofing system spanning up to

Fig. 7.4 Profiled glass sections
3 m. A large radius curve is possible as well as the normal straight butt jointed system and the joints are sealed with one-part translucent silicone. The standard double-glazed system has a $U$-value of 2.8 W/m² K, but this can be enhanced to 1.8 W/m² K by the use of low-emissivity coated glass. Amber or blue tinted versions are available for solar control or aesthetic reasons. The double-glazed system produces a sound reduction within the 100–3200 Hz range of typically 40 dB.

The incorporation of 16 mm thick translucent aerogel insulated polycarbonate panels into profiled trough sections significantly increases the thermal insulating properties of the units. Airborne sound insulation is enhanced, particularly at frequencies below 500 Hz. Light transmission through the translucent aerogel panels is approximately 50%, but UV transmission is zero. (Aerogels are described in Chapter 13, Insulating Materials, page 347.)

**Glass blocks**

Glass blocks for non-load-bearing walls and partitions are manufactured by casting two half-blocks at 1050°C, joining them together at 800°C, followed by annealing at 560°C. The void space is partially evacuated, giving a $U$-value of 2.5 W/m² K. The standard blocks (Fig. 7.5) are 115, 190, 240 and 300 mm square with thicknesses of 80 and 100 mm, although rectangular and circular blocks are also available. Walls may be curved as illustrated in Fig. 7.5. The variety of patterns, offering differing degrees of privacy, include clear, frosted, Flemish, reeded, and crystal designs with colours ranging from blue, green and grey to pink and gold. Blocks with solar reflective glass or incorporating white glass fibres offer additional solar control; colour may be added to either the edge coating or the glass itself. Special blocks are also available to form corners and ends also for ventilation.

For exterior and fire-retarding applications natural or coloured mortar (2 parts Portland cement, 1 part lime, 8 parts sand) is used for the jointing. Walls may be straight or curved; in the latter case, the minimum radius varies according to the block size and manufacturer’s specification. Vandal- and bullet-proof blocks are available for situations requiring higher security. For interior use, blocks may be laid with sealant rather than mortar. Glass blocks are now available in preformed panels for speedier installation.

Glass blocks, jointed with mortar, give a fire resistance of 60 minutes with respect to integrity and either
15 (F15), 30 (TF30) or 60 (TF60) minutes with respect to insulation according to BS EN 1364-1: 1999. Fire stop blocks (TF30) are manufactured from 26 mm rather than 8 mm glass to offer the increased fire protection. Sound reduction over the 100–3150 Hz range is typically 40–42 dB for standard blocks, but up to 45–49 dB for the TF30 blocks. Visible light transmission ranges downwards from 85 to 77% depending on the pattern and block size, but this is reduced to 60% for coloured glass.

**Glass pavers**

Glass pavers are cast as either single-layer blocks or shells, and as hollow blocks where insulation is required. The standard sizes are 120, 150 and 190 mm square, with a depth of 55, 60 or 80 mm for single-layer shells and 80, 100 and 160 mm for double-layer insulating blocks. Single-layer square blocks range in size from 90 to 300 mm with thicknesses typically between 18 and 38 mm. Surfaces may be clear or textured for non-slip action. On-site installation requires reinforced concrete construction allowing a minimum 30 mm joint between adjacent blocks and appropriate expansion joints around panels. Precast panels offer a high standard of quality control and rapid installation. They may also be used to create architectural features such as naturally illuminated domes or archways.

**CELLULAR OR FOAMED GLASS**

The manufacture of cellular or foamed glass as an insulation material is described in Chapter 13.

**Sheet products**

**STANDARD FLOAT GLASS SIZES**

The standard thicknesses for float glass are 3, 4, 5, 6, 8, 10, 12, 15, 19 and 25 mm to maximum sheet sizes of 3 × 12 m. Thicker grades are available to smaller sheet sizes. (The \( U \)-value for standard 6 mm float glass is 5.7 W/m² K.)

**TRADITIONAL BLOWN AND DRAWN GLASSES**

Traditional blown and drawn glasses are available commercially both clear and to a wide range of colours. Drawn glass up to 1600 × 1200 mm is optically clear but varies in thickness from 3 to 5 mm. It is suitable for conservation work where old glass requires replacement. Blown glass contains variable quantities of air bubbles and also has significant variations in thickness giving it an antique appearance. Where laminated glass is required, due to variations in thickness, these traditional glasses can only be bonded to float glass with resin.

**CURVED SHEET GLASS**

BS 952-2: 1980 refers to standard glass curvatures. Curved glass can be manufactured by heating annealed glass to approximately 600°C, when it softens and sags to the shape of the supporting mould. Sheets up to 3 × 4 m or 2 × 5 m can be bent with curvature in either one or two directions. Thicknesses between 4 and 19 mm are standard and tight curvatures can be produced for architectural feature glass. Patterned, textured, tinted, clear white and pyrolytic coated solar control glasses can all be curved by this technique. Bent glass can subsequently be sand-blasted, toughened, or laminated incorporating coloured interlayers if required.

**SELF-CLEANING GLASS**

Self-cleaning glass has an invisible hard coating 15 μm thick, which incorporates two special features. The surface incorporating titanium dioxide is photocatalytic, absorbing ultra-violet light, which with oxygen from the air, breaks down or loosens any organic dirt on the surface. Additionally, the surface is hydrophilic, causing rainwater to spread evenly over the surface, rather than running down in droplets, thus uniformly washing the surface and preventing any unsightly streaks or spots appearing when the surface dries. Self-cleaning glass has a slightly greater mirror effect than ordinary float glass, with a faint blue tint. It is available as normal annealed glass also in toughened or laminated form. The surface coating, which reduces the transmittance of the glass by about 5%, is tough, but as with any glass, can be damaged by scratching. A blue solar control version, suitable for conservatory roofs, reduces the solar heat gain by approximately 60% depending on the thickness of glass used and its combination in a double-glazed system.

**CLEAR WHITE GLASS**

Standard float glass is slightly green due to the effect of iron oxide impurities within the key raw material sand.
However, clear white glass can be produced, at a greater cost, by using purified ingredients. The light transmittance of clear white glass is approximately 2% greater than that of standard glass for 4 and 6 mm glazing. Unlike standard glass, which appears green at exposed polished edges, clear white glass is virtually colourless. Clear white glass is normally required for the external leaf in double-glazing units within the A rating of the Window Energy Rating (WER) classification.

**PATTERNED GLASS**

A wide range of 3, 4, 6, 8 and 10 mm patterned glass is commercially available, offering a range of obscuration factors depending on the depth and design of the pattern as illustrated in Fig. 7.6. The classification of obscuration factors differs between manufacturers with 1 (lowest) to 5 (highest) for Pilkington and 1 (highest) to 10 (lowest) for Saint Gobain. The degree of privacy afforded by the various glasses is dependent not only on the pattern but also on the relative lighting levels on either side and the proximity of any object to the glass. The maximum stock sheet size is 3300 × 1880 mm.

Patterned glass is manufactured to BS EN 572-5: 2004, from a ribbon of molten cast glass, which is passed through a pair of rollers one of which is embossed. Certain strong patterns, such as *Cotswold* or *reeded*, require client choice of orientation, whereas the more flowing designs may need appropriate matching. Patterned glasses may be toughened, laminated or incorporated into double-glazing units for thermal, acoustic or safety considerations; a limited range is available in bronze tinted glass. Wired patterned glass (BS EN 572-6: 2004) is available with a 12.5 or 25 mm square mesh.

**SCREEN-PRINTED GLASS**

White or coloured ceramic frit is screen printed onto clear or tinted float glass, which is then toughened and heat soaked, causing the ceramic enamel to fuse permanently into the glass surface. Standard patterns (Fig. 7.7) or individual designs may be created, giving the required level of solar transmission and privacy. Screen printed glass, which is colourfast and abrasion-resistant, is usually installed with the printed side as the inner face of conventional glazing. The lace pattern on the John Lewis, Leicester, store is printed on two layers of glass to give views out when viewed frontally from within the retail outlet, but when viewed obliquely from street level the patterns are displaced, increasing the privacy. The pattern (Fig. 7.8) draws its inspiration from Leicester’s history of textiles and weaving, the John Lewis tradition of quality fabrics and the translucency of Indian saris worn by the local Asian population.

**DECORATIVE ETCHED AND SANDBLASTED GLASS**

Acid etched glass, 4 and 6 mm in thickness, is available to a maximum sheet size of 2140 × 1320 mm with a small range of patterns (Fig. 7.9). These glasses have a low obscuration factor and should not be used in areas of high humidity, as condensation or water causes temporary loss of the pattern. Etched glasses need to be handled carefully on site, as oil, grease and finger marks are difficult to remove completely. Etched glasses may be toughened or laminated; when laminated, the etched side should be outermost to retain the pattern effect, and when incorporated into double glazing, the etched glass forms the inner leaf with the etched face towards the air gap. As with embossed glass, pattern matching and orientation is important. Similar visual effects can be achieved by sandblasting techniques, although the surface finish is less smooth. Patterns may be clear on a frosted background, or the reverse, depending on the aesthetic effect and level of privacy required. Additionally, glass with both a textured and etched finish is available.

**DECORATIVE COLOURED GLASS**

Traditional coloured glass windows constructed with lead came s, soldered at the intersections and wired to saddle bars, are manufactured from uniform pot, surface flashed or painted glasses. For new work, additional support is afforded by the use of lead came s with a steel core and non-corroding saddle bars of bronze or stainless steel should be used. The basilica at Brioude, France (Fig. 7.10), is enhanced by a complete set of modernistic coloured glass windows, installed in 2008, ranging in colours from soft blues to brilliant red and orange.

A three-dimensional effect is achieved by fixing with ultraviolet sensitive adhesive, coloured bevelled glass to clear or coloured sheet glass, the thin edges (1.5–2.5 mm) being covered with adhesive lead strip. Such effects can also be simulated by the use of coloured polyester or vinyl film and lead strip acrylic bonded to a single sheet of clear glass. The base glass
Fig. 7.6 Selection of patterned (texture) glasses with appropriate obscuration factors. Copyright © images reproduced by permission of Pilkington Plc.
Fig. 7.7 Screen printed glass

may be toughened or laminated as appropriate, and the decorative coloured glass laminate can be incorporated into standard double-glazing units. The effect of coloured glass can also be achieved by using a coloured polyvinyl butyral interlayer within laminated glass. Various colour effects and tones can be achieved by combining up to four different-coloured interlayers and, if required, a variety of patterns including spots, stripes, squares and dots. These laminated coloured and patterned glasses have the same impact resistance and acoustic insulation as standard clear laminated glass of the same dimensions.

GEORGIAN WIRED GLASS

Georgian wired glass (BS EN 572-3: 2004) is produced by rolling a sandwich of a 13 mm electrically welded steel wire mesh between two ribbons of molten glass. This produces the standard cast 7 mm sheet, suitable when obscuration is required. For visual clarity the cast product is subsequently ground with sand and water and then polished with jeweller’s rouge to 6 mm sheet (Fig. 7.11). Both the cast and polished grades have a light transmission of 80%. Wired glass is not stronger than the equivalent thickness of annealed glass; however, when cracked, the pieces remain held together.

On exposure to fire, the wire mesh dissipates some heat, but ultimately Georgian glass will crack, particularly if sprayed with water when hot. However, the wire mesh holds the glass in position, thus retaining its integrity and preventing the passage of smoke and flame. Accidental damage may cause the breakage of the glass, but again it is retained in position by the mesh, at least until the wires are affected by corrosion.

Georgian wired glass is available in sheet sizes up to 1985 × 3500 mm (cast) and 1985 × 3300 mm (polished). It is easily cut and can be laminated to other glasses but cannot be toughened. Standard Georgian glass is not considered to be a safety glass to BS 6206: 1981, which defines three classes with decreasing impact resistance down from Class A to Class C. However, certain laminates or products with increased wire thickness do achieve the impact resistance standards for safety glass to BS 6206: 1981 and should be marked accordingly. They may therefore be used in locations requiring safety glass according to Part N of the Building Regulations and BS 6262-4: 2005.

TOUGHENED GLASS

Toughened glass (Fig. 7.12) is up to four or five times stronger than standard annealed glass of the same thickness. It is produced by subjecting preheated annealed glass at 650°C to rapid surface cooling by the application of jets of air. This causes the outer faces to be set in compression with balancing tension forces within the centre of the glass. As cracking within glass commences with tensile failure at the surface, much greater force can be withstood before this critical point is reached.

Toughened glass cannot be cut or worked, and therefore all necessary cutting, drilling of holes and grinding or polishing of edges must be completed in advance of the toughening process. In the roller hearth horizontal
Fig. 7.8 Screen printed glass facade — John Lewis Department Store, Leicester. *Architects:* Foreign Office Architects. *Photographs:* Arthur Lyons
Fig. 7.9 Selection of etched glasses — Oriel™ collection. Copyright © images reproduced by permission of Pilkington Plc.
toughening process, some bow, roller wave distortions and end edge sag may develop but these will be within narrow tolerances; however, they may be observed in the more reflective glasses when viewed from the outside of a building. In the vertical toughening process the sheet is held by tongs, which leave slight distortions where they have gripped the glass.

Toughened glass will withstand considerable extremes of temperature and sudden shock temperatures. If broken, it shatters into small granules which are not likely to cause the serious injuries associated with the accidental breakage of annealed glass. The form of breakage for toughened glass is defined in BS EN 12600: 2002. To be classified as a safety glass toughened glass must be tested and marked according to required standard BS 6206: 1981. When toughened glass is specified for roof glazing, balustrades and spandrel panels, it is subjected to a heat soaking at 290°C, a process which is destructive to any substandard units. This removes the low risk of spontaneous breakage of toughened glass on site, caused by the presence of nickel sulphide inclusions within the material. All standard float, coated, rough cast and some patterned glasses can be toughened. Toughened glass is defined by the standard BS EN 12150-1: 2000.

**HEAT-STRENGTHENED GLASS**

Heat-strengthened glass is manufactured by a process similar to toughening, but with a slower rate of cooling which produces only half the strength of toughened glass. On severe impact, heat-strengthened glass breaks into large pieces like annealed glass, and therefore alone is not a safety glass. It does not require heat soaking to prevent the spontaneous breakage which occurs occasionally with toughened glass. Heat-strengthened glass is frequently used in laminated glass where the residual strength after fracture gives some integrity to the laminate. Typical applications include locations where resistance to wind pressure is necessary, such as the upper storeys and corners of high buildings, and also in spandrels where there is an anticipated higher risk of thermal cracking. Modern applications of laminated heat-strengthened glass include the use of 12 mm thick 1.5 × 3 m roofing panels developed by Arup in
Fig. 7.12 Process for manufacturing toughened glass

Flintholm, Copenhagen. The residual strength of the laminate prevents the glazing falling out of the frame if broken, reducing the hazard to users of the transport interchange building.

LAMINATED GLASS

Laminated glass (Fig. 7.13) is produced by bonding two or more layers of glass together with a plastic interlayer of polyvinyl butyral (PVB) sheet or a polymethyl methacrylate low-viscosity resin. The low-viscosity resin is more versatile as it allows for the manufacture of curved laminates or the incorporation of patterned glasses. The lamination process greatly increases the impact resistance over annealed glass of the same thickness. Furthermore, on impact, whilst the glass laminations crack, they do so without splintering or disintegration, being held together by the interlayer. The form of breakage for laminated glass is defined in BS EN 12600: 2002. Laminated glass can be defined as a safety glass provided it achieves the appropriate class standard to BS 6206: 1981.

The all-glass facade of the Prada Store in Tokyo by Herzog & de Meuron (Fig. 7.14) is manufactured from rhomboidal panes of PVB-bonded laminated glass; some units are flat, but others are curved with either the convex or concave surface to the outside giving rise to varied colours by reflection. The perimeters of the double-glazed units are necessarily flat to ensure correct positioning and sealing.

Fig. 7.13 Laminated glass — Glazing resistant to manual attack and bullet-resistant glazing

The impact resistance of laminated glass may be increased by the use of thicker interlayers, increasing the numbers of laminates, or by the inclusion of polycarbonate sheet. Typically, glazing resistant to manual attack (BS 5544: 1978) has two or three glass laminates, while bullet-resistant glass (BS EN 1063: 2000) has four or more glass laminates depending on the anticipated calibre and muzzle velocity. To prevent spalling, the rear face of bullet-resistant glass may be sealed with a scratch-resistant polyester film, and for fire protection, Georgian wired glass may be incorporated. Laminated glasses made from annealed glass can be cut and worked after manufacture.

Specialist properties for X-ray or ultraviolet light control can be incorporated into laminated glasses by appropriate modifications to the standard product.
The latter reduces transmissions in the 280–380 nm wavelength ranges, which cause fading to paintings, fabrics and displayed goods.

The incorporation of specialist film interlayers within laminated glass offers further diversity. Interlayers may have variable optical and thermal transmission properties, by incorporating photochromic, thermochromic, thermotropic or electrochromic materials. Alternatively, the interlayer may diffract the incident light through specified angles, as within prisms and holograms. Thus, within a deep room, natural light can be refracted up to a white ceiling for dispersion further back within the space.

**PLASTIC FILM LAMINATES**

A range of transparent and translucent plastic films is readily applied internally or externally to modify the properties of glass. These include patterned films to create privacy, manifestation films to prevent people accidentally walking into clear glass screens or doors and reflective films to reduce solar gain and glare. Safety films, as applied to overhead glazing on the Waterloo train terminal (Fig. 5.1, Chapter 5, page 165), remove the risk of injury from falling glass when nickel sulphide inclusions cause the spontaneous failure of toughened glass. Similarly, security films ensure that glass damaged by accidental impact or vandalism remains in place.

**FIRE-RESISTANT GLASS**

The ability of a particular glass to conform to the criteria of integrity and insulation within a fire is a measure of its fire resistance (Table 7.1). However, to achieve a specified performance in fire it is necessary to ensure that the appropriate framing, fixings and glass have all been used, as fire resistance is ultimately dependent on the whole glazing system and not the glass alone.

The European specification (BS EN 13501-2: 2003) for the fire resistance of a material or an assembly is classified by its performance against the criteria: integrity (E), insulation (I), radiation (W) and also, not normally relevant to glass, load bearing capacity (R). The standard time periods are 15, 20, 30, 45, 60, 90, 120, 180 and 240 minutes.

**Typical classifications:**

- E30 integrity only 30 minutes
- EW30 integrity and radiation protection for 30 minutes
- EI30 integrity and insulation for 30 minutes
- E60EI30 integrity 60 minutes and insulation 30

**Non-insulating glass**

Integrity-only non-insulating glass products will prevent the passage of flame, hot gases and smoke, but will allow heat transmission by radiation and conduction; thus, ultimately further fire spread may occur through the ignition of secondary fires. Intense radiation through glass areas may render adjacent escape routes impassable.

Georgian wired glass offers up to a 120 minutes’ fire resistance rating with respect to integrity, depending on the panel size and fixings. If the glass cracks within a fire, its integrity is retained, as the wire mesh prevents loss of the fractured pieces. Georgian wired glass is cheaper than insulated fire-resistant glasses and may be cut to size on site.
### Table 7.1  Typical fire resistance properties of glass

<table>
<thead>
<tr>
<th>Type</th>
<th>Thickness (mm)</th>
<th>E integrity (min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Georgian wired glass—cast</td>
<td>7</td>
<td>30–120</td>
</tr>
<tr>
<td>Georgian wired glass—polished</td>
<td>6</td>
<td>30–120</td>
</tr>
<tr>
<td>Toughened standard glass</td>
<td>6–19</td>
<td>30–60 (90 in steel)</td>
</tr>
<tr>
<td>Toughened borosilicate glass</td>
<td>6–12</td>
<td>30–120</td>
</tr>
<tr>
<td>Glass blocks</td>
<td>80</td>
<td>30–60</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Type</th>
<th>Thickness (mm)</th>
<th>EI integrity and insulation (min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Laminated intumescent glass</td>
<td>7</td>
<td>20</td>
</tr>
<tr>
<td></td>
<td>19</td>
<td>45</td>
</tr>
<tr>
<td></td>
<td>27</td>
<td>60</td>
</tr>
<tr>
<td></td>
<td>54 (double-glazing unit)</td>
<td>120</td>
</tr>
<tr>
<td>Laminated gel interlayer glass</td>
<td>20–22</td>
<td>30</td>
</tr>
<tr>
<td></td>
<td>32</td>
<td>60</td>
</tr>
<tr>
<td></td>
<td>38</td>
<td>90</td>
</tr>
<tr>
<td></td>
<td>55</td>
<td>120</td>
</tr>
</tbody>
</table>

**Note:**

Fire resistance data are significantly dependent on glass thickness, glazing size, aspect ratio and the metal, timber, plastic or butt-jointed glazing system.

Toughened calcium-silica-based glasses can achieve 90 minutes’ fire resistance with respect to integrity. The glazing remains intact and transparent, but will break up into harmless granules on strong impact if necessary for escape. Toughened glass cannot be cut or worked after manufacture.

Boro-silicate glass, with a low coefficient of expansion, is more resistant to thermal shock than standard annealed glass and does not crack on exposure to fire. It can be thermally strengthened to increase its impact resistance. Certain ultra-heat-resistant ceramic glasses have a zero coefficient of thermal expansion. As a result, they can resist temperatures up to 1000°C and the thermal shock of a cold water spray when heated by fire.

### Insulating glass

Insulating glasses are manufactured from float glass laminated with either intumescent or gel materials. Intumescent laminated glass has clear interlayers which, on exposure to fire, expand to a white opaque material, inhibiting the passage of conductive and radiant heat (Fig. 7.15). The glass layers adjacent to the fire crack but retain integrity owing to their adhesion with the interlayers. The fire resistance, ranging between 30 and 120 minutes for insulation and integrity, depends on the number of laminations, usually between 3 and 5. To avoid the green tint associated with thick laminated glass, a reduced-iron-content glass may be used to maintain optimum light transmission. For exterior use, the external grade has an additional glass laminate with a protective ultraviolet filter interlayer. Laminates

**Fig. 7.15**  Fire-resistant glass with intumescent material laminates
may be manufactured with tinted glass or combined with other patterned or solar control glasses. Insulating glass is supplied cut to size and should not be worked on site. Double-glazed units with two leaves of intumescent laminated glass give insulation and integrity ratings of 120 minutes; alternatively, units may be formed with one intumescent laminate in conjunction with specialist solar control glasses. The fire-resistant laminated glasses conform to the requirements of BS 6206: 1981, in respect of Class A impact resistance. Typical systems for integrity and insulation are 15 mm thickness for EI30 and 27 mm for an external EI60 rating.

Gel insulated glasses are manufactured from laminated toughened glass with the gel layer sandwiched between two or more glass layers. In the event of fire, the gel interlayer, which is composed of a polymer-containing aqueous inorganic salt solution, absorbs heat by the evaporation of water and produces an insulating crust. The process is repeated layer by layer. Depending on the thickness of the gel layer, fire resistance times of 30, 60 or 90 minutes are achieved.

Partially insulating glass

Partially insulating glass, consisting of a 10 mm triple laminate of float glass with one intumescent interlayer and one polyvinyl butyral layer, offers a modest increase in fire resistance over non-insulating glass. Surface treatments can also increase the heat reflectance of both glass faces.

ENERGY CONSERVATION

The Building Regulations (Approved Document Part L1A: Conservation of fuel and power in new dwellings [2006 edition]) requires that all new dwellings have predicted carbon dioxide emissions – Dwelling Emission Rate (DER) – not greater than the Target Emission Rate (TER) determined for the equivalent notional building of the same size and shape. The TER, expressed as the quantity of CO₂ in kilogram emitted per square metre of floor area per year, is calculated for dwellings of less than or equal to 450 m² total floor area by the Standard Assessment Procedure (SAP 2005 edition). For all dwellings above 450 m² floor area, the Simplified Building Energy Model (SBEM) calculation is required. The following reference figures are area-weighted average limits for guidance. However, for most building enhanced specifications are likely to be required in order to achieve the required carbon dioxide TER.

<table>
<thead>
<tr>
<th>Component</th>
<th>U-value (W/m² K)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wall</td>
<td>0.35</td>
</tr>
<tr>
<td>Floor</td>
<td>0.25</td>
</tr>
<tr>
<td>Roof</td>
<td>0.25</td>
</tr>
<tr>
<td>Window/roof window/door</td>
<td>2.2</td>
</tr>
</tbody>
</table>

The Building Regulations also require provision to prevent excessive summer solar gains causing high internal temperatures. This can be achieved by the appropriate use of window size and orientation, solar control with shading or specialist glazing systems, ventilation and high thermal capacity. The Building Regulations set limits on services performance and envelope airtightness (including sample pressure testing), and, in addition, they require each new building to be supplied with guidance on its energy-efficient operation.

The TER method gives a good degree of flexibility in relation to the design by allowing trade-offs between different energy-saving factors. The benefits of useful solar radiation gains may also be taken into account.

The Building Regulations (Approved Document Part L2A: Conservation of fuel and power in new buildings other than dwellings [2006 edition]) requires the predicted carbon dioxide emissions – Building Emission Rate (BER) to be no greater than the Target Emission Rate (TER) calculated by the Simplified Building Energy Model (SBEM) or other approved software tools. The area-weighted average limits for elements are set at the same standards as for new dwellings. However, the appropriate BER, rather than individual element U-values, is required for compliance.

Building Regulations Approved Documents Part L1B and L2B refer to work on existing dwellings and other buildings, respectively. Guidance is quoted on the reasonable provision and standards for new extensions and replacement of existing thermal elements, but again full compliance in all aspects of the regulations is required.

Standards for new elements in existing buildings:

<table>
<thead>
<tr>
<th>Component</th>
<th>U-value (W/m² K)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wall</td>
<td>0.28</td>
</tr>
<tr>
<td>Floor</td>
<td>0.22</td>
</tr>
<tr>
<td>Pitched roof – insulation at ceiling level</td>
<td>0.16</td>
</tr>
<tr>
<td>Pitched roof – insulation at rafter level</td>
<td>0.16</td>
</tr>
<tr>
<td>Flat roof – integral insulation</td>
<td>0.16</td>
</tr>
</tbody>
</table>
Double and triple glazing

Whenever the internal surface of exterior glazing is at a lower temperature than the mean room surface temperature and the internal air temperature, heat is lost by a combination of radiation exchange at the glass surfaces, air conduction and air convection currents inside and out, also by conduction through the glass itself. This heat loss can be reduced considerably by the use of multiple glazing with air, partial vacuum or inert gas fill (Fig. 7.16).

Gas-filled glazing units

Double glazing reduces the direct conduction of heat by the imposition of an insulating layer of air between the two panes of poorly insulating glass. The optimum air gap is between 16 and 20 mm, as above this value convection currents between the glass panes reduce the insulating effect of the air. Argon is now frequently used as the filling agent as it has a lower thermal conductivity than air, thus reducing heat transfer by conduction. The standard is 90% argon, because of the difficulties in ensuring that all air has been flushed out of the void space. The use of krypton or even xenon within a 16 mm double-glazing gap in conjunction with low-emissivity glass can achieve a $U$-value of 0.8 W/m² K. These rarer inert gases are now commercially available. (thermal conductivities $\lambda$ air = 0.025, $\lambda$ argon = 0.017, $\lambda$ krypton = 0.009,

Fig. 7.16  Mechanism of heat loss through single and double glazing (after Button and Pye: 1993)
### Table 7.2 U-values for single- and multiple-glazing systems—
typical centre pane values

<table>
<thead>
<tr>
<th>Glass system</th>
<th>U-value (W/m² K)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single clear glass</td>
<td>5.8</td>
</tr>
<tr>
<td>Double clear glass</td>
<td>2.8</td>
</tr>
<tr>
<td>Double clear glass with argon fill</td>
<td>2.7</td>
</tr>
<tr>
<td>Double clear glass with hard low-emissivity coating</td>
<td>1.7</td>
</tr>
<tr>
<td>Double clear glass with hard low-emissivity coating and argon fill</td>
<td>1.5</td>
</tr>
<tr>
<td>Double clear glass with soft low-emissivity coating and argon fill</td>
<td>1.4</td>
</tr>
<tr>
<td>Double clear glass with soft low-emissivity coating and argon fill</td>
<td>1.2</td>
</tr>
<tr>
<td>Triple clear glass with two low-emissivity coatings and two argon fills</td>
<td>0.8</td>
</tr>
<tr>
<td>Triple clear glass with two low-emissivity coatings and two xenon fills</td>
<td>0.4</td>
</tr>
</tbody>
</table>

Note:
The data relates to 4 mm standard glass and 16 mm spacing.
Argon is 90% filling.
Overall window ratings will differ considerably according to the perimeter materials and construction system.

$\lambda_{\text{xenon}} = 0.005 \text{ W/m K}$; BS EN ISO 10456: 2007.

Similar further reductions in conducted heat can be achieved by the incorporation of an additional air space within triple glazing. Thin low-emissivity films suspended within the cavity can further reduce the U-values of double-glazing units to as low as 0.6 W/m² K. Typical centre pane U-values are shown in Table 7.2.

**Vacuum glazing units**
Vacuum glazing units incorporating a gap of only 0.2 mm have the same level of thermal efficiency as standard gas-filled low-emissivity systems. The sheets of clear float glass and low-emissivity glass are kept apart by 0.5 mm diameter micropacers at 20 mm centres, which are only visible on close inspection. The vacuum is created through a hole in the inner pane, which is permanently sealed with a 15 mm cover. Units can be manufactured up to $2.4 \times 1.35$ m, and can be fitted as replacements for existing single glazing. The centre pane U-value for this vacuum system is 1.4 W/m² K.

**Window energy ratings**
The British Fenestration Rating Council (BFRC) system of window energy rating (WER) bands for complete units, based on the range A (best) to G (poorest), gives guidance to specifiers on energy efficiency. The ratings take into account a combination of the three key factors which affect performance; namely, thermal efficiency for heat retention (U-value), solar heat gain (g-value between zero and 1) and heat loss by air leakage (L-factor). The solar heat gain and U-values relate to the whole unit, not just the glazed areas. The leakage rate is taken for average conditions. It is determined as $\text{m}^3$/h/m² at a pressure of 50 Pa and then scaled down by a factor of 20 to a realistic level. It is then converted into an equivalent heat loss (W/m² K) to be added to the heat gains and losses. The window energy rating (WER) bands are colour coded from dark green (A) through light green (C) to red (F) and brown (G) and they measure the overall energy performance of the windows as the total annual energy flow (kW hour/m²/year). Windows in the A band will give a small positive energy contribution to the building, whilst C band windows contribute an energy loss of the order of 15 kW hour/m²/year. The scheme, supported by the Glass and Glazing Federation, is based on a standard window of $1.40 \times 1.25$ m with a central mullion, one fixed light and one opening light to ensure equity of comparison between different products.

The industry standard for domestic windows is currently the C rating, and this will be required by the Building Regulations around 2010. However, manufacturers are developing the higher rated units at less premium costs in response to client demands. A major advancement in double-glazing units has been the development of warm edge spacer bar technology and the increased thermal efficiency of frames. Materials now used for spacers include thermoplastics and GRP. In some cases steel reinforcement previously used in PVC-U frames is omitted or reduced to improve thermal performance, but this may diminish the frame strength.

Windows for domestic housing are dominated by PVC-U and timber systems, but aluminium and steel frames also have a significant market share. Additionally composite windows, typically with aluminium for the exterior and timber on the interior, combine the advantages of both materials including aesthetic and maintenance properties; they also have the potential for the installation of a wide range of sealed double- or triple-glazing units for thermal and acoustic efficiency.

An appropriately insulated and sealed PVC-U framed system with double glazing, using 4 mm clear white glass for the outer pane, 4 mm vacuum sputtered soft-coated low-emissivity glass as the inner pane, PVC-U thermal inserts, 20 mm spacing with 90% argon fill, can achieve the highest A rating. The typical combination of factors to give the A rating would
be an overall window $U$-value of 1.4 W/m$^2$ K with a solar factor ($g$) of 0.45 and effective air leakage ($L_o$) of 0.0 W/m$^2$ K. An A+ rating ($U$-value 0.8 W/m$^2$ K) can be achieved with inert-gas-filled triple-glazing set in multi-chamber reinforced PVC-U frames.

Relationship between BFRC window ratings and annual heat loss per year:

<table>
<thead>
<tr>
<th>WER rating</th>
<th>Energy loss per year (kW.hour/m$^2$/year)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>greater than zero (positive energy gain)</td>
</tr>
<tr>
<td>B</td>
<td>10 to less than 0 (small energy loss)</td>
</tr>
<tr>
<td>C</td>
<td>20 to less than 10</td>
</tr>
<tr>
<td>D</td>
<td>30 to less than 20</td>
</tr>
<tr>
<td>E</td>
<td>50 to less than 30</td>
</tr>
<tr>
<td>F</td>
<td>70 to less than 50</td>
</tr>
<tr>
<td>G</td>
<td>less than 70 (high energy loss)</td>
</tr>
</tbody>
</table>

Figure 7.17 illustrates a typical British Fenestration Rating Council average C energy rating.

**Low-emissivity glass**

Low-emissivity glasses are manufactured from float glass by the application of a transparent low-emissivity coating on one surface. The coating may be applied either online, within the annealing lehr at 650°C, as a pyrolytic hard coat, or off-line after glass manufacture by magnetic sputtering under vacuum which produces a softer coat. On-line manufactured low-emissivity glasses and some off-line products may be toughened after coating; also off-line low-emissivity coatings may be applied to previously toughened glass. The on-line surface coating is more durable and is not normally damaged by careful handling.

Low-emissivity glass functions by reflecting back into the building the longer-wavelength heat energy associated with the building’s occupants, heating systems and internal wall surfaces, whilst allowing in the transmission of the shorter-wavelength solar energy (Fig. 7.18). The incoming solar energy is absorbed by the internal walls and re-radiated as longer-wavelength energy, which is then trapped by the low-emissivity coating on the glass.

Low-emissivity coatings can reduce by three quarters the radiant component of the thermal transfer between the adjacent surfaces within double glazing. The reduction in emissivity of standard uncoated glass, from 0.84 to below 0.16, gives a decrease in $U$-value from 2.8 W/m$^2$ K for standard double glazing to 1.7 W/m$^2$ K with low-emissivity glass. Frequently, low-emissivity glass is protected in use within sealed
double-glazed units. The outer leaf in the double-glazing system may be clear or any other specialist glass for security or solar control. Pyrolytic low-emissivity coatings are suitable for incorporation into secondary glazing for existing windows. The emissivities of low-E coatings range for hard coats from 0.15 to 0.20 and for soft coatings from 0.04 to 0.10.

Following the changes in the requirements for energy conservation, low-emissivity glass double glazing has effectively become the standard for all new building works, as in the Swiss Re building in central London (Fig. 7.19).

**Double-glazing units**

Hermetically sealed double-glazing units are usually manufactured with aluminium or thin stainless steel spacers which incorporate moisture-adsorbing molecular sieve or silica gel and are sealed with polyisobutylene, polyurethane, polysulphide or epoxy-sulphide. The primary seal is backed up by a secondary seal, usually a two-part silicone to prevent leakage, and is covered with a protective cap (Fig. 7.20). Greater thermal efficiency, to reduce cold bridging and the risk of condensation, is afforded by thermoplastic or GRP spacers coated with a thin foil of aluminium or stainless steel to prevent gas loss. Timber frames offer good insulation. PVC frames use multi-chamber systems and plastic or minimal steel reinforcement to reduce thermal conduction. Aluminium and steel frames require the inclusion of thermal breaks to reduce the risk of surface condensation and significant heat loss. Table 7.3 shows the relationship between centre pane and overall window $U$-values for a sample of timber, PVC-U, aluminium and steel frame systems.

**SOLAR CONTROL GLASSES**

Solar control glasses offer a modified passage of light and heat energy compared to clear glass of the same thickness (Fig. 7.21). A descriptive code indicates the relative quantities of light and heat transmitted for a single sheet of a particular glass (e.g. 49/58 for 6 mm bronze glass) and this can be related to the equivalent data for clear float glass (87/83 for 6 mm clear glass). However, solar control glasses are always used within an insulating glass unit (IGU), so the overall transmittance of light and heat is important in specification (Table 7.4). The method of solar control is a combination of heat absorption and heat reflection by the outer pane of the IGU. The absorption of some of the incident solar radiation by the outer pane
causes the glass to warm. The glass then dissipates this absorbed heat mainly to the outside due to external air movement. The outer pane of the IGU also reflects some solar energy dependent on any coatings applied online during the float process or subsequently as sputtered surface treatment. Solar control glasses are typically grey, bronze, green, blue or silver, and may be toughened or laminated for use in hazardous areas. Additionally, double-glazed units can be augmented by adjustable blinds or louvres. One system incorporates magnetically adjustable Venetian blinds into the 18 or 20 mm sealed double-glazing units. Solar control reflective film is available in a range of grades, thickness and colours for retrofitting onto existing glazing.

**ACOUSTIC CONTROL**

The level of sound reduction by glazing is influenced by the mass of the glass and the extent of air leakage around the opening lights. Sound insulation for single glazing follows the mass law – doubling the glass thickness reduces sound transmission by approximately 4 dB. Toughened, patterned and wired glass of the same thickness respond as for plain glass, but laminated glasses based on a 1 mm layer of polymethyl methacrylate (PMMA) or a thick polyvinyl butyral (PVB) interlayer (one or two 0.4 mm layers) have enhanced sound insulation properties. The plastic interlayers, because they are soft material,

<table>
<thead>
<tr>
<th>Glass separation (mm)</th>
<th>Timber</th>
<th>PVC-U</th>
<th>Aluminium</th>
<th>Steel</th>
</tr>
</thead>
<tbody>
<tr>
<td>Centre pane U-value (W/m² K)</td>
<td>1.2</td>
<td>1.2</td>
<td>1.2</td>
<td>1.2</td>
</tr>
<tr>
<td>Standard unit (W/m² K)</td>
<td>1.8</td>
<td>1.7</td>
<td>1.9</td>
<td>1.9 – 2.5</td>
</tr>
<tr>
<td>High-performance unit (W/m² K)</td>
<td>1.5</td>
<td>1.4</td>
<td>1.5*</td>
<td>1.7 – 2.4**</td>
</tr>
</tbody>
</table>

Notes:
- Timber, PVC-U and aluminium window U-values are based on the standard size of 1.48 × 1.23 m with a central mullion, one fixed and one opening light. Units combine soft coat low-emissivity glass (0.05) and 90% argon fill.
- High-performance units have enhanced edge spacers.
- Aluminium windows are slim line with a polyamide thermal break.
- Building Regulations Part L require a maximum average U-value of 2.2 W/m² K for metal and 2.0 W/m² K for timber and PVC-U.
- *Uses 20 mm separation and a centre pane U-value of 1.18 W/m² K.
- **Uses 8 mm krypton or 6 mm xenon gap.
Table 7.4  Characteristics of typical solar control glasses

<table>
<thead>
<tr>
<th>Colour</th>
<th>Manufacturer’s code</th>
<th>Light transmittance</th>
<th>Light reflectance</th>
<th>Solar radiant heat transmission</th>
<th>Shading coefficient</th>
</tr>
</thead>
<tbody>
<tr>
<td>Green</td>
<td>75/79</td>
<td>0.67</td>
<td>0.12</td>
<td>0.48</td>
<td>0.55</td>
</tr>
<tr>
<td>Bronze</td>
<td>49/58</td>
<td>0.44</td>
<td>0.08</td>
<td>0.48</td>
<td>0.55</td>
</tr>
<tr>
<td>Grey</td>
<td>43/58</td>
<td>0.39</td>
<td>0.07</td>
<td>0.46</td>
<td>0.53</td>
</tr>
<tr>
<td>Blue</td>
<td>50/27</td>
<td>0.50</td>
<td>0.19</td>
<td>0.28</td>
<td>0.33</td>
</tr>
<tr>
<td>Silver</td>
<td>50/30</td>
<td>0.49</td>
<td>0.39</td>
<td>0.31</td>
<td>0.36</td>
</tr>
<tr>
<td>Clear</td>
<td>70/40</td>
<td>0.70</td>
<td>0.10</td>
<td>0.43</td>
<td>0.49</td>
</tr>
<tr>
<td>Clear</td>
<td>30/17</td>
<td>0.30</td>
<td>0.26</td>
<td>0.19</td>
<td>0.21</td>
</tr>
<tr>
<td>Standard clear float glass</td>
<td>0.87</td>
<td>0.08</td>
<td>0.83</td>
<td></td>
<td>0.95</td>
</tr>
</tbody>
</table>

Note: 6 mm solar control glass, 16 mm space and 6 mm standard glass inner pane.

change the frequency response of the composite sheet in comparison with the same weight of ordinary glass and also absorb some of the sound energy.

Double glazing for sound insulation should be constructed with the component glasses differing in thickness by at least 30% to reduce sympathetic resonances; typically 6 and 10 mm would be effective. Where the passage of speech noise (630–2000 Hz) is to be reduced, double-glazing units filled with sulphur hexafluoride have been used, but this is detrimental to the attenuation of traffic and other low-frequency noises within the range 200–250 Hz; also it is now considered environmentally inappropriate. A typical high-performance double-glazing unit, giving a U-value of 1.3 W/m² K and a sound reduction of 35 dB, would be constructed from 6.4 mm laminated glass with a 15.5 mm argon-filled cavity and a 4 mm inner pane of low-emissivity glass. For enhanced sound insulation an air gap of at least 100 mm is required, with the economical optimum being 200 mm. The reveals should be lined with sound-absorbing material such as fibreboard, to reduce reverberation within the air space. All air gaps must be fully sealed with opening lights closed by multipoint locking systems and compressible seals. An air gap corresponding to only 1% of the window area can reduce the efficiency of sound insulation by 10 dB.

COLOURED ENAMELLED GLASS

Opaque or translucent enamelled glass spandrel panels for curtain walling can be manufactured to match or contrast with the range of vision area solar control glasses. Manufactured from toughened heat-soaked glass for impact and thermal shock resistance, they may be single or double glazed, with integral glass fibre or polyurethane foam insulation and an internal finish. Panels are colourfast and scratch-resistant, and may be manufactured from plain, screen-printed or decorative glass.

SPECIALIST GLASSES

One-way observation glass

Where unobserved surveillance is required, one-way observation mirror glass can be installed. In order to maintain privacy, the observer must be at an illumination level no greater than one seventh that of the observed area and wear dark clothing. From the observed area the one-way observation glass has the appearance of a normal mirror. One-way observation glass is available annealed, toughened or laminated.

Mirror glass

Standard mirror glass is manufactured by the chemical deposition onto float glass of a thin film of silver, from aqueous silver and copper salt solutions. The film is then protected with two coats of paint or a plastic layer. A recent development is the production of mirror glass by chemical vapour deposition within the float glass process. Mirror glass is produced by the online application of a three-layer coating of silicon–silica–silicon, which acts by optical interference to give the mirror effect. Mirror glass manufactured by this process is less prone to deterioration and may be more easily toughened, laminated or bent than traditional mirror glass.

Anti-reflection glass

Treatment of standard float glass can reduce surface reflection from 0.09 to 0.025, thus increasing the transmittance. The coating is applied equally to both faces.
Although used mostly for the protection of displayed art works, this material can be used for interior display windows and dividing screens, and also to reduce multiple reflections from the surfaces of double-glazed units, where both sheets of glass must be anti-reflective.

**Alarm glass**

Glass containing either a ceramic loop or a series of straight wires can be incorporated into an intruder alarm system, which is activated when the glass is broken. Usually, the alarm glass conductive circuit is incorporated into laminated glass on the inside of the outer ply, or fixed to the inner face of the toughened outer pane within a double-glazing system.

**Electrically heatable laminated safety glass**

Electrically heatable laminated safety glass incorporates fine electrically conducting wires which may be switched on when there is the risk of condensation. Typical applications are in areas of high humidity, such as swimming pools, kitchens and glass roofs, particularly when there are significant differences between the internal and external temperatures. Power consumption ranges from 100 W/m² for homes to 500 W/m² for industrial applications depending on the internal environment and external ambient conditions. Electrical connections are made at the perimeter of the glass.

**Dichroic glass**

Dichroic glass has a series of metal oxide coatings which create optical interference effects. These cause the incident light to be split into the spectral colours which, depending on the angle of incidence of the light, are either reflected or transmitted. This effect can be used to create interesting colour patterns, which vary with both the movement of the sun and the observer. One material produces a range of iridescent colours from blue through purple to violet.

**Sphere glass and cracked glass**

Sphere glass consists of an array of glass hemispheres, typically 4 to 8 mm in diameter, fixed to standard toughened glass, and is suitable for use in feature walls, partitions and ceilings. The hemispheres and the base material each may be clear glass, standard or custom coloured. Cracked glass is also a feature material manufactured from a laminate of toughened glass between layers of annealed glass. After curing, the laminate is machined causing the toughened glass to crack, giving the desired aesthetic effect. It is available in a range of colours and mirror finish.

**Antibacterial glass**

Antibacterial glass kills the majority of bacteria which fall onto its surface. The action is due to the presence of silver ions in the surface layer of glass. The silver ions interact with the bacteria and inhibit their division metabolism. This type of glass has potential for reducing the number of hospital-acquired bacterial infections, which is currently a significant health problem.

**Electromagnetic radiation-shielding glass**

Electromagnetic radiation-shielding glass can be used to protect building zones containing magnetically stored data from accidental or deliberate corruption by external electric fields. For maximum security, the conducting laminates within the composite glass should be in full peripheral electrical contact with the metal window frames and the surrounding wall surface screening.

**X-ray protection glass**

X-ray protection glass contains 70% lead oxide, which produces significant shielding against ionising radiation. The glass is amber in colour due to the high lead content. A 6 mm sheet of this lead glass has the equivalent shielding effect of a 2 mm lead sheet against $\gamma$ radiation.

**Sound-generating glass**

Terfenol-D is a magnetostrictive material, which when stimulated by a magnetic field, expands and contracts rapidly producing a large physical force. If a device containing Terfenol-D is attached to the smooth surface of glass and an audio input is fed into the system, then the whole sheet of glass will vibrate, acting as a loudspeaker. Thus, shop windows can be turned into loudspeakers, producing across their surface a uniform sound which can be automatically controlled to just greater than the monitored street noise level, thus avoiding sound pollution. Two devices, appropriately positioned, will generate stereo sound. Magnetostrictive devices will operate similarly on any flat rigid surfaces such as tabletops, work surfaces and rigid
partitions. Terfenol-D is named from the metallic elements iron, terbium and dysprosium from which it is manufactured.

**Light-emitting diode (LED) illuminated glass**

Light-emitting diodes laminated between two sheets of glass and a PVB interlayer can be illuminated by the electrical conductivity of the surface oxide film on special glass, thus eliminating any visible electrical connections. The technology has been used in high-profile buildings in Dubai, California and Neuchâtel, Switzerland. White and colour points of light activated by complex switching sequences are all attainable.

**Manifestation of glass**

Where there is a risk that glazing might be unseen, and thus cause a hazard to the users of a building, particularly large areas at entrances and in circulation spaces, the presence of the glass should be made clear with solid or broken lines, decorative features or company logos at heights between 850 and 1000 mm and also between 1400 and 1600 mm above floor level. In such circumstances, the risk of impact injury must be reduced by ensuring that the glass is either robust, protected, in small panes, or breaks safely. Critical locations with a risk of human impact (BS 6262-4: 2005) are clear glazed panels from floor level to 800 mm, and also floor level to 1500 mm for glazed doors and glazed side panels within 300 mm of doors. Figure 7.22 illustrates a typical example of glass manifestation.

**VARIABLE TRANSMISSION (SMART) GLASSES**

Variable transmission or *smart* glasses change their optical and thermal characteristics under the influence of light (photochromic), heat (thermochromic) or electric potential (electrochromic). These glasses offer the potential of highly responsive dynamic climate control to building facades. These smart materials including thermotropic products are also available as plastic laminates for incorporation into laminated glass systems.

**Photochromic glass**

Photochromic glasses incorporate silver halide crystals, which are sensitive to ultraviolet or short-wave visible light. The depth of colour is related directly to the intensity of the incident radiation and is fully reversible. For use in buildings, these materials have the disadvantage that they respond automatically to changes in solar radiation, rather than to the internal environment within the building.

**Thermochromic glass**

Thermochromic glasses change in transmittance in response to changes in temperature. Like photochromic glass, these materials have the disadvantage of responding to local conditions, rather than to the requirements of the building’s internal environment.

**Electrochromic glass**

Electrochromic glasses change their transmittance in response to electrical switching and are therefore likely to become the basis of *smart* windows. Electrochromic multilayer thin-film systems become coloured in response to an applied low voltage, and are then cleared by reversal of the electric potential. The depth of colouration is dependent on the magnitude...
of the applied d.c. voltage. Optically stable materials, which exhibit electrochromism, are the oxides of tungsten, nickel and vanadium. Electrochromic thin film systems may be laminated to any flat sheet glass.

**Electro-optic laminates**

Electrically operated vision-control glass (*Priva-lite*) consists of a laminated system of glass and polyvinyl butyral layers containing a polymer-dispersed liquid crystal layer, which can be electrically switched from transparent to white/translucent for privacy.

**Intelligent glass**

Conventional glass coatings reduce both light and heat transmission. However, a coating based on tungsten-modified vanadium dioxide allows visible light through at all times, but reflects infrared radiation at temperatures above 29°C. Thus, at this temperature, further heat penetration through the glass is blocked. Therefore the intelligent glass, which has a slight yellow/green colour, admits useful solar gain in cooler conditions but cuts out excessive infra-red solar gain under hot conditions.

**Intelligent glass facades**

An intelligent glass building facade changes its physical properties in response to sensors detecting the external light and weather conditions, thus reducing the energy consumption necessary to maintain the appropriate internal environment. Therefore intelligent facades have ecological significance in reducing global greenhouse emissions and also in reducing operational building costs to clients and users.

Truly intelligent facades capitalise on the incident solar energy striking the facade of a building, adapt the skin functionality to the appropriate thermal control and solar protection and, in addition, may generate electricity through photovoltaic cell systems. Solar control may be provided by switched electrochromic glass or by using laminated prismatic or holographic films which deflect the solar radiation according to its angle of incidence. Additionally, intelligent facades respond to air flows or ground heat sources to ensure appropriate and responsive ventilation. This function is usually achieved by the use of a double-skin facade, which acts as a ventilation cavity. During the heating season the double skin can prewarm the incoming fresh air, and when cooling is required it can remove, by convection, built-up excess heat from the double-glazing unit. Furthermore, excess heat energy can be stored for redistribution when required.

Smart shading systems respond to reduce excess incident solar radiation. Electrically controlled louvres or blinds located between two glass panes open and close according to either a solar detector or a range of weather sensing devices. Such mechanical systems, including the prototype iris diaphragm shading devices used by Jean Nouvel in the Institute de Monde Arabe in Paris, demand considerable maintenance for continued operation.

**Glass supporting systems**

Modern glazing facades are sophisticated as illustrated in Fig. 7.23. The fixing of glazing and particularly solar control glasses should be sufficiently flexible to allow for tolerances and thermal movements. A minimum edge clearance of at least 3 mm is required for single-glazing and 5 mm for double-glazing units. Edge cover
should be sufficient to cope with the design wind loading, with a minimum normally equal to the glass or unit thickness to ensure a neat sight line. Glass thickness should be checked for suitability against predicted wind speeds, modified appropriately by consideration to the effects of local topography, building height and size of the glazing component.

The Pilkington Planar System (Fig. 7.24) offers the designer a flush and uninterrupted facade of glass. The only fixings to be seen on the external facade are the countersunk bolt heads. The system which can be used for single, double or triple glazing, vertically or sloping, is designed such that each glazing unit is separately supported by the mullion system, so there is no restriction on the height of the building. Thermal and wind movement is taken up by the fixing plate, which is sufficiently flexible to allow some rotation of the glass. In the double-glazed system the units are principally supported by the outer pane. Glass-to-glass butt joints are sealed with silicone.

The Planar Triple Glazing System (Fig. 7.25), with one solar control, two low-emissivity glazings and two 16 mm airspaces, can achieve a $U$-value of 0.8 W/m² K. Insulating glass units are manufactured from three panes of toughened and heat-soaked glass, with thicknesses between 4 and 19 mm according to the unit size.

The Financial Times building in Docklands, London (Fig. 7.26), is designed with a long clear facade sandwiched between two aluminium-clad solid ends. The glazed section, $96 \times 16$ m, consists of a single-glazed suspended toughened-glass wall, bolted by circular plate assemblies to external aerofoil forms and intermediate cantilevered arms. This creates a wall of uninterrupted glass which is striking by day and transparent at night when the illuminated printing presses can be seen clearly.

**STRUCTURAL GLASS**

Glass columns are frequently used as fins to restrain excessive deflection caused by wind and other lateral loads to glass facades. The fixings between the facade glazing and fin units are usually stainless steel clamps bolted through preformed holes in the toughened or laminated glass, although silicone adhesives can also be used. BS EN 13022: 2006 indicates that where exposed to UV light the outer seal of any structural glazing unit should be sealed with silicone. Furthermore, BS EN 15434: 2006 indicates that where exposed to UV light, silicone adhesive is the only structural adhesive appropriate for fixing glazing units. Typically, storey-height fins are 200–300 mm wide in 12–15 mm toughened glass, fixed into aluminium or stainless steel shoes to the floor and/or glazing head. A soft interlayer between the metal fixing and the glass is incorporated to prevent stress concentrations on the glass surface and to allow for differential thermal movement between the glass and metal.

Glass is strong in compression and therefore an appropriate material for load-bearing columns and walls, provided that the design ensures sufficient strength, stiffness and stability. Generally, consideration of buckling is the critical factor, although safety factors must be considered in relation to robustness and protection against accidental damage. Column sections need not be rectangular, as, for example, a cruciform section manufactured from laminated toughened glass gives both efficient and elegant solution.

Glass beams are usually manufactured by laminating toughened glass. Typically, a $4 \times 600$ mm deep beam manufactured from three 15 mm toughened glass laminates could carry a load of over 5 tonnes, thus supporting at 2 m centres a 4 m span glass roof.
Fig. 7.25  Planar triple-glazing system. Diagram: Courtesy of Pilkington Plc.
Fig. 7.26 Glazing system — Financial Times, Docklands, London. Architects: Grimshaw Architects. Photographs: Courtesy of Grimshaw Architects, Jo Reid and John Peck
Glass beams can be jointed to glass columns by mortice and tenon jointing fixed with adhesive.

Single-storey all-glass structures, such as the small pavilions at the Gateshead Millennium Bridge (Fig. 7.27), entrance foyers and cantilevered canopies have been constructed using a combination of laminated and toughened glass walls, columns and beams. Usually, metal fixings have been used, but where the purity of an all-glass system is required, high-modulus structural adhesives are used for invisible fixing.

Where clear double-glazed structural units are required, standard aluminium spacers can be replaced by glass spacers sealed with clear silicone, although the edges may require etching to conceal the necessary desiccant. It should be remembered that all-glass constructions require careful design consideration in relation to excessive solar gain and other environmental factors.

Where glass is used as a load-bearing element it must conform to the structural and fire performance requirements of the relevant parts of BS EN 1365. The requirement is usually for 60 minutes of integrity and insulation, but this may be reduced to EI30 for residential buildings no higher than 5 m.

Structural double-glazing units filled with aerogel can be considered as alternatives for facades and roofing. A 70 mm wide cavity filled with aerogel gives a $U$-value of 0.25 W/m K, and 30 mm filled cavity
produces a $U$-value of 0.54 W/m K. Solar and light transmittance through 70 mm of nanogel is 19% and 49% for a 30 mm filled cavity. Structural requirements would include a toughened laminated inner pane.

**Glass staircases, balustrades and floors**

Other structural applications for glass include stairs, walkways, floors, balustrades and canopies. For stairs, laminated glass, either annealed or toughened, with a polymer interlayer may be used. The overall thickness depends upon the span and fixings, but a typical system would be a laminate of 25 and 10 mm, with the thicker layer uppermost. Glass is, however, slippery when wet, even if treated by sandblasting, chemically bonded polymer resin or grooved, and should not normally be considered for external locations. An all-glass staircase in Notting Hill, London (Fig. 7.28), uses glass risers and treads joined with structural silicone. The complete stair unit is supported on one side by a masonry wall but on the other by a full height laminated glass wall, to which it is fixed only with metal brackets and structural silicone. The steps incorporate sand-blasted dots to prevent slipping.

Glass flooring, usually in panels up to a maximum of 1 m², is used for effect in domestic and commercial locations. Coloured or clear panels may be illuminated from below to create interesting features. Floor thicknesses depend on the anticipated loading and fire regulations, but are normally within the range from 29 mm (19 + 10 mm) for domestic environments to 50 mm (25 + 25 mm) for commercial situations.
Surfaces will normally be etched or sandblasted for safety. The Chicago Sear’s Tower has 40 mm thick glass floor balconies projecting 1.2 m out from the 103rd floor, to give a view down onto the city.

Free-standing glass balustrades are usually manufactured from heat-soaked toughened glass ranging in thickness from 12 to 25 mm depending on their height and anticipated loading. Features may include curvature and fired-on screen print decoration.

Glass canopies for commercial buildings may be manufactured from laminates of toughened glass, which gives double security against the risk of dangerous failure. Recent innovations also include the structural use of glass rods in web compression members, and glass tubes in structural compression elements.

Structural glass must conform to the relevant parts of BS EN 1365 in relation to load-bearing properties and fire performance.

### Glazing checklist

Perhaps more than any other building component glazing is expected to perform many functions. It is therefore necessary to ensure that all factors are taken into consideration in the specification of glasses. It is evident that many of the environmental control factors are closely interrelated and the specifier must check the consequences of a design decision against all the parameters. Many of these factors are illustrated by the feature roof to the Great Court of the British Museum in London (Fig. 7.29).

**Key functions:**

- view in and out by day and night;
- visual appearance by day and by night – colour and reflectivity;
- energy-conscious balance between daylight and artificial lighting;
- sky and reflected glare;
- overheating and solar control;
- shading;
- passive solar gain and energy efficiency;
- thermal comfort, U-values and condensation;
- ventilation;
- acoustic control;
- security – impact damage, vandalism and fire spread.

### References

**FURTHER READING**


BS 4873: 2009 Aluminium alloy window and door sets. Specification.


BS 5051 Bullet-resisting glazing:


BS 5516 Patent glazing and sloping glazing for buildings:

BS 5544: 1978 Specification for anti-bandit glazing (glazing resistant to manual attack).

BS 5713: 1979 Specification for hermetically sealed flat double glazing units.

BS 5821 Methods for rating the sound insulation in buildings and of building elements:

BS 6100 Building and civil engineering terms:


BS 6262 Glazing for buildings:

BS 6375 Performance of windows and doors:
  Part 1: 2009 Classification for weathertightness and guidance on selection and specification.
  Part 2: 2009 Classification for operation and strength characteristics and guidance on selection and specification.
Part 3: 2009 Classification for additional performance characteristics and guidance on selection and specification.
BS 6510: 2005 Steel-framed windows and glazed doors.
BS 7412: 2007 Specification for windows and door sets made from unplasticised polyvinyl chloride (PVC-U) extruded hollow profiles.
BS 7950: 1997 Specification for enhanced security performance of casement and tilt/turn windows in domestic applications.
BS 8000 Workmanship on building sites:
BS 8213 Windows, doors and rooflights:
pr BS ISO 11479 Glass in building. Coated glass:
BS EN 357: 2004 Glass in building. Fire resistant glazed elements with transparent or translucent products.
BS EN 572 Glass in building. Basic soda lime silicate glass products:
  Part 5: 2004 Patterned glass.
  Part 7: 2004 Wired or unwired channel shaped glass.
  Part 8: 2004 Supplied and final cut sizes.

BS EN 1036 Glass in building. Mirrors from silver-coated float glass for internal use:
BS EN 1051 Glass in building. Glass blocks and glass pavers:
BS EN 1096 Glass in building. Coated glass:
  Part 2: 2001 Class A, B and S coatings.
  Part 3: 2001 Class C and D coatings.
BS EN 1279 Glass in building. Insulating glass units:
  Part 2: 2002 Requirements for moisture penetration.
  Part 4: 2002 Methods of test for the physical attributes of edge seals.
BS EN 1364 Fire resistance tests for non-loadbearing elements:
  Part 2: 1999 Ceilings.
BS EN 1365 Fire resistance tests for loadbearing elements:
  Part 4: 1999 Columns.
  Part 5: 2004 Balconies and walkways.
  Part 6: 2004 Stairs.
BS EN 1634 Fire resistance and smoke control tests for door, shutter and openable window assemblies:
Part 3: 2004 Smoke control test for door and shutter assemblies.
BS EN 1748 Glass in building. Special basic products:
BS EN 1863 Glass in building. Heat strengthened soda lime silicate glass:
BS EN ISO 10077 Thermal performance of windows, doors and shutters:
BS EN ISO 10456: 2007 Building materials and products. Hygrothermal properties. Tabulated design values and procedures for determining the declared and design thermal values.
BS EN 12150 Glass in building. Thermally toughened soda lime silicate safety glass:
BS EN 12337 Glass in building. Chemically strengthened soda lime silicate glass:
BS EN ISO 12543 Glass in building. Laminated glass and laminated safety glass:
BS EN 12567 Thermal performance of windows and doors. Determination of thermal transmittance:
  Part 2: 2005 Roof windows and other projecting windows.
BS EN 12608: 2003 Unplasticised polyvinyl chloride (PVC-U) profiles for the fabrication of windows and doors.
BS EN 13022 Glass in building. Structural sealant glazing:
  Part 1: 2006 Glass products for structural sealant glazing systems for supported and unsupported monolithic and multiple glazing.
BS EN 13024 Glass in building. Thermally toughened borosilicate safety glass:
BS EN 13363 Solar protection devices combined with glazing:
BS EN 13501 Fire classification of construction products and building elements:
  Part 1: 2007 Classification using data from reaction to fire tests.
  Part 2: 2007 Classification using data from fire resistance tests.
BS EN 14178 Glass in building. Basic alkaline earth silicate glass products:
BS EN 14179 Glass in building. Heat soaked thermally toughened soda lime silicate safety glass:
BS EN 14321 Glass in building. Thermally toughened alkaline earth silicate safety glass:
BS EN 14600: 2005  Doorsets and openable windows with fire resisting and/or smoke control characteristics. Requirements and classification.
pr EN 15269: 2008  Extended application of test results from fire-resistance and/or smoke control for door, shutter and openable window assemblies.
BS EN 15434: 2006  Glass in building. Product standard for structural and/or UV resistant sealant.
CP 153  Windows and rooflights:
PD 6512  Use of elements of structural fire protection:

BUILDING RESEARCH ESTABLISHMENT PUBLICATIONS

BRE Digests

BRE Digest 346  The assessment of wind loads:
  Part 1: 1992  Background and method.
  Part 4: 1992  Terrain and building factors and gust peak factors.

BRE Digest 453: 2000  Insulating glazing units.
BRE Digest 497: 2005  Factory glazed windows (Parts 1 & 2).
BRE Digest 498: 2006  Selecting lighting controls.

BRE Good building guide

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BRE IP 3/98  Daylight in atrium buildings.
BRE IP 2/02  Control of solar shading.
BRE IP 3/02  Whole life performance of domestic automatic window controls.
BRE IP 11/02  Retrofitting solar shading.
BRE IP 17/03  Impact of horizontal shading devices on peak solar gains through windows.
BRE IP 1/05  Impact standards for glass.

BRE Report

BR 443: 2006  Conventions for U-value calculations.

ADVISORY ORGANISATIONS

British Glass Manufacturers Confederation, 9 Churchill Way, Chapeltown, Sheffield, South Yorkshire S35 2PY, UK (0114 2901850).
Glass & Glazing Federation, 44-48 Borough High Street, London SE1 1XB, UK (0870 042 4255).
Plastics Window Federation, Federation House, 85-87 Wellington Street, Luton, Bedfordshire LU1 5AF, UK (01582 456147).
Steel Window Association, The Building Centre, 26 Store Street, London WC1E 7BT, UK (020 7637 3571).
CERAMIC MATERIALS

Introduction

Ceramic materials, manufactured from fired clay, have been used in construction since at least 4000 BC in Egypt, and represent the earliest manufactured building materials. Whilst the strict definition of ceramics includes glass, stone and cement, this chapter deals only with the traditional ceramics based on clays. The variety of traditional ceramic products used within the building industry arises from the wide range of natural and blended clays used for their production. The roof of the spectacular Sydney Opera House (Fig. 8.1) is surfaced with white ceramic tiles which reflect the changing light associated with the time of day.

CLAY TYPES

Clays are produced by the weathering of igneous rocks, typically granite, which is composed mainly of feldspar, an alumino-silicate mineral. Clays produced within the vicinity of the parent rock are known as primary clays. They tend to be purer materials, less plastic and more vulnerable to distortion and cracking on firing. Kaolin (Al₂O₃·2SiO₂·2H₂O), which is the purest clay, comes directly from the decomposition of the feldspar in granite. Secondary clays, which have been transported by water, have a higher degree of plasticity, and fire to a buff or brown colour depending on the nature and content of the incorporated oxides. Generally, secondary clays, laid down by the process of sedimentation, have a narrower size distribution and their particulate structure is more ordered.

The most common clay minerals used in the manufacture of building materials are kaolin, illite (a micaceous clay) and montmorillonite, a more plastic clay of variable composition. Clay crystals are generally hexagonal in form, and in pure kaolin, the crystals are built up of alternating layers of alumina and silica (Fig. 8.2). However, in illite and montmorillonite clays, the variable composition produced by sedimentation produces more complex crystal structures.

Ball clays are secondary clays containing some organic matter which is burnt off during the firing process; they tend to have a fine grain size which makes them plastic. When fired alone they have a high shrinkage and produce a light grey or buff ceramic, but they are usually blended into other clays such as kaolin to make a workable clay. Terracotta clays contain significant proportions of iron oxide which gives rise to the characteristic red colour on firing. Whilst the major clay materials used in the manufacture of ceramics are kaolin, illite, feldspar and ball clay; chalk, quartz and other minor constituents are frequently incorporated to produce required ceramic properties on firing.

WATER IN CLAY

Moist clay contains both chemically and physically bonded water. It is the latter which permeates between the clay particles, allowing them to slide over each other during the wet forming processes. As the formed clay slowly dries out before firing, a small proportion of the residual physically bonded water holds the clay in shape. On firing, the last of the physically bonded water is removed as the temperature exceeds 100°C.

MANUFACTURING PROCESSES

Clay products are formed by either wet or dry processes. In the former case, the artefacts after extrusion
must be dried slowly prior to firing, allowing for shrinkage without cracking. Where a high level of dimensional accuracy is required, as in most wall and floor tiles, a dry process is used in which powdered clay is compressed into the required form. The standard BS EN 14411: 2006 defines all ceramic tiles according to whether they are produced by extrusion or dry pressing, in addition to their water absorption properties, irrespective of their end use.

During firing of the clay, as the temperature is gradually increased, the majority of the chemically bonded water is removed by 500°C. At 800°C, carbonaceous matter has been burnt off as carbon dioxide, and the sintering process commences, at first producing a highly porous material. As the temperature is further raised towards 1200°C, the alumina and silica components recrystallise to form mullite. With an additional increase in firing temperature, a more glassy ceramic
is produced due to further recrystallisation, and if the firing temperature reaches 1300°C, any remaining free silica is recrystallised. In the presence of potassium or sodium salts vitrification occurs giving an impervious product (Fig. 8.3).

**Ceramic products**

**FIRECLAY**

A range of clays, predominantly blends of alumina and silica, high in silica (40–80%) and low in iron oxide (2–3%), produce fireclay refractory products, which will withstand high temperatures without deformation. Dense products have high flame resistance, whilst the insulating lower-density products are suitable for flue linings. White glazed fireclay is typically used for urinals, floor channels, industrial and laboratory sinks.

**BRICK CLAYS**

**Bricks and roofing tiles**

Bricks can be manufactured from a wide range of clays, the principal ones being Keuper marl, Etruria marl, Oxford clay, London clay, Coal measure shale, Weald and Gault clays with some production from alluvial and fireclay deposits. The composition of the clay varies widely depending on the type, but typically contains 40% to 65% silica, 10% to 25% alumina and 3% to 9% iron oxide. The loss on firing may reach 17% in the case of clay containing high levels of organic matter. The production of bricks is described in detail in Chapter 1.

Glazed bricks are manufactured in a wide range of high-gloss, uniform or mottled colours. Colour-fast glazed bricks offer a low-maintenance, frost- and vandal-resistant material suitable for light-reflecting walls. Standard and purpose-made specials can be manufactured to order. Normal bricklaying techniques are appropriate but to reduce the visual effect of the mortar joints they may be decreased from the standard 10 to 6 mm. For conservation work, in order to match new to existing, it may be necessary to fire the glazed bricks a second time at a reduced temperature to simulate the existing material colour.

Roofing tiles are made from similar clays to bricks, such as Etruria marl, but for both hand- and machine-made tiles, the raw materials have to be screened to a
finer grade than for brick manufacture. Traditional red, brown, buff, brindled or ‘antique’ ceramic roofing tiles are unglazed with a plain or sanded finish. Whilst most interlocking clay tiles can be used to a minimum pitch of 22.5°, one terracotta red UK product with full interlocking may be used down to only 15°. Where bright colours are required, high- and low-gloss pantiles are available in a range of strong colours, or to individual specification. For plain tiles, a range of standard fittings is produced for hips, valleys, eaves, ridges, verges, internal and external angles, as shown in Fig. 8.4. Tiles are usually shrink-wrapped for protection and ease of handling on site.

Marl floor tiles
Certain floor tiles are also manufactured from Etruria marl. Firing to 1130°C produces sufficient vitrification to give a highly durable, chemical- and frost-resistant product, with water absorption limited to less than 3%. This corresponds to the lowest category (Group 1) of water absorption to BS EN 14411: 2006. Medium water absorption (Group 2) is defined as between 3% and 10%, and Group 3 refers to water absorption greater than 10%. Where high slip resistance is required, a studded profile or carborundum (silicon carbide) grit may be incorporated into the surface (Fig. 8.5). Figure 8.6 illustrates the appropriate use of ceramic floor tiles.

TERRACOTTA
In order to produce intricately detailed terracotta building components, the clay has to be more finely divided than is necessary for bricks and roof tiles. The presence of iron oxide within the clay causes the buff, brown or red colouration of the fired product. During the latter part of the nineteenth century many civic buildings were constructed with highly decorative terracotta blocks. The material was used because it was cheaper than stone, durable and could be readily moulded. The blocks, which were usually partly hollowed out to facilitate drying and firing, were filled with concrete during construction.

Modern terracotta blocks may still be supplied for new work or refurbishment as plain ashlar, profiled or with sculptural embellishments. Terracotta may be used as the outer skin of cavity wall construction or as 25–40 mm thick cladding hung with stainless steel mechanical fixings. The production of terracotta blocks requires the manufacture of an oversize model (to allow for shrinkage), from which plaster moulds are made. Prepared clay is then pushed into the plaster mould, dried under controlled conditions and finally fired. Traditional colours together with greens and blues and various textures are produced. For refurbishment work existing terracotta can, subject to natural variations, usually be colour matched. In addition to cladding units, terracotta clay is also used in the manufacture of terracotta floor tiles and an extensive range of decorative ridge tiles and finials (Fig. 8.7). Standard terracotta building blocks made by extrusion have been described in Chapter 2.

Terracotta rainscreen cladding
Rainscreen cladding is the external weathering element to multi-layer rainscreen wall systems. The rainscreen facade is drained and back-ventilated to protect the structural wall from the adverse effects of the sun, wind and rainwater. The design of the joints and the cavity between the facade and the structure result in an equalisation of air pressure between the cavity and the exterior, thus inhibiting the drive of airborne moisture across the cavity. In some systems a vertical EPDM gasket seal is located in the vertical joints. A breather membrane is usually fixed to the structure before the rainscreen system is applied.

Rainscreen systems are appropriate for masonry, concrete, timber-frame and concrete-frame construction. A grid of vertical or horizontal aluminium extrusions is fixed to the facade, creating a minimum air gap of 25 mm. The rainscreen units are then clipped to the support system. The rainscreen cladding units may be manufactured in terracotta or from a wide range of other materials including stone laminate, stainless steel, copper, aluminium or zinc. Rainscreen units are shaped to shed water out of the open-drained joints, and individual units may be removed for maintenance or repair. A range of colours and dimensions is available in terracotta units to create the required aesthetic effect. Terracotta rainscreen cladding (Fig. 8.8) is fire resistant and durable requiring virtually no maintenance except occasional cleaning.

FAIENCE
Faience is glazed terracotta, used either as structural units or in the form of decorative slabs applied as cladding. It was popular in the nineteenth century and was frequently used in conjunction with polychrome brickwork on the facades of buildings such as public
Fig. 8.4 Roof tiles—feature tiles and plain tile fittings
houses. Either terracotta may be glazed after an initial firing to the biscuit condition or the slip glaze may be applied prior to a single firing. The latter has the advantage that it reduces the risk of the glaze crazing, although it also restricts the colour range. Faience, with an orange-peel texture, is available with either a matt or gloss finish and in plain or mottled colours. It is a highly durable material unaffected by weathering, frost or ultraviolet light, but strong impacts can chip the surface causing unsightly damage.
STONEWARE

Stoneware is manufactured from secondary plastic clays, typically fireclays blended with an added flux such as feldspar. On firing to between 1200°C and 1300°C, the material vitrifies, producing an impermeable ceramic product with high chemical resistance. The majority of unglazed vitrified clay pipes are stoneware. For most purposes push-fit polypropylene couplings are used, which allow flexibility to accommodate ground movement; however, if required, traditional jointed socket/spigot drainage goods are also available in stoneware.

Large stoneware ceramic panels up to 1.2 m square and 8 mm in thickness are manufactured as cladding units for facades. The units, which are colour fast, frost- and fire-resistant may be uniform in colour or flecked and glazed or unglazed. Fixing systems are exposed or hidden; the open joint system offers rear ventilation, allowing any moisture diffusing from the supporting wall to be dissipated by natural air movement.

Stoneware floor tiles

Stoneware is also used in the manufacture of some floor tiles. The high firing temperature gives a product of low porosity, typically less than 3%. In one manufacturing process a granular glaze is applied to the tiles within the kiln to produce an impervious vitreous finish. The standard BS EN 14411: 2006 classifies six levels of abrasion resistance for glazed floor tiles. Class 0 tiles are not recommended for use on floors. Class 5 tiles have the maximum resistance to severe pedestrian traffic over sustained periods.

EARTHENWARE

Earthenware is produced from a mixture of kaolin, ball clay and flint, with in some cases, feldspar as a flux. The material when fired at 1100°C is porous and requires a glaze to prevent water absorption. In the manufacture of traditional glazed drainage goods, the salt glaze is produced by adding damp common salt to the kiln during the firing process. The salt decomposes to form sodium oxide, which then reacts with silica and alumina on the surface of the clay component to produce the salt glaze which is impermeable to moisture.

EARTHENWARE WALL TILES

Wall tiles (Fig. 8.9) are generally manufactured from earthenware clay to which talc (magnesium silicate)
or limestone (calcium carbonate) is added to ensure a white burning clay. To prepare the clay for manufacturing wall tiles by the dry process, the components, typically a blend of china clay (kaolin), ball clay and silica sand together with some ground recycled tiles, are mixed with water to form a slip. This is sieved, concentrated to a higher density slip and then dried to a powder by passage down a heated tower at 500°C. The clay dust, which emerges with a moisture content of approximately 8%, is then pressed into tiles. A glaze is required to both decorate and produce an impermeable product and this may be applied before a single firing process or after the tiles have been fired at 1150°C to the biscuit stage in a tunnel kiln. Either the unfired or biscuit tiles are coated with a slip glaze followed by firing under radiant heat for approximately 16 hours. Damaged tiles are rejected for recycling, and the quality-checked tiles are packaged for dispatch. Standard sizes are 108 × 108 mm, 150 × 150 mm, 200 × 150 mm, 200 × 200 mm and 250 × 200 mm.

**VITREOUS CHINA**

Vitreous china, used for the manufacture of sanitary ware, has a glass-like body which limits water absorption through any cracks or damage in the glaze to 0.5%. It is typically manufactured from a blend of kaolin (25%), ball clay (20%), feldspar (30%) and quartz (25%). For large units such as WCs and wash basins, a controlled drying-out period is required before firing to prevent cracking. Glaze containing metallic oxides for colouration is applied before firing to all visually exposed areas of the components.

**Vitreous china floor tiles**

Vitreous china is also used in the manufacture of some floor tiles due to its impermeable nature. Unglazed floor tiles may be smooth, alternatively studded or ribbed to give additional non-slip properties. Standard sizes are 100 × 100 mm, 150 × 150 mm, 200 × 200 mm and 300 × 300 mm, with thickness usually in the range 8–13 mm. For lining swimming pools, additional protection against water penetration is given by the application of a glaze.

**REPRODUCTION DECORATIVE TILES**

Reproduction moulded ceramic wall tiles, encaustic tiles with strong colours burnt into the surface, and geometrical floor tiles can be manufactured to match existing units with respect to form, colour and texture for restoration work. Some manufacturers retain both the necessary practical skills and appropriate detailed drawings to ensure high-quality conservation products, which may be used to replace lost or seriously damaged units. There is also an increasing demand for reproduction decorative tiles in new-build work.
Fig. 8.11  Ceramic mosaic finish and detail — Concert Hall, Tenerife. Architect: Santiago Calatrava. Photographs: Arthur Lyons
MOSAICS
Mosaics in glazed or unglazed porcelain are hard wearing, frost-proof and resistant to chemicals. Unglazed mosaics may be used for exterior use and other wet areas such as swimming pools, where good slip resistance is important. Mosaics are usually supplied attached to paper sheets for ease of application. Figure 8.10 illustrates a formal mosaic floor, while Fig. 8.11 shows the broken tile mosaic finish used by Calatrava on the Tenerife Concert Hall, following the technique developed by Gaudi.

CERAMIC GRANITE
Ceramic granite is a blend of ceramic and reconstituted stone, manufactured from a mixture of feldspar, quartz and clay. The components are crushed, graded, mixed and compressed under very high pressure, followed by firing at 1200°C. The material is produced in 20 and 30 mm slabs, which can be cut and polished to produce a hard shiny finish with the appearance of natural marble or granite, suitable for worktops. Colours range from ochre, off-white and grey to green and blue depending on the initial starting materials.

References
FURTHER READING

STANDARDS
BS 65: 1991 Specification for vitrified clay pipes, fittings, and ducts, also flexible mechanical joints for use solely with surface water pipes and fittings.
BS 493: 1995 Airbricks and gratings for wall ventilation.
BS 1188: 1974 Ceramic wash basins and pedestals.
BS 1196: 1989 Clayware field drain pipes and junctions.
BS 3402: 1969 Quality of vitreous china sanitary appliances.
BS 5385 Wall and floor tiling:
  Part 1: 2009 Code of practice for the design and installation of internal ceramic and natural stone wall tiling and mosaics in normal conditions.
BS 5504 Specification for wall hung WC pan:
BS 5506 Specification for wash basins:
  Part 3: 1977 Wash basins (one or three tap holes), materials, quality, design and construction.
BS 8000 Workmanship on building sites:
BS EN 295 Vitrified clay pipes and fittings and pipe joints for drains and sewers:
  Part 2: 1991 Quality control and samples.
  Part 4: 1995 Requirements for special fittings, adaptors and compatible accessories.
  Part 5: 1994 Requirements for perforated vitrified clay pipes.
  Part 7: 1996 Requirements for vitrified clay pipes and joints.
  Part 10: 2005 Performance requirements.
BS EN 539 Clay roofing tiles for discontinuous laying. Determination of physical characteristics:
  Part 2: 2006 Test for frost resistance.
BS EN 997: 2003 WC pans and WC suites with integral trap.
BS EN 1457: 1999 Chimneys clay/ceramic flue liners.
BS EN ISO 10545 Ceramic tiles:
Part 7: 1999 Resistance to surface abrasion for glazed tiles.
Part 8: 1996 Determination of linear thermal expansion.
Part 15: 1997 Determination of lead and cadmium given off by glazed tiles.
Part 16: 2000 Determination of small colour differences.
BS EN 12808 Grouts for tiles. Requirements, evaluation of conformity, classification and designation:
BS EN 13502: 2002 Requirements and test methods for clay/ceramic flue terminals.
BS EN 13888: 2009 Grout for tiles.
BS EN 14411: 2006 Ceramic tiles. Definitions, classification, characteristics and marking.
BS EN 14437: 2004 Determination of the uplift resistance of installed clay or concrete tiles for roofing.

BUILDING RESEARCH ESTABLISHMENT PUBLICATIONS

BRE Digests

BRE Digest 467: 2002 Slate and tile roofs: avoiding damage from aircraft wake vortices.
BRE Digest 486: 2004 Reducing the effect of climate change by roof design.
BRE Digest 499: 2006  Designing roofs for climate change. Modifications to good practice guidance.

**BRE Good building guides**

BRE GBG 28: 1997  Domestic floors - repairing or replacing floors and flooring.
BRE GBG 64: 2005  Tiling and slating pitched roofs (Parts 1, 2 and 3).

**ADVISORY ORGANISATIONS**

British Ceramic Confederation, Federation House, Station Road, Stoke-on-Trent ST4 2SA, UK (01782 744631).

CERAM Research, Queens Road, Penkhull, Stoke-on-Trent, Staffordshire ST4 7LQ, UK (01782 764428).
Clay Pipe Development Association Ltd., Copsham House, 53 Broad Street, Chesham, Buckinghamshire HP5 3EA, UK (01494 791456).
Clay Roof Tile Council, Federation House, Station Road, Stoke-on-Trent, Staffordshire ST4 2SA, UK (01782 744631).
Tile Association, Forum Court, 83 Copers Cope Road, Beckenham, Kent BR3 1NR, UK (020 8663 0946).
Introduction

The term stone refers to natural rocks after their removal from the earth’s crust. The significance of stone as a building material is illustrated by widespread prehistoric evidence and its sophisticated use in the early civilisations of the world, including the Egyptians, the Incas of Peru and the Mayans of Central America.

Geologically, all rocks can be classified into one of three groups: igneous, metamorphic or sedimentary, according to the natural processes by which they were produced within or on the earth’s surface.

For construction work, stone specification to BS EN 12440: 2008 requires the scientific petrological family as defined within the standard BS EN 12670: 2002, the typical colour range and the place of origin. The latter should be as detailed as possible, including the location of the quarry, its nearest town, the region and the country.

Types of stone

IGNEOUS ROCKS

Igneous rocks are the oldest, having been formed by the solidification of the molten core of the earth or magma. They form about 95% of the earth’s crust, which is up to 16 km thick. Depending on whether solidification occurred slowly within the earth’s crust or rapidly at the surface, the igneous rocks are defined as plutonic or volcanic, respectively. In the plutonic rocks, slow cooling from the molten state allowed large crystals to grow, which are characteristic of the granites. Volcanic rocks such as pumice and basalt are fine-grained and individual crystals cannot be distinguished by eye; thus, the stones are visually less interesting. Dolerites, formed by an intermediate rate of cooling, exhibit a medium-grained structure.

Apart from crystal size, igneous rocks also vary in composition according to the nature of the original magma, which is essentially a mixture of silicates. High-silica-content magma produces acid rocks (e.g. granite) whilst low silica content forms basic rocks (e.g. basalt and dolerite). Granites are mainly composed of feldspar (white, grey or pink), which determines the overall colour of the stone but they are modified by the presence of quartz (colourless to grey or purple), mica (silver to brown), or hornblende (dark coloured). The basic rocks such as dolerite and basalt in addition to feldspar contain augite (dark green to black) and sometimes olivine (green). Although basalt and dolerite have not been used widely as building stones they are frequently used as aggregates, and cast basalt is now being used as a reconstituted stone.

Granites

Most granites are hard and dense, and thus form highly durable building materials, virtually impermeable to water, resistant to impact damage and stable within industrial environments. The appearance of granite is significantly affected by the surface finish which may be sawn, rough punched, picked, fine tooled, honed or polished. It is, however, the highly polished form of granite which is most effective at displaying the intensity of the colours and reflectivity of the crystals. Additionally, granites may be flamed to a spalled surface, produced by the differential expansion of the various crystalline constituents. Many recent buildings
have combined the polished and flamed material to create interesting contrasts in depth of colour and texture. Grey and pink granites are quarried in Scotland, the North of England, Devon and Cornwall, but a wide variety of colours including black, blue, green, red, yellow and brown are imported from other countries (Fig. 9.1; Table 9.1). Because of the high cost of quarrying and finishing granite, it is frequently used as a cladding material (40 mm externally or 20 mm internally) or alternatively cast directly onto concrete cladding units. Granite is available for flooring and for hard landscaping including pavings, sets and kerbs. Polished granite is also used as a kitchen countertop material due to its strength, durability and high-quality finish.
Table 9.1  UK and imported granites

<table>
<thead>
<tr>
<th>Colour</th>
<th>Name</th>
<th>Country of origin</th>
</tr>
</thead>
<tbody>
<tr>
<td>UK</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Light grey</td>
<td>Merrivale, Devon</td>
<td>England</td>
</tr>
<tr>
<td>Silver grey</td>
<td>De Lank &amp; Hantergantick, Cornwall</td>
<td>England</td>
</tr>
<tr>
<td>Light and dark pink to brownish red</td>
<td>Shap</td>
<td>England</td>
</tr>
<tr>
<td>Pink</td>
<td>Peterhead</td>
<td>Scotland</td>
</tr>
<tr>
<td>Pale to deep red</td>
<td>Ross of Mull</td>
<td>Scotland</td>
</tr>
<tr>
<td>Grey</td>
<td>Aberdeen</td>
<td>Scotland</td>
</tr>
<tr>
<td>Black</td>
<td>Hillend</td>
<td>Scotland</td>
</tr>
<tr>
<td>Black</td>
<td>Baltmoss</td>
<td>Scotland</td>
</tr>
<tr>
<td>Imported</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Red with black</td>
<td>Balmoral Red</td>
<td>Finland</td>
</tr>
<tr>
<td>Red</td>
<td>Bon Accord Red</td>
<td>Sweden</td>
</tr>
<tr>
<td>Black</td>
<td>Bon Accord Black</td>
<td>Sweden</td>
</tr>
<tr>
<td>Red</td>
<td>Virgo Granite</td>
<td>Sweden</td>
</tr>
<tr>
<td>Dark red with blue to purple quartz</td>
<td>Rose Swede</td>
<td>Sweden</td>
</tr>
<tr>
<td>Grey</td>
<td>Grey Royal</td>
<td>Norway</td>
</tr>
<tr>
<td>Grey</td>
<td>Sardinian Grey</td>
<td>Sardinia</td>
</tr>
<tr>
<td>Yellow</td>
<td>Nero Tijuca</td>
<td>Brazil</td>
</tr>
<tr>
<td>Beige/brown</td>
<td>Juparana</td>
<td>Brazil</td>
</tr>
<tr>
<td>Blue</td>
<td>Blue Pearl</td>
<td>Norway</td>
</tr>
<tr>
<td>Green/black</td>
<td>Emerald Pearl</td>
<td>Norway</td>
</tr>
<tr>
<td>Pink to red</td>
<td>Torcicola</td>
<td>Brazil</td>
</tr>
<tr>
<td>Beige/brown</td>
<td>Giallio Veneziano</td>
<td>Brazil</td>
</tr>
</tbody>
</table>

* Available only in limited quantities.

Cast basalt

Basalt is a fine-grained stone nearly as hard as granite. It can be melted at 2400°C and cast into tile units which are deep steel grey in colour. A slightly patterned surface can be created by swirling the molten basalt within the mould. Annealing in a furnace produces a hard virtually maintenance-free, shiny textured surface flecked with shades of green, red and bronze. Larger cast units for worktops, in either a honed or a polished finish, can be cut to size.

SEDIMENTARY ROCKS

Sedimentary rocks are produced by the weathering and erosion of older rocks. In the earliest geological time these would have been the original igneous rocks, but subsequently other sedimentary and metamorphic rocks too will have been reworked. Weathering action by water, ice and wind breaks the rocks down into small fragments, which are then carried by rivers and sorted into size and nature by further water action. Most deposits are laid down in the oceans as sedimentary beds of mud or sand, which build up in layers, become compressed and eventually are cemented together by minerals such as calcium carbonate (calcite), quartz...
Table 9.2  Typical UK sandstones and their characteristics

<table>
<thead>
<tr>
<th>Name</th>
<th>Colour</th>
<th>Source</th>
<th>Characteristics</th>
</tr>
</thead>
<tbody>
<tr>
<td>Doddington</td>
<td>Purple/pink</td>
<td>Northumberland</td>
<td>Fine- to medium-grained</td>
</tr>
<tr>
<td>Darley Dale—Stancliffe</td>
<td>Buff</td>
<td>Derbyshire</td>
<td>Fine-grained</td>
</tr>
<tr>
<td>Birchover gritstone</td>
<td>Pink to buff</td>
<td>Derbyshire</td>
<td>Medium- to coarse-grained</td>
</tr>
<tr>
<td>York stone</td>
<td>Buff, fawn, grey, light brown</td>
<td>Yorkshire</td>
<td>Fine-grained</td>
</tr>
<tr>
<td>Mansfield stone</td>
<td>Buff to white</td>
<td>Nottinghamshire</td>
<td>Fine-grained</td>
</tr>
<tr>
<td>Hollington</td>
<td>Pale pink, dull red pink with</td>
<td>Staffordshire</td>
<td>Fine- to medium-grained</td>
</tr>
<tr>
<td></td>
<td>darker stripe</td>
<td></td>
<td></td>
</tr>
<tr>
<td>St. Bees</td>
<td>Dark red</td>
<td>Cumbria</td>
<td>Fine-grained</td>
</tr>
<tr>
<td>Blue Pennant</td>
<td>Dark grey/blue</td>
<td>Mid-Glamorgan</td>
<td>Fine-grained</td>
</tr>
</tbody>
</table>

(silica), iron oxide or dolomite (magnesium and calcium carbonate) remaining in the ground water. The natural bedding planes associated with the formation of the deposits may be thick or thin but are potentially weak; this is used to advantage in the quarrying process. In masonry, to obtain maximum strength and durability, stones should be laid to their natural bed except for cornices, cills and string courses which should be edge-bedded. Face-bedded stones will tend to delaminate (Fig. 9.2). When quarried, stones contain *quarry sap* and may be worked and carved more easily than after exposure to the atmosphere.

Sandstones

Deposits of sand cemented together by calcium carbonate, silica, iron oxide and dolomite produce calcareous, siliceous, ferruginous and dolomitic sandstones, respectively. Depending on the nature of the original sand deposit, the sandstones may be fine or coarse in texture. Sandstones range in colour from white, buff and grey through to brown and shades of red depending on the natural cement; they are generally frost-resistant. Some common UK sandstones are listed in Table 9.2, and examples are illustrated in Figure 9.3. The recent Chester Song School (Fig. 9.4) was built in Lochabriggs red sandstone to empathise with the old red sandstone of the cathedral. Some sandstones imported from India have fern fossils embedded within the strata (Fig. 9.5). Typical finishes are sawn, split faced and clean rubbed, although a range of tooled finishes including broached and droved, can also be selected (Fig. 9.6). For cladding, sandstone is normally 75–100 mm thick and fixed with non-ferrous cramps and corbels. Sandstones are quarried in Scotland, the North of England, Yorkshire and Derbyshire; they include the old and new red sandstones, York Stone and Millstone Grit. Sandstone is also imported from Spain and Italy from where *Pietra Serena* is sourced.

**Calcareous sandstone**

Calcareous sandstones are not durable in acid environments, which may cause the slow dissolution of the natural calcium carbonate cement of the stone. Pure calcite is white, so these sandstones are generally white in colour.

**Siliceous sandstone**

Siliceous sandstones are predominantly grains of silica (sand) cemented with further natural silica, and are therefore durable even in acid environments. Siliceous sandstones are generally grey in colour.

**Ferruginous sandstone**

Ferruginous sandstones are bound with oxides of iron which may be brown, ochre or red. They are generally durable.

**Dolomitic sandstone**

Dolomitic sandstones are bound with a mixture of magnesium and calcium carbonates, and therefore do not weather well in urban environments. They are generally off-white and buff in colour.

Limestones

Limestones consist mainly of calcium carbonate, either crystallised from solution as calcite or formed from accumulations of fossilised shells deposited by various sea organisms (Fig. 9.7). They are generally classified according to their mode of formation. Many colours are available ranging from off-white, buff, cream, grey and blue. Limestones are found in England in a belt from Dorset, the Cotswolds, Oxfordshire, and Lincolnshire to Yorkshire. Limestone is also imported from Ireland, France and Portugal to widen the palette of colours. Some common UK limestones are listed.
Fig. 9.3 Examples of UK sandstones. Photographs: Natural stone images supplied by Stancliffe Stone
in Table 9.3, and examples of limestones and ironstone are illustrated in Figure 9.8. The standard finishes are fine rubbed, fine dragged and split faced, although tooled finishes are also appropriate. Externally, limestones must not be mixed with or located above sandstones, as this may cause rapid deterioration of the sandstone.

*Oölitic limestone*

Oölitic limestones are formed by crystallisation of calcium carbonate in concentric layers around small
fragments of shell or sand, producing spheroidal grains or ooliths. The ooliths become cemented together by the further deposition of calcite to produce the rock. Typically, the ooliths are up to 1 mm in diameter, giving a granular texture to the stone, which may also incorporate other fossils. Oolitic limestones are very workable and include Bath stone, Portland stone. Clipsham stone and Ketton stone have been widely used at Oxford and Cambridge, respectively, including the recent Queen’s Building of Emmanuel College, Cambridge (Fig. 9.9), which is built of load-bearing Ketton limestone, with appropriately massive columns and flat voussoir arches to the colonnade and window openings. Lime mortar is used to ensure an even spreading of the load between stones. In the case of Foundress Court, Pembroke College, Cambridge (Fig. 9.10), the Bath stone (Monks Park) is built three storeys up from ground level as a well-detailed cladding, with restraint back to the load-bearing blockwork inner skin. The flexibility of lime mortar is used to reduce the number of visible movement joints.

**Organic limestone**

Organic limestones are produced in bedded layers from the broken shells and skeletal remains of a wide variety of sea animals and corals. Clay is frequently incorporated into organic limestones and this adversely affects the polish, which can otherwise be achieved on the cut stone.

---

**Table 9.3  Typical UK limestones and their characteristics**

<table>
<thead>
<tr>
<th>Name</th>
<th>Colour</th>
<th>Source</th>
<th>Characteristics</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ancaster</td>
<td>Cream to buff</td>
<td>Lincolnshire</td>
<td>Oolitic limestone—variable shell content; freestone available</td>
</tr>
<tr>
<td>Bath Stone</td>
<td>Pale brown to light cream</td>
<td>Avon</td>
<td>Oolitic limestone</td>
</tr>
<tr>
<td>- Westwood Ground</td>
<td></td>
<td></td>
<td>Coarse-grained—buff coloured</td>
</tr>
<tr>
<td>- Monks Park</td>
<td></td>
<td></td>
<td>Fine-grained—buff coloured</td>
</tr>
<tr>
<td>Clipsham</td>
<td>Buff to cream</td>
<td>Rutland</td>
<td>Medium-grained oolitic limestone with shells; some blue stone; Best-quality stone is durable</td>
</tr>
<tr>
<td>Doulting</td>
<td>Pale brown</td>
<td>Somerset</td>
<td>Coarse textured; fossils uncommon</td>
</tr>
<tr>
<td>Hopton Wood</td>
<td>Cream or grey</td>
<td>Derbyshire</td>
<td>Carboniferous limestone containing many attractive fossils; may be polished</td>
</tr>
<tr>
<td>Ketton</td>
<td>Pale cream to buff and pink</td>
<td>Lincolnshire</td>
<td>Medium-grained oolitic limestone; even-textured; durable stone</td>
</tr>
<tr>
<td>Portland Stone</td>
<td>White</td>
<td>Dorset</td>
<td>Exposed faces weather white, Protected faces turn black.</td>
</tr>
<tr>
<td>- Roach</td>
<td></td>
<td></td>
<td>Coarse open-textured shelly stone; weathers very well</td>
</tr>
<tr>
<td>- Whitbed</td>
<td></td>
<td></td>
<td>Fine-grained—some shell fragments; durable stone</td>
</tr>
<tr>
<td>- Basebed</td>
<td></td>
<td></td>
<td>Fine-grained with few shells; suitable for carving</td>
</tr>
<tr>
<td>Purbeck</td>
<td>Blue/grey to buff</td>
<td>Dorset</td>
<td>Some shells; durable stone</td>
</tr>
</tbody>
</table>
Crystallised limestone
When water containing calcium bicarbonate evaporates, it leaves a deposit of calcium carbonate. In the case of hot springs the material produced is travertine, and in caves, stalactites and stalagmites or onyx-marble result.

Dolomitic limestone
Dolomitic limestones have had the original calcium carbonate content partially replaced by magnesium carbonate. In general, this produces a more durable limestone, although it is not resistant to heavily polluted atmospheres.

METAMORPHIC ROCKS
Metamorphic rocks are formed by the recrystallisation of older rocks, when subjected to intense heat or pressure or both, within the earth’s crust. Clay is metamorphosed to slate, limestone to marble and sandstone to quartzite.
Slate

Slate is derived from fine-grained sand-free clay sediments. The characteristic cleavage planes of slate were produced when the clay was metamorphosed, and frequently, they do not relate to the original bedding planes. Slate can be split into thin sections (typically 4–10 mm for roofing slates) giving a natural riven finish, or it may be sawn, sanded, fine rubbed, honed, polished, flame textured or bush hammered. A range of distinctive colours is available: blue/grey, silver grey and green from the Lake District, blue, green, grey and plum red from North Wales and grey from Cornwall. Slate is also imported from Ireland (grey/green), Canada (blue/grey), France (blue/grey), India (blue/grey), China (blue/green/grey), Brazil (grey/green/plum) and blue/black from Spain, which is the world’s largest producer of the material. Slate is strong, acid- and frost-resistant, lasting up to 400 years as a roofing material. The minimum recommended pitch for slate roofing is $20^\circ$ under sheltered or moderate exposure and $22.5^\circ$ under severe exposure, and these situations require the use of the longest slates (460, 560 or 610 mm). Where thick slates (up to 20 mm in thickness) are used for a roof pitch of less than $25^\circ$, it should be noted that the slates lie at a significantly lower pitch than the rafters. Fixing nails should be of copper or aluminium. Slate is also used for flooring, cladding, copings, cills and stair treads. When used as a cladding material it should be fixed with non-ferrous fixings or cast directly onto concrete cladding units.

Roofing and external cladding slates satisfy the requirements for the Class A1 characteristic reaction to fire performance, without the need for testing.

Recycled roofing slates, particularly Welsh slate, are generally available in a range of sizes and are appropri-
ate for both conservation work and new build where an immediate weathered appearance is required. Welsh slates have a good reputation for durability, making the recycled product a viable option. Certain regional slates, such as Swithland in Leicestershire, are only available as recycled products. This particular type of slate has a single top nail fixing, and unlike most roof- ing slates which are of a uniform size, it is graded from large slates at the eaves to smaller units at the ridge (Fig. 9.11).

Reconstituted slate
Reconstituted slate for roofing is manufactured from slate granules and inert filler, mixed with a thermosetting resin and cast into moulds to give a natural riven slate finish. Certain products incorporate glass- fibre reinforcement and offer a wider range of colours than are available in natural slate. Some interlocking slates may be used down to a pitch of 17.5°, whilst double-lap simulated natural slates can be used down to a pitch of 20° depending on the degree of exposure. Reconstituted slate is also manufactured in glass-fibre reinforced cement (GRC) as described in Chapter 11.

Marble
Marble is metamorphosed limestone in which the calcium carbonate has been recrystallised into a mosaic of approximately equal-sized calcite crystals. The process, if complete, will remove all traces of fossils, the size of the crystals being largely dependent on the duration of the process. Some limestones which can be polished are sold as marble, but true marble will not contain any fossilised remains. Calcite itself is white, so a pure marble is white and translucent. The colours and vein- ing characteristics of many marbles are associated with impurities within the original limestone; they range from red, pink, violet, brown, green, beige, cream and white to grey and black. Marble is attacked by acids; therefore, honed, rather than highly polished surfaces, are recommended for external applications. Marbles are generally hard and dense, although fissures and veins sometimes require filling with epoxy resins. Most
Fig. 9.11 Slate roofs

Marbles used within Britain are imported from Europe as indicated in Table 9.4; a selection is illustrated in Figure 9.12.

For external cladding above first floor level 40 mm thick slabs are used, although 20 mm may be appropriate for internal linings and external cladding up to first floor level. Fixing cramps and hooks should be in stainless steel, phosphor bronze or copper. Floor slabs, to a minimum thickness of 30 mm, should be laid on a minimum 25 mm bed. Marble wall and bathroom floor tiles are usually between 7 and 10 mm in thickness.
Reconstituted marble
Reconstituted marble is manufactured from marble chippings and resin into tiles and slabs for use as floor and wall finishes. The material has the typical colours of marble but without the veining associated with the natural material.

Quartzite
Quartzite is metamorphosed sandstone. The grains of quartz are recrystallised into a matrix of quartz, producing a durable and very hard-wearing stone used mainly as a flooring material. The presence of mica allows the material to be split along smooth cleavage planes, producing a riven finish. Quartzite is mainly imported from Norway and South Africa and is available in white, grey, grey-green, blue-grey and ochre colours.

Table 9.4  A selection of imported marbles

<table>
<thead>
<tr>
<th>Colour</th>
<th>Name</th>
<th>Country of origin</th>
</tr>
</thead>
<tbody>
<tr>
<td>White</td>
<td>White Carrara/Sicilian</td>
<td>Italy</td>
</tr>
<tr>
<td>White</td>
<td>White Pentelicon</td>
<td>Greece</td>
</tr>
<tr>
<td>Cream</td>
<td>Perlato</td>
<td>Sicily</td>
</tr>
<tr>
<td>Cream</td>
<td>Travertine</td>
<td>Italy</td>
</tr>
<tr>
<td>Beige</td>
<td>Botticino</td>
<td>Sicily</td>
</tr>
<tr>
<td>Pink</td>
<td>Rosa Aurora</td>
<td>Portugal</td>
</tr>
<tr>
<td>Red</td>
<td>Red Bilbao</td>
<td>Spain</td>
</tr>
<tr>
<td>Brown</td>
<td>Napoleon Brown</td>
<td>France</td>
</tr>
<tr>
<td>Green</td>
<td>Verde Alpi</td>
<td>Italy</td>
</tr>
<tr>
<td>Black</td>
<td>Belgian Black</td>
<td>Belgium</td>
</tr>
<tr>
<td>Black with white veins</td>
<td>Nero Marquina</td>
<td>Algeria</td>
</tr>
</tbody>
</table>

ALABASTER
Alabaster is naturally occurring gypsum or calcium sulphate. Historically, it has been used for building as in the Palace of Knossos, Crete, but in the UK its use has been mainly restricted to carved monuments and ornaments. The purest form is white and translucent, but traces of iron oxide impart light brown, orange or red colourations.

Stonework
TRADITIONAL WALLING
Dressed stone may be used as an alternative to brick or block in the external leaf of standard cavity construction. Limestone and sandstone are the most frequently used for walling, but slate is also used where it is available locally. Although random rubble and
hand-dressed stone can be supplied by stone suppliers, sawn-bedded (top and bottom) stones are generally the most available. These are normally finished split faced, pitch faced, fine rubbed or sawn. The standard sizes are 100 or 105 mm on bed, with course heights typically 50, 75, 100, 110, 125, 150, 170, 225 and 300 mm (Fig. 9.13). Stones may be to a particular course length, e.g. 300 mm or 450 mm, although they are frequently to random lengths. Quoin blocks, window and door surrounds, cills and other components are often available as standard. In ashlar masonry, the stones are carefully worked and finely jointed. Stones within horizontal courses are of the same height and are perfectly rectangular in elevation. Joints are generally under 6 mm in width.

The mortar for stone masonry should be weaker than the stone selected. For porous limestones and sandstones, crushed stone aggregate is frequently used as the aggregate in the mortar, typically in a 1:3:12 mix of Portland cement, lime putty and crushed stone. For ashlar Bath stone a typical mix would be 1:2:8 cement, lime and stone dust. Dense sandstones may be bonded with a stronger 1:1:6 mix, and granite a 1:2 or 1:3 Portland cement to fine aggregate mix. Jointing should generally be to a similar texture and colour as that of the dressed stone itself, and should be slightly recessed to emphasise the stones rather than the joints. In ashlar work, in which accurately cut squared stones are used, a matching 5 mm flush joint is appropriate.

The David Mellor Cutlery Factory, Hathersage, Derbyshire (Fig. 9.14) illustrates the use of traditionally detailed Derbyshire stone as loadbearing masonry worked in conjunction with precast concrete quoins and padstones. The building takes its form from the base of an old gasholder which provides its foundations.

**Gabions**

Gabions are wire cages filled with crushed rocks or recycled concrete rubble. They are frequently used in civil engineering applications as retaining walls, and are simply stacked to the required height either vertically or to an appropriate incline. Compressive loads
Fig. 9.14 Load-bearing stone masonry — David Mellor Cutlery Factory, Hathersage, Derbyshire. Architects: Hopkins Architects. Photographs: Arthur Lyons
are transmitted through the stones or concrete rubble, and any spreading movement is restrained by the tensile forces within the wire cage. Normally, the cages are of heavy gauge woven or welded steel mesh, which may be zinc, aluminium/zinc alloy, or PVC coated, but for use in load-bearing building applications, such as walls, stainless steel should be used. Gabions are now being used as significant components in building construction, where the particular rugged aesthetic is required (Fig. 9.15). Gabions may be delivered on site
filled or flat packed for filling and fastening, usually with a helical binder in alloy coated or stainless steel. A range of sizes is available based mainly on a metre module.

As an alternative, softer aesthetic, landscape-retaining structures may be constructed from a combination of stones, steel mesh and timber (Fig. 9.16).

**STONE CLADDING**

For the majority of large commercial buildings, stone is used as a cladding material mechanically fixed to the structural system. The strength of the stone largely determines the appropriate cladding panel thickness. For granites, marbles and slate 40 mm slabs are usual for external elevations above ground floor level, but for the softer limestones and sandstones a minimum thickness of 75 mm is frequently recommended. However, at heights of less than 3.7 m above ground level, thinner sections are permissible provided that they are of sufficient strength not to suffer distortion and failure. The standard (BS 8298: 1994) gives details of stone thicknesses for external cladding and internal lining. Fixings (Fig. 9.17) must be manufactured from stainless steel or non-ferrous metal and must be sized to sustain the dead load of the cladding together with applied loads from wind and maintenance equipment. Movement joints are required to accept the differential structural movements of the frame and the thermal and moisture movements of the cladding. Horizontal compression joints of 15 mm minimum should be located at each floor level; vertical movement joints of 10 mm should be at approximately 6 m centres. Polysulphides, polyurethanes and silicones are used as joint sealants, although non-staining silicones should be used on stones which darken by absorption of silicone fluid. Stone cladding systems should ideally be protected from impact damage at ground level by the design detailing.
Concrete-backed stone cladding

An alternative approach to traditional stone cladding is the use of an integral stone veneer on concrete cladding panels. Stone is fixed to the concrete with a series of non-corroding dowels inclined in opposite directions, creating a mechanical fixing, not dependent on the bond between stone and cast concrete. With limestone, a stone veneer of 50 mm is appropriate. The concrete should be cast with appropriate reinforcement and fixings for attachment to the building structure.

Stone-faced masonry blocks

Stone-faced concrete blocks are manufactured to the standard size of $440 \times 215$ mm, and faced with a 10 mm veneer of polished marble or granite, fixed with a frost-resistant adhesive. A range of matching special shapes including quoins, end blocks and lintels is available, formed with mitred-stone corner joints.

Lightweight stone cladding

Thin-section stone (approximately 6 mm) can be bonded to lightweight backing materials to reduce the dead weight of stone cladding (Fig. 9.18). The reduction in dead load is significant compared to thick-stone sections which would require traditional stone cladding techniques. One such material, originally used in the aerospace industry, is a sandwich panel consisting of a core of honeycomb aluminium faced with glass-fibre reinforced epoxy resin skins. The polished stone facing is bonded to one face with epoxy resin, to create a lightweight stone-finished panel, which, if detailed appropriately, has all the visual qualities associated with solid stone masonry.

Rainscreen stone cladding

Ventilated rainscreen stone cladding systems usually incorporate natural stone backed with lightweight concrete and/or insulation and fixed to an aluminium support system. Stone of 8–10 mm is typically bonded to 19 mm lightweight concrete. External and internal corner special units are prefabricated to match the facade.

Deterioration of stone

The main agencies causing the deterioration of stone are soluble salt action, atmospheric pollution, frost, corrosion of metal components and poor design or workmanship.

SOLUBLE SALT ACTION

If moisture containing soluble salts evaporates from the surface of stonework, then the salts will be left on the surface either as white efflorescence or as crystals within the porous surface layer. If the wetting and
drying cycles continue, the crystalline material builds up within the pores to the point at which the pressure produced may exceed the tensile strength of the stone, causing it to crumble. The actual pore size significantly influences the durability of individual stones, but generally the more porous stones, such as limestone and sandstone, are susceptible to soluble salt action.

**ATMOSPHERIC POLLUTION**

Stones based on calcium carbonate are particularly vulnerable to attack by acid atmospheric pollutants. Sulphur dioxide in the presence of water and oxygen from the air produces sulphuric acid, which attacks calcium carbonate to produce calcium sulphate. Limestones and calcareous sandstones are vulnerable to attack. In the case of limestone, the gypsum (calcium sulphate) produced on the surface is slightly soluble, and on exposed surfaces, it gradually washes away leaving the eroded limestone clean. In unwashed areas, the surface becomes blackened with soot producing a hard crust, which eventually blisters exposing powdered limestone. Magnesian limestones react similarly, except that in some cases the recrystallisation of magnesium sulphate under the blackened crust causes a more serious cavernous decay of the stone. Calcareous sandstones, when rain-washed, gradually decay to powder; however, in unwashed areas they produce a hard crust in which the pores are blocked with gypsum. The crust eventually fails due to differential thermal expansion. Dolomitic sandstones are less vulnerable to acid attack, unless they contain a significant proportion of vulnerable calcite. Silicious sandstones, which are not attacked directly by atmospheric acids, can be damaged by the calcium sulphate washings from limestone, which then cause crystallisation damage to the sandstone surface. Marble, which is essentially calcium carbonate, is also affected by atmospheric acids. Any polished surface is gradually eroded; however, as marble is generally non-porous, crystallisation damage is unusual, and limited to sugaring in some cases.

**FROST ACTION**

Frost damage occurs in the parts of a building which become frozen when very wet, such as copings, cornices, string courses, window hoods and cills. Frost causes the separation of pieces of stone, but it does not produce powder as in crystallisation attack. Generally, limestones and magnesian limestones are more vulnerable to frost damage than sandstones. Marble, slate and granite used in building are normally unaffected by frost due to their low porosities.

**CORROSION OF METALS**

Rainwater run-off from copper and its alloys can cause green colour staining on limestones. Iron and steel produce rust staining which is difficult to remove from porous stones. Considerable damage is caused by the expansion of iron and steel in stonework caused by corrosion. All new and replacement fixings should normally be manufactured from stainless steel or non-ferrous metals.

**FIRE**

Fire rarely causes the complete destruction of stonework. In the case of granite, marble and most sandstones, the surfaces may be blackened or spall. Limestones are generally unaffected by fire, although the paler colours may turn permanently pink due to the oxidation of iron oxides within the stone. Reigate stone, a calcareous sandstone, is also resistant to heat, but it is not a durable stone for exterior use.

**PLANTS**

Generally, large plants including ivy should be removed from old stonework; however, Virginia creeper and similar species are not considered harmful. Lichens may contribute to deterioration of limestones, and affected stonework should be treated. Damp north-facing walls and sloping sandstone surfaces are vulnerable to developing algae and lichen growth.

**Maintenance of stonework**

**CLEANING**

External granite, marble and slate claddings require regular washing with a mild detergent solution. Particularly, highly polished external marble should be washed at least twice per year to prevent permanent dulling of the surface. Limestone, where it is not self-cleaned by rainwater, should be cleaned with a fine water spray and brushing, removing only deposit and not the gypsum-encrusted surface. However, the washing of limestone may cause a ginger staining or efflorescence as the stone dries out and risk the possible corrosion of embedded ferrous cramps, so water
quantities should be adequately controlled. Sandstone is usually cleaned mechanically by abrasive blasting or chemical cleaning. Abrasive blasting with sand or grit is satisfactory for hard stones but can seriously damage soft stone and moulded surfaces. Hydrofluoric acid and sodium hydroxide (caustic soda) are used in the chemical cleaning of sandstones, but both are hazardous materials which need handling with extreme care by specialist contractors.

**STONE PRESERVATION**

Generally, coatings such as silicone water repellents should only be applied to stonework following expert advice and testing. Silicone treatment may in certain cases cause a build-up of salt deposits behind the treated layer, eventually causing failure. Silicone treatment should not be applied to already decayed stone surfaces. Polymeric silanes can be used to consolidate decaying stone. The silane is absorbed up to 50 mm into the stone, where it polymerises stabilising the stone but without significantly changing its external appearance. Different silane compositions offer a range of solidification and water-repellent properties which must be matched to the substrate. Generally, such treatment is appropriate for small artefacts which are in immediate danger of loss if left untreated.

**Cast stone**

The appearance of natural stones, such as Bath, Cotswold, Portland and York, can be recreated using a mixture of stone dust and natural aggregates with cement. In certain cases, iron oxide pigments may also be added to match existing stonework as required. Many architectural components such as classical columns, capitals, balustrades and porticos are standard items (Figs. 9.19 and 9.20), but custom-made products may be cast to designers’ specifications as illustrated in the facade of the Thames Water building at Reading (Fig. 9.21). High-quality finishes are achieved by the specialist manufacturers, and cast stone often surpasses natural stone in terms of strength and resistance to moisture penetration. Cast stone may be homogeneous, or, for reasons of economy, may have the facing material intimately bonded to a backing of concrete, in which case the facing material should be at least 20 mm thick. The standard BS 1217: 2008 requires that untreated steel reinforcement should have at least

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**Fig. 9.19** Typical cast stone units

**Fig. 9.20** Cast stone classical Ionic column
40 mm cover from any visual faces or a minimum of 30 mm if galvanised. Corrosion-resistant metals (e.g. stainless steel, bronze or copper) require at least 10 mm cover from any exposed face. Most masonry units are designed to be installed with 5 or 6 mm joints, and locating holes for dowel joints should be completely filled. Mortars containing lime are recommended rather than standard sand and cement (Table 9.5). Careful workmanship is required to prevent staining of the cast stone surfaces with mortar, as it is difficult to remove. Cast stone should weather in a similar manner to the equivalent natural stone.

**Table 9.5  Recommended grades of mortar for cast stonework**

<table>
<thead>
<tr>
<th>Exposure</th>
<th>Masonry cement : sand</th>
<th>Plasticised cement : sand</th>
<th>Cement : lime : sand</th>
</tr>
</thead>
<tbody>
<tr>
<td>Severe</td>
<td>1 : 4½</td>
<td>1 : 6</td>
<td>1 : 1 : 6</td>
</tr>
<tr>
<td>Moderate</td>
<td>1 : 6</td>
<td>1 : 8</td>
<td>1 : 2 : 9</td>
</tr>
</tbody>
</table>

The process is used for the repetitive casting of smaller components, which can be removed from the mould immediately after compaction, allowing many units to be made each day. The wet cast stone system is used for the manufacture of larger units, which remain in the mould for 24 hours, and may incorporate anchor fixings and more complex reinforcement.

**References**

**FURTHER READING**


### STANDARDS

| BS 5080 | Structural fixings in concrete masonry: |
| BS 5385 | Wall and floor tiling: |
| BS 5628 | Code of practice for the use of masonry: |
| Part 2: 2005 | Structural use of reinforced and prestressed masonry. |
| BS 5642 | Sills and copings: |
| BS 6100 | Building and civil engineering. Vocabulary: |
| BS 7533 | Pavements constructed with clay, natural stone or concrete pavers: |
| Part 4: 2006 | Construction of pavements of precast concrete flags or natural stone slabs. |
| Part 7: 2002 | Construction of pavements of natural stone setts and cobbles. |
| Part 8: 2003 | Structural design of lightly trafficked pavements of precast concrete and natural stone flags. |
| Part 10: 2004 | Structural design of trafficked pavements constructed of natural stone setts. |
| Part 12: 2006 | Structural design of trafficked pavements using concrete paving flags and natural stone slabs. |
| BS 8000 | Workmanship on building sites: |
| Part 11 | Code of practice for wall and floor tiling: |
| Sec. 11.1: 1989 | Ceramic tiles, terrazzo tiles and mosaics. |
| Sec. 11.2: 1990 | Natural stone tiles. |
| BS 8221 | Code of practice for cleaning and surface repair of buildings: |


BS 8298: 1994 Code of practice for design and installation of natural stone cladding and lining:
   pr Part 2: 2007 Traditional handset external cladding.
   pr Part 4: 2007 Rainscreen and stone on metal frame cladding systems.

BS EN 771 Specification for masonry units:
   Part 5: 2003 Manufactured stone masonry units.
   Part 6: 2005 Natural stone masonry units.

BS EN 772 Methods of test for masonry units:

BS EN 1341: 2001 Slabs of natural stone for external paving. Requirements and test methods.


BS EN 1343: 2001 Kerbs of natural stone for external paving. Requirements and test methods.


BS EN 12326 Slate and stone products for discontinuous roofing and cladding:

BS EN 12370: 1999 Natural stone test methods. Determination of resistance to salt crystallisation.

BS EN 12371: 2001 Natural stone test methods. Determination of frost resistance.


BS EN 12407: 2007 Natural stone test methods. Petrographic examination.

BS EN 12440: 2008 Natural stone. Denomination criteria.


BS EN 14579: 2004 Natural stone test methods. Determination of sound speed propagation.


DD CEN/TS 15209: 2008 Tactile paving surface indicators produced from concrete, clay and stone.

BUILDING RESEARCH ESTABLISHMENT PUBLICATIONS

BRE Digests


BRE Digest 449: 2000 Cleaning exterior masonry (Parts 1 and 2).

BRE Digest 467: 2002 Slate and tile roofs: avoiding damage from aircraft wake vortices.


BRE Good building guide


BRE Information papers

BRE IP 6/97 External cladding using thin stone.
BRE IP 7/98 External cladding—how to determine the thickness of natural stone panels.
BRE IP 17/98 Use of lightweight veneer stone claddings.
BRE IP 18/98 Stone cladding panels—in situ weathering.
BRE IP 9/99 Cleaning exterior masonry.
BRE IP 10/00 Flooring, paving and setts.
BRE IP 10/01 Lightweight veneer stone cladding panels.

BRE Reports


BR 84: 1986 The building sandstones of the British Isles, E. Leary.

ADVISORY ORGANISATIONS

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National Federation of Terrazzo, Marble & Mosaic Specialists, PO Box 2843, London W1A 5PG, UK (0845 609 0050).
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PLASTICS

Introduction

The plastics used in the construction industry are generally low-density non-load-bearing materials. Unlike metals, they are not subject to corrosion, but they may be degraded by the action of direct sunlight, with a corresponding reduction in mechanical strength. Many plastics are flammable unless treated; the majority emit noxious fumes in fires. Approximately 20% of plastics produced within the UK are used by the building industry. Polyvinyl chloride (PVC), which has a high embodied energy content, accounts for 40% of this market share, predominantly in pipes, but also in cladding, electrical cable insulation, windows, doors and flooring applications. Foamed plastics for thermal and acoustic insulation are formulated as either open or closed-cell materials, the latter being resistant to the passage of air and water.

In terms of their chemical composition plastics form a diverse group of materials which have chain-like molecular structures composed of a large number of small repeat units. Whilst some materials such as rubber and cellulose derivatives are based on natural products, the majority of plastics are produced from petrochemical products. The manufacture of polythene, which dates back to 1933, involves the polymerisation of ethylene monomer, a colourless gas, which under high pressure at 200°C, is converted into the clear polymer, polyethylene or polythene (Fig. 10.1).

Polymerisation

In the production of polythene the small molecular units of ethylene are joined end to end by an addition polymerisation process to produce the long-chain macromolecules. A similar process converts vinyl chloride into polyvinyl chloride (PVC) (Fig. 10.2), styrene monomer into polystyrene and tetrafluoroethylene into polytetrafluoroethylene (PTFE).

While the molecular backbones of plastics are predominantly composed of chains of carbon atoms, variations occur, particularly when the polymerisation process involves the elimination of water between adjacent monomer units. Thus, in the case of condensation
polymerisation (Fig. 10.3), oxygen or nitrogen atoms are incorporated into the backbone of the macromolecular chains as in the polyesters (resins) and polyamides (nylons).

**BRANCHED CHAINS**

Depending on the conditions during the polymerisation process, the polymer chains produced may be linear or branched. In the case of polythene, this affects the closeness of packing of the chains and therefore the bulk density of the material. Thus high-density polythene (HDPE) (s.g. 0.97), which is relatively stiff, has few branched chains compared to low-density polythene (LDPE) (s.g. 0.92), which is softer and waxy (Fig. 10.4).

**COPOLYMERS**

Where two or more different monomers are polymerised together, the product is a copolymer. The properties of the copolymer are significantly dependent on whether the two components have joined together in alternating, random or block sequences (Fig. 10.5).

More complex plastics can be produced for their specific physical properties by combining several components. Thus, acrylonitrile butadiene styrene (ABS) is produced by grafting styrene–acrylonitrile onto a preformed butadiene latex of a carefully controlled particle size.

**CRYSTALLINITY**

In the initial manufactured state, most polymers consist of amorphous randomly orientated molecular chains. However, if the plastic material is stretched in one direction, such as during the drawing of spun fibres, it causes an alignment of the molecular chains, leading to partial formation of crystalline regions and an associated anisotropy (Fig. 10.6). Crystalline regions may also be produced during the solidification of simple polymers such as polyethylene, but they will be limited in their extents due to the general entanglement of the molecular chains.

**GLASS TRANSITION TEMPERATURE**

In the molten state, the individual molecular chains of a plastic material move freely relative to each other, allowing the material to be moulded within the various forming processes used for the manufacture of components. As the temperature of melted plastic material

![Fig. 10.3 Condensation polymerisation](image)

![Fig. 10.4 Straight- and branched-chain plastics](image)

![Fig. 10.5 Random, alternate and block copolymers](image)
is lowered, the freedom of movement of the molecular chains is reduced; gradually the plastic becomes more viscous, until eventually it solidifies at its characteristic melting point temperature. However, even when solid, most plastics remain rubbery or flexible, due to rotations within the individual molecular chains. As the temperature is lowered further, the material will eventually become rigid and brittle, as movement can no longer take place within the individual molecular units. The temperature at which a particular plastic changes from flexible to rigid is defined as its characteristic glass transition temperature. Depending on the nature of the particular plastic material this may be above or below normal ambient temperatures. Further, the glass transition temperature for a particular plastic can be significantly changed by, for example, the addition of plasticisers, characterised by the differences in physical properties between PVC-U (unplasticised) and PVC (plasticised polyvinyl chloride).

**Polymer types**

Polymers are normally categorised in respect of their physical properties as either thermoplastic, thermosetting or elastomeric.

**THERMOPLASTICS**

Thermoplastics soften on heating and reset on cooling. The process is reversible and the material is unaffected by repeating the cycle, provided that excessive temperatures, which would cause polymer degradation, are not applied. Many thermoplastics are soluble in organic solvents, whilst others swell by solvent absorption. Thermoplastics are usually produced initially in the form of small granules for subsequent fabrication into components.

**THERMOSETTING PLASTICS**

Thermosetting plastics have a three-dimensional cross-linked structure, formed by the linkage of adjacent macromolecular chains (Fig. 10.7). Thermosets are not softened by heating, and will only char and degrade if heated to high temperatures. Thermosets are usually produced from a partially polymerised powder or by mixing two components, such as a resin and a hardener. The resin is essentially the macromolecular component and the hardener cross-links the liquid resin into the thermoset plastic. Curing for epoxy resin adhesives and polyesters as in glass-fibre reinforced polyester (GRP) occurs at room temperature, while for phenolic and formaldehyde-based resins, a raised temperature and pressure are required. Thermosets, because of their three-dimensional structure, are usually solvent resistant and harder than thermoplastics.

**ELASTOMERS**

Elastomers are long-chain polymers in which the naturally helical or zig-zag molecular chains are free to
straighten when the material is stretched, and recover when the load is removed. The degree of elasticity depends on the extensibility of the polymeric chains. Thus natural rubber is highly extensible, but when sulphur is added, the vulcanisation process increasingly restricts movement by locking together adjacent polymer chains (Fig. 10.8). For most uses some cross-linking is required to ensure that an elastomeric material returns to its original form when the applied stress is removed.

Additives

PLASTICISERS

Plasticisers are frequently incorporated into plastics to increase their flexibility. The addition of the plasticiser separates the molecular chains, decreasing their mutual attraction. Thus unplasticised PVC (PVC-U) is suitable for the manufacture of rainwater goods, window units and glazing, whereas plasticised PVC is used for flexible single-layer roof membranes, tile and sheet floor coverings and electrical cable insulation. Loss of plasticiser by migration can cause eventual embrittlement of plasticised PVC components.

PIGMENTS AND STABILISERS

Dyes and pigments may be added to the monomer or polymer. Stabilisers are added to absorb ultraviolet light which otherwise would cause degradation. For example, organotin compounds are used in clear PVC sheet to preferentially absorb incident ultraviolet light, in order to prevent degradation by the elimination of hydrogen chloride.

Degradation of plastics

The degradation of plastics is most frequently attributed to the breakdown of the long molecular chains (Fig. 10.9), or in the case of PVC, the loss of plasticiser. Polymeric molecular chains may be broken by the effect of heat, ultraviolet light or ozone, or by a combination of any of these factors, thus reducing their average molecular chain length. Discolouration occurs through the production of molecular units with double bonds, usually causing a yellowing of the plastic. Surface crazing and stress cracks may develop where degradation has caused cross-linking, resulting in embrittlement of the surface.

Where plasticiser is lost by migration from PVC, the glass transition temperature is gradually raised, so eventually the material becomes brittle at ambient temperatures. Typically, high-boiling point oils such as dibutyl phthalate and dioctyl phthalate are incorporated into the original PVC, but these gradually evaporate leaving the surface vulnerable to cracking and shrinkage.

Properties of plastics

FIRE

All plastics are combustible, producing noxious fumes and smoke (Table 10.1). Carbon monoxide
Degradation of plastics is produced by most organic materials, but in addition, plastics containing nitrogen, such as polyurethane foam, generate hydrogen cyanide, and PVC produces hydrochloric acid. Some plastics, particularly acrylics and expanded polystyrene, have a high surface spread of flame and produce burning droplets; however, others when treated with fire retardant are difficult to ignite and some are self-extinguishing.

**STRENGTH**

Although plastics have a good tensile strength-to-weight ratio, they also have a low modulus of elasticity which renders them unsuitable for most load-bearing situations, the only exception being glass-fibre-reinforced polyester which has been used for some limited load-bearing applications. Generally, thermoplastics soften at moderate temperatures and are subject to creep under ambient conditions.

**THERMAL AND MOISTURE MOVEMENT**

The thermal expansion of most plastics is high. The expansion of GRP is similar to that of aluminium, but most other plastics have larger coefficients of linear expansion. For this reason, attention must be paid to careful detailing to allow for adequate thermal movement, particularly where weather exclusion is involved. Most plastics are resistant to water absorption, and therefore do not exhibit moisture movement. (Typical coefficients of linear expansion are polythene (HD) \((110–130) \times 10^{-6} \text{C}^{-1}\), polypropylene \(110 \times 10^{-6} \text{C}^{-1}\), ABS \((83–95) \times 10^{-6} \text{C}^{-1}\), PVC \((40–80) \times 10^{-6} \text{C}^{-1}\), GRP \((20–35) \times 10^{-6} \text{C}^{-1}\).)

<table>
<thead>
<tr>
<th>Material</th>
<th>Behaviour in fire</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thermoplastics</td>
<td></td>
</tr>
<tr>
<td>Polythene, Polypropylene</td>
<td>Melts and burns readily</td>
</tr>
<tr>
<td>Polyvinyl chloride</td>
<td>Melts, does not burn easily, but emits smoke and hydrogen chloride</td>
</tr>
<tr>
<td>PTFE/ETFE</td>
<td>Does not burn, but at high temperatures evolves toxic fumes</td>
</tr>
<tr>
<td>Polymethyl methacrylate</td>
<td>Melts and burns rapidly, producing droplets of flaming material</td>
</tr>
<tr>
<td>Polystyrene</td>
<td>Melts and burns readily, producing dense black smoke and droplets of flaming material</td>
</tr>
<tr>
<td>ABS copolymer</td>
<td>Burns readily</td>
</tr>
<tr>
<td>Polyurethane</td>
<td>The foam burns readily producing highly toxic fumes including cyanides and isocyanates</td>
</tr>
<tr>
<td>Thermosetting plastics</td>
<td></td>
</tr>
<tr>
<td>Phenol formaldehyde, Melamine formaldehyde</td>
<td>Resistant to ignition, but produce noxious fumes including ammonia</td>
</tr>
<tr>
<td>Urea formaldehyde</td>
<td>Burns producing smoke, but flame-retarded grades are available</td>
</tr>
<tr>
<td>Glass-reinforced polyester (GRP)</td>
<td></td>
</tr>
<tr>
<td>Elastomers</td>
<td>Burns readily producing black smoke and sulphur dioxide</td>
</tr>
<tr>
<td>Rubber</td>
<td>Better fire resistance than natural rubber</td>
</tr>
<tr>
<td>Neoprene</td>
<td></td>
</tr>
</tbody>
</table>
**Plastics forming processes**

Depending on the nature of the product, plastics may be formed by either continuous or batch processes. With thermoplastics, frequently a two-stage process is most appropriate in which the raw materials, supplied by the primary manufacturer as powder or granules, are formed into an extrusion or sheet which is then reformed into the finished product. However, thermosetting plastics must be produced either from a partially polymerised material or directly from the resin and hardener mix in a single-stage process. Most foamed plastics are either blown with internally generated gas, or produced by a vacuum process, which reduces reliance on the previously used environmentally damaging CFCs and HCFCs.

**CONTINUOUS PROCESSES**

**Extrusion**

Plastic granules are fed continuously into the heated barrel of a screw extruder, which forces the molten thermoplastic through an appropriately shaped die to produce rod, tube or the required section (Fig. 10.10). Products include pipes, rainwater goods and fibres.

**Film blowing**

As a molten thermoplastic tube is produced in the extrusion process, air is blown in to form a continuous cylindrical plastic sheet, which is then rolled flat and trimmed to produce a folded sheet. Adjustment of the applied air pressure controls the sheet thickness.

**Calendering**

Sheet thermoplastic materials may be produced from plastics granules by compression and fusion between a series of heated rollers. Laminates may be produced by heating together two or more thermoplastic sheets, and during this process, sheet reinforcement material may be incorporated.

**BATCH PROCESSES**

**Injection moulding**

Thermoplastic granules are melted in a screw extruder to fill a ram which injects the plastic into an appropriate mould. After cooling, the components are removed from the mould and trimmed as necessary. The process is low cost and rapid. A series of moulds can be attached to the injection moulding machine to ensure continuity of production (Fig. 10.11). Thermosetting polymers can be injection moulded by initial forming at a low temperature followed by heating of the mould to cross-link the liquid plastic.

**Compression moulding**

In the compression moulding process for thermosetting resins, the appropriate quantity of uncross-linked resin powder is subjected to pressure and heat within the mould. When the polymer has melted and cross-linked, the mould can be opened and the component removed.

**Pressing**

Pressing is used to form thermoplastic sheet into components. The sheet plastic is initially heated to...
softening point and then pressed between an appropriately shaped pair of dies.

**Vacuum forming and blow moulding**

During vacuum forming, thermoplastic sheet is heated over a mould which is then evacuated through a series of fine holes, drawing the soft plastic into the appropriate form. In the similar process, blow moulding, positive air pressure is applied inside a molten polymer tube which is expanded into the shape of the mould.

**RAPID PROTOTYPING**

New techniques in computer-aided manufacturing enable prototype components to be manufactured to very close tolerances from three-dimensional computer-aided design (CAD) solid modelling images. This has implications not only for the design of building components but also for the manufacture of architectural models.

The systems are based on the successive build-up of very thin layers of solid material to the exact pattern of layered CAD sections. Various lay-up systems have been developed for the deposition of plastic layers. These range from a fine nozzle, to using laser technology to accurately polymerise viscous resin in very thin layers and the use of adhesive-backed paper cut by laser to the required section shapes. Each system produces a highly accurate three-dimensional solid, over a period of several hours depending on the product size. Where any part of the build-up of the solid object needs support during manufacture, the system automatically produces additional material in a weak form. This can be broken away easily after the whole object is complete, and in the case of laser/resin production finally cured. In all these manufacturing processes the build-up layers are extremely thin, so smooth and accurate surfaces are achieved.

The reverse of this process allows prototype complex-shape components or small-scale architectural models to be turned into accurate three-dimensional CAD files, using a delicate probe mechanism which senses all over the object’s surfaces. This allows the designer to generate CAD files for highly complex three-dimensional forms, which would be virtually impossible to draw directly into a CAD system.

**Plastics in construction**

The broad range of thermoplastic, thermosetting and elastomeric plastics are collated into families in Fig. 10.12. Typical uses in construction are listed in Table 10.2.

(Glass-fibre reinforced polyester is described in Chapter 11, foamed plastics as insulation materials in Chapter 13 and plastics used primarily as adhesives in Chapter 14.)

**THERMOPLASTICS**

**Polythene (polyethylene)**

Polyethylene (PE) is one of the cheapest plastics and is available in a range of grades of differing densities and physical properties. Low-density polyethylene (LDPE) has a softening point of 90°C and high-density polyethylene (HDPE) softens at 125°C.

<table>
<thead>
<tr>
<th>Grades of polyethylene</th>
<th>Density (kg/m³)</th>
</tr>
</thead>
<tbody>
<tr>
<td>LDPE</td>
<td>900–939</td>
</tr>
<tr>
<td>VLDPE</td>
<td>880–915</td>
</tr>
<tr>
<td>LDPE</td>
<td>916–925</td>
</tr>
<tr>
<td>MDPE</td>
<td>926–940</td>
</tr>
<tr>
<td>HDPE</td>
<td>941–970</td>
</tr>
<tr>
<td>HMWPE</td>
<td>947–950</td>
</tr>
<tr>
<td>UHMWPE</td>
<td>930–935</td>
</tr>
<tr>
<td>PEX</td>
<td>926–970</td>
</tr>
</tbody>
</table>

Polyethylene is resistant to chemicals and tough at low temperatures, but is rapidly embrittled by ultraviolet light unless carbon black is incorporated. Polyethylene burns and has a relatively high coefficient of thermal expansion. Low-density polyethylene is used widely for damp-proof membranes, damp-proof courses and vapour barriers. High-density polyethylene, which is stiffer than the low-density material, is used for tanking membranes to basements. Ultra-high molecular weight polyethylene (melting point 130°C) has chemical inertness and good environmental stress crack resistance to solvents, oils and detergents. Linear low-density polyethylene, used for pipes and cable jacketing, is easily extruded and has good impact resistance.
Polyethylene is used for the production of cold-water cisterns, but is only suitable for cold-water plumbing applications due to its high thermal expansion. Mains water pressure pipes, which are manufactured from medium-density (PE80) and ultra-high molecular weight (PE100) polyethylene, require a significant wall thickness due to the relatively low tensile strength of polyethylene.

Cross-linked polyethylene (PEX), manufactured by the action of peroxide catalyst on normal polyethylene, is used for domestic hot water and underfloor heating systems as it can withstand operating temperatures up to 90°C. In underfloor heating systems, an interlayer of aluminium is incorporated into the PEX pipe to prevent the ingress of oxygen which would cause corrosion of steel components within the system. Certain barrier composite pipes can operate at temperatures up to 95°C.

The standard BS EN ISO 11542-1: 2001 describes ultra-high molecular weight polyethylene.

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**Polypropylene**

Polypropylene (PP), with a softening point of 150°C, is slightly stiffer than polyethylene, to which it is closely related chemically. Like polyethylene, it is resistant to chemicals and susceptible to ultraviolet light, but unlike polyethylene, it becomes brittle below 0°C. However, the block copolymer with ethylene does have improved low-temperature impact resistance. Polypropylene is used for pipes, drainage systems, water tanks, DPCs, connecting sleeves for clay pipes and WC cisterns. Polypropylene fibres are used in fibre-reinforced cement to produce an increase in impact resistance over the equivalent unreinforced material. Polypropylene permanent shuttering for concrete is frequently a double-layer system incorporating polystyrene insulation and radon protection where necessary. Certain breather membranes used for tile underlay and timber-frame construction are manufactured from multilayer systems incorporating polypropylene with polyethylene and glass-fibre...
**Table 10.2  Typical uses of plastics in construction**

<table>
<thead>
<tr>
<th>Material</th>
<th>Examples of plastics in construction</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Thermoplastics</strong></td>
<td></td>
</tr>
<tr>
<td>Polythene</td>
<td>DPC, DPM, vapour checks, roof sarking</td>
</tr>
<tr>
<td>(low density LD)</td>
<td>Cold water tanks, cold water plumbing</td>
</tr>
<tr>
<td>(high density HD)</td>
<td>Hot and cold water plumbing</td>
</tr>
<tr>
<td>(cross-linked PEX)</td>
<td>Pipework and fittings, cold water plumbing, drainage systems, water tanks, WC cisterns, DPCs, fibres in fibre-reinforced concrete</td>
</tr>
<tr>
<td>Polypropylene (PP)</td>
<td>Hot and cold-water pipework and fittings</td>
</tr>
<tr>
<td>Polyvinyl chloride (PVC-U)</td>
<td>Rainwater goods, drainage systems, cold water, underground services, window and door frames, conservatories, garage doors, translucent roofing sheets, cold water tanks</td>
</tr>
<tr>
<td>(PVC-UE)</td>
<td>Claddings, barge boards, soffits, fascias, window boards</td>
</tr>
<tr>
<td>(PVC)</td>
<td>Tile and sheet floor coverings, single-ply roofing, cable insulation, electrical trunking systems, sarking, tensile membrane structures, glazing to flexible doors, door seals, handrail coatings, vinyl-film finishes to timber products</td>
</tr>
<tr>
<td>(CPVC)</td>
<td>Hot and cold water systems, window and door frames</td>
</tr>
<tr>
<td>ETFE</td>
<td>Inflated systems for translucent wall and roof membranes</td>
</tr>
<tr>
<td>PTFE</td>
<td>Sealing tape for plumbing, tensile membrane structures, low-friction movement joints</td>
</tr>
<tr>
<td>Polymethyl methacrylate</td>
<td>Baths, shower trays, kitchen sinks, glazing, roof lights, luminaires</td>
</tr>
<tr>
<td>Polycarbonate</td>
<td>Vandal-resistant glazing, spa baths, kitchen sinks</td>
</tr>
<tr>
<td>Polystyrene</td>
<td>Bath and shower panels, decorative expanded polystyrene tiles</td>
</tr>
<tr>
<td>ABS copolymer</td>
<td>Pipes and fittings, rainwater goods, drainage systems, shower trays</td>
</tr>
<tr>
<td>Nylons</td>
<td>Electrical conduit and trunking, low-friction components—hinges, brush strips for sealing doors and windows, carpet tiles and carpets, shower curtains</td>
</tr>
<tr>
<td><strong>Thermosetting plastics</strong></td>
<td></td>
</tr>
<tr>
<td>Phenol formaldehyde</td>
<td>Decorative laminates</td>
</tr>
<tr>
<td>Melamine formaldehyde</td>
<td>Laminates for working surfaces and doors, moulded electrical components, WC seats</td>
</tr>
<tr>
<td>Urea formaldehyde</td>
<td>Decorative laminates</td>
</tr>
<tr>
<td>Glass-reinforced polyester (GRP)</td>
<td>Cladding and roofing panels, simulated cast-iron rainwater goods, cold-water tanks, spa baths, garage doors, decorative tiles and panels</td>
</tr>
<tr>
<td><strong>Elastomers</strong></td>
<td></td>
</tr>
<tr>
<td>Rubber</td>
<td>Flooring, door seals, anti-vibration bearings</td>
</tr>
<tr>
<td>Neoprene</td>
<td>Glazing seals, gaskets</td>
</tr>
<tr>
<td>EPDM</td>
<td>Glazing seals, gaskets, single-ply roofing systems</td>
</tr>
<tr>
<td>Butyl rubber</td>
<td>Sheet liners to water features and land-fill sites</td>
</tr>
<tr>
<td>Nitrile rubber</td>
<td>Tile and sheet flooring</td>
</tr>
</tbody>
</table>

reinforcement. Such products are wind and watertight, but vapour-permeable. Many geotextiles for soil stabilisation are manufactured as a mat material from non-woven heat-bonded polypropylene continuous fibres. The material may be reinforced by woven polyester fibres. Polypropylene is highly fatigue resistant and therefore used for integral hinges in lightweight components.

The standard BS EN ISO 1873-1: 1995 describes the designation system for polypropylene.

**Polybutylene**

Polybutylene (polybutene-1 PB) is used for pipework as an alternative to copper. It has the advantage of flexibility and the very smooth internal surface is resistant to the build-up of scale and deposits. It can withstand continuous operating temperatures up to 82°C. As polybutylene is slightly permeable to oxygen, some pipe manufacturers incorporate an ethyl vinyl alcohol copolymer (EVOH) barrier as an interlayer.
The standard BS EN ISO 8986-1: 1999 describes the designation system for polybutylene.

Polyvinyl chloride

Polyvinyl chloride is the most widely used plastic material in the construction industry. It is available both in the unplasticised form (PVC-U) and as the plasticised product (PVC). In both forms polyvinyl chloride is combustible giving off noxious hydrogen chloride fumes; however, the unplasticised form tends to burn only with difficulty. PVC begins to soften at 75°C, and therefore cannot be used for hot water systems, although chlorinated PVC (CPVC) can be used at higher temperatures. PVC is soluble in certain organic solvents which, therefore, can be used for the solvent welding of joints, but PVC is unaffected by acids and alkalis.

Plasticised PVC is extensively used in the manufacture of floor coverings, either as individual tile units or as continuously jointed sheet. It is also the most widely used material for single-layer roofing systems, due to its durability, colour range and ease of application. It also offers an alternative to bitumen sheet for sarking. Plasticised PVC is the standard for electrical cable insulation, and many small building components are made from injection-moulded PVC.

PVC-U

PVC-U is widely used for rainwater goods, usually in white, grey, black or brown, and similarly for soil and waste pipes. It is also used colour-coded for underground water, gas, electrical and telecommunications systems. PVC-U is used extensively for the manufacture of extruded window frames, door frames and conservatories, usually incorporating sealed double-glazing units. Where insufficient rigidity is achieved with the PVC-U alone, steel inserts within the extruded sections give strength and provide additional protection against forced entry. PVC-U is used in the manufacture of translucent, transparent and coloured profiled sheeting for domestic structures such as carports and conservatories, where an economical product is required, although eventually the products discolour and craze due to the effects of direct ultraviolet light. Current research suggests that PVC-U may be recycled several times without loss of performance. The recycled material is generally ground into granular form and reused in the co-extrusion of profile sections or in the production of 100% recycled PVC-U cavity closers.

PVC-UE

Extruded cellular unplasticised PVC (PVC-UE) is used for cladding, fascias, soffits, window boards, barge boards and other components of uniform section. It is manufactured by the co-extrusion of a high-impact PVC-U surface material over a core of closed-cell PVC foam. The foaming agent is usually sodium bicarbonate. The high stiffness-to-weight ratio arises from the combination of a dense outer skin and a cellular core. Both the cellular core and the wearing surface are stabilised with metallic additives to prevent degradation and discolouration. Contact with bitumen should be avoided. The material will char and melt in fire, but with a limited surface spread of flame. The material is described within the standard BS 7619: 1993.

Tensile membrane structures

PVC-coated polyester is the standard material used for tensile membrane structures and canopies. The durability depends directly on the degree of translucency; and at 15% transmission, a period of 15 years can be reasonably expected. At greater levels of translucency, the expected serviceable lifetime is considerably reduced; however, fluoropolymer lacquers to the top surface of the fabric enhance durability. While white fabric is the standard, coloured and patterned membranes are available according to client requirements. PVC-coated polyester membranes are a cheaper alternative to PTFE-coated fabrics, but are not non-combustible. PVC-coated polyester is more flexible than PTFE-coated fibreglass and is therefore the preferred material for temporary structures which may be folded for transport and storage. The thermal insulation afforded by single-layer tensile membrane roofs is negligible.

Tensile membrane structures are manufactured from a set of tailored panels stitched or welded together. They are usually tensioned by wires or rods running through edge pockets, or by fixing directly to structural elements. Accurate tensioning is required to generate the correct form and resistance to wind and snow loads. The use of double curvature within the panel elements imparts structural rigidity to the overall membrane structure. Damage by accident or vandalism can usually be repaired on site.

Polytetrafluoroethylene

Polytetrafluoroethylene (PTFE) coated glass-fibre woven fabrics are used for permanent tensile membrane structures. In a fire, PTFE gives off toxic
combustion products, but only at temperatures above which any fabric would have already failed and vented the heat and smoke. With a fire rating of Class 0, PTFE-coated glass-fibre tensile membranes are more expensive than the Class 1 rated PVC-coated polyesters, but are generally more durable with an anticipated lifespan in excess of 20–25 years. The low-friction PTFE surface has good self-cleaning properties.

The Inland Revenue Amenity Building in Nottingham (Fig. 10.13) and the Millennium Dome at Greenwich are roofed with PTFE (Teflon)-coated glass-fibre tensile membranes. The translucent fabric gives well-lit internal spaces during the daytime, and striking glowing surfaces at night. In the Nottingham building, the membrane roof is suspended from four steel columns, and is linked to the fixed structure below by inflatable elements which absorb any movement. The 321 m high Burj Al Arab seven-star hotel in Dubai (Fig. 10.14) is clad on one face with a PTFE-coated glass-fibre tensile membrane. It was the first building to use this material orientated vertically.

PTFE tape has a very low coefficient of friction and a high melting point. It is therefore ideal for use as a sealing tape for threaded joints in water and gas pipes. It is also used to form sliding joints in large structures.

The standard BS EN ISO 12086-1: 2006 describes the designation system for fluoropolymers.

**Ethylene tetrafluoroethylene**

Ethylene tetrafluoroethylene copolymer (ETFE) is used as a translucent foil for low-pressure pneumatic metal-framed building envelope cushions. The fluorocopolymer has the advantages over glass that when used to form two- to five-layer air cushion systems, it offers higher thermal insulation with greater transparency to UV light. ETFE is strong, shatter proof, half the cost and only one-hundredth the weight of the equivalent glass, thus offering significant economies to the required structural supporting system. ETFE with an anticipated life span of 25 years, can withstand maintenance loads, can be easily repaired and is recyclable. It has been used very effectively on the galvanised-tubular steel space-frame envelope for the biomes (domes sheltering plants from around the world) at the Eden Centre, Cornwall (Fig. 10.15). The structure is formed from an icosahedral geodesic outer layer, with a combination of hexagons, pentagons and triangles as the inner layer of the three-dimensional space frame. Only a small pumping system, powered by photovoltaic cells, is required to maintain the air-fill of the ETFE cushions. At the National Space Centre at Leicester, also designed by Grimshaw Architects, the ETFE cushion clad tower houses the main space rocket exhibits (Fig. 10.16).

If automatic smoke venting of an ETFE atrium is required, electrical wiring can be incorporated into the cushion frames, which release the cushions, except at one point, in case of fire. An enclosed atrium is thus turned into a fully open light well.

Transmitted light levels through ETFE cushions can be constantly adjusted by the use of partially printed internal layers within the cushions, which can be moved closer or further apart by changing the pumped air pressure, thus modifying the shadowing effect. Interesting patterns can be created by the use of coloured cushions, whilst aluminium-coated foils will give a highly reflective effect with reduced sunlight penetration.

**Polymethyl methacrylate**

Acrylic or polymethyl methacrylate (PMMA) is available in a wide variety of translucent or transparent, clear or brightly coloured sheets. It softens at 90°C, and burns rapidly with falling droplets of burning material. Stress crazing may occur where acrylic has been shaped in manufacture and not fully annealed, but generally the material is resistant to degradation by ultraviolet light. Acrylic is frequently used for decorative signs, rooflights and light fittings. Baths and shower trays are manufactured from acrylic as a lighter alternative to cast iron and ceramics. Although not resistant to abrasion, scratches can usually be polished out with proprietary metal polish. Polymethyl methacrylate from industrial and commercial sources can be fully recycled when clean and separated from other waste materials.

The standard BS ISO 8257-1: 1987 describes the designation system for polymethyl methacrylate.

**Polycarbonate**

Polycarbonates (PCs) are used as vandal-resistant glazing, due to their high impact resistance, good optical transparency and low ignitability. Polycarbonate glazing is used extensively for rooflights, roof domes, smoking shelters, car ports, covered walkways and road barriers. Polycarbonate blocks offer a lightweight alternative to traditional cast glass blocks. Proprietary extruded cellular systems of double-,
Fig. 10.13  Tensile membrane — Inland Revenue Amenity Building, Nottingham. Architects: Hopkins Architects. Photographs: Courtesy of Martine Hamilton Knight
Fig. 10.14  Teflon-coated glass fibre membrane — Burj Al Arab Hotel, Dubai. Architects: WS Atkins. Photograph: Athur Lyons

triple-, five- and seven-walled polycarbonates offer combined thermal insulation and vandal-resistant properties. The protective outer surface prevents ultraviolet degradation for 10 years, and sections may be curved on site within the limits of the manufacturers’ specifications.

The standard BS EN ISO 7391-1: 2006 describes the designation system for polycarbonate.

**Acrylonitrile butadiene styrene**

Acrylonitrile butadiene styrene (ABS) plastics are a range of complex terpolymers manufactured by combining together the two copolymers, styrene–acrylonitrile and butadiene–styrene. ABS plastics are relatively expensive but tough and retain their strength at low temperatures. They are used to manufacture moulded components, rainwater and drainage goods. A special ABS solvent cement is required for solvent welding.

**Nylons**

Nylons, usually nylon 66 or nylon 6, are used for the manufacture of small components where low friction is required. Nylons are tough and strong but tend to be embrittled and become powdery on prolonged exposure to sunlight. Carpet tiles in nylon 66 are durable and hard wearing.

**Kevlar**

*Kevlar* (polyparabenzamide) fibres are produced by extrusion of a cold solution of the polymer into a cylinder at 200°C, which causes the solvent to evaporate. The resulting fibres are stretched by a drawing process, which aligns the polymer molecules along the fibres to produce a very high modulus material, used in ropes and composite plastics.

**THERMOSETTING PLASTICS**

**Phenol formaldehyde**

Phenol formaldehyde (PF) was the original, and remains the cheapest, thermosetting resin. Currently, its main use is in the production of laminates by the hot pressing of layers of resin-impregnated paper, fabric or glass fibre. The cured resin is brown, but heat-resistant laminates for working surfaces and wallboards are laminated with a decorative printed paper film and coated with a clear melamine formaldehyde finish. Phenol formaldehyde is resistant to ignition, but produces a phenolic smell on burning.

**Urea formaldehyde**

Urea formaldehyde (UF) is similar to phenol formaldehyde except that because it is clear it can be produced in a range of colours including white. It is used in the manufacture of electrical components and other moulded components such as WC seats. Urea formaldehyde is resistant to ignition, but produces a fishy smell on burning. Urea formaldehyde foam is no longer used for cavity wall insulation.

**Melamine formaldehyde**

Melamine formaldehyde (MF) is available clear and in a wide range of colours. When heat cured, it is hard wearing, durable and resistant to heat, and is therefore used as the surface laminate over the cheaper brown phenol formaldehyde layers in the production of working surface and wallboard laminates. Melamine formaldehyde is resistant to ignition, but produces a fishy smell on burning.
ELASTOMERS

Natural rubber

Natural rubber is harvested from the species *Hevea brasiliensis*, in Africa, South America and Malaysia. The white latex is predominantly cis-polyisoprene, a macromolecule containing some double bonds within the carbon chain. It is these double bonds which permit cross-linking with sulphur when natural rubber is heated under pressure in the vulcanisation process. Natural rubber is usually reinforced with carbon and treated with antioxidants to prevent degradation. It is used for flooring and in antivibration bearings for buildings and large structures. Natural rubber has the advantage that it is a renewable resource and also the rubber trees absorb carbon dioxide during growth.

Uncontaminated natural rubber can be recycled several times without severe degradation.

Neoprene

Neoprene (polychloroprene), unlike natural rubber, is resistant to chemical attack, and is therefore used for glazing seals and gasket systems. It is available only in black.

EPDM

Unlike neoprene, ethylene propylene diene monomer (EPDM) can be obtained in any colour, and is characterised by high elongation and good weathering resistance to ultraviolet light and ozone. It is therefore taking over from neoprene as the key material for
gaskets and weather strips. EPDM is extensively used in single-ply roofing systems.

**Butyl rubber**

A copolymer of isobutylene and isoprene, this material has good chemical and weathering resistance. It is used as liners to landfill sites and decorative water features.

**Nitrile rubber**

Nitrile rubber is formed by the copolymerisation of acrylonitrile and butadiene. It is water and oil resistant, so it is frequently used in structural movement joints which may be subject to surface oil.

**COMPOSITE PLASTICS**

Composite plastic materials, such as glass-fibre reinforced polyester (Chapter 11), have physical properties which differ significantly from those of the individual component materials. An increasing variety of composite plastics are reaching the construction industry, driven by the demand for product diversity and in some cases recycling.

**Wood plastic composites**

Wood plastic composites (WPC) encompass a range of materials incorporating polymers such as polyethylene, polypropylene and polyvinyl chloride blended with natural fibres and saw dust. The plastics materials may be virgin or recycled. The natural fibres are usually wood chips, but hemp, sisal, jute or rice may also be used. The materials are defined in the publications DD CEN/TS 15534: 2007 and BRE Digest 480: 2004.

For the standard production of wood plastic composites, saw mill wood chips are dried to 2% or 3% moisture and ground down through a hammer mill to wood fibres (<5 mm). Wood flour may be incorporated as a filler giving bulk to the product. The prepared wood-fibre material and any filler are mixed with lubricants, UV stabilisers and pigments into the molten polymer, either in a batch or continuous process. The components are then formed by injection moulding for items such as architectural mouldings. Extrusion or pultrusion, a combination of extrusion and pulling, are both used for continuous sections such as window profiles and decking.

Wood plastic composites for outdoor products such as decking, fencing and garden furniture can be manufactured using a significant proportion of recycled polymer and scrap wood, which reduces the quantities of these materials in the waste stream.

Wood plastic composites are generally less stiff than wood and have lower strengths, but they are normally resistant to rot and insect attack. Fire resistance is similar to that of wood of the same density, but is improved by the incorporation of flame and smoke retardants during the manufacturing process. The materials are generally coloured grey or brown to reduce the effects of fading. Thermal movement is large and appropriate expansion joints are necessary.

**Corian**

Corian is a composite of natural minerals, pigments and acrylic polymer, which combine together to produce a highly durable and tough material, available in a wide range of colours. A typical composition is 33% acrylic polymer and 67% aluminium trihydrate. The proprietary product is frequently used for kitchen and
other countertops as it can be moulded into complex forms and inconspicuously joined into single units with the base plastic composite. Thicknesses range from 6 to 19 mm in sheets up to 930 × 3650 mm.

**LIGHT-CONDUCTING PLASTICS**

A range of light- and colour-sensitive materials has been developed by embedding a matrix of light-conducting plastic channels into a substrate of either concrete or acrylic polymer. Each of the light-conducting channels, which operate like fibre optics, gathers light or shadow from one end and transfers it to the other end, creating scintillation or darkening, respectively. Overall this creates an optical rippling effect as an object or light passes over the surface. The material, according to the substrate, can be used as floor tiles, walls, partitions, facades or tabletop surfaces. In each case the surface is seen to respond to object movement or changes in light intensity and colour. Units may be individual tiles or larger panels and are available in a range of standard or custom colours.

**VARIABLE-COLOUR PLASTIC FILMS**

Dichroic plastic films cause the observed colour and opacity to change depending on the viewpoint and also on the direction and intensity of the light source. For example, one dichroic sheet changes between green, gold and orange, and another between purple and blue. These films can be used to make the external envelope or internal environment of buildings appear active.

Thermochromic pigments in fibre-reinforced plastic sheet cladding change colour with the temperature, allowing the building to visually respond to outside temperature. The thermochromic pigment is incorporated into the core fibres and the gel coat to gain maximum effect.

**Recycling of plastics**

The increasing use of plastics in our everyday lives has led to a concern regarding large accumulation of wastes, which can only be resolved by extensive recycling. Many plastics are slow to degrade in landfill sites, and, as many are based on products from the petrochemical industry, this finite resource should not be wasted.
than landfill disposal. Currently, some domestic plastic bottle waste is recycled into loft insulation.

**RECYCLED PLASTIC LUMBER**

Mixed domestic plastic waste is cut up into small flakes, melted at 200°C into a grey viscous liquid and cast into moulds to produce structural components. The product, *polywood*, has been used to create a 7.5 m span lightweight bridge with a capacity of 30 tonnes in America. Recycled plastic was used to construct the I-beams and other structural sections. Recycled plastic lumber has the advantage over timber that it requires no maintenance or treatment with noxious chemicals and its use reduces the demand on landfill sites. Polywood is light, although more dense than timber; however, it suffers from creep. Also, it has greater thermal movement and lower stiffness (modulus of elasticity) than timber. Prior to its recent structural use, polywood has been used for decking, fencing, garden furniture and various marine applications. One UK product incorporating 90% thermoplastic recycled material is extruded into wood-coloured profiles for cladding, decking, windows, doors, conservatories, sheds and similar products normally associated with timber.

**BLENDED RECYCLED PLASTIC**

Crushed and ground plastic waste, when mixed with resin and reagents causing an exothermic reaction, produces a liquid plastic which can be poured and cast into moulds. The material sets quickly and cures after several hours to a strong, waterproof and durable product, which has the potential to be an alternative to both timber and concrete. The product is based largely on recycled domestic plastic which otherwise would have been sent to landfill sites.

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GLASS-FIBRE REINFORCED PLASTICS, CEMENT AND GYPSUM

Introduction

Composite materials, such as the glass-fibre reinforced materials glass-fibre reinforced polyester (GRP), glass-fibre reinforced cement (GRC) and glass-fibre reinforced gypsum (GRG), rely for their utility on the advantageous combination of the disparate physical properties associated with the individual component materials. This is possible when a strong bond between the glass fibres and the matrix material ensures that the two materials within the composite act in unison. Thus polyester, which alone has a very low modulus of elasticity, when reinforced with glass fibres produces a material which is rigid enough for use as a cladding material. Cement, which alone would be brittle, when reinforced with glass fibres can be manufactured into thin, impact-resistant sheets. Similarly, glass-fibre reinforcement in gypsum considerably increases its impact and fire resistance.

Glass fibres

The glass fibres for GRP and GRG are manufactured from standard E-glass as shown in Fig. 11.1. Molten glass runs from the furnace at 1200°C into a fore-hearth, and through a spinneret of fine holes from which it is drawn at high speed down to approximately 9 μm in thickness. The glass fibres are coated in size and bundled before winding up on a collet. Subsequently, the glass-fibre ‘cake’ is either used as continuous rovings or cut to 20–50 mm loose chopped strand. Glass-fibre rovings may be manufactured into woven mats; chopped strand mats are formed with organic binder.

Glass-fibre reinforced plastics

The standard matrix material for glass-fibre reinforced plastics is polyester resin, although other thermosetting resins including phenolic, epoxy and polyurethane may be used. Glass fibres as continuous rovings or chopped strand are used for most purposes; however, the highest-strength products are obtained with woven glass fabrics and unidirectionally aligned fibres. The proportion of glass fibres ranges widely from 20% to 80% by weight depending on the strength required. Enhanced performance can be achieved by using the more expensive S-grade high-strength and high-modulus glass fibres used mainly in the aerospace industry. Alternative higher tensile strength fibres include the polyaramids such as Kevlar and carbon fibres, but these are considerably more expensive than glass.

FABRICATION PROCESS

A major investment in the manufacture of GRP cladding panels lies within the production of the high-quality moulds. These are usually made from timber, but steel or GRP itself may also be used. Moulds are reused, sometimes with minor variations (e.g. the insertion of a window void within a wall unit), as many
times as possible to minimise production costs. The number of different mould designs for any one building is therefore kept to a minimum, and this may be reflected in the repetitiveness of the design.

In the fabrication process, the mould is coated with a release agent to prevent bonding and associated damage to the finished panel surface. A gel coat, which ultimately will be the weathering surface, is applied to a finished thickness of 0.25–0.4 mm. Early examples of GRP without sufficient gel coat have weathered to a rough surface with consequent exposure of the glass fibres; however, modern gel coats when applied to the correct thickness are durable. The subsequent fabrication involves the laying-up of layers of glass fibres and polyester resin to the required thickness, usually with either sprayed rovings or chopped strand mat. Reinforcement and fixings, normally in aluminium due to similarities in coefficients of expansion, may be incorporated and areas requiring additional strength can be thickened as appropriate by the laying-up process. Plastic foam insulation may be encapsulated to give the required thermal properties. Curing may take up to 2 weeks, after which the unit is stripped from the mould, trimmed around the edges and fitted out.

**PHYSICAL PROPERTIES AND DESIGN CONSIDERATIONS**

The choice of GRP, for example as a cladding panel, imparts its own aesthetic on a building design. The high strength-to-weight ratio of GRP allows for the use of large panel units, but cost constraints in the mould-making reduce the number of panel variations to a minimum. Curved edges to panels and openings are preferred to reduce stress-raising points at very sharp corners. The high thermal expansion coefficient of GRP demands careful detailing of movement joints and their appropriate sealing where necessary with components that retain their flexibility. In some cases the high expansion can be resolved by the use of profiled forms, which also impart strength. Colour fading
and yellowing of GRP panels have been problems; however, recent products with ultraviolet light protection are more colour-fast. Slightly textured finishes are generally more durable than smooth ones for exposure to full direct sunlight. GRP can be manufactured with fire-resistant additives; the phenolic resins have the advantage of lower flammability and smoke emissions. Long-term creep precludes the use of GRP as a significant load-bearing material, although single-storey structures, two-storey mobile units and structural overhead walkways are frequently constructed from the material. GRP is vandal resistant and can be laminated sufficiently to be bullet resistant. Where both surfaces are to be exposed, the material can be pressed between the two halves of a die.

USES OF GLASS-FIBRE REINFORCED PLASTICS

The lightweight properties of GRP make it an eminently suitable material for the manufacture of large cladding panels and custom-moulded structures as illustrated in the belfry and spire of St James, Piccadilly, London (Fig. 11.2). Finishes may be self-coloured or incorporate a natural stone aggregate finish. In addition, GRP is frequently used for the production of architectural features such as barge boards, dormer windows, classical columns and entrance canopies (Fig. 11.3). GRP may be pigmented to simulate various timbers, slate, Portland or Cotswold stone and lead or copper. It is also used to produce a wide range of small building components including baths, valley troughs, flat roof edge trim and water drainage systems. In addition, a wide range of composite cladding panels is manufactured from glass-fibre reinforced resins incorporating stone granules within the core of the material. These products, which are impact and fire resistant, are available with a granular stone, painted or gel-coat finish.

CARBON-FIBRE AND ARAMID COMPOSITES

Carbon fibres, which were originally developed for the aerospace industry, combine strength and stiffness with low weight, but have poor impact resistance. They are produced from polyacrylonitrile fibres by controlled oxidation at 250°C followed by carbonisation at 2600°C in an inert atmosphere. Three grades, high strength (HS), high modulus (HM) and intermediate modulus (IM) are produced. Carbon fibres, like glass fibres, are available as woven material, chopped strand or continuous filament. Carbon fibres have a small negative coefficient of expansion along the fibre axis; thus, composite materials of zero thermal expansion can be produced.

Aramids are aromatic polyamide liquid crystalline polymers, with high strength-to-weight ratio in tension, but poorer properties under compression or bending. Impact resistance is greater than that for carbon fibres. Aramid fibres, typically Kevlar, are produced by spinning the continuous fibre from solution. A variety of products is available with a range of modulus, elongation and impact resistance properties. Aramid composites exhibit good abrasion resistance. Carbon and aramid fibres may be combined in a composite material where strength, stiffness and impact resistance are all required.

Although polyester resins may be used as the matrix material for either carbon or aramid fibres, usually these more expensive fibres are incorporated into
higher-performance epoxy resins. In addition to the standard GRP laying-up production process, pultrusion and preimpregnation are used for manufacturing carbon-fibre-reinforced components. Pultrusion, a combination of extruding and pulling (Fig. 11.4), is used for making continuous profiles which may be either solid or hollow. Preimpregnation involves coating the continuous fibre or woven fibre fabric with a mixture of resin and curing agent, which can be stored frozen at this stage, then thawed and moulded into shape when required. By using low-temperature moulding processes large and complex structures can be fabricated for the construction industry.

For externally bonded reinforcement to concrete structures, pultruded carbon-fibre reinforced plates may be bonded to the concrete with a thixotropic epoxy resin. Alternatively, woven carbon fibre mat is wrapped around the concrete and pasted on with epoxy resin. The required level of reinforcement can be achieved by building up the appropriate thickness of epoxy resin saturated carbon fibre mat. The technique can also be applied to reinforcing steel, masonry, timber or cast iron. Although usually used for remedial work, this type of reinforcement could also be considered for new-build elements.

**RECYCLING FIBRE-REINFORCED POLYMERS**

According to 2004 data, the UK fibre-reinforced polymer (FRP) industry produces 240,000 tonnes per year, of which 11% is used within the construction industry. Currently, the majority of waste fibre-reinforced polymers are disposed into landfill sites. Whilst melting and remoulding is possible for thermoplastic FRPs, this is not applicable to thermoset GRP.

One solution for GRP is to grind the waste into powder and use this material in conjunction with other binders; however, this process is difficult where embedded metal fixings are incorporated into the original components. The recyclate powder can be blended with other recycled plastics to produce GRP/plastic lumber, which can be used for lightly loaded piles, decking, fencing and similar applications. This material can be cut and worked like the natural timber which it replaces. Alternatively, ground GRP can be incorporated into particleboard to make GRP-reinforced wood particleboard, which has enhanced mechanical properties compared to the standard grade (P5) of particleboard used for domestic flooring. However, when energy costs, transportation and other factors are considered, the ecological balance towards recycling fibre-reinforced polymers may be dependent on future considerations of recycling at the initial design stage.

Alternative potential processes for recycling GRP include thermal or chemical processes to recover the fibres, although the quality of the fibres is reduced.
in the process. Incineration with energy recovery is a viable option for GRP waste, but currently most incineration sites burn material to reduce volume rather than to generate electricity. GRP has a high calorific content compared to most domestic refuse, which on balance means that less domestic waste can be incinerated and therefore more sent to landfill.

**Glass-fibre reinforced cement**

Glass-fibre reinforced cement (GRC) is a material that was developed in the early 1970s by the Building Research Establishment. The standard material is produced from a mixture of alkali-resistant glass fibres with Portland cement, sand aggregate and water. Admixtures such as pozzolanas, superplasticisers and polymers are usually incorporated into the mix to give the required fabrication or casting properties. The breakthrough in the development of the material was the production of the alkali-resistant (AR) glass fibres, as the standard E-glass fibres, which are used in GRP and GRG, corrode rapidly in the highly alkaline environment of hydrated cement. Alkali-resistant glass, in addition to the sodium, silicon and calcium oxide components of standard E-glass, contains zirconium oxide. Alkali-resistant glass fibres, which have been improved by progressive development, are manufactured under the trade name Cem-FIL. The addition to GRC mixes of metakaolin, a pozzolanic material produced by calcining china clay at 750–800°C, prevents the development of lime crystals around the glass fibres. In the unblended GRC this leads to some gradual loss of strength. Standard grey GRC has the appearance of sheet cement and is non-combustible.

**MANUFACTURE OF ALKALI-RESISTANT GLASS FIBRES**

Silica, limestone and zircon are melted in a furnace; the alkali-resistant glass produced is drawn into fibres of 14 or 20 μm diameter and rolled into cakes for subsequent use as continuous rovings or for conversion into chopped strand. The process is comparable to that for standard E-glass fibres. Cem-FIL glass tissue with a fine texture is also available.

**CEMENT MATRIX**

Portland cement, strength class 42.5 or 42.5R (rapid early strength), is normally used. Portland cement will produce a grey finish, but white Portland cement or added pigments may be used to give different effects. However, with the use of pigments care must be taken to ensure uniformity of colour. Washed sand and fly ash (pulverised-fuel ash) are the usual aggregates, but crushed marble, limestone or granite can be used when a particular exposed aggregate finish is required.

**FABRICATION PROCESSES**

Fibre-reinforced cement components may be formed either by using a spray-gun, which mixes the glass fibres with a slurry of cement as it sprays directly into the mould, or by premixing a blend of cement, sand, water, admixtures and glass fibres before casting. Moulds similar to those required for the production of GRP components are used. Extrusion and injection moulding techniques are applicable for linear or small components, and bagged pre-blended mixes can be used for on-site applications.
Sprayed glass-fibre reinforced cement

Spray techniques, which may be manual (Fig. 11.5) or robotic, are used to build up the required thickness, usually between 10 and 20 mm. During spraying, the gun chops the fibres into 25–40 mm lengths, depositing a uniform felt of fibres and mortar into the mould. A typical sprayed mix would contain 5% glass fibres, 36% Portland cement, 36% washed sand, 11% additives/polymer and 12% water. The curing of GRC is relatively slow, with 95% strength developed after 7 days.

Premixed glass-fibre reinforced cement

It is normal to premix the cement, sand, water and admixtures, then add the chopped fibres. A typical mix would contain up to 3.5% of 12 mm fibres in a sand : cement mix of 0.5 : 1 with a water/cement ratio of 0.35. The mix is then cast and vibrated, or pressed into form for smaller components. For renderings, a glass-fibre content of between 1% and 2% is appropriate. A recent development involves the direct spraying of the premixed material.

Properties of Glass-fibre Reinforced Cement

Appearance

While standard GRC has the appearance of cement, a wide diversity of colours, textures and simulated materials can be manufactured. A gloss finish should be avoided, as it tends to craze and show any defects or variations. The use of specific aggregates followed by grinding can simulate marble, granite, terracotta, etc., while reconstructed stone with either a smooth or tooled effect can be produced by the action of acid etching. An exposed aggregate finish is achieved by the use of retardants within the mould, followed by washing and brushing. Applied finishes, which are usually water-based synthetic latex emulsions, can be applied to clean, dust-free surfaces.

Moisture and thermal movement

GRC exhibits an initial irreversible shrinkage followed by a reversible moisture movement of approximately 0.2%. The coefficient of thermal expansion is within the range \((7–20) \times 10^{-6}\) °C\(^{-1}\), typical for cementitious materials.

Thermal conductivity

The thermal conductivity of GRC is within the range 0.21–1.0 W/m K. Double-skin GRC cladding panel units usually incorporate expanded polystyrene, mineral wool or foamed plastic insulation. Cold bridging should be avoided where it may cause shadowing effects.

Durability

GRC is less permeable to moisture than normal concrete, so it has good resistance to chemical attack; however, unless manufactured from sulphate-resisting cement, it is attacked by soluble sulphates. GRC is unaffected by freeze/thaw cycling.

Impact resistance

GRC exhibits a high impact resistance but toughness and strength does decrease over long periods of time. However, the incorporation of metakaolin \((2SiO_2\cdotAl_2O_3)\) into the mix appears to improve the long-term performance of the material.
USES OF GLASS-FIBRE REINFORCED CEMENT

GRC is used extensively for the manufacture of cladding and soffit panels because it is lightweight and easily moulded (Fig. 11.6). It is used in conservation work as a replacement for natural stone and in architectural mouldings, including sophisticated decorative screens within countries of the Middle East. It is used as permanent formwork for concrete, fire-resistant partitioning and in the manufacture of small components including slates, tiles and decorative ridge tiles. Glass-fibre reinforced cement slates are manufactured to simulate the texture and colour of natural slate. Some manufacturers incorporate blends of other non-asbestos natural and synthetic fibres together with pigments and fillers to produce a range of coloured products with glossy, matt or simulated riven finishes.

Glass-fibre reinforced gypsum

Glass-fibre reinforced gypsum (GRG) combines the non-combustibility of gypsum plaster with the reinforcing strength of glass fibres. Products contain typically 5% of the standard E-glass fibres, which considerably improve impact as well as fire resistance. Commercial GRG products are available as standard panels, encasement systems for the fire protection of steel and decorative wall panels. As with all gypsum products, standard GRG should not be used in damp conditions or at temperatures regularly over 50°C.
Fig. 11.8 Decorative GRG plaster components

GLASS-FIBRE REINFORCED GYPSUM BOARDS

The standard boards, available in a range of thicknesses from 5 to 15 mm, are manufactured with a glass-fibre reinforced gypsum core and glass-fibre tissue immediately below the gypsum faces. The material is suitable for a wide range of applications including wall linings, ceilings and protected external positions such as roof soffits. The material can be easily cut on site and fixed with nails or screws; in addition, owing to the effect of the glass-fibre reinforcement, it can be curved to fit, for example, barrel-vault ceilings. The minimum radius of curvature depends on the board thickness. The material has a smooth off-white finish; joints should be taped before finishing board plaster is applied.

For the fire protection of steelwork, high-performance, Class 0 board thicknesses of 15, 20, 25 and 30 mm are available. Depending on the steel section factor (Hp/A m⁻¹), with double layers and staggered joints, up to 120 minutes’ fire resistance can
be achieved (Fig. 11.7). With specialist steel fixings to the structural steelwork, up to 180 minutes’ fire protection is achieved with glass-fibre reinforced gypsum boards.

Decorative glass-fibre reinforced gypsum boards and components

Decorative boards manufactured with a range of motifs can be used as dado or wall panels (Fig. 11.8). Dabs of sealant are used initially to fix panels to existing walls and allow adjustment to a flush finish. Panels may be painted after jointing.

Ceiling tiles manufactured from GRG are available in a wide range of designs, including plain, textured, patterned, open or closed-cell surface and with square, tapered or bevelled edges. The standard size is usually 600 × 600 mm, although some manufacturers produce a wide range of units. GRG has good fire-resistant properties; it is non-combustible to BS 476 Part 4: 1970, Class 1 surface spread of flame to BS 476 Part 6: 1989, up to Class 0 to Building Regulations Section E15 and Euroclass A2-s1, d0 (no significant contribution to the fire load, no evolution of smoke or flaming droplets). It does not emit noxious fumes in fire. Acoustic tiles with enhanced sound absorption and attenuation properties are normally part of the standard range, which may also include Imperial sizes for refurbishment work.

A wide range of small decorative components is manufactured in GRG, including cornices, coving, ceiling centres, corbels, columns, arches and niches.


STANDARDS

BS 476 Fire tests on building materials and structures:
  Part 7: 1997 Method of test to determine the classification of the surface spread of flame of products.
BS 4592 Industrial type flooring and stair treads:
BS 5544: 1978 Specification for anti-bandit glazing (glazing resistant to manual attack).

References

FURTHER READING

BS EN 1013-2: 1999  Light transmitting profiled sheet for single skin roofing (GRP).
BS EN 1170  Precast concrete products. Test method for GRC:
  Part 2: 1998  Measuring the fibre content in fresh GRC.
  Part 3: 1998  Measuring the fibre content in sprayed GRC.
  Part 8: 2008  Cyclic weathering type test.
BS EN 1447: 2009  Plastics piping systems. Glass reinforced thermoplastics (GRP) pipes.
BS EN 1796: 2006  Plastics piping systems for water supply. GRP based on unsaturated polyester resin (UP).
BS EN 13121-3: 2008  GRP tanks and vessels for use above ground. Design and workmanship.
BS EN 14845  Test methods for fibres in concrete:

BUILDING RESEARCH ESTABLISHMENT PUBLICATIONS

BRE Digests
BRE Digest 405: 2000  Polymer composites in construction.
BRE Digest 442: 1999  Architectural use of polymer composites.
BRE Digest 480: 2004  Wood plastic composites and plastic lumber.

BRE Good repair guide
BRE GRG 34: 2003  Repair and maintenance of FRP structures.

BRE Information papers
BRE IP 7/99  Advanced polymer composites in construction.
BRE IP 19/01  The performance of fibre cement slates.
BRE IP 10/03  Fibre reinforced polymers in construction: durability.
BRE IP 11/03  Fibre reinforced polymers in construction: predicting weathering.
BRE IP 2/04  Wood plastic composites: market drivers and opportunities in Europe.
BRE IP 4/04  Recycling fibre reinforced polymers in the construction industry.
BRE IP 5/04  Fibre reinforced polymers in construction.

BRE Reports

ADVISORY ORGANISATIONS
British Plastics Federation, 6 Bath Place, Rivington Street, London EC2A 3JE, UK (020 7457 5000).
Glassfibre Reinforced Concrete Association, Concrete Society, 4 Meadows Business Park, Station Approach, Camberley, Surrey GU17 9AB, UK (01276 607140).
Introduction

Plastering, based on lime, was brought to Britain by the Romans. In Britain, it was originally used to strengthen and seal surfaces and, in the case of combustible materials to afford some fire protection, but by the eighteenth century its value as a decorative finish had been appreciated. The use of gypsum plaster both as a sealant and as a decorative material by the Minoan civilisation is well documented, and current UK practice is now based on gypsum (hydrated calcium sulphate), rather than lime. Gypsum is mined from geological deposits produced by the gradual evaporation of lakes containing the mineral; there are extensive reserves within the UK, mainly in the North of England, but also in the East Midlands.

Historically, fibrous materials have been used to reinforce plaster and particularly to control shrinkage in lime plaster. Traditionally, ox, horse and goat hair were the standard materials; however, straw, hemp and jute have also been used. The earliest lightweight support for plasters was interwoven hazel twigs, but by the fifteenth century split timber laths were common. The modern equivalent is the use of galvanised and stainless steel expanded metal.

Gypsum plaster

MANUFACTURE OF GYPSUM PLASTER

Rock gypsum is mined, crushed and ground to a fine powder. The natural mineral may be white or discoloured pale pink, grey or brown due to small quantities of impurities, which do not otherwise affect the product. On heating to temperatures in the range 130–170°C, water is driven off the hydrated gypsum; the type of plaster produced is largely dependent on the extent of this dehydration process.

\[
\begin{align*}
\text{CaSO}_4 \cdot 2\text{H}_2\text{O} & \xrightarrow{130^\circ C} \text{CaSO}_4 \cdot \frac{1}{2}\text{H}_2\text{O} \\
\text{hydrated gypsum} & \quad \text{hemi-hydrate} \\
\text{170}^\circ C & \quad \text{anhydrous gypsum}
\end{align*}
\]

Gypsum plasters including those for special purposes are defined in the standard BS EN 13279-1: 2008. Gypsum plasterboards are defined in BS EN 520: 2004.

TYPES OF PLASTER

Plaster of Paris

Plaster of Paris is produced by driving off three quarters of the water content from natural hydrated gypsum. Plaster of Paris sets very quickly on the addition of water, and is therefore often used as a moulding material.

Retarded hemi-hydrate gypsum plaster

A majority of plasters in current use within construction are based on retarded hemi-hydrate gypsum. The addition of different quantities of a retarding agent, usually keratin, is used to adjust the setting time (usually between 1.5 and 2 hours) for different products. Other additives and admixtures include fillers, fibres, lime, lightweight aggregates, pigments, retarders and plasticisers.
Types of gypsum binders and plasters (BS EN 13279-1: 2008):

<table>
<thead>
<tr>
<th>Notation</th>
<th>Designation</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Gypsum binder for further processing</td>
</tr>
<tr>
<td>A1</td>
<td>Gypsum binders for direct use</td>
</tr>
<tr>
<td>A2</td>
<td>Gypsum binders for direct use on site</td>
</tr>
<tr>
<td>A3</td>
<td>Gypsum binders for further processing</td>
</tr>
<tr>
<td>B</td>
<td>Gypsum plaster</td>
</tr>
<tr>
<td>B1</td>
<td>Gypsum building plaster</td>
</tr>
<tr>
<td>B2</td>
<td>Gypsum-based building plaster (minimum 50% gypsum)</td>
</tr>
<tr>
<td>B3</td>
<td>Gypsum-lime building plaster (&gt;5% lime)</td>
</tr>
<tr>
<td>B4</td>
<td>Lightweight gypsum building plaster (inorganic or organic aggregates)</td>
</tr>
<tr>
<td>B5</td>
<td>Lightweight gypsum-based building plaster</td>
</tr>
<tr>
<td>B6</td>
<td>Lightweight gypsum-lime building plaster</td>
</tr>
<tr>
<td>B7</td>
<td>Enhanced surface hardness gypsum plaster</td>
</tr>
<tr>
<td>C</td>
<td>Gypsum plaster for special purposes</td>
</tr>
<tr>
<td>C1</td>
<td>Gypsum plaster for fibrous plasterwork</td>
</tr>
<tr>
<td>C2</td>
<td>Gypsum mortar</td>
</tr>
<tr>
<td>C3</td>
<td>Acoustic plaster</td>
</tr>
<tr>
<td>C4</td>
<td>Thermal insulation plaster</td>
</tr>
<tr>
<td>C5</td>
<td>Fire protection plaster</td>
</tr>
<tr>
<td>C6</td>
<td>Thin-coat plaster, finishing product</td>
</tr>
<tr>
<td>C7</td>
<td>Finishing product</td>
</tr>
</tbody>
</table>

Undercoat and one-coat plasters
The main constituents of undercoat and one-coat plasters are retarded hemi-hydrate gypsum, with lightweight aggregates for the lightweight products, together with small quantities of limestone, anhydrite (anhydrous gypsum), clay and sand. In addition, other materials are incorporated to adjust the product specification and setting time, which normally ranges between 1 and 2 hours. Thus, lime is added to undercoat plaster, and for backgrounds of high suction, a water retention agent is also required. For example, browning is suitable for use on backgrounds with moderate or high suction and a good mechanical key (e.g. brickwork and blockwork), but on smooth surfaces or low suction backgrounds such as concrete, an initial bonding coat is required. For higher impact resistance, cement and granulated blast furnace slag are incorporated, and for a one-coat plaster, limestone is added. Typical applications for walls would be 11 mm for undercoats with a finish coat of 2 mm, or a single one-coat application of 13 mm. The undercoat for ceilings would normally be 8 mm with a 2 mm finish coat. A maximum total plaster thickness of 25 mm is recommended.

Finish-coat plasters
For finish-coat plasters, like undercoat plasters, the main constituent is retarded hemi-hydrate gypsum, but with a small addition of lime to accelerate the set. The lightweight products contain exfoliated vermiculite. Finish coats on masonry substrates are usually 2 mm in thickness, and board finish plaster is normally applied to 2–3 mm.

Lightweight plasters
Lightweight plasters usually contain inorganic expanded perlite or exfoliated vermiculite, but organic lightweight aggregates may also be used. The range of densities and associated thermal conductivities is listed in Table 12.1.

<table>
<thead>
<tr>
<th>Density (kg/m³)</th>
<th>Thermal conductivity (W/m K)</th>
</tr>
</thead>
<tbody>
<tr>
<td>600</td>
<td>0.18</td>
</tr>
<tr>
<td>700</td>
<td>0.22</td>
</tr>
<tr>
<td>800</td>
<td>0.26</td>
</tr>
<tr>
<td>900</td>
<td>0.30</td>
</tr>
<tr>
<td>1000</td>
<td>0.34</td>
</tr>
<tr>
<td>1100</td>
<td>0.39</td>
</tr>
<tr>
<td>1200</td>
<td>0.43</td>
</tr>
<tr>
<td>1300</td>
<td>0.47</td>
</tr>
<tr>
<td>1400</td>
<td>0.51</td>
</tr>
<tr>
<td>1500</td>
<td>0.56</td>
</tr>
</tbody>
</table>

BACKGROUNDS FOR PLASTER
Plaster bonds to the background by a combination of mechanical key and adhesion. Backgrounds should be clean, dry and free from other contamination and the specification of the plaster should be appropriate to the suction of the background surface. Where possible, as in the case of brickwork, a good mechanical key should be obtained by raking out the joints. On hard, low-suction materials such as smooth concrete and ceramic tiles, a polyvinyl acetate (PVA) or proprietary bonding agent should be applied. Similarly, to control the high suction in substrates such as aerated concrete blocks, a bonding agent can be applied or the substrate wetted prior to the application of plaster. Plaster can, however, be applied directly to dense aggregate concrete blocks without prior wetting. Where two or more coats of plaster are applied, the undercoats should be scratched to ensure good subsequent bonding. Gypsum plasters, if applied correctly, do not shrink or crack on
drying out and subsequent coats can be applied in quick succession.

**PLASTERBOARD**

Plasterboard consists of a gypsum core bonded to strong paper liners. Most wallboards have one light surface for direct decoration or plaster skim and one grey surface. The decorative surface may be either tapered or squared. The standard board sizes are 1200 and 900 mm wide to coordinate with timber or metal stud partitioning systems. Plasterboard may be cut with a saw or scored and snapped. Nail fixings should be driven in straight, leaving a shallow depression but without fracturing the paper surface. Alternatively boards may be screwed. Standard thicknesses are 12.5, 15 and 19 mm, although 9.5 mm board can also be obtained.

Only the moisture-resistant grades of plasterboard (Type H) normally require the application of a bonding agent before plastering. These have a water-resistant core and treated liners, so may be used in moist and humid conditions such as kitchens or bathrooms and behind external finishes such as vertical tiling and weatherboarding, or in external sheltered positions protected from direct rain.

Fire-resistant boards (Type F), available in 12.5 and 15 mm thicknesses and reinforced with glass fibres, offer increased fire resistance over standard gypsum boards. Fire-resistant boards are colour-coded pink. Impact-resistant boards are also reinforced with glass fibres and have a high-strength paper liner. Glass-fibre reinforced gypsum is described in Chapter 11 Glass-fibre reinforced plastics, cement and gypsum.

Sound insulation boards, colour-coded blue, have a modified gypsum core, making them heavier than standard wallboards. The extra weight enhances the sound attenuation by up to 5 dB \( R_w \) compared to standard gypsum board. When used in conjunction with Robust Details, sound insulation boards provide enhanced acoustic performance. The Approved Document—Building Regulations Part E—Resistance to the Passage of Sound, requires a surface mass of 10 kg/m\(^2\) for plasterboard separating and internal walls also floors.

The heavy 19 mm gypsum planks, produced to a width of 600 mm, are used in walls, ceilings and floors to comply with the requirements of the Building Regulations when constructed according to Robust Details.

Boards are available finished with PVC, backed with aluminium foil or laminated to insulation (expanded polystyrene, extruded polystyrene, phenolic foam or mineral wool) for increased thermal properties. (The thermal conductivity of standard plasterboard is 0.19 W/m K.)

**Types of plasterboards (BS EN 520: 2004):**

<table>
<thead>
<tr>
<th>Type</th>
<th>Designation</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Gypsum plasterboard with a face suitable for a gypsum finish coat or decoration</td>
</tr>
<tr>
<td>H</td>
<td>Gypsum plasterboard with a reduced water absorption rate</td>
</tr>
<tr>
<td>E</td>
<td>Gypsum sheathing board for external walls but not permanently exposed to weather conditions</td>
</tr>
<tr>
<td>F</td>
<td>Gypsum plasterboard with enhanced core cohesion at high temperatures</td>
</tr>
<tr>
<td>P</td>
<td>Gypsum baseboard to receive gypsum plaster</td>
</tr>
<tr>
<td>D</td>
<td>Gypsum plasterboard with a controlled density</td>
</tr>
<tr>
<td>R</td>
<td>Gypsum plasterboard with enhanced strength</td>
</tr>
<tr>
<td>I</td>
<td>Gypsum plasterboard with enhanced surface hardness</td>
</tr>
</tbody>
</table>

Note:
Some boards may combine more than one designation.

**Plasterboard systems**

Plasterboard non-load-bearing internal walls may be constructed using proprietary metal stud systems or as traditional timber stud walls. Where appropriate, acoustic insulation should be inserted within the void spaces. Dry lining to masonry may be fixed with dabs of adhesive, alternatively with metal or timber framing. Plasterboard suspended ceiling systems are usually supported on a lightweight steel framework fixed directly to either concrete or timber. Convex and concave surfaces can be achieved. Sound transmission through existing upper-storey timber-joist floors can be reduced by a combination of resiliently mounted plasterboard and mineral wool insulation. Compliance with the acoustic requirements of the Building Regulations Approved Document E for domestic building can be achieved using Robust Details. Components for plasterboard metal-framing systems are defined in BS EN 14195: 2005.

**Plasterboard ceiling tiles**

Plasterboard ceiling tiles are available in a range of smooth, textured and perforated finishes to produce various levels of sound-insulating and sound-absorbing properties. The standard tiles are 600 \( \times \) 600 mm, for fixing to metal sub-framing. Fire classification is Class 0 and Euroclass A2-s1, d0.
Fibre-reinforced gypsum boards

Fibre-reinforced gypsum boards are manufactured with either natural or glass fibres. Glass-fibre reinforced gypsum (GRG) is described in Chapter 11.

Natural fibre-reinforced gypsum boards are manufactured from cellulose fibres, frequently from recycled paper, or wood fibres within a matrix of gypsum. The boards may be uniform with dispersed fibres or laminated with woven or non-woven gypsum-reinforced sheets encasing a perlite and gypsum core. Boards are impact and fire-resistant and easily fixed by nails, screws, staples or adhesive as a dry-lining system to timber, metal framing or masonry. Standard boards are 1200 × 2400 mm with thicknesses normally in the range 12.5–18 mm. Joints are filled or taped and corners beaded as for standard plasterboard products. A composite board of fibre-reinforced gypsum and expanded polystyrene offers enhanced insulation properties. Gypsum boards with fibrous reinforcement are defined in BS EN 15283: 2008. (The thermal conductivity of gypsum board containing 13% wood fibres is 0.24 W/m K.)

Types of fibre-reinforced plasterboard (BS EN 15283: 2008):

<table>
<thead>
<tr>
<th>Type</th>
<th>Designation</th>
</tr>
</thead>
<tbody>
<tr>
<td>GM</td>
<td>Gypsum board with mat reinforcement</td>
</tr>
<tr>
<td>GM-H1, GM-H2</td>
<td>Gypsum board with mat reinforcement and reduced water absorption rate</td>
</tr>
<tr>
<td>GM-I</td>
<td>Gypsum board with mat reinforcement and enhanced surface hardness</td>
</tr>
<tr>
<td>GM-R</td>
<td>Gypsum board with mat reinforcement and enhanced strength</td>
</tr>
<tr>
<td>GM-F</td>
<td>Gypsum board with mat reinforcement and enhanced core cohesion at high temperatures</td>
</tr>
<tr>
<td>GF</td>
<td>Gypsum fibre board</td>
</tr>
<tr>
<td>G-H</td>
<td>Gypsum fibre board with reduced water absorption rate</td>
</tr>
<tr>
<td>GF-W1, GF-W2</td>
<td>Gypsum fibre board with reduced surface water absorption rate</td>
</tr>
<tr>
<td>GF-D</td>
<td>Gypsum fibre board with enhanced density</td>
</tr>
<tr>
<td>GF-I</td>
<td>Gypsum fibre board with enhanced surface hardness</td>
</tr>
<tr>
<td>GF-R1, GF-R2</td>
<td>Gypsum fibre board with enhanced strength</td>
</tr>
</tbody>
</table>

SPECIAL PLASTERS

Renovating plaster

Renovating plaster is used where walls have been stripped of existing plaster during the successful installation of a new damp-proof course. Renovating plasters contain aggregates which promote surface drying when they are applied to structures with residual moisture, but they should not be used in permanently damp locations below ground level. Renovating plaster should also not be used where masonry is heavily contaminated with salts, such as in buildings not originally built with damp-proof courses, and on the brickwork of chimney breasts. Renovating plasters contain a fungicide to inhibit mould growth during the drying-out process.

Projection plaster

Projection plaster is sprayed onto the background from a projection machine as a continuous ribbon which flows sufficiently for the ribbons to coalesce. The plaster should be built up to the required thickness, ruled to an even surface then flattened and trowelled to a flat surface. As with all plastering the process should not be carried out under freezing, excessively hot or dry conditions. A typical application to masonry would be 13 mm and should not exceed 25 mm.

Acoustic plaster

Acoustic plaster has a higher level of sound absorption than standard gypsum plaster owing to its porosity and surface texture. Aluminium powder is added to the wet plaster mix to produce fine bubbles of hydrogen gas, which remain trapped as the plaster sets giving it a honeycomb structure. One form of acoustic plasterboard consists of perforated gypsum plasterboard, which may be backed with a 100 mm glass wool sound-absorbing felt.

X-ray plaster

X-ray plaster is retarded hemi-hydrate plaster containing barium sulphate (barytes) aggregate. It is used as an undercoat plaster in hospitals, etc., where protection from X-rays is required. Typically, a 20 mm layer of X-ray plaster affords the same level of protection as a 2 mm sheet of lead, provided that it is free of cracks.

Textured plaster

Textured plaster is frequently applied to plasterboard ceilings. A variety of different patterns and textures
can be achieved. The textured surface may be left as a natural white finish or painted as required.

**Fibrous plaster**

Fibrous plaster is plaster of Paris reinforced with jute, sisal, hessian, glass fibres, wire mesh or wood laths. It is used for casting in moulds, ornate plasterwork, such as fire surrounds, decorative cornices, dados, friezes, panel mouldings, corbels and centrepieces for ceilings in both restoration and new work. The reinforcement material may be elementary in the form of random fibres or sheet material, or complementary as softwood laths or lightweight steel sections. Fibrous plaster is described in the standards BS EN 13815: 2006 and BS EN 15319: 2007.

**Phase change material plaster**

An alternative method of stabilising internal room temperatures, rather than using thermal mass, is to incorporate phase change materials (PCMs) into the building fabric. One approach is to use a layer of gypsum plaster containing 26–30% by volume of phase change material. A commercial system uses wax encapsulated within 3 µm particles of PVA, which are formed as a dispersion, and then dried to a 0.1–0.3 mm powder for mixing into gypsum. The wax undergoes a phase change at 23°C or 26°C, absorbing heat as it melts and releasing it as it solidifies. A 30 mm layer of phase change gypsum plaster has an equivalent thermal stabilising effect to a mass of 180 mm of concrete or 230 mm of brickwork, and a 15 mm PCM gypsum board is equivalent to 100 mm of concrete. The boards may be used for ceilings or wall linings as appropriate, and the phase change temperature should be specified. Current costs are of the order of 10 times that of standard plasterboard.

**ACCESSORIES FOR PLASTERING**

**Beads**

Angle and stop beads are manufactured from galvanised or stainless perforated steel strip or expanded metal. They provide a protected, true straight arris or edge with traditional plastering to masonry or for thin-coat plasterboard. Proprietary systems are manufactured similarly from perforated galvanised or stainless steel to form movement joints in dry lining systems (Fig. 12.1). These components are defined in BS EN 14353: 2007.

**Scrim and jointing tape**

Scrim, an open-weave material, is used across joints between plasterboards and in junctions between plaster and plasterboard. Both self-adhesive glass-fibre mesh and traditional jute scrim are available. For the prevention of thermal movement cracking at plasterboard butt joints, 50 mm paper tape bedded into the plaster skim is often more effective than self-adhesive scrim. The paper-based tape and bedding compounds for plasterboards are defined in BS EN 14353: 2007.

**Coves and cornices**

Decorative coves and cornices are manufactured from gypsum plaster encased in a paper liner. In some cases the gypsum is reinforced with glass fibres. The components (Fig. 12.2) can be cut to size with a saw, and are normally fixed with proprietary adhesives. The standard sizes of cove are 100 and 127 mm.

**GYPSUM FLOOR SCREED**

Gypsum interior floor screed, manufactured from a mixture of hemi-hydrate gypsum, limestone and less than 2% cement, may be used as an alternative to a traditional sand and cement screed, provided that a floor covering is to be used. The material is self-smoothing and may be pumped. It is laid on a polythene membrane to a minimum thickness of 35 mm for floating screeds, and may be used over under-floor heating systems. When set, the hard plaster has a minimum 28 day compressive strength of 30 MPa.

**FIRE RESISTANCE OF PLASTER MATERIALS**

Gypsum products afford good fire protection within buildings due to their basic chemical composition. Gypsum, hydrated calcium sulphate (CaSO$_4$·2H$_2$O) as present in plaster and plasterboard, contains nearly 21% water of crystallisation. When exposed to a fire this chemically combined water is gradually expelled in the form of vapour. It is this process which absorbs the incident heat energy from the fire, considerably reducing the transmission of heat through the plaster, thus protecting the underlying materials. The process of dehydrating the gyspsum commences on the face adjacent to the fire, and immediately the dehydrated material, because it adheres to the unaffected gyspsum, acts as an insulating layer slowing down further dehydration. Even when all the water of crystallisation has been expelled, the remaining anhydrous gyspsum
continues to act as an insulating layer whilst it retains its integrity. The inclusion of glass fibres into gypsum plasterboards increases the cohesiveness of the material within fires. Gypsum binders and gypsum plasters are classified as Class A1 (no contribution to fire) if they contain less than 1% of organic material.

**RECYCLING GYPSUM PRODUCTS**

A large proportion of gypsum for plasterboard (over 1 million tonnes per annum) is produced as a by-product of the removal of sulphur dioxide from the flue gases of coal-fired electricity-generating stations. A slurry of pulverised limestone absorbs the sulphur dioxide, producing high-purity calcium sulphate. However, the supply of this synthetic gypsum (known as desulphogypsum, DSG) is dependent on the continuing generation of electricity by coal-fired stations also the sulphur content of the coal used. Low-sulphur coal imported from Australia may reduce the production of desulphogypsum.

Waste of plasterboard on site is high, ranging from 10% to 25% of the delivered material. Disposal in
landfill sites is very limited, as gypsum, in association with biodegradable materials, can produce toxic hydrogen sulphide. Gypsum products must therefore be deposited in biodegradable-free locations. Currently, the industry maximum recycled gypsum content for plasterboard is 18%. The paper liners for plasterboard are made from 97% recycled paper and cardboard. One manufacturer will collect all of its own surplus gypsum board, ceiling tiles, coving and GRG from site for recycling into new products.

The publication PAS 109: 2008 gives the specification for the production of recycled gypsum from waste plasterboard.

Lime plaster

Hydraulic lime plaster is suitable for interior application, particularly on earth structures and unfired clay walls. It is usually applied in two or three coats – the best quality work requiring the three-coat system. In this case a 13 mm coat of coarse stuff containing 5 mm sand (lime : sand, 1 : 2.5) is followed when dry, with a similar thickness of a 1: 3 mix and a thin final coat of between 1 : 1 and 1 : 2 lime to sand. Other additions, including horsehair and cow dung, may be added to improve the setting properties of the lime plaster.

Calcium silicate boards

Calcium silicate boards are manufactured from silica with lime and/or cement, usually incorporating cellulose fibres or softwood pulp and mica or exfoliated vermiculite filler, to produce a range of densities. The high-density material is laminated under steam and pressure, whilst the lower-density material is produced by rolling followed by curing in an autoclave. Calcium silicate boards, like gypsum boards are non-combustible. The material is grey or off-white in colour, easily worked and nailed. Calcium silicate boards are durable, moisture-, chemical- and impact-resistant with dimensional stability and a good strength-to-weight ratio. They are available with a range of smooth or textured factory finishes for interior or exterior use and also laminated to extruded polystyrene for enhanced insulation properties. Standard thicknesses include 4.5, 6.0, 9.0 and 12.0 mm, although thicknesses up to 60 mm are available in the vermiculite lightweight boards used for fire protection, giving up to 240 minutes’ resistance. Typical applications include wall, roof and partition linings, suspended ceilings, fascias, soffits, weatherboarding and fire protection to structural steelwork. External cladding boards may be finished with a sprayed or trowelled render to produce a seamless finish. (The thermal conductivities of calcium silicate boards are usually within the range 0.13–0.29 W/m K depending on their composition.)

References

FURTHER READING


STANDARDS

BS 476 Fire tests on building materials and structures:
Part 7: 1997 Method of test to determine the classification of the surface spread of flame of products.

BS 5270 Bonding agents for use with gypsum plaster and cement:

BS 6100 Building and civil engineering. Vocabulary:

BS 7364: 1990 Galvanized steel studs and channels for stud and sheet partitions and linings using screw fixed gypsum wallboards.

BS 8000 Workmanship on building sites:


BS EN 520: 2004 Gypsum plasterboards. Definitions, requirements and test methods.

BS EN 998 Specification for mortar for masonry:

BS EN 12859: 2008 Gypsum blocks. Definitions, requirements and test methods.


BS EN 13055-1: 2002 Lightweight aggregates for concrete, mortar and grout.

BS EN 13139: 2002 Aggregates for mortar.

BS EN 13279 Gypsum binders and gypsum plasters:
  Part 1: 2008 Definitions and requirements.
  Part 2: 2004 Test methods.

BS EN 13501 Fire classification of construction products and building elements:
  Part 1: 2007 Classification using data from reaction to fire tests.
  Part 2: 2007 Classification using data from fire resistance tests.

BS EN 13658 Metal laths and beads. Definitions, requirements and test methods:

BS EN 13815: 2006 Fibrous gypsum plaster casts. Definitions, requirements and test methods.

BS EN 13914 Design, preparation and application of external rendering and internal plastering:
  Part 2: 2005 Design considerations and essential principles for internal plastering.

BS EN 13915: 2007 Prefabricated gypsum plasterboard panels with a cellular paperboard core.

BS EN 13950: 2005 Gypsum plasterboard thermal/acoustic insulation composite panels. Definitions, requirements and test methods.


BS EN 13964: 2004 Suspended ceilings. Requirements and test methods.

BS EN 14190: 2005 Gypsum, plasterboard products from reprocessing. Definitions, requirements and test methods.

BS EN 14195: 2005 Metal framing components for gypsum plasterboard systems. Definitions, requirements and test methods.


BS EN 14246: 2006 Gypsum elements for suspended ceilings. Definitions, requirements and test methods.

BS EN 14353: 2007 Metal beads and feature profiles for use with gypsum plasterboards. Definitions, requirements and test methods.

BS EN 14496: 2005 Gypsum-based adhesives for thermal/acoustic insulation composite panels and plasterboards.

BS EN 14566: 2008 Mechanical fasteners for gypsum plasterboard systems. Definitions, requirements and test methods.


BS EN 15254 Extended application of results from fire-resistance tests:

BS EN 15283 Gypsum boards with fibrous reinforcement:

BS EN 15303 Design and application of plasterboard systems on frames:
BS EN 15318: 2007  Design and application of gypsum blocks.
BS EN 15319:2007  General principles of design of fibrous (gypsum) plasterworks.

BUILDING RESEARCH ESTABLISHMENT PUBLICATIONS

BRE Good building guides
BRE GBG 65: 2005  Plastering and internal rendering.
BRE GBG 70: 2007  Plasterboard; types and their applications (Parts 1, 2 and 3).

BRE Good repair guide

ADVISORY ORGANISATION

Gypsum Products Development Association, PO Box 35084, London NW1 4XE, UK (020 7935 8532).
INSULATION MATERIALS

Introduction

With the increasing emphasis on energy-conscious design and the broader environmental impact of buildings, greater attention is necessarily being focused on the appropriate use of thermal and sound insulation materials, for both new build and refurbishment work. Currently, 27% of UK carbon emissions are from domestic buildings and 17% from non-domestic buildings. The UK government has set the target that new homes will be net zero carbon from 2016. To achieve this target, the energy efficiency of new homes will need to increase by 25% in 2010 and by 44% in 2013 relative to the 2006 standards. The equivalent target date for net zero carbon for new non-domestic buildings is 2019. The energy conservation criteria within the Approved Document of the UK Building Regulations Part L—Conservation of fuel and power, are outlined in Chapter 7 Glass (page 246).

Thermal and sound insulation materials

To consider the relative efficiency of insulating materials, the thermal conductivities (λ, W/m K) are quoted at the standard 10°C to allow direct comparisons. U-values would not illustrate direct comparability owing to the varying thicknesses used and the wide variety of combinations of materials typically used in construction.

In considering acoustic control, distinction is made between the reduction of sound transmitted directly through the building components and the attenuation of sound reflected by the surfaces within a particular enclosure. Furthermore, transmitted sound is considered in terms of both impact and airborne sound. Impact sound is caused by direct impact onto the building fabric which then vibrates, transmitting the sound through the structure; it is particularly significant in the case of intermediate floors. Airborne sound waves, from the human voice and sound-generating equipment, cause the building fabric to vibrate, thus transmitting the sound. Airborne sound is particularly critical in relation to separating walls and is significantly increased by leakage at discontinuities within the building fabric, particularly around unsealed openings. Flanking sound is transmitted between rooms via the adjoining elements rather than directly through the separating wall. The reduction in sound energy passing through a building element is expressed in decibels (dB). The doubling of the mass of a building component reduces the sound transmission by approximately 5 dB; thus, sound-insulating materials are generally heavy structural elements. However, the judicious use of dissipative absorbers within walls can reduce the reliance for sound absorption on mass alone. Noise may be transmitted through service installations, so consideration should be given to the use of acoustic sleeves and linings as appropriate.

For dwellings, the Approved Document of the UK Building Regulations Part E (2003)—‘Resistance to the Passage of Sound’ requires protection against sound from adjacent buildings and within buildings including internal walls and floors. The use of Robust Details in housing can ensure adherence to the regulations and eliminate the requirement for pre-completion sound testing to demonstrate compliance. The Building Regulations list minimum criteria for the attenuation of airborne and impact sound.
Table 13.1  Typical sound absorption coefficients at 125, 500 and 2000 Hz for various building materials

<table>
<thead>
<tr>
<th>Material</th>
<th>Absorption coefficient</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>125 Hz</td>
</tr>
<tr>
<td>Concrete</td>
<td>0.02</td>
</tr>
<tr>
<td>Brickwork</td>
<td>0.05</td>
</tr>
<tr>
<td>Plastered solid wall</td>
<td>0.03</td>
</tr>
<tr>
<td>Glass 6 mm</td>
<td>0.1</td>
</tr>
<tr>
<td>Timber boarding, 19 mm over air space against solid backing</td>
<td>0.3</td>
</tr>
<tr>
<td>Wood wool slabs, 25 mm, on solid backing, unplastered</td>
<td>0.1</td>
</tr>
<tr>
<td>Fibreboard, 12 mm on solid backing</td>
<td>0.05</td>
</tr>
<tr>
<td>Fibreboard, 12 mm over 25mm air space</td>
<td>0.3</td>
</tr>
<tr>
<td>Mineral wool, 25 mm with 5% perforated hardboard over</td>
<td>0.1</td>
</tr>
<tr>
<td>Expanded polystyrene board, 25 mm over 50 mm airspace</td>
<td>0.1</td>
</tr>
<tr>
<td>Flexible polyurethane foam, 50 mm on solid backing</td>
<td>0.25</td>
</tr>
</tbody>
</table>

The absorption of sound at surfaces is related to the porosity of the material. Generally, light materials with fibrous or open surfaces are good absorbers, reducing ambient noise levels and reverberation times, whereas smooth hard surfaces are highly reflective to sound (Table 13.1). Sound absorption is measured on a 0–1 scale, with 1 representing total absorption of the sound.

**FORMS OF INSULATION MATERIALS**

Thermal and sound insulation materials may be categorised variously according to their appropriate uses in construction, their physical forms or their material origin. Many insulating materials are available in different physical forms, each with their appropriate use in building. Broadly, the key forms of materials could be divided into:

- structural insulation materials;
- rigid and semi-rigid sheets and slabs;
- loose fill, blanket materials and applied finishes;
- aluminium foil.

However, within this grouping, it is clear that certain materials spread over two or three categories. Insulation materials are therefore categorised according to their composition, with descriptions of their various forms, typical uses in construction and, where appropriate, fire protection properties. Materials are initially divided into those of inorganic and organic origin, respectively.

The broad range of non-combustible insulating materials are manufactured from ceramics and inorganic minerals including natural rock, glass, calcium silicate and cements. Some organic products are manufactured from natural cork or wood fibres but materials developed by the plastics industry predominate. In some cases these organic materials offer the higher thermal insulation properties but many either are inflammable or decompose within fire. Cellular plastics include open- and closed-cell materials. Generally, the closed-cell products are more rigid and have better thermal insulation properties and resistance to moisture, whereas the open-cell materials are more flexible and permeable. Aluminium foil is considered as a particular case as its thermal insulation properties relate to the transmission of radiant rather than conducted heat. Typical thermal conductivity values are indicated in Table 13.2.

**Inorganic insulation materials**

**FOAMED CONCRETE**

The manufacture of foamed concrete is described in Chapter 3.

Foamed concrete with an air content in the range 30–80% is a fire- and frost-resistant material. Foamed concrete can be easily placed without the need for compaction but it does exhibit a higher drying shrinkage than dense concrete. It is suitable for insulating under floors and on flat roofs where it may be laid to a fall of up to 1 in 100. (Thermal conductivity ranges from 0.10 W/m K at a density of 400 kg/m³ to 0.63 W/m K at a density of 1600 kg/m³.)

**LIGHTWEIGHT AGGREGATE CONCRETE**

Lightweight concrete blocks and in situ concrete are discussed in Chapters 2 and 3, respectively. Lightweight
Table 13.2  Typical thermal conductivity values for various building materials

<table>
<thead>
<tr>
<th>Material</th>
<th>Thermal conductivity (W/m K)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aerogel</td>
<td>0.018</td>
</tr>
<tr>
<td>Phenolic foam</td>
<td>0.018–0.031</td>
</tr>
<tr>
<td>Polyurethane foam (rigid)</td>
<td>0.019–0.025</td>
</tr>
<tr>
<td>Foil-faced foam</td>
<td>0.020</td>
</tr>
<tr>
<td>Polyisocyanurate foam</td>
<td>0.023–0.025</td>
</tr>
<tr>
<td>Extruded polystyrene</td>
<td>0.025–0.027</td>
</tr>
<tr>
<td>Expanded PVC</td>
<td>0.030</td>
</tr>
<tr>
<td>Mineral wool</td>
<td>0.031–0.040</td>
</tr>
<tr>
<td>Glass wool</td>
<td>0.031–0.040</td>
</tr>
<tr>
<td>Expanded polystyrene</td>
<td>0.033–0.040</td>
</tr>
<tr>
<td>Cellulose (recycled paper)</td>
<td>0.035–0.040</td>
</tr>
<tr>
<td>Flax</td>
<td>0.037</td>
</tr>
<tr>
<td>Sheep’s wool</td>
<td>0.037–0.039</td>
</tr>
<tr>
<td>Rigid foamed glass</td>
<td>0.037–0.048</td>
</tr>
<tr>
<td>Urea-formaldehyde foam</td>
<td>0.038</td>
</tr>
<tr>
<td>Hemp wool</td>
<td>0.040</td>
</tr>
<tr>
<td>Corkboard</td>
<td>0.042</td>
</tr>
<tr>
<td>Coconut fibre boards</td>
<td>0.045</td>
</tr>
<tr>
<td>Perlite board</td>
<td>0.045–0.050</td>
</tr>
<tr>
<td>Fibre insulation board</td>
<td>0.050</td>
</tr>
<tr>
<td>Straw bales</td>
<td>0.050</td>
</tr>
<tr>
<td>Exfoliated vermiculite</td>
<td>0.062</td>
</tr>
<tr>
<td>Thatch</td>
<td>0.072</td>
</tr>
<tr>
<td>Wood wool slabs</td>
<td>0.077</td>
</tr>
<tr>
<td>Medium-density fibreboard (MDF)</td>
<td>0.10</td>
</tr>
<tr>
<td>Foamed concrete (low density)</td>
<td>0.10</td>
</tr>
<tr>
<td>Lightweight to dense concrete</td>
<td>0.10–1.7</td>
</tr>
<tr>
<td>Compressed straw slabs</td>
<td>0.10</td>
</tr>
<tr>
<td>Softwood</td>
<td>0.13</td>
</tr>
<tr>
<td>Oriented strand board (OSB)</td>
<td>0.13</td>
</tr>
<tr>
<td>Hardboard</td>
<td>0.13</td>
</tr>
<tr>
<td>Particleboard/plywood</td>
<td>0.14</td>
</tr>
<tr>
<td>Gypsum plasterboard</td>
<td>0.19</td>
</tr>
<tr>
<td>Bituminous roofing sheet</td>
<td>0.19</td>
</tr>
<tr>
<td>Cement bonded particleboard</td>
<td>0.23</td>
</tr>
<tr>
<td>Unfired clay blocks</td>
<td>0.24</td>
</tr>
<tr>
<td>Calcium silicate boards</td>
<td>0.29</td>
</tr>
<tr>
<td>GRC—lightweight</td>
<td>0.21–0.5</td>
</tr>
<tr>
<td>GRC—standard density</td>
<td>0.5–1.0</td>
</tr>
<tr>
<td>Mastic asphalt</td>
<td>0.5</td>
</tr>
<tr>
<td>Calcium silicate brickwork</td>
<td>0.67–1.24</td>
</tr>
<tr>
<td>Clay brickwork</td>
<td>0.65–1.95</td>
</tr>
<tr>
<td>Glass—sheet</td>
<td>1.05</td>
</tr>
</tbody>
</table>

Notes:
Individual manufacturers’ products may differ from these typical figures.

Concrete materials offer a range of insulating and load-bearing properties, starting from 0.10 W/m K at a crushing strength of 2.8 MPa. Resistance to airborne sound in masonry walls is closely related to the mass of the wall. However, any unfilled mortar joints which create air paths will allow significant leakage of sound. In cavity walls, again mass is significant, but additionally, to reduce sound transmissions the two leaves should be physically isolated, with the exception of any necessary wall ties, to comply with the Building Regulations.

GYPSUM PLASTER

Plasterboard thermal linings will increase the thermal response in infrequently heated accommodation; the effect can be enhanced with metallised polyester-backed boards, which reduce radiant, as well as transmitted, heat loss. The addition of such linings for either new or upgrading existing buildings reduces the risk of thermal bridging at lintels, etc. (The thermal conductivity of gypsum plaster is typically 0.16 W/m K.)

Sound transmission through lightweight walls can be reduced by the use of two layers of differing thicknesses of gypsum plasterboard (e.g. 12.5 and 19 mm) as these resonate at different frequencies. The addition of an extra layer of plasterboard attached to existing ceilings with resilient fixings can reduce sound transmission from upper floors particularly if an acoustic quilt can also be incorporated.

WOOD WOOL SLABS

Wood wool (WW) slabs manufactured from wood fibres and cement (Chapter 4) are both fire and rot resistant. Wood wool products are described in BS EN 13168: 2008. With their combined load-bearing and insulating properties, wood wool slabs are suitable as a roof decking material, which may be exposed, painted or plastered to the exposed lower face. Wood wool slabs offer good sound absorption properties due to their open-textured surface, and this is largely unaffected by the application of sprayed emulsion paint. Acoustic insulation for a pre-screeded 50 mm slab is typically 30 dB. (The thermal conductivity of wood wool is typically 0.077 W/m K.)

MINERAL WOOL

Mineral wool (MW) is manufactured from volcanic rock (predominantly silica, with alumina and
magnesium oxide), which is blended with coke and limestone and fused at 1500°C in a furnace. The melt runs onto a series of rotating wheels which spin the droplets into fibres; they are then coated with resin binder and water-repellent mineral oil. The fibres fall onto a conveyor belt, where the loose mat is compressed to the required thickness and density, then passed into an oven where the binder is cured; finally, the product is cut into rolls or slabs. New binders are bio-polymers rather than phenolic resins, thereby reducing the embodied energy. Mineral wool is non-combustible, water-repellent and rot proof, and contains no CFCs or HCFCs.

Mineral wool is available in a range of forms depending on its degree of compression during manufacture and its required use:

- loose for blown cavity insulation;
- mats for insulating lofts, lightweight structures and within timber framed construction;
- batts (slabs) for complete cavity fill of new masonry;
- semi-rigid slabs for partial cavity fill of new masonry;
- rigid slabs for warm pitched roof and flat roof insulation;
- rigid resin-bonded slabs for floor insulation;
- weather-resistant boards for inverted roofing systems;
- dense pre-painted boards for exterior cladding;
- ceiling tiles.

The mats and board materials may be faced with aluminium foil to enhance their thermal properties. Roof slabs may be factory cut to falls or bitumen faced for torch-on bitumen membrane-roofing systems. Floor units are coated with paper when they are to be directly screeded. A resilient floor can be constructed with floor units manufactured from mineral wool slabs, with the fibres orientated vertically rather than horizontally, bonded directly to tongued and grooved flooring-grade chipboard.

The thermal conductivity of mineral wool products for internal use ranges typically between 0.031 and 0.039 W/m K at 10°C, although products for external use have higher conductivities.

Mineral wool can be used effectively to attenuate transmitted sound. In lightweight construction, acoustic absorbent quilts are effective in reducing transmitted sound through separating walls when combined with double plasterboard surfaces and a wide airspace, as well as in traditional timber joist floors when combined with a resilient layer between joists and floor finish. Pelletised mineral wool can be used for pugging between floor joists to reduce sound transmission, and is particularly appropriate for upgrading acoustic insulation during refurbishment.

Mineral wool, due to its non-combustibility, is used for the manufacture of fire stops to prevent fire spread through voids and cavities, giving fire resistance ratings between 30 and 120 minutes. Mineral wool slabs give typically between 60 minutes and 4 hours fire protection to steel. Similar levels of protection can be achieved with sprayed-on mineral wool, which may then be coated with a decorative finish.

Ceiling tiles for suspended ceilings manufactured from mineral wool typically provide Class 1 Spread of Flame to BS 476 Part 7 (1997) and Class 0 to Part 6 (1989) on both their decorative and back surfaces. The thermal conductivity of mineral wool suspended ceiling tiles is typically within the range 0.052–0.057 W/m K. Sound attenuation of mineral wool ceiling tiles usually lies within the range 34–36 dB, but depending on the openness of the tile surface, the sound absorption coefficient may range from 0.1 for smooth tiles, through 0.5 for fissured finishes to 0.95 for open-cell tiles overlaid with 20 mm mineral wool.

**GLASS WOOL**

Glass wool is made by the Crown process (Fig. 13.1), which is similar to the process used for mineral wool. A thick stream of glass flows from a furnace into a forehearth and by gravity into a rapidly rotating steel alloy dish, punctured by hundreds of fine holes around its perimeter. The centrifugal force expels the filaments which are further extended into fine fibres by a blast of hot air. The fibres are sprayed with a bonding agent and then sucked onto a conveyor to produce a mat of the appropriate thickness. This is cured in an oven to set the bonding agent, and then finally cut, trimmed and packaged.

Glass wool is non-combustible, water-repellent and rot proof, contains no CFCs or HCFCs and is available in a range of product forms:

- loose for blown cavity wall insulation;
- rolls, either unaced or laminated between kraft paper and polythene, for roofs, within timber-frame construction, internal walls and within floors;
- semi-rigid batts with water-repellent silicone for complete cavity fill of new masonry;
• rigid batts for partial cavity fill within new masonry;
• compression-resistant slabs for solid concrete or beam and slab floors;
• a laminate of rigid glass wool and plasterboard for dry linings;
• PVC-coated rigid panels for exposed factory roof linings.

(The thermal conductivity of glass wool products ranges typically between 0.031 and 0.040 W/m K at 10°C.)

The sound- and fire-resistant properties of glass wool are similar to those of mineral wool. Glass wool sound-deadening quilts, which have overlaps to seal between adjacent units, are used to reduce impact sound in concrete and timber floating floors. Standard quilts are appropriate for use in lightweight partitions and over suspended ceilings.

Resin-bonded glass wool treated with water repellent is used to manufacture some ceiling tiles which meet the Class 0 fire spread requirements of the Building Regulations (BS 476: Parts 6 and 7) and also offer sound absorption to reduce reverberant noise levels.

**CELLULAR OR FOAMED GLASS BLOCKS**

Cellular or foamed glass (CG) is manufactured from a mixture of crushed glass and fine carbon powder, which on heating to 1000°C, causes the carbon to oxidise, creating bubbles within the molten glass. The glass is annealed, cooled and finally cut to size. The black material is durable, non-combustible, easily worked, and has a high compressive strength. It is water resistant due to its closed-cell structure, impervious to water vapour and contains no CFCs. Cellular glass products are described in BS EN 13167: 2008.

Cellular glass slabs are appropriate for roof insulation, including green roofs and roof-top car parks owing to their high compressive strength. The slabs are usually bonded in hot bitumen to concrete screeds, profile metal decking or reinforced bitumen membrane on timber roofing. Foamed glass is suitable for floor insulation under the screed and may be used internally, externally or within the cavity of external walls. Externally, it may be rendered or tile hung and internally finished with plasterboard or expanded metal and conventional plaster. (The thermal conductivity of cellular glass is within the range 0.037–0.048 W/m K at 10°C, depending on the grade.)

**EXFOLIATED VERMICULITE**

Exfoliated vermiculite (EV) is manufactured by heating the natural micaceous mineral. The material, containing up to 90% air by volume, is used as a loose fill for loft insulation and within a cementitious spray produces a hard fire protection coating for exposed structural steelwork. Where thicknesses over 30 mm are required, application should be in two coats. The product has a textured surface finish which may be exposed internally or painted in external applications. Depending on the thickness of application and the ratio (Hp/A) between exposed surface area and steel cross-section (Chapter 5), up to 240 minutes’ fire protection may be obtained. Vermiculite is used for certain demountable fire stop seals where services penetrate...
through fire compartment walls. The standards BS EN 14317 Parts 1 and 2 describe coated and hydrophobic vermiculite and premixed vermiculite with binders. (The thermal conductivity of exfoliated vermiculite is 0.062 W/m K. Within lightweight aggregate concrete a thermal conductivity of typically 0.11 W/m K can be achieved.)

EXPANDED PERLITE

Expanded perlite (EPB) is manufactured by heating natural volcanic rock minerals. It is used for loose and bonded in situ insulation for roofs, ceilings, walls and floors also as preformed boards. The expanded perlite products are described in BS EN 13169: 2008 and the loose fill material in BS EN 14316 Parts 1 and 2. (The thermal conductivity of expanded perlite boards is 0.05 W/m K.)

CALCIUM SILICATE

Calcium silicate (CS), which is described in Chapter 12, has the advantage of good impact resistance and is very durable. Various wallboards are manufactured with calcium silicate boards laminated to extruded polystyrene. (Calcium silicate typically has a thermal conductivity of 0.29 W/m K.)

GLASS AND MULTIPLE GLAZING

The thermal and sound insulation effects of double and triple glazing and the use of low-emissivity glass are described in Chapter 7.

AEROGEL

Aerogels are extremely lightweight, hydrophobic amorphous silica materials with densities as low as 3 kg/m³ ($\rho_{\text{air}} = 1.2$ kg/m³). They are manufactured by solvent evaporation from silica gel under reduced pressure. Aerogels are highly porous with typically 95–97% and even 99.8% air space, but significantly the pore size of 20 nm is so small that it is less than the mean free path of nitrogen and oxygen in the air. This prevents the air particles moving and colliding with each other, which would normally give rise to gas phase heat conduction. With only 3–5% solid material, heat conduction in the solid phase is very limited. When used to fully fill the cavity in glazing units, 0.5–4 mm aerogel granules prevent the movement of air, thus reducing heat transfer by convection currents. Limited heat transfer can therefore only occur across the glazing unit by radiation.

Light transmission through aerogel is approximately 80% per 10 mm thickness, giving a diffuse light and eliminating the transmission of ultraviolet light. Airborne sound transmission is reduced particularly for lower frequencies of less than 500 Hz. The material is hydrophobic and therefore resistant to mould growth.

Polycarbonate glazing units filled with aerogel are available as 10 and 16 mm panels, over a range of sizes to fit profiled glass trough sections (Chapter 7) or as translucent rooflights or wall panels (e.g. 1220 × 3660 mm). One proprietary cladding and roofing system uses aluminium framing to support a sandwich panel of fibreglass sheets separated by aerogel. The panels, 1200 × 3600 mm or 1500 × 3000 mm (maxima), have $U$-values of 0.28 W/m² K. (The thermal conductivity of aerogel silica is typically 0.018 W/m K.)

Organic insulation materials

The use of straw bales is described in Chapter 17 with other recycled products.

CORK PRODUCTS

Cork is harvested from the cork oak ($Quercus suber$) on a 9-year (or more) cycle and is therefore considered to be an environmentally friendly material. For conversion into boards, typically used for roof insulation, cork granules are expanded, then formed under heat and pressure into blocks using the natural resin within the cork. The blocks are trimmed to standard thicknesses or to a taper to produce falls for flat roofs (Fig. 13.2). For increased thermal insulation properties, the cork may be bonded to closed-cell polyurethane or polyisocyanurate foam. In this case the laminate should be laid with the cork uppermost. Cork products are unaffected by the application of hot bitumen in
flat roofing systems. Expanded cork (ICB) is described in BS EN 13170: 2008. (The thermal conductivity of corkboard is 0.042 W/m K.)

**SHEEP’S WOOL**

Sheep’s wool is a very efficient renewable resource insulation material, with a low conductivity that compares favourably to other fibrous insulants. It is available in grey batts ranging in thickness from 50 and 75 to 100 mm thick. Wool is a hygroscopic material, that is, it reversibly absorbs and releases water vapour, and this effect is advantageous when it is used for thermal insulation. When the building temperature rises, wool releases its moisture causing a cooling effect in the fibre and thus a reduced flow of heat into the building. In winter the absorption of moisture warms the material. This evolution of heat helps to prevent interstitial condensation in construction cavities by maintaining the temperature of the fibres above the dew point, and also effectively reduces the heat loss from the building.

Wool is safe to handle only requiring gloves and a dust mask as minimal protection. It causes no irritation except in the rare cases of people with a specific wool allergy. Wool batts, which contain 85% wool and 15% polyester to maintain their form, can easily be cut with a sharp knife or torn to size. Wool is potentially susceptible to rodents, which may use it as a nesting material if it is accessible, but the batts are treated with an insecticide to prevent moth or beetle attack and with an inorganic fire retardant.

Wool batt insulation is suitable for ventilated loft applications between joists or rafters and for timber frame construction. It should be installed with a vapour-permeable breather membrane on the cold side, and kept clear of any metal chimneys or flues. Wool also acts as an effective acoustic insulating material. (The thermal conductivity of wool batts is 0.039 W/m K.)

**CELLULOSE INSULATION**

Cellulose insulation is manufactured from shredded recycled paper. It is treated with borax for flammability and smouldering resistance; this also makes it unattractive to vermin and resistant to insects, fungus and dry rot. Unlike mineral fibre and glass-fibre insulation it does not cause skin irritation during installation. Recycled cellulose has a low embodied energy compared to mineral and glass-fibre insulation, and when removed from a building it may be recycled again or disposed of safely without creating toxic waste. (Treatment with the inorganic salt, borax, ensures that cellulose insulation conforms to BS 5803 Part 4: 1985—Fire Test Class 1 and Smoulder Test Class B2.)

Cellulose insulation may be used directly from bags for internal floors and also lofts where the required eaves ventilation gap must be maintained. For other cavities, including sloping roof voids, the material is dry injected under pressure completely filling all spaces to prevent air circulation. In breathing walls, cellulose insulation is filled inside a breathing membrane, which allows the passage of water vapour through to the outer leaf of the construction. Cellulose may be damp-sprayed in between wall studs before the wall is closed. Cellulose is a hygroscopic material, which under conditions of high humidity absorbs water vapour and then releases it again under dry conditions. Cellulose is an effective absorber of airborne sound. (The thermal conductivity of cellulose is 0.035 W/m K in horizontal applications and 0.038–0.040 W/m K in walls.)

**RECYCLED PLASTIC**

Loft insulation material is manufactured from recycled plastic bottles into a non-woven, clean material (Fig. 13.3). The product contains approximately 85–90% of recycled material, and unlike the equivalent glass fibre or mineral wool, has no loose fibres which may cause irritation. The material is available in rolls of thicknesses from 100 to 200 mm and has a thermal conductivity of 0.043 W/m K.

**FLAX, HEMP AND COCONUT FIBRE**

As the demand for sustainable insulating materials increases, products derived from renewable flax, hemp and coconut fibres are becoming available. Flax
Recycled plastic insulation is suitable for ventilated or breathing constructions. The batts may be used in ceilings and walls, and rolls in lofts, suspended floors and walls. Flax is treated with borax for fire and insect resistance and bonded with potato starch, giving a moisture-absorbing, non-toxic product with good thermal and acoustic insulation properties. (The thermal conductivity of flax is 0.037 W/m K.)

Chopped hemp fibres, treated with borax for fire resistance, are used to produce insulation batts, and also as loose fill for floors and roofs. Hemp is also blended with either wood fibres or waste cotton and 15% polyester fibres to form insulation batts. Hemp, a very tough material, is used in the manufacture of certain particleboards in Germany and generally for paper production. Hempcrete is described in Chapter 3. (The thermal conductivities of hemp insulation products are in the range 0.038–0.040 W/m K.)

Coconut fibre thermal and acoustic insulation boards have the advantage of natural rot resistance. They are available in a range of thicknesses from 10 to 25 mm, and typical uses include ceiling and floor insulation including under-screed applications. (The thermal conductivity of coconut fibre is 0.045 W/m K. The sound reduction for a typical 18 mm under-screed application is 26 dB.)

**FIBRE INSULATION BOARD**

The manufacture of insulation board (WF) or softboard, which is a low-density wood fibre building board, is described in Chapter 4. Standard grades of insulation board should only be used in situations where they are not in contact with moisture, or at risk from the effects of condensation. Insulating board is used for wall linings and may be backed with aluminium foil for increased thermal insulation. Wood fibre insulation products are described in BS EN 13171: 2008.

Insulating board may be impregnated with inorganic fire retardants to give a Class 1 Surface Spread of Flame to BS 476 Part 7 or finished with plasterboard to give a smooth Class 0 fire rated surface. The Euroclass fire performance rating under the conditions specified in BS EN 13986: 2004 for 9 mm untreated high-density medium board of 600 kg/m³ is Class D-s2, d0 for non-floor use. For untreated low-density medium board of 400 kg/m³ the equivalent rating is Class E, pass for non-floor use and for 9 mm untreated softboard of 250 kg/m³ the rating is Class E, pass for non-floor use.

Exposed insulation board has good sound-absorbing properties due to its surface characteristics. Standard 12 mm lining softboard has a noise reduction coefficient of 0.42, although this is increased to 0.60 for the 24 mm board.

Bitumen-impregnated insulating board, with its enhanced water-resistant properties, is used as a thermal insulation layer on concrete floors. The concrete floor slab is overlaid with polythene, followed by bitumen-impregnated insulation board and the required floor finish such as particleboard. In the upgrading of existing suspended timber floors, a loosely laid layer of bitumen-impregnated insulating board under a new floor finish can typically reduce both impact and airborne sound transmission by 10 dB. Bitumen-impregnated insulating board is frequently used in flat-roofing systems as a heat protective layer to polyurethane, polystyrene, or phenolic foams prior to the application of the hot bitumen waterproof membrane. It is also used for sarking in pitched roofs. (The thermal conductivity of insulating board is typically 0.050 W/m K.)

**EXPANDED POLYSTYRENE**

Expanded polystyrene (EPS) is a combustible material, which, in fire, produces large quantities of noxious black smoke, although Type A, with a flame retardant additive is not easily ignitable. Expanded polystyrene, a closed-cell product, is unaffected by water, dilute acids and alkalis but is readily dissolved by most organic solvents. It is rot and vermin proof, and
free of formaldehyde, CFCs and HCFCs. Expanded polystyrene is described in BS EN 13163: 2008.

**Polystyrene beads**

Expanded polystyrene beads are used as loose fill for cavity insulation. To prevent subsequent slippage and escape through voids, one system bonds the polystyrene beads by spraying them with atomised PVA adhesive during injection process, although other processes leave the material loose. Walls up to 12 m in height can be insulated by this type of system. Polystyrene bead insulation should not be used where electrical wiring is present in the cavity, as the polystyrene gradually leaches the plasticiser out from plastic cables, causing their embrittlement which could lead to problems later if the cables are subsequently moved. Polystyrene bead aggregate cement is used to form an insulating sandwich core in concrete cladding panel systems.

**Expanded polystyrene boards**

Expanded polystyrene rigid lightweight boards are used for thermal insulation and four standard grades are available (Table 13.3). The standard material is classified as Euroclass F in relation to fire, but certain flame retardant modified boards are classified as Euroclass E. (Grades range from A1 and A2 through to E.) Load-bearing expanded polystyrene for impact sound insulation properties is designated as type EPS T to BS EN 13163: 2008.

The boards, which are manufactured by fusing together pre-foamed beads under heat and pressure, can easily be cut, sawn or melted with a hot wire. Polystyrene boards provide thermal insulation for walls, roofs and floors. In addition, polystyrene may be cast into reinforced concrete, from which it is easily removed to create voids for fixings.

In cavity wall insulation, a 50 mm cavity may be retained to prevent the risk of water penetration, with proprietary wall ties fixing the boards against the inner leaf. Alternatively, with a fully filled cavity system, the boards may be slightly moulded on the outer surface to shed any water back onto the inside of the external masonry leaf. Interlocking joints prevent cold bridging, air leakage and water penetration at the board joints. In upgrading existing walls, external expanded polystyrene insulation should be protected by suitably supported rendering or tile hanging. Weatherproof rendering should tolerate thermal and moisture movement and be frost resistant to minimise maintenance. For internal wall insulation, expanded polystyrene can be used in conjunction with 12.5 mm plasterboard either separately or as a laminate. Expanded polystyrene is used to give thermal insulation in ground floors. It may be laid below or above the oversite slab; in the latter case, it may be screeded or finished with chipboard. Composite floor panels manufactured from expanded polystyrene and oriented strand board are suitable for beam and block floors while proprietary systems offer thermal insulation to prestressed concrete beam and reinforced concrete screed floors. Expanded polystyrene boards reduce impact and airborne sound transmission through intermediate floors.

Expanded polystyrene is suitable for thermal insulation in flat and pitched roofs. For flat roofs it may be cut to falls. Where hot bitumen products are to be applied, the expanded polystyrene boards must be protected by an appropriate layer of bitumen-impregnated fibreboard, perlite board or corkboard. In metal deck applications the insulating layer may be above or below the purlins, whereas in traditional pitched roofs expanded polystyrene panels are normally installed over the rafters. Expanded polystyrene, although a closed-cell material, acts as a sound absorber, provided it is installed with an air gap between it and the backing surface. It particularly absorbs sound at low frequencies and may be used in floors and ceilings. It is, however, less effective than the open-cell materials,
such as flexible polyurethane foam. Expanded polystyrene incorporating graphite is grey in colour and has enhanced thermal insulation properties as it absorbs infrared radiation and reflects heat. (The thermal conductivity of expanded polystyrene is in the range 0.033–0.040 W/m K depending on the grade. The graphite product has a thermal conductivity of 0.033 W/m K.)

**EXTRUDED POLYSTYRENE**

Extruded polystyrene (XPS) is normally manufactured by a vacuum process, although some is blown with CFCs or carbon dioxide. It is slightly denser and therefore slightly stronger in compression than expanded polystyrene but has a lower thermal conductivity. It has a closed-cell structure with very low water-absorption and vapour-transmission properties. It is produced either with or without a polystyrene skin. Extruded polystyrene is described in BS EN 13164: 2008. It is available with densities ranging from 20 to 40 kg/m³. Extruded polystyrene is widely used for cavity wall and pitched roof insulation. Because of its high resistance to water absorption, extruded polystyrene may be used for floor insulation below the concrete slab and on inverted roofs where its resistance to mechanical damage from foot traffic is advantageous. Extruded polystyrene is also available laminated to tongued and grooved moisture-resistant flooring grade particle-board for direct application to concrete floor slabs, and laminated to plasterboard as a wallboard. (The thermal conductivity of extruded polystyrene is typically 0.025–0.027 W/m K.)

**EXPANDED PVC**

Plasticised PVC open, partially open and closed-cell foams are manufactured as flexible or rigid products within the density range 24–72 kg/m³. The rigid closed-cell products provide low water permeability and are self-extinguishing in fire. Expanded PVC boards are used in sandwich panels and for wall linings. The low-density open-cell material has particularly good acoustic absorbency and can be used to reduce sound transmission through unbridged cavities and floating floors. (The thermal conductivity of expanded PVC is typically 0.030 W/m K.)

**POLYISOCYANURATE FOAM**

Polyisocyanurate foam (PIR) is usually blown with pentane. It is used as a roof insulation material, since it is more heat resistant than other organic insulation foams, which cannot be directly hot-bitumen bonded. Polyisocyanurate is also appropriate for use in wall and floor insulation. PIR is combustible (BS 476 Part 4) with a Class 1 Surface Spread of Flame (BS 476 Part 7) but is more fire resistant than polyurethane foam and it can be treated to achieve Class 0 rating. Polyisocyanurate foam boards have a fire classification of Class F, but the Building Regulations permit their use in wall cavities within masonry leaves of at least 75 mm thickness (BS 4841: 2006). Polyisocyanurate tends to be rather friable and brittle. Certain proprietary systems for insulated cavity closers use PVC-U-coated polyisocyanurate insulation. Such systems offer a damp-proof barrier and can assist in the elimination of cold bridging, which sometimes causes condensation and mould growth around door and window openings. (The thermal conductivity of polyisocyanurate foam is usually in the range 0.023–0.025 W/m K.)

**POLYURETHANE FOAM**

Rigid polyurethane (PUR) is a closed-cell foam currently manufactured using carbon dioxide, pentane or hydrofluorocarbons. The pentane or hydrofluorocarbons remain trapped within the closed cells enhancing the thermal performance, but carbon dioxide diffuses out. Certain polyurethanes are modified with polyisocyanurates. Polyurethane foam products are defined in BS EN 13165: 2008. Rigid polyurethane is a combustible material producing copious noxious fumes and smoke in fire, although a flame-resistant material is available. Polyurethane boards have a fire classification of Class F, but the Building Regulations permit their use in wall cavities within masonry leaves of at least 75 mm thickness (BS 4841: 2006). Polyurethane is used to enhance the thermal insulation properties of concrete blocks either by filling the void spaces in hollow blocks or by direct bonding onto the cavity face. Roofboards, in certain systems pre-bonded to bitumen roofing sheet, are suitable for mastic asphalt and reinforced bitumen membrane roofing systems. Owing to the temperature stability of polyurethane no additional protection from the effects of hot bitumen application is required; the durability of the material also makes it suitable for use in inverted roofs. Laminates with foil or kraft paper are available. Factory-manufactured double-layer profiled-metal sheeting units are frequently filled with rigid polyurethane foam due to its good adhesive and thermal insulation properties. Polyurethane laminated to plasterboard is used as a wallboard.
When injected as a pre-mixed two-component system into cavity walls polyurethane adheres well to the masonry, foaming and expanding in situ to completely fill the void space. It has been used in situations where the cavity ties have suffered serious corrosion, and where additional bonding between the two leaves of masonry is required, but it is not now widely used as a cavity insulation material. However, two-component polyurethane foam spray is effective at closing gaps around service voids, eliminating cold bridges, under tile insulation and for filling inaccessible locations.

Flexible polyurethane foam is an open-cell material offering good noise absorption properties. It is therefore used in unbridged timber-frame partitions, floating floors and duct linings to reduce noise transmission. Polyurethane foams are resistant to fungal growth, aqueous solutions and oils, but not to organic solvents. (The thermal conductivity of rigid polyurethane foam is usually in the range 0.019–0.023 W/m K, at a nominal density of 32 kg/m³. Flexible polyurethane foam typically has a thermal conductivity of 0.048 W/m K.)

**UREA-FORMALDEHYDE FOAM**

Urea-formaldehyde foam (UF) was used extensively in the 1980s for cavity wall insulation, but it can shrink after installation, creating fissures which link the outer and inner leaves. Occasionally, in conditions of high exposure, this had led to rain water penetration. After installation the urea-formaldehyde foam emits formaldehyde fumes, which have in certain cases entered buildings, causing occupants to suffer from eye and nose irritation. The problem normally arises only if the inner leaf is permeable and a cavity greater than 100 mm is being filled. Recent advances claim to have reduced formaldehyde emissions but all installations must be undertaken to the stringent British Standard BS 5618: 1985. The Health & Safety Executive advises against the use of urea formaldehyde where the inner leaf of the cavity wall is porous or has unsealed connections with the interior. (The thermal conductivity of urea-formaldehyde foam is typically 0.038 W/m K.)

**PHENOLIC FOAM**

Phenolic foams (PF), which have very low thermal conductivities, are used as alternatives to rigid polyurethane and polyisocyanurate foams, where a self-extinguishing, low smoke emission material is required. Phenolic foams are produced with densities in the range 35–200 kg/m³. The closed-cell material is expanded with pentane, iso-propyl chloride, hydrofluorocarbons or a mixture of these agents. Phenolic foam is described in BS EN 13166: 2008. Wallboard laminates with plasterboard offer good thermal insulation properties due to the very low thermal conductivity of phenolic foam, compared to polyurethane or extruded polystyrene. Phenolic foams are stable up to a continuous temperature of 120°C. (The thermal conductivity of phenolic foam in the density range 35–60 kg/m³ is typically 0.020 W/m K, although the open-cell material has a thermal conductivity of 0.031 W/m K.)

**Aluminium foil**

Aluminium foil is frequently used as an insulation material in conjunction with organic foam or insulating gypsum products. It acts by a combination of two physical effects. First, it reflects back incident heat due to its highly reflecting surface. Second, owing to its low emissivity, the re-radiation of any heat that is absorbed is reduced. Thin aluminium reflective foil insulation can be inserted between studs, joists or rafters, leaving a 25 mm air gap on either side. In addition to insulation it acts as an air infiltration and vapour barrier.

**THERMO-REFLECTIVE INSULATION PRODUCTS**

Proprietary quilt systems incorporating multi-layers of aluminium foil, fibrous materials and cellular plastics act as insulation by reducing conduction, convection and radiation (Fig. 13.4). A range of these thermoreflective insulation products is manufactured using different combinations of thin plastic foam, plastic bubble sheet, non-woven fibrous wadding with plain and reinforced aluminium foil. Thus, a 10 mm sandwich system consisting of four layers of aluminium foil alternating with three layers of polyethylene bubble sheet can achieve the thermal insulation effect equivalent to that of approximately 100 mm mineral wool. A 30 mm thick system, composed of 19 intermixed layers of aluminium foil, foam and wadding, can achieve the thermal efficiency as roof insulation equivalent to that of 210 mm of mineral wool. Higher levels of insulation are achieved by systems composed of larger numbers of layers and increased overall thicknesses. (The thermal conductivity of foil-faced foam is typically 0.020 W/m K.)
Chlorofluorocarbons in foamed plastics

Until relatively recently, rigid polyurethane and polyisocyanurate foams were blown with chlorofluorocarbons (CFCs). However, due to the worldwide concern over the effects of these gases on the ozone layer, the use of CFCs was phased out in favour of reduced ozone depletion potential (ODP) blowing agents such as the partially halogenated alkanes (PHAs), usually hydrochlorofluorocarbons (HCFCs), which themselves have now virtually been eliminated in favour of hydrofluorocarbons (HFCs) and hydrocarbons with zero ozone depletion potential. Carbon dioxide can be used as a blowing agent, but it produces less dimensionally stable products. Specifically, HFC blown polyurethane foam has better insulating properties (thermal conductivity – 0.019 W/m K) than the equivalent foam blown by carbon dioxide (thermal conductivity – 0.022 W/m K).

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Introduction

Although used in relatively small quantities compared with the load-bearing construction materials, sealants, gaskets and adhesives play a significant role in the perceived success or failure of buildings. A combination of correct detailing and appropriate use of these materials is necessary to prevent the need for expensive remedial work. Many manufacturers are producing solvent-free sealants and adhesives as environmentally friendly alternatives to the traditional solvent-based systems.

Sealants

Sealants are designed to seal the joints between adjacent building components while remaining sufficiently flexible to accommodate any relative movement. They may be required to exclude wind, rain and airborne sound. A wide range of products is available matching the performance characteristics of the sealant to the requirements of the joint. Incorrect specification or application, and poor joint design or preparation are likely to lead to premature failure of the sealant. The standard (BS EN ISO 11600: 2003) classifies sealants into type G for glazing applications and type F (façade) for other construction joints. For both types, classes are defined by movement capability, modulus and elastic recovery (Fig. 14.1).

Key factors in specifying the appropriate sealant are:

- matching the nature and extent of movement to an appropriate sealant;
- appropriate joint design, surface preparation and sealant application;
- service life of the sealant.

Relative Movement within Buildings

The most common causes of movement in buildings are associated with settlement, dead and live load effects including wind loading, fluctuations in temperature, changes in moisture content and, in some cases, the deteriorative effects of chemical or electrolytic action. Depending on the prevailing conditions, the various effects may be additive or compensatory.

Settlement

Settlement is primarily associated with changes in loadings on the foundations during the construction process, although it may continue for some time, frequently up to 5 years, after the construction is complete. Subsequent modifications to a building or its contents may cause further relative movement. Settlement is usually slow and in one direction, creating a shearing effect on sealants used across the boundaries.

Thermal Movement

All building materials expand and contract to some degree with changes in temperature. For timber the movement is low, but moderate for glass, steel, brick, stone and concrete, and relatively high for plastics and aluminium. Such thermal movements are accentuated by the effects of colour, insulation and thickness of
the material. Dark materials absorb solar radiation and heat more quickly than light reflective materials. Also, well-insulated claddings respond quickly to changes in solar radiation, producing rapid cyclical expansion movements, whereas heavy construction materials respond more slowly but will still exhibit considerable movements over an annual cycle. Typical thermal movements are shown in Table 14.1.

**Moisture movement**

Moisture movement falls into two categories: irreversible movements as new materials acclimatise to the environment, and reversible cyclical movements due to climatic variations. Many building materials, especially concrete and mortars, exhibit an initial contraction during the drying-out process. Incorrectly seasoned timber will also shrink but new bricks used too quickly after manufacture will expand. After these initial effects, all materials which absorb moisture will expand and contract to varying degrees in response to changes in their moisture content. Depending on climatic conditions, moisture and thermal movements may oppose or reinforce each other. Typical irreversible and reversible moisture movements are shown in Table 14.2.

**Loading and deterioration**

Movements associated with live loads, such as machinery, traffic and wind, can cause rapid cyclical movements within building components. The deterioration of materials, such as the corrosion of steel or sulphate attack on concrete, is often associated with irreversible expansion, causing movement of adjacent components. Concrete structures may exhibit creep, which is gradual permanent deformation under load, over many years.

**TYPES OF SEALANT**

There are three distinct types of sealant, plastic, elastoplastic and elastic, each of which exhibit significantly different properties which must be matched to the appropriate application (BS 6213: 2000).

**Plastic sealants**

Plastic sealants, which include general-purpose mastics, allow only a limited amount of movement, but when held in a deformed state they stress-relax. Elastic recovery is limited to a maximum of 40%. Plastic sealants dry by the formation of a surface skin, leaving liquid material encased to retain flexibility. However, with time, the plastic core continues to harden; thus, durability is related to the thickness of the material used. Plastic sealants (Code P to BS EN ISO 11600: 2003) are more suitable for locations in which the majority of movement is irreversible (BS 6213: 2000).

**Oil-based mastics**

For oil-based mastics a 10 mm depth is required for optimum durability with a typical life expectancy of 2–10 years. The effects of ultraviolet degradation
Table 14.1 Thermal movements of building materials

<table>
<thead>
<tr>
<th>Material</th>
<th>Typical thermal movement (mm/m for 85°C change)</th>
<th>Coefficient of linear expansion per °C × 10⁻⁶</th>
</tr>
</thead>
<tbody>
<tr>
<td>Masonry</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Concrete—standard aggregates</td>
<td>1.2</td>
<td>10–14</td>
</tr>
<tr>
<td>Calcium silicate brickwork</td>
<td>1.2</td>
<td>8–14</td>
</tr>
<tr>
<td>Concrete blockwork</td>
<td>1.0</td>
<td>6–12</td>
</tr>
<tr>
<td>Concrete—aerated</td>
<td>0.7</td>
<td>8</td>
</tr>
<tr>
<td>Concrete—limestone aggregate</td>
<td>0.6</td>
<td>7–8</td>
</tr>
<tr>
<td>Clay brickwork</td>
<td>0.5–0.7</td>
<td>5–8</td>
</tr>
<tr>
<td>GRC</td>
<td>1.0</td>
<td>7–12</td>
</tr>
<tr>
<td>Plaster</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dense plaster</td>
<td>1.5</td>
<td>18–21</td>
</tr>
<tr>
<td>Lightweight plaster</td>
<td>1.4</td>
<td>16–18</td>
</tr>
<tr>
<td>Metals</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Zinc (along roll)</td>
<td>2.7</td>
<td>32 (22 across roll)</td>
</tr>
<tr>
<td>Lead</td>
<td>2.5</td>
<td>29</td>
</tr>
<tr>
<td>Aluminium</td>
<td>2.0</td>
<td>24</td>
</tr>
<tr>
<td>Titanium zinc</td>
<td>1.8</td>
<td>20–22</td>
</tr>
<tr>
<td>Copper</td>
<td>1.4</td>
<td>17</td>
</tr>
<tr>
<td>Stainless steel (austenitic)</td>
<td>1.4</td>
<td>18</td>
</tr>
<tr>
<td>Stainless steel (ferritic)</td>
<td>0.8</td>
<td>10</td>
</tr>
<tr>
<td>Terne-coated stainless steel</td>
<td>1.4</td>
<td>17</td>
</tr>
<tr>
<td>Structural steel</td>
<td>1.0</td>
<td>12</td>
</tr>
<tr>
<td>Stone and glass</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Glass</td>
<td>0.9</td>
<td>9–11</td>
</tr>
<tr>
<td>Slate</td>
<td>0.9</td>
<td>9–11</td>
</tr>
<tr>
<td>Granite</td>
<td>0.8</td>
<td>8–10</td>
</tr>
<tr>
<td>Sandstone</td>
<td>0.8</td>
<td>7–12</td>
</tr>
<tr>
<td>Marble</td>
<td>0.4</td>
<td>4–6</td>
</tr>
<tr>
<td>Limestone</td>
<td>0.3</td>
<td>3–4</td>
</tr>
<tr>
<td>Plastics</td>
<td></td>
<td></td>
</tr>
<tr>
<td>ABS</td>
<td>8.0</td>
<td>83–95</td>
</tr>
<tr>
<td>PVC (rigid)</td>
<td>6.0</td>
<td>40–80</td>
</tr>
<tr>
<td>GRP</td>
<td>3.0</td>
<td>18–35</td>
</tr>
<tr>
<td>Timber</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Wood (along grain)</td>
<td>0.5</td>
<td>4–6</td>
</tr>
</tbody>
</table>

Note:
Typical thermal movements of building materials in use calculated for a temperature variation of 85°C (e.g. −15 to +70°C) (measured in mm/m)

are reduced by painting. Typical uses include sealing around window and door frames in traditional low-rise building. Oil-based mastics are not suitable for use with PVC-U windows. (The typical movement accommodation for oil-based mastics is 10%).

Butyl sealants
Butyl sealants are plastic but with a slightly rubbery texture. They are used in small joints as a gap filler and general-purpose sealant where oil-based mastics would dry too rapidly. Life expectancy is between 10 and 20 years if they are protected from sunlight by painting, but only up to 5 years in exposed situations. (The typical movement accommodation for butyl sealants is 10%).

Acrylic sealants
Water-based acrylic sealants are frequently used for internal sealing such as between plaster and new windows. The solvent-based acrylic sealants are durable for up to 20 years, with good adhesion to slightly contaminated surfaces. They accommodate only limited movement but produce a good external seal around windows, for both new and remedial work. (The typical movement accommodation for water- and solvent-based acrylic sealants is 15% and 20%, respectively.)

Polymer/bitumen sealants
Solvent-based bitumen sealants are generally suitable for low-movement joints in gutters and flashings. Hot-poured bitumen is used for sealing movement joints
in asphalt and concrete floor slabs, although compatibility with any subsequent floor coverings should be verified.

Linseed oil putty
Traditional putty contains a mixture of linseed oil and inorganic fillers (BS 544: 1969), which sets by a combination of aerial oxidation of the oil and some absorption into the timber. A skin is produced initially, but the mass ultimately sets to a semi-rigid material. Application is with a putty knife onto primed timber. For application to steel window frames, non-absorbent hardwoods and water-repellent preservative-treated softwoods, non-linseed oil putty is appropriate. Linseed oil putty should be painted within 2 weeks, whereas metal casement putty may be left 3 months before painting.

Elastoplastic sealants
Elastoplastic sealants will accommodate both slow cyclical movements and permanent deformations. A range of products offer appropriately balanced strength, plastic flow and elastic properties for various applications.

Polysulphide sealants
Polysulphide sealants are available as one- or two-component systems. The one-component systems have the advantage that they are ready for immediate use. They cure relatively slowly by absorption of moisture from the atmosphere, initially forming a skin and fully curing within 2–5 weeks. One-component systems are limited in their application to joints up to 25 mm in width, but their ultimate performance is comparable to that of the two-component materials. Typical uses include structural movement joints in masonry, joints between precast concrete or stone cladding panels and sealing around windows. The two-component polysulphide sealants require mixing immediately before use and fully cure within 24–48 hours. They are more suitable than one-component systems for sealing joints which are wider than 25 mm, have large movements, or are subject to vandalism during setting. Uses include sealing joints within concrete and brickwork cladding systems and also within poorly insulated lightweight cladding panels. Polysulphides have a life expectancy of 20–25 years. (The typical movement accommodation for polysulphide sealants is up to 25% for one-part systems and up to 30% for two-part systems.)

Elastic sealants
Elastic sealants are appropriate for sealing dynamic joints where rapid cyclic movement occurs. They are often sub-classified as low- or high-modulus depending on their stiffness. Low-modulus sealants should be used where joints are exposed to long periods of compression or extension and where the substrate material is weak. Elastic sealants are categorised as Code E to BS EN ISO 11600: 2003.

Polyurethane sealants
Polyurethane sealants are available as one- or two-component systems. The products are highly elastic but surfaces should be carefully prepared and usually primed to ensure good adhesion. Polyurethane sealants have good abrasion resistance and durability is good, ranging from 20 to 25 years. Typical applications are joints within glazing, curtain walling, lightweight cladding panels and floors. (The typical movement accommodation for polyurethane sealants is between 10% and 30% depending on the modulus.)

Silicone sealants
Silicone sealants are usually one-component systems which cure relatively quickly in air, frequently with the evolution of characteristic smells such as acetic acid. Generally, silicone sealants adhere well to metals and glass, but primers may be necessary on friable or porous surfaces, such as concrete or stone. Silicone sealants may cause staining on stonework which is difficult to remove. High-modulus silicone sealants are resilient. Typical applications include glazing and curtain-wall systems, movement joints in ceramic tiling and around sanitary ware. Low-modulus silicone sealants are very extensible and are appropriate for use in joints subject to substantial thermal or moisture movement. Typical applications are the perimeter sealing of PVC-U and aluminium windows and also cladding systems. Silicone sealants are durable with life expectancies within the range 25–30 years. (The typical movement accommodation for silicone sealants ranges from 20% to 70% depending on the modulus.)

Epoxy sealants
Epoxy sealants are appropriate for stress-relieving joints where larger movements in compression than in tension are anticipated. Typical applications include floor joints and the water sealing of tiling joints within swimming pools. Epoxy sealants have a life expectancy of 10–20 years. (The typical movement accommodation of epoxy sealants is within the range 5–15%).
JOINT DESIGN

There are three forms of joint; butt, lap and fillet (Fig. 14.2). However, only butt and lap joints will accommodate movement. Generally, lap joints, in which the sealant is stressed in shear, will accommodate double the movement of butt joints in which the sealant is under tension or compression. Furthermore, lap joints tend to be more durable as the sealant is partially protected from the effects of weathering. However, lap joints are generally more difficult to seal than butt joints. Frequently, joints are made too narrow, either for aesthetic reasons or due to miscalculation of component tolerances. The effect is that extent of movement is excessive in proportion to the width of sealant, causing rapid failure.

To correctly control the depth of the sealant and to prevent it adhering to the back of the joint, a compressible back-up material, usually rectangular or round closed cell polyethylene, is inserted (Fig. 14.3). The polyethylene acts as a bond-breaker by not adhering to the sealant. Where the joint is filled with a filler board, such as impregnated fibreboard or corkboard, a plastic bond-breaker tape or closed-cell polyethylene strip should be inserted. Normally, the depth of the sealant should be half the width of the joint for elastic and elastoplastic sealants and equal to the width of the joint for plastic sealants. A minimum depth of 6 mm is normally required. The minimum width of the joint is calculated from the maximum total relative movement (TRM) to be accommodated and the movement accommodation factor (MAF), that is, the extensibility of the sealant to be used. Where insufficient depth is available to insert a polyethylene foam strip, a tape bond-breaker should be inserted at the back of the joint.

Minimum joint width calculation (BS 6093: 2006):

Total relevant movement (TRM) = 3 mm  
Movement accommodation factor (MAF) = 25%  
Width of sealant to accommodate movement = \( \frac{3}{0.25} + 3 = 15 \text{ mm} \)

In order to obtain good adhesion, the joint surfaces should be prepared by the removal of contaminants, loose material or grease and by the application of a primer if specified by the sealant manufacturer. Most sealants are applied directly by gun application,
although tooled, poured and tape/strip sealants are also used. Tooling helps to remove air bubbles entrained in two-component mixes; if left, air bubbles would reduce the durability of the seal. Externally, recessed cladding joints show less staining than flush joints, although the usual finish is a slightly concave surface. Where stonework is being sealed, non-staining silicone sealants must be used to prevent the migration of plasticiser into the stone, which could cause discolouration. Sealants to floor joints need to be tough, and therefore wider to accommodate the necessary movements and recessed to prevent mechanical damage. Alternatively, proprietary mechanical jointing systems should be used.

**Colour matching**

While most sealants, except the black bituminous products, are available in white, translucent, greys and browns, the silicone sealants appropriate for use around kitchen and bathroom units are available in a wide range of colours. For these purposes, fungicides are often included within the formulation.

**FIRE-RESISTANT SEALANTS**

Many fire-resistant sealants are based on the use of intumescent materials which expand copiously in fire. The intumescent components commonly used are either ammonium phosphate, hydrated sodium silicate or intercalated graphite (layers of water and carbon) and these are incorporated into the appropriate sealant. Intumescent oil-based mastics and acrylic sealants are suitable for sealing low-movement joints around fire check doors. For the fire-resistant sealing of structural movement joints, fire-resistant grades of low-modulus silicone, two-part polysulphide and acrylic sealants are available. Maximum fire resistance is obtained if the sealant is applied to both faces of the joint, with mineral wool or glass-fibre insulation in the void space. Four hours’ fire resistance with respect to both integrity and insulation can be achieved for a 20 mm wide movement joint within 150 mm concrete (BS 476-20: 1987). The low-modulus silicone is appropriate for sealing fire-resisting screens, curtain walls, claddings and masonry subject to movement. The two-part polysulphide is designed for use in concrete and masonry fire-resisting joints. Acrylic sealants are appropriate over a wide range of materials but where timber is involved an allowance must be made for its loss by charring.

Intumescent fillers manufactured from acrylic emulsions with inert fillers and fire-retardant additives can be applied either by gun or by trowel to fill voids created around service ducts within fire-resistant walls. Four hours of fire resistance can be achieved with these materials. Intumescent tapes are appropriate for application within structural movement joints. Most intumescent sealants are now *low-smoke* and evolve no halogenated products of combustion in fire situations. (The typical movement accommodation for intumescent acrylic sealants is 15%.)

**FOAM SEALANTS**

Compressible strips of closed-cell PVC and polyethylene, or open-cell polyurethane foams, coated on one or both edges with pressure-sensitive adhesive, are used to seal thermal movement and differential settlement joints, gaps around window and door frames, and in air-conditioning ductwork. Strips may be uniform in section or profiled for particular applications. Aerosol-dispersed polyurethane foam is widely used as an all-purpose filler. It is available either as foam or as expanding foam, and acts as an adhesive, sealant, filler and insulator.

**CONCRETE JOINT FILLERS AND SEALANTS**

Concrete joint fillers for use in pavements are specified by the standards BS EN 14188: 2004, Parts 1 and 2 for hot and cold application sealants, respectively. Sealants for cold application are classified as single-component systems (S) or multi-component systems (M) and subdivided into self-levelling (sl) or non-sag (ns) types. An additional classification A, B, C or D relates to increasing level of resistance to chemicals. Standard hot-applied joint sealants are classified as elastic (high extension) Type N1, and normal (low extension) Type N2. Where fuel resistance is also required, the higher specification grades F1 and F2 are necessary.

**Gaskets**

Gaskets are lengths of flexible components of various profiles, which may be solid or hollow and manufactured from either cellular or non-cellular materials. They are held in place either by compression or encapsulation into the adjacent building materials, and maintain a seal by pushing against the
two surfaces (Fig. 14.4). Typical applications include the weather sealing of precast cladding units and facade systems. Within precast concrete, glass-fibre reinforced polyester (GRP) or glass-fibre reinforced cement (GRC) cladding units, the gaskets are typically inserted into recessed open-drained joints. The gaskets therefore act as a rain barrier, but because they do not necessarily fit tightly along their full length, they can be backed up by compressed cellular foam wind penetration seals. Gaskets should not be either stretched or crammed in during insertion as they will subsequently shrink leaving gaps or pop out causing failure.

In glazing and related curtain walling systems, gaskets may be applied as capping seals, retained by appropriate profiles within the mullions and transoms; alternatively, the gaskets may be recessed within the joints of the glazing system to give narrower visual effect to the joint. Some glazing gaskets of H- or U-sections are sealed with a zipper or filler strip which is inserted in the profile, compressing the material into an air- and watertight seal. Gaskets and weatherstripping for use on doors, windows and curtain walling are classified by a letter and digit code which defines the use and key physical properties of the particular product, enabling appropriate specification (Table 14.3).

The standard materials for gaskets used in construction are neoprene which is highly elastic, ethylene propylene diene monomer (EPDM) which has better weathering characteristics than neoprene, and silicone rubbers which are highly resistant to ultraviolet light, operate over a wide range of temperatures and are available in almost any colour. Cruciform section gaskets of polychloroprene rubber are suitable for vertical joints between precast concrete panels.

Waterstops for embedding into in situ concrete for sealing movement and construction joints are manufactured in PVC or rubber according to the required movement (Fig. 14.5). Sections are available in long extruded lengths and factory-produced intersections. Applications include water-containing structures and water exclusion from basements. Waterstops placed centrally within concrete will resist water pressure from either side, but externally positioned waterstops, not encased below the concrete slab or within permanent concrete shuttering, will only resist water pressure from the outer face.

Proprietary systems offer watertight expansion jointing for horizontal surfaces, such as roof car parks and pedestrian areas. Systems usually combine complex aluminium or stainless steel profiles with extruded synthetic rubber inserts. Materials can withstand high loads, with good resistance to bitumen and salt water.

Dry glazing strips are based on elastomeric polymers, typically EPDM or butyl rubber. Usually, the synthetic rubber strip has a self-adhesive backing which adheres to the rebate upstand. With external

<table>
<thead>
<tr>
<th>Letter (G or W)</th>
<th>Digit 2</th>
<th>Digit 3</th>
<th>Digit 4</th>
<th>Digit 5</th>
<th>Digit 6</th>
</tr>
</thead>
<tbody>
<tr>
<td>Category</td>
<td>Working range (mm)</td>
<td>Compression force (KPa)</td>
<td>Working temperature range (°C)</td>
<td>Deflection recovery (%)</td>
<td>Recovery after ageing (%)</td>
</tr>
<tr>
<td>Gasket (G)</td>
<td>9 grades identified</td>
<td>9 grades identified</td>
<td>6 grades identified</td>
<td>8 grades identified</td>
<td>8 grades identified</td>
</tr>
<tr>
<td>Weatherstripping (W)</td>
<td>(1–9)</td>
<td>(1–9)</td>
<td>(1–6)</td>
<td>(0–7)</td>
<td>(1–8)</td>
</tr>
</tbody>
</table>
beading, the dry glazing strip can also be applied to each bead, which is then fixed with suitable compression to ensure a good seal to the glass. The performance requirements and classification for gaskets and weather-stripping for doors, windows and curtain walling are described in the standard BS EN 12365-1:2003.

Adhesives

**TYPES OF ADHESIVE**

The traditional adhesives based on animal and vegetable products have largely been superseded by synthetic products manufactured by the polymer industry, except for casein, manufactured from skimmed milk, which is currently used as a timber adhesive (BS EN 12436:2002). The range of adhesives is under constant development and particular applications should always be matched to manufacturers’ specifications. Special notice should be taken of exclusions where materials and adhesives are incompatible, and also to safety warnings relating to handling and the evolution of noxious fumes or flammable vapours. Adhesives are more efficient when bonding components subject to shear forces rather than direct tension. They are least efficient against the peeling stresses. Most adhesives have a *shelf life* of 12 months when stored unopened under appropriate conditions. The *pot life* after mixing the two-component systems ranges from a few minutes to several hours.

**Tile adhesives**

The standard BS EN 12004:2007 classifies adhesives for tiles into three types: cementitious (C), dispersion (D) and reaction resin (R). Each of these types may have further characteristics defined by classes relating to enhanced adhesive properties, faster setting, reduced slip or extended open time (the time between spreading the adhesive and applying the tiles) (Table 14.4). Dispersion adhesives are the ready-for-use aqueous polymer dispersions, whilst the reaction resin adhesives are one- or two-component systems which set by chemical reaction. Tile adhesives with no more than 1% organic material are classified for fire resistance as Class A1 or A1fl.

**Ceramic wall tile adhesives**

Wall tile adhesives are usually polyvinyl acetate (PVA), acrylic- or cement-based compositions. The standard PVA thin-bed adhesives, typically to 3 mm, will only tolerate moisture, whereas the thin-bed water-resistant acrylic-based adhesives are suitable for fixing wall

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**Table 14.4 Classification of tile adhesives by composition and properties (BS EN 12004: 2007)**

<table>
<thead>
<tr>
<th>Classification</th>
<th>Composition and properties</th>
</tr>
</thead>
<tbody>
<tr>
<td>Type C</td>
<td>Cementitious adhesive — hydraulic binding resin</td>
</tr>
<tr>
<td>Type D</td>
<td>Dispersion adhesive — aqueous organic polymer resin</td>
</tr>
<tr>
<td>Type R</td>
<td>Reaction resin adhesive — one or two component synthetic resin</td>
</tr>
<tr>
<td>Class 1</td>
<td>Normal adhesive</td>
</tr>
<tr>
<td>Class 2</td>
<td>Improved adhesive</td>
</tr>
<tr>
<td>Class F</td>
<td>Fast setting adhesive</td>
</tr>
<tr>
<td>Class T</td>
<td>Reduced slip adhesive</td>
</tr>
<tr>
<td>Class E</td>
<td>Extended open time adhesive</td>
</tr>
<tr>
<td>Class S1</td>
<td>Deformable adhesive</td>
</tr>
<tr>
<td>Class S2</td>
<td>Highly deformable adhesive</td>
</tr>
</tbody>
</table>
tiles and mosaics in damp and wet conditions associated, for example, with domestic showers. Some acrylic-based products evolve ammonia on setting. The water-resistant cements and polymer-modified cement products are appropriate for both internal and external use and can usually be applied with either thin or thick bedding. The polymer-modified cement adhesives are also suitable for fixing marble, granite and slate tiles up to 15 mm thick. For chemical resistance thin-bed epoxy resin-based adhesives are available. In all cases the substrate must be sound with new plaster, brickwork, concrete, fully dried out for 2–6 weeks. Plasterboard and timber products must be adequately fixed at 300 mm centres horizontally and vertically to ensure rigidity. In refurbishment work, flaking or multi-layered paint should be removed and glazed surfaces made good. Where the tile adhesive is classified as waterproof either acrylic- or cement-based, it may be used as the grouting medium. Alternatively, equivalent waterproof grouting is available in a wide range of colours to blend or contrast with the wall tiles. Epoxy-resin tile grout is available for very wet conditions.

**Ceramic floor tile adhesives**

The majority of ceramic floor tile adhesives are cement based, used either as thick bed (up to 25 mm) or thin bed according to the quality of the substrate. Standard products are suitable for fixing ceramic tiles, quarries, brick slips, stone and terrazzo to well dried out concrete or cement/sand screed. Where suspended timber floors are to be tiled, they must be well ventilated and strong enough to support the additional dead load. An overlay of 12 mm exterior grade plywood, primed with bonding agent and screwed at 200 mm centres, may be necessary. In refurbishment work, flaking or multi-layered paint should be removed and glazed surfaces made good. Where the refurbishment work, flaking or multi-layered paint should be removed and glazed surfaces made good. Where the tile adhesive is classified as waterproof either acrylic- or cement-based, it may be used as the grouting medium. Alternatively, equivalent waterproof grouting is available in a wide range of colours to blend or contrast with the wall tiles. Epoxy-resin tile grout is available for very wet conditions.

**Contact adhesives**

Contact adhesives based on polychloroprene rubber, either in organic solvents or in aqueous emulsions, are normally suitable for bonding decorative laminates, and other rigid plastics such as PVC and ABS to timber, timber products and metals. The adhesive is usually applied on both surfaces; the solvent or emulsion is allowed to become touch dry, prior to bringing the two surfaces into contact, when an immediate strong bond is produced. The aqueous emulsion products can also be suitable for fixing sealed cork and expanded polystyrene and have the advantage that no fumes are evolved. Expanded polystyrene tiles may be adversely affected by solvent-based formulations.

**Vinyl floor tile and wood block adhesives**

Most vinyl floor tile and wood block adhesives are based on either rubber/bitumen, rubber/resin or modified bitumen emulsions. In all cases it is essential that the sub-floor is dry, sound, smooth and free from any contamination which would affect the adhesion. Where necessary cement/acrylic or cement/latex floor levelling compound should be applied to concrete, asphalt or old ceramic tiled floors. Some cement/latex materials evolve ammonia during application.

**Wood adhesives**

Wood joints generally should be in close contact with a gap of less than 0.15 mm, but so-called gap-filling adhesives satisfactorily bond up to 1.3 mm. Polyvinyl acetate (PVA) wood glues are widely used for most on-site work and in the factory assembly of mortice and tenon joints for doors, windows and furniture. The white emulsion sets to a colourless translucent thermoplastic film, giving a bond of similar strength to the timber itself, but insufficient for bonding load-bearing structural members. Components should be clamped in position for up to 12 hours to ensure maximum bonding, although this may be reduced by increasing the temperature. Waterproof PVA adhesives, which partially cross-link on curing, are suitable for protected external use but not immersion in water. PVA adhesives generally retain their strength up to 60°C and do not discolour the timber, except by contact with ferrous metals.

The thermosetting wood resins are mainly two-component systems based on phenolic compounds, such as urea, melamine, resorcinol or phenol, which
cure with formaldehyde to produce load-bearing adhesives (BS EN 301: 2006). Most formulations require the mixing of the resin and hardener, but a pre-mixed dry powder to which water is added is also available. Structural resin-based adhesives are designated for exterior or high-temperature exposure (Type 1) or protected use (Type 2). Melamine formaldehyde adhesives will not resist prolonged exposure to weathering. Urea formaldehyde adhesives are generally moisture resistant or for interior use only. Certain timber fire-retardant and preservative treatments reduce the efficiency of timber adhesives, although generally those based on phenol formaldehyde/resorcinol formaldehyde are unaffected.

Wallpaper adhesives

Standard wallpaper adhesives are based on methyl cellulose, a white powder which is water soluble giving a colourless solution. For fixing the heavier papers and decorative dado strips, polyvinyl acetate is an added component. Cold water starch is also available as both a wall sizing agent and wallpaper adhesive. Most wallpaper pastes contain fungicide to inhibit mould growth. The standard BS 3046: 1981 describes five types of adhesive ranging from low solids to high wet and dry strength with added fungicide.

Epoxy resin adhesives

Epoxy resins are two-component, cold-curing adhesives which produce high-strength, durable bonds. Most require equal quantities of the resin and hardener to be mixed and various formulations are available giving curing times ranging from minutes to hours. Strong bonds can be obtained to timber, metal, glass, concrete, ceramics and rigid plastics. Epoxy resins may be used internally or externally and they are resistant to oils, water, dilute acids, alkalis, and most solvents except chlorinated hydrocarbons. Epoxy resins are frequently used for attaching stainless steel fixings into stone and brick slips prior to their casting into concrete cladding panels. Epoxy flooring adhesives may be used for bonding vinyl floor finishes in wet service areas and to metal surfaces.

Cyanoacrylate adhesives

Cyanoacrylates are single-component adhesives which bond components held in tight contact within seconds. A high tensile bond is produced between metals, ceramics, most plastics and rubber. The curing is activated by adsorbed moisture on the material surfaces, and only small quantities of the clear adhesive are required. The bond is resistant to oil, water, solvents, acid and alkalis but does not exhibit high impact resistance. A range of adhesive viscosities is manufactured to match to particular applications.

Hot-melt adhesives

Hot-melt adhesives for application by glue gun are usually based on the thermoplastic copolymer, ethylene vinyl acetate (EVA). Formulations are available for joining materials to either flexible or rigid substrates. Generally, the adhesive should be applied to the less easily bonded surface first (e.g. the harder or smoother surface) and then the two components should be pressed together for at least one minute. Where metals are to be bonded they should be pre-warmed to prevent rapid dissipation of the heat. Similar adhesives are used in iron-on edging veneers for plastic- and wood-faced particleboard.

Bitumen sheet roofing adhesives

Bitumen adhesives are available for hot application, emulsion or in hydrocarbon solvent for the cold bonding bituminous sheet roofing. The adhesives should be poured and spread by trowel to avoid air pockets, which may cause premature delamination of the sheet from the substrate. Excess bitumen should be removed as it may stain adjacent materials.

Plastic pipe adhesives

Solvent-based vinyl resin adhesives are used for bonding PVC-U and ABS pipes and fittings. The adhesive is brush-applied to both components which are then united and slightly rotated to complete the seal. Curing is rapid but in cold water supply systems, water pressure should not be applied for several hours.

Gap-filling adhesive

Gun grade gap-filling adhesives, either solvent-borne rubber/synthetic rubber resins with filler reinforcement or solvent-free systems, are versatile in their applications. They are generally formulated to bond timber, timber products, decorative laminates, sheet metals, PVC-U and rigid insulating materials (except polystyrene), to themselves and also to brickwork,
blockwork, concrete, plaster and GRP. Typical applications include the fixing of decorative wall panels, dado rails, architraves and skirting boards without nailing or screwing. Surfaces to be bonded must be sound and clean, but the gap-filling properties of the products can allow fixing to uneven surfaces. The materials have good immediate adhesion, and can allow the components to be adjusted into position.

**PVA bonding agent and sealant**

Polyvinyl acetate (PVA) is a versatile material which will act not only as an adhesive as described but also as a bonding agent or surface sealant. As a bonding agent it will bond cement screeds, rendering and plaster to suitable sound surfaces without the requirement for a good mechanical key. PVA will seal porous concrete surfaces to prevent dusting.

**References**

**FURTHER READING**


**STANDARDS**

BS 476 Fire tests on building materials and structures:


BS 544: 1969 Linseed oil putty for use in wooden frames.

BS 1203: 2001 Hot-setting phenolic and aminoplastic wood adhesives.

BS 2499 Hot-applied joint sealant systems for concrete pavements:


BS 3712 Building and construction sealants:


BS 4255 Rubber used in preformed gaskets for weather exclusion from buildings:


BS 5212 Cold applied joint sealants for concrete pavements:


BS 5270 Bonding agents for use with gypsum plaster and cement:


BS 5385 Wall and floor tiling:

BS 8000 Workmanship on building sites:
BS ISO 16938 Building construction. Determination of staining of porous substrates by sealants used in joints:
  Part 2: 2008 Test without compression.
BS EN 204: 2001 Classification of thermoplastic wood adhesives for non-structural applications.
BS EN 301: 2006 Adhesives, phenolic and aminoplast, for load-bearing timber structures. Classification and performance requirements.
BS EN 302 Adhesives for load-bearing timber structures:


BS EN 12365: 2003 Building hardware. Gaskets and weather stripping for doors, windows shutters and curtain walling:


BS EN 12765: 2001 Classification of thermosetting wood adhesives for non-structural applications.

BS EN 12860: 2001 Gypsum based adhesives for gypsum blocks. Definitions, requirements and test methods.

BS EN 13022 Glass in building. Structural sealant glazing:


BS EN 13415: 2002 Adhesives. Test of adhesives of floor coverings.


BS EN 14187 Cold applied joint sealants. Test methods:

BS EN 14188 Joint fillers and sealants:
  Part 4: 2009 Specification for primers to be used with joint sealants.

pr EN 14496: 2002 Gypsum based adhesives for thermal/acoustic insulation composite panels and plasterboards.

BS EN 14680: 2006 Adhesives for non-pressure plastic piping systems. Specifications.

pr EN 14815: 2003 Adhesives, phenolic and aminoplast for finger-joints in lamellae for loadbearing timber structures.

BS EN 14840: 2005 Joint fillers and sealants. Test methods for preformed joint seals.


BS EN 15275: 2007 Structural adhesives. Characterisation of anaerobic adhesives.

BS EN 15416: 2006 Adhesives for load bearing timber structures other than phenolic and aminoplast:

BS EN 15425: 2005 Adhesives, one component polyurethane, for load bearing timber structures. Classification and performance requirements.

BS EN 15434: 2006 Glass in building. Product standard for structural and/or UV resistant sealant.

BS EN 15466 Primers for cold and hot applied joint sealants:

BS EN 15497: 2009 Finger jointed structural timber.

pr BS EN 15651 Sealants for joints in building construction. Definitions, requirements and evaluation of conformity:

BUILDING RESEARCH ESTABLISHMENT
PUBLICATIONS

BRE Digests

BRE Digest 463: 2002  Selecting building sealants with ISO 11600.
BRE Digest 469: 2002  Selecting gaskets for construction joints.

BRE Information paper

BRE IP 12/03  VOC emissions from flooring adhesives.

ADVISORY ORGANISATION

British Adhesives and Sealants Association, 5 Alderson Road, Worksop, Nottinghamshire S80 1UZ, UK (01909 480888).
Introduction

As colour is an important factor in the description of paints, wood stains and varnishes, the key elements of the British Standards, Natural Color System®, RAL and Pantone systems and the Colour Palette Notation are described, although other colour systems including Munsell are also used within the construction industry. Colour is a key feature of architectural design, as illustrated in Fig. 15.1 by the vivid spectrum colour sequence in Barajas Airport, Madrid, by Rogers Stirk Harbour and Partners, and Estudio Lamela.

Colour

BRITISH STANDARDS SYSTEM

The British Standards BS 5252: 1976 (236 colours) and BS 4800: 1989 (100 colours) define colour for building purposes and paints, respectively. A specific colour is defined by the BS 4800 framework with a three-part code consisting of hue (two digits, 00–24), greyness (letter A to E) and weight (two further digits) (Fig. 15.2). Hue is the attribute of redness, yellowness, blue-ness, etc., and the framework consists of 12 rows of hue in spectral sequence plus one neutral row. Greyness is a measure of the grey content of the colour at five levels from the maximum greyness Group A, to clear Group E. The third attribute, weight, is a subjective term which incorporates both lightness (reflectivity to incident light) and greyness. Within a given column, colours have the same weight, but comparisons between columns in different greyness groups should only be made in respect of lightness. The framework has up to 8 columns of equal lightness in each greyness group commencing with the highest lightness. Thus, any colour is defined through the system by its

Fig. 15.1 Colour feature—Barajas Airport, Madrid. Architects: Rogers Stirk Harbour + Partners. Photograph: AENA / Manuel Renau
three-part code, for example Magnolia is yellow-red 08, nearly grey B and low weight 15 (i.e. 08 B 15), Midnight 20 C 40 and Plum 02 C 39. The draft Standard pr BS 4800: 2008 includes an additional 22 colours giving a total of 122 colours including black and white.

**NATURAL COLOR SYSTEM®©**

The Natural Color System®© (NCS) was developed by the Scandinavian Colour Institute in the 1980s and modified in 1995 with a second edition in which extra colours were added and some removed leaving a total of 1950 colours. It is a colour language system which can describe any colour by a notation, communicable in words without the need for visual matching. It has been used by architects, builders and designers who need to coordinate colour specification across a broad range of building products. A range of materials can be colour-referenced using the system; these include wall, floor and ceiling tiles, carpets, fabrics, wall coverings, flexible floor finishes, paints, architectural ironmongery and metalwork, sanitary fittings, laminates and furniture.

The Natural Color System®© is based on the assumption that for people with normal vision there are six pure colours; yellow, red, blue, green, white and black. The four colours, yellow, red, blue and green are arranged around the colour circle, which is then subdivided into 10% steps. For example, yellow changes to red through orange, which could be described as Y50R (yellow with 50% red) (Fig. 15.3). In order to
Fig. 15.3 Natural Color System®©. Images: Courtesy of the Scandinavian Colour Institute AB, www.sci-sweden.se
superimpose the black/white variation and also intensity of colour, each of the forty 10% steps around the colour circle may be represented by colour triangles, with the pure colour at the perimeter apex and the vertical axis illustrating blackness/whiteness. A colour may be therefore described as having 10% blackness and 80% chromatic intensity. The full colour specification thus reads 1080-Y50R for an orange with 10% blackness, 80% chromatic intensity at yellow with 50% red.

The system allows for a finer subdivision of the colour circle, and this is necessary to define any colour and make direct comparisons with colours defined within the British Standards system. Thus, Magnolia (BS 08 B 15) is 0606-Y41R (6% blackness, 6% chromatic intensity on a yellow with 41% red); Plum (BS 02 C 39) is 5331-R21B and Midnight (BS 20 C 40) is 7415-R82B in the Natural Color System®

RAL COLOUR COLLECTION

The RAL Colour Collection is used significantly within the building industry for defining the colours of finishes, particularly to plastics and metals, but also to materials such as glazed bricks. Typical applied finishes include acrylics, polyesters and polyurethane as well as some paints and lacquers. The RAL system, established in Germany in 1925, has developed through several phases. It commenced with 40 colours, subsequently, many were added and others removed, leaving 170 standard colour shades. Because of its development, the RAL system (designated RAL 840-HR) does not have a systematic order of colours with equal steps between shades. Colours are defined by four digits, the first being the colour class (1 yellow, 2 orange, 3 red, 4 violet, 5 blue, 6 green, 7 grey, 8 brown and 9 black/white) and the further three digits relate only to the sequence in which the colours were filed. An official name is also applied to each standard RAL colour (e.g. RAL 1017 Saffron Yellow, RAL 5010 Gentian Blue, RAL 6003 Olive Green). Some additional colours have been added to the RAL classic collection, giving over 200 colours. The collection for matt shades is designated RAL 840-HR and that for glossy shades is RAL 841-GL. A CD of 195 of the RAL classic colours is available for computer applications giving the colour specifications in RGB (red/green/blue), HLC (hue/lightness/chroma) and offset printing format.

RAL DESIGN SYSTEM

Unlike the RAL Colour Collection which has only a limited selection of standard colours, the RAL Design System has 1624 colours arranged in a colour atlas based on a three-dimensional colour space defined by the coordinates of hue, lightness and chroma. Hue is the attribute of colour, for example red, blue or yellow. Lightness ranges from black to white, and chroma is the saturation or intensity of the colour. The system is equivalent to the HLS (hue, lightness, saturation) system which is used alongside RGB (red, green, blue) in many computer colour systems. The RAL Design System is similar to the Natural Color System®, except that it is based on a mathematical division of the whole visible wavelength spectrum, rather than the visually assessed four standard colours; yellow, red, blue and green.

The colour spectrum is therefore divided into mostly 10° steps around a circle. Each step, illustrated on a page of the associated colour atlas, represents a particular hue. For each hue on the colour atlas page, samples illustrate lightness decreasing from top to bottom and intensity or saturation increasing from the inside to the outside. A colour is therefore coded with the three numbers relating to hue, lightness and chroma, for example 70 75 55. The standard RAL colour collection numbers do not fit neatly into the RAL design system coding but any colour can be defined, thus Saffron Yellow (RAL 1017) becomes 69.9 75.6 56.5. However, as the number defining the hue is not exactly 70, the colour Saffron Yellow will not appear on the atlas page. Computer programs generating colour through the attributes of hue, lightness and chroma can immediately formulate colours according to this system. The electronic version of RAL illustrates the three-dimensional colour atlas, offers 1900 standard colours and links proposed colours to the nearest standard RAL colour. The system was revised in 2007 by the addition of 26 new pastel tones and the removal of some less used darker colours.

COLOUR PALETTE NOTATION SYSTEM

The Colour Palette (Dulux) notation system is based on the three factors: hue, light reflectance value (LRV) and chroma (Fig. 15.4). The hue or colour family is derived from eight divisions of the spectrum, each of which is sub-divided into a further 100 (00–99) divisions to give a precise colour within a particular hue.
Example

**30BB 08/263**

**Hue**

The colour family

**Light Reflectance Value (LRV)**

The lightness or darkness of the colour.

The higher the number the lighter the colour.

**Chroma**

The intensity of the colour.

The higher the number the more intense the colour.

---

*Fig. 15.4 Colour Palette Notation. Images: Courtesy of Imperial Chemical Industries Plc. (Dulux Master Palette and Colour Palette are trademarks of ICI Plc)*
Hue families:

RR  magenta to red
YR  red to orange
YY  orange to yellow to lime
GY  lime to green
GG  green to turquoise
BG  turquoise to blue
BB  blue to violet
RB  violet to magenta

Light reflectance value (LRV) is a measure of lightness or darkness on a 00–99 scale, with light colours having a high two-digit number. Thus, most pastel shades have a light reflectance value between 75 and 83 and the majority of colours fall within the range 04 (very dark) to 83 (very light).

Chroma is a measure of the saturation or strength of the colour measured in 1000 steps from 000 to 999, with high numbers indicating high saturation or intensity.

Thus, a strong bright lemon colour would be specified as:

\[
37YY \ 61/ \ 877 \\
hue \ \ LRV \ \ chroma
\]

(where LRV is light reflectance value)

The Master Palette (Dulux) is based on the same system as the Colour Palette using the three factors: hue, light reflectance value and chroma. The hue or colour family is derived from the same eight divisions of the natural spectrum, with the addition of neutral (NN). Each hue is then sub-divided into a further five sections (10, 30, 50, 70, 90), except YY and YR which have 10 and 7 sections, respectively. These figures represent the percentage movement across the spectrum, for example in RR – from magenta towards red. Thus, hues are described by two digits and two letters (e.g. 50RR). The light reflectance value (LRV) and chroma are the same as in the Colour Palette notation system.

A deep emerald green colour would be described as:

\[
10GG \ 15/ \ 346 \\
hue \ \ LRV \ \ chroma
\]

PANTONE

The Pantone Matching System is frequently used by architects and interior designers for specifying colours for plastics, fabrics and paint. Colours are normally sampled on paper swatches in either coated (C), uncoated (U) or matte (M) finish. Each specific colour is defined by name and a three- or four-digit number and an appropriate finish, for example Pantone 185 C.

Within Pantone there are various sub-sets, including one specifically for paints and interiors. This set is defined by two digits followed by a dash, four digits and a letter suffix. The letters TCX and TPX differentiate between editions printed on cotton or paper, respectively. Each colour also has a name (e.g. Pantone 15-1247 TCX Tangerine).

The Pantone Goe system, a recent development, has 2058 colours arranged chromatically, defined by a three-part hyphenated numbering system. The first number (1–165) defines the colour family, the second number (1–5) defines the page within the family, and the final number defines the position on the page (1–7). A suffix C or U defines coated or uncoated paper (e.g. Pantone 105-5-3 C).

Pantone colours may only be approximately matched on an RGB screen or in standard CMYK printing, but by using hexachrome printing, which includes orange and green, the vibrancy of the Pantone colours can normally be reproduced. The Pantone Formula Guide indicates which colours are achievable by RGB and CMYK screen and printing systems.

VISUAL COMPARISON OF PAINT COLOURS

The European Standard for the comparison of paint colours (BS EN ISO 3668: 2001) is based on observation under specified illumination and viewing conditions (either natural diffuse daylight or scientifically specified artificial light). The colour-matching process is based on an assessment of the differences in hue, chroma and lightness between test panels and reference colour standards.

Cross referencing between colour systems

The NCS system can supply digital cross references to the 237 colours in the British Standard BS 5252, to 189 of the 194 colours in RAL and also to the 1150 Colour Matte Finish and 1500 Glossy Finish Collection colours of the Munsell system. A printed cross reference is available between the NCS system and the European system CMYK (cyan, magenta, yellow, black) used for colour printing. Translation tables from NCS through RGB (computer screen) to CMYK (printing) are available for working with CAD systems on standard operating systems.
PAINTS

The British Standard BS 6150: 2006 gives a detailed description of paint systems and their appropriate applications to the range of building materials used within construction. The standard describes pretreatments for substrates, also the maintenance and deterioration of paint finishes. The choice of a paint system is based on a combination of factors including exposure conditions, function, substrate, constraints, initial cost and maintenance.

COMPONENTS OF PAINTS

Paints consist of a blend of components, each with their specific function. Commonly, these include the binder (or medium), solvent, base, extenders, pigments and driers, although other additives may be incorporated into specialist paints.

The binder solidifies to produce the paint film. Traditionally, the binder was natural linseed oil, which set by gradual oxidation on exposure to air. However, linseed oil has now largely been replaced by alkyd resins which oxidise in air, or vinyl and acrylic resins which solidify by drying. To ensure adequate fluidity of the paint during application by brushing or spraying, either water or organic solvents (hydrocarbons, ketones or esters) are incorporated; paint thinners have the same effect. The base material, usually white titanium dioxide, produces the required opacity, although the body of the paint may be increased by the incorporation of inert extenders such as silica, calcium carbonate, china clay or barytes. Colouring materials are frequently a mixture of organic and inorganic dyes and pigments. Driers which induce the polymerisation of the binder ensure a rapid drying process.

Changes in legislation and environmental concerns have led to the development of paints with reduced levels of volatile organic compounds (VOCs). Mainly this has been through the increased use of water-borne rather than solvent-borne paints. In some respects water-borne paints have the advantage. They have low odour emissions, brushes can be cleaned with water and they tolerate damp surfaces. However, they are not ideal for external use in cold and wet conditions. Other developments have been towards high-solids paints, which have low solvent content and therefore very low VOC emissions.

A further trend is towards the use of natural paints based on plant oils, casein, cellulose and mineral compositions. These are free of the high embodied energy materials such as petrochemical products. Natural paints consume considerably less energy in manufacture, and are environmentally friendly in application and less problematic in waste management; however, they are not suitable for all applications. Typically, casein emulsions, available in a range of colours, are suitable for internal application only.

PAINT SYSTEMS

Coats within a paint system perform specific tasks. Usually, a complete system would require primer, undercoat and finishing coat, although in the case of new external materials, four coats may be appropriate.

Primers

The primer must adhere well to the substrate, offer protection from deterioration or corrosion and provide a good base for the undercoat. To ensure adhesion, the substrate surface must be free of loose or degraded material. Appropriate systems are indicated in Table 15.1. For use on timber, primers may be oils, alkyd resins or acrylic emulsions, frequently with titanium oxide. Aluminium wood primer is recommended for resinous woods and to seal aged creosoted and bitumen-coated surfaces. For the corrosion protection of ferrous metals, primers incorporate zinc-rich compounds within oils or alkyd resins. The use of lead-based paints in the UK is subject to the Control of Lead at Work Regulations, 2002. The newly developed low-VOC coatings offer temporary protection against the corrosion of structural steelwork as either pre-fabrication or post-fabrication primers. Alternatively, acrylated rubber paints which form a physical barrier over steel may be used as primers. For non-ferrous metals, zinc phosphate primers are frequently used. The application of primers suitable to ferrous metals may cause increased corrosion on non-ferrous substrates, particularly aluminium. Masonry paints are usually based on alkyd or acrylic resins with titanium oxide; where surfaces are likely to be alkaline, such as new plaster, brickwork or concrete, alkali-resisting primer should be used.

Undercoats

Undercoats provide cover and a good base for the finishing coat. Most undercoats are based on alkyd resins or acrylic emulsions.
Table 15.1  Recommended primers for various substrates

<table>
<thead>
<tr>
<th>Primer</th>
<th>Suitable substrates and conditions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Timber</td>
<td></td>
</tr>
<tr>
<td>Alkyd primer (solvent-borne)</td>
<td>Softwoods and hardwoods, interior and exterior</td>
</tr>
<tr>
<td>Acrylic primer (water-borne)</td>
<td>Softwoods and hardwoods</td>
</tr>
<tr>
<td>Aluminium wood primer</td>
<td>Resinous softwoods and hardwoods</td>
</tr>
<tr>
<td>Preservative primer</td>
<td>Exterior use, contains fungicides</td>
</tr>
<tr>
<td>Plaster and masonry</td>
<td></td>
</tr>
<tr>
<td>Acrylic primer sealer</td>
<td>Loose, friable surfaces</td>
</tr>
<tr>
<td>Alkali-resisting primer</td>
<td>Plaster, cement and concrete</td>
</tr>
<tr>
<td>Ferrous metals</td>
<td></td>
</tr>
<tr>
<td>Pre-treatments</td>
<td>Improve adhesion of paint systems</td>
</tr>
<tr>
<td>Zinc phosphate</td>
<td>Steel, iron, galvanised steel. Good rust inhibitor</td>
</tr>
<tr>
<td>Zinc-rich primer</td>
<td>Steel. Two-component system. Must be a blast-cleaned surface</td>
</tr>
<tr>
<td>Chromate primer</td>
<td>Ferrous and non-ferrous metals</td>
</tr>
<tr>
<td>Epoxy primer</td>
<td>Steel. Two-component system containing aluminium powder.</td>
</tr>
<tr>
<td>Acrylated rubber</td>
<td>Steel, iron, galvanised steel. Must be a full acrylated rubber system</td>
</tr>
<tr>
<td>Non-ferrous metals</td>
<td></td>
</tr>
<tr>
<td>Zinc phosphate</td>
<td>Aluminium</td>
</tr>
<tr>
<td>Acrylated rubber</td>
<td>Aluminium. Must be a full acrylated rubber system</td>
</tr>
<tr>
<td>Acrylic metal primer</td>
<td>Aluminium, copper, lead, brass. Quick-drying water-based primer</td>
</tr>
</tbody>
</table>

**Finishing coats**

Finishing coats provide a durable and decorative surface. Some gloss, eggshell and satin finishes are still based on oils and alkyd resins, although water-borne products are increasingly becoming predominant. Some water-borne gloss finishes tend to be visually softer and are more moisture permeable than the traditional solvent-borne hard glosses. However, they have the advantage of quick drying without the evolution of solvent odour; generally they are more durable and do not yellow on ageing. Matt and silk finishes are usually vinyl or acrylic emulsions.

**Special paints**

**LIGHT REFLECTING WHITE PAINT**

White paints with lower light absorbency and greater reflectivity can enhance the distribution of light within room enclosures, thus reducing the energy input required to achieve appropriate lighting levels. Products are available in matt and eggshell finishes.

**MULTICOLOUR PAINTS**

Multicolour paints incorporating flecks give a hard-wearing surface which may be glazed over to ease the removal of graffiti. Application is with a spray gun, which can be adjusted to change the pattern and texture of the fleck. This type of paint system may be applied to most dust- and grease-free internal surfaces.

**Broken-colour paints**

Broken-colour effects, reflecting the traditional processes of graining, marbling, ragging and stippling, are once again popular. Most modern broken-colour effects require a base coat, applied by brush or roller, which is then overpainted, with a clear coloured glaze. The glaze is then patterned or distressed to create the desired effect. One proprietary system uses a special rag-roller, which flails the wet finish coat giving random partial exposure of the darker first coat. Alternative finishes include metallic, pearlescent and graining effects. An iridescent finish produces a two-tone shimmer effect by optical interference of the reflected light. Water-based acrylic glazes are virtually odour-free and are touch dry within two hours.

**ACRYLATED RUBBER PAINTS**

Acrylated rubber paints are suitable for internal and external applications exposed to chemical attack or wet and humid atmospheric conditions. Acrylated rubber paints are tending to replace chlorinated rubber coatings which rely on carbon tetrachloride solvent, now considered environmentally damaging. Acrylated rubber paints may be applied to metal or masonry by either brushing or spraying. Usually, a film of dry thickness 100 μm is applied compared to 25–30 μm for most standard paint products.

**HEAT-RESISTING PAINT**

Aluminium paint, which has a lustrous metallic finish, is resistant to temperatures up to 230–260°C. A dry film thickness of 15 μm is typical. Acrylated rubber paints can usually be used satisfactorily to 100°C.
FLAME-RETARDANT PAINTS

Flame-retardant paints emit non-combustible gases when subject to fire, the usual active ingredient being antimony oxide. Combustible substrates such as plywood and particleboard can be raised to Class 1 (BS 476 Part 7) surface spread of flame. Products include matt, semi-gloss and gloss finishes, and may be applied by brush, roller or spray.

INTUMESCENT COATINGS

Thin-film intumescent coatings, typically 1 or 2 mm in thickness, offer fire protection to structural steel without noticeable visual effect. In the event of fire, the thin coating expands up to 50 times to form a layer of insulating foam. The carbonaceous material in the coating, typically starch, is charred, whilst the heat also causes the release of acids. These act to produce large volumes of non-inflammable gases which blow up the charring starch within the softened binder into an insulating cellular carbon layer. Coatings may be applied to give 30, 60 or 120 minutes’ fire protection. Intumescent emulsion paints or clear varnishes are appropriate for use on timber, although, where timber has been factory impregnated with a flame-retardant salt, the compatibility of the intumescent coating and flame retardant must be verified.

FUNGICIDE PAINTS

Fungicide paints, for application in areas where mould growth is a recurrent problem, usually contain a blend of fungicides to give high initial activity and a steady long-term performance. The latter can be achieved with fungicide constituents of low solubility which are gradually released to the surface during the lifetime of the paint. Matt acrylic finishes are available in a range of colours.

ENAMEL PAINTS

Enamel paints based on polyurethane or alkyd resins give highly durable, impact-resistant, easily cleaned, hard-gloss surfaces. Colours tend to be strong and bright, suitable for machinery and plant in interior and exterior locations.

MICACEOUS IRON OXIDE PAINTS

Micaceous iron oxide paints have good resistance to moisture on structural steelwork, iron railings, etc. due to the mica plates which reduce permeability to moisture vapour. A dry film thickness of 45–50 μm is typical, thus requiring longer drying times than standard paint products. Micaceous iron oxide paints should be applied over an appropriate metal primer.

MASONRY PAINTS

Smooth and sand-textured masonry paints are suitable for application to exterior walls of brick, block, concrete, stone or renderings. Where fine cracks are present, these can often be hidden using the sand-textured material. Usually, masonry paints contain fungicides to prevent discolouration by moulds and algae. Acrylic resin-based products are predominantly water-based; however, fast-drying solvent-based systems are also produced. Mineral silicate paints form a crystalline protective layer over the masonry surface, which tends to be more durable than the organic finishes from synthetic resins.

WATER-REPELLENT AND WATERPROOF PAINTS

Silicone water-repellent paints can be applied to porous surfaces including brick, concrete, stone and renderings to prevent damp penetration. Such treatment does not prevent rising damp, but will allow the continued evaporation of moisture within the masonry. Two-pack epoxy waterproofing systems may be applied to sound masonry surfaces to provide an impervious coating. Typical applications are to rooms where condensation causes the blistering of normal paint films; also, in basements and solid external walls where penetrating water is a problem, provided that a good bond can be achieved between substrate and epoxy resin. Bituminous paints provide a waterproof finish to metals and masonry, also they may be used as a top dressing to asphalt or for renovating bitumen membrane roofing. Aqueous bitumen coatings, if fully protected against physical damage, can provide a vertical membrane where the external ground level is higher than the internal floor level.

EPOXY PAINTS

Epoxy ester paint coatings are highly resistant to abrasion and spillages of oils, detergents or dilute aqueous chemicals. They are therefore frequently used as finishes to concrete, stone, metal or wood in heavily trafficked workshops and factories. Many are produced
as two-pack systems requiring mixing immediately before application.

**ANTI-GRAFFITI PAINTS**

The main purpose of anti-graffiti paints is to aid easy removal of graffiti. Paints may be sacrificial or permanent. The sacrificial paints, usually based on wax emulsions, acrylates or biopolymers, are easily removed by jets of hot water. The permanent anti-graffiti paints based on tough fluoro-acrylics or two-component polyurethane systems are resilient to chemical and abrasive cleaning techniques. The permanent anti-graffiti paints are more successful on smooth surfaces. Multi-colour finishes help to disguise any discolouration.

**ANTI-MICROBIAL COATINGS**

Anti-microbial wall and ceiling coatings, incorporating silver ions, can reduce bacteria levels within hospitals and hygienic environments.

**Natural wood finishes**

Natural wood finishes include wood stains, varnishes and oils. Wood stains are pigmented resin solutions which penetrate into the surface and may then build up a sheen finish. Varnishes are unpigmented resin solutions which are intended to create a surface film. Timber preservatives are described in Chapter 4.

**WOOD STAINS**

Most wood stain systems for exterior use include a water- or solvent-based preservative basecoat which controls rot and mould growth. Typical formulations include zinc or copper naphthenate, dichlorofluanid, tri-(hexylene glycol) bborate and disodium octaborate tetrahydrate. Wood stain finishes are either low-, medium- or high-build systems, according to the particular application. They usually contain iron oxide pigments to absorb the ultraviolet light which otherwise causes the surface degradation of unprotected timber. Generally, for rough-sawn timber, deeply penetrating wood stains are appropriate, whereas for smooth-planed timber a medium or high-build system gives the best protection from weathering. Products are based on acrylic and/or alkyd resins.

For sawn timber, both organic solvent-based and water-based materials are available, usually in a limited range of colours. Solvent-based low-build products, which are low in solids, penetrate deeply, leaving a water-repellent matt finish, enhance the natural timber grain and are suitable for timber cladding. Deep penetration should eliminate the risk of flaking or blistering on the surface. Medium- and high-build products for exterior joinery offer the choice of semi-transparency to allow the grain to be partially visible, or opaque colours for uniformity. Products are available in a wide range of colours with matt or gloss finishes. The first coat both penetrates and adheres to the surface, whilst the second coat provides a continuous microporous film which is both permeable to moisture vapour and water repellent, thus reducing the moisture movement of the timber. Additional coats should be applied to end grain. The coating, typically 30–40 µm thick, should remain sufficiently flexible to accept natural timber movements. Low-VOC products based on water-borne emulsions or high-solids solvent-borne resins are now generally available.

**VARNISHES**

Traditional varnishes are combinations of resins and drying oils, but most products are now based on modified alkyd resins. Polyurethane varnishes are available in matt, satin or gloss finishes, based on either water- or solvent-based systems. The solvent-based systems produce the harder and more durable coatings up to 80 µm thick, suitable for exterior woodwork. Products either retain the natural wood colour, enhance it, or add colour. Screening agents to protect timber from the effects of ultraviolet light are normally included in the formulations. Urethane-modified alkyd resins are suitable for interior use, and have the advantage of high resistance to scuffing and hot liquids. External weathering causes eventual failure by flaking and peeling as light passing through the varnish gradually degrades the underlying wood surface. For example, hardwood doors decorated with polyurethane varnish, protected from rain and direct sunlight by a porch, should have extended periods between maintenance. End grain should be sealed to prevent trapped moisture encouraging the development of staining fungi.

Wood finishes entirely composed of natural products are also available. These are based on blends of beeswax, larch resin, vegetable oils, and may incorporate minerals and earth pigments for colour and opacity.
OILS

Oils such as teak oil are used mainly for internal applications. Formulations based on natural oils for exterior use are high in solids, producing an ultraviolet-resistant, microporous finish which may be transparent or opaque. The finish, which should not flake or crack, may be renovated by the application of a further coat.

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ENERGY-SAVING MATERIALS AND COMPONENTS

Introduction
The trend towards increasingly energy-conscious design has resulted in a greater focus on energy-saving materials and components. These include photovoltaics and solar collectors, which turn the sun’s energy into electricity and hot water, respectively. Light tubes and wind catchers are energy-saving devices which can make modest reductions in the energy consumption of buildings in the context of a holistic energy-efficient strategy. Micro-wind turbines, within the appropriate context, can generate modest quantities of electricity. Rainwater harvesting systems can eliminate or reduce mains potable water consumption for certain less critical domestic uses.

Photovoltaics
Photovoltaics (PVs) are silicon-based devices which, under sunlight, generate a low-voltage direct electric current. The quantity of electricity produced is directly related to the intensity of incident solar radiation or irradiance (W/m²). Both direct and diffuse sunlight are effective, although the intensity of direct sunlight is typically 10-fold that of an overcast sky, and the efficiency of energy conversion is around 15%. Photovoltaic cells are connected in series to generate a higher voltage. The supply is then passed through an inverter to convert the direct current into more usable alternating current at the standard voltage. The electricity generated may then be used within the building or sold back into the national supply if generation exceeds the demand.

Photovoltaic units are manufactured from a sandwich of at least two variants of mono- or polycrystalline silicon (Fig. 16.1). These n- and p-type (negative and positive) silicon crystals generate electricity at their interface under solar (photon) radiation. Cells are arranged in rectangular modules ranging from 0.3 to 1.5 m². A typical unit of one square metre could produce 150 W of electrical power under bright sunshine of 1000 W/m².

The cells are usually laminated with a protective layer of glass, backed with metal sheeting and mounted on a steel frame. However, translucent systems built into glass double-glazing units or flexible units faced with a plastic cover are available. Mono- or polycrystalline silicon modules are usually blue or black in colour. The mono-crystalline modules are a uniform colour, whereas the poly-crystalline units have a sparkling surface. Other colours can be achieved, but with a reduced level of efficiency.

The alternative amorphous thin-film silicon (TFS) cell modules are matt red, orange, yellow, green, blue or black in colour, and can be laminated into glass or mounted on a flexible plastic backing. These systems have significantly lower levels of efficiency than the crystalline cells in good light conditions, but are more efficient in poor light conditions. Thin-film amorphous silicone photovoltaics are much cheaper to produce than the crystalline systems which ameliorates their low efficiency at 8%. Hybrid units combining mono-crystalline and thin-film technologies give good output over the range of light conditions. It is
anticipated that thin-film silicon photovoltaics will increase their share of the market from 14% in 2008 to 30% by 2015.

Recent developments include the production of cadmium telluride (CdTe) photovoltaics with an efficiency of 10%. The newer materials including copper indium selenide (CIS) and copper indium gallium selenide (CIGS) can produce 11% energy conversion. Multi-layer thin composites can achieve 20% conversion rates and the expensive gallium arsenide (GaAs) multi-function cells can achieve 40% efficiency, by absorbing the majority of the solar spectrum including infrared and ultraviolet light.

More total energy can be obtained by tracking the sun to optimise the orientation of the photovoltaic cells or by using mirrors to increase the solar radiation received. However, these systems involving moving components are more expensive and will require maintenance.

Photovoltaic systems are normally supplied as panels, but PV cladding, slates and glazing systems are also available. The location and tilt angle of PV installations are critical in respect of maximising output. The maximum efficiency in the northern hemisphere is gained from a south orientation with a tilt from the horizontal equal to the geographic latitude minus 20° (Fig. 16.2). Thus, for London, at a latitude of 51°, the optimum tilt is 31° from the horizontal. However, in urban areas, the effects of the immediate environment must be taken into account when assessing the available solar energy. Shadowing and inter-reflection from adjacent buildings, together with local regular weather patterns, affect the total annual solar energy, which ultimately determines the electrical output.

Photovoltaic units, particularly those manufactured from crystalline silicon, decrease in performance with increased temperature, and any roof, curtain walling or rainscreen cladding system should be naturally ventilated to maintain efficiency. The use of PVs should be an integral part of the energy strategy for the building.

A 1 m² crystalline photovoltaic unit operating efficiently will generate about 100 kWh per year. The pay-back time for such a unit would be of the order of 10 years including the energy of manufacture, and most installations will last between 20 and 30 years. One appropriately sized system offers a maximum 1.2 kW output, giving 1000 kWh per year, roughly one fifth of the average UK annual domestic consumption.

PHOTOVOLTAIC ROOFING AND CLADDING MATERIALS

Photovoltaic slates (Fig. 16.3) and tiles, which have the general appearance of fibre-cement slates and shiny plain tiles, respectively, can be used on suitably orientated roofs as an ecological alternative to standard roofing, subject to appropriate planning consent. The lower part of the individual slates comprise
photovoltaic cells which are connected into a standard photovoltaic system. The tiles are 2.1 m strips of material, marked in units to resemble conventional roof tiles and overlapped to give the required visual effect. To obtain 1 kW of power under optimum conditions, 10 m² of slate or 16 m² of tiling is required depending on the lap. Photovoltaic roof tiles can be coordinated with matching solar panel tiles for hot water systems, giving a fully integrated appearance to a tile roof.

High-performance roofing membranes are available incorporating laminated flexible amorphous thin-film silicone cells. Rainscreen cladding, curtain walling and sunshade louvres incorporating photovoltaic systems may be more appropriate for tall buildings with relatively small roof areas. Thin-film photovoltaic modules
can be produced to sizes up to $2.2 \times 2.6\,\text{m}$ for facade and roofing applications.

**PHOTOVOLTAIC GLAZING**

Amorphous silicon photovoltaic cells integrated into the glass of laminated or double-glazing systems can be a source of electrical energy. The glazing may be semi-transparent retaining visual contact with the exterior or opaque in various colours. The electrical connections are made at the perimeter of the units within the frame system. Cell spacing can be tuned to give the optimum balance between electricity generation and daylight transmission.

**Solar collectors**

The two standard types of solar collectors are the flat-plate and the evacuated-tube systems. Flat-plate collectors consist of a metal heat-absorbing plate, closely bonded to copper water pipes which transport the heated water to a storage system. The maximum efficiency is achieved using a low-emissivity matt black absorbing plate, which limits the loss of energy through re-radiation from the hot surface. A low iron-content double-glazed cover which admits the maximum quantity of short-wave energy protects the absorbing plate and retains the entrapped heat. The underside of the pipework is insulated with fibreglass or polyisocyanurate foam to prevent heat loss to the aluminium casing and the underlying roof structure or support system.

Evacuated-tube collectors (Fig. 16.4) consist of a double-layer glass tube, with a vacuum between the two layers. The outer glass is clear, admitting light and heat with minimal reflection. The inner tube is coated to absorb the maximum quantity of radiation. The heat from the inner tube is transferred in a sealed unit vapourising and condensing system to a heat exchanger within the main the liquid flow which is circulating to the heat storage system. Evacuated-tube collectors are substantially more expensive than flat-plate collectors, but are more efficient if angled correctly and will produce higher temperatures.

Flat-plate solar collectors may be located in any unshaded location, at ground level or attached to buildings. The best orientation is directly towards the mid-day sun, but a variation of up to $15^\circ$ east or west will have little adverse effect. The optimum tilt from horizontal for solar hot water collectors for maximum all-year-round efficiency equals the location’s latitude. However, for increased winter efficiency, when solar gains are at a premium, the tilt from horizontal should be increased by $10^\circ$, to pick up more energy at lower sun altitudes. Solar hot water systems are heavy, and must be fixed securely to suitable substrates. On tiled- or slated-pitched roofs an air gap should allow for the clear passage of rainwater and melting snow.

Solar collector systems with the appearance of standard plain concrete roofing tiles can be integrated into a tiled roof without significant visual effect. Furthermore, solar collector tiles can be matched to photovoltaic tiles giving a fully integrated appearance to a solar-responsive tile roof.

Hot water from the solar collector is usually circulated through an indirect system to a solar storage tank (Fig. 16.5). This acts as a heat store of preheated water to be fed into a standard hot water cylinder system, where the temperature can then be boosted from a boiler to the required level. Circulation may be through either a gravity thermosyphon system operated by hot water convection with the storage tank located above the collector, or a pumped system, in which case the tank may be below the collector. The circulated water must contain antifreeze and a rust inhibitor. An alternative direct system feeds tap water directly into the solar collector, but scaling and corrosion of the pipework can be problematic. A $5\,\text{m}^2$ solar collector panel will heat $250\,\text{l}$ of hot water per day, which is a typical four-person family demand.

Solar-energy district-heating plants in Europe, including Scandinavia, contribute significantly to a direct reduction in the energy requirements for
small-town domestic hot-water systems. Water is preheated by large arrays of solar collectors before the local conventional-fuel heating system tops up the temperature to the required domestic level. Furthermore, solar heating systems, in conjunction with large underground heat storage tanks, can significantly reduce winter energy consumption by preheating the water supplies during periods when direct solar gain is ineffective.

**Solar air heating systems**

Solar air heating systems applicable to commercial buildings consist of steel wall or roof panels which absorb solar energy and transfer the heat to internal air channels created by the sheeting profile. Convection currents take the warmed air into the building as required, supplementing the main heating system.

**Light pipes**

Light pipes or tubes transmit direct sunlight and natural daylight from roof level into the building space below (Fig. 16.6). At roof level an acrylic self-cleaning dome admits light into a highly reflective pipe which transmits it down to a white translucent dome at ceiling level where the light is diffused into the space below. The mirror-finish aluminium tube can be of any length including offsets, although the quantity of light transmitted is typically reduced by 3% for each metre length and by 8% for each bend.

Standard light pipe diameters range from 200 to 600 mm, although larger sizes up to 1500 mm are available for commercial applications. The systems should be free of condensation and not cause winter heat loss or summer solar gain to the building enclosure. Rectangular units, similar in appearance to standard or flush fitting conservation rooflights, are also available,
and the ceiling unit can be a square diffuser to integrate into suspended ceiling systems.

A 330 mm diameter system will typically deliver between 100 W from a winter overcast sky and 400 W under full summer sun, for a straight tube not exceeding 3 m in length. Such systems can offer energy-saving solutions to existing buildings, and may be considered as one element within a fully integrated lighting strategy for new-build.

A more sophisticated system combines the functions of both a light pipe and a wind catcher to admit natural daylight and ventilation into internal spaces poorly served by normal external glazing.

**SUNLIGHT TRANSPORTING SYSTEMS**

Sunlight received on the roof by collecting panels can be transported to any internal location within 20 m, using 30 mm diameter fibre optic cable bundles terminated by light-emitting luminaires. The output is a combination of diffuse and parallel light.

**Heat pump systems**

Heat pumps operate by withdrawing low-temperature heat from a source and rejecting it at a higher temperature. The standard process uses a refrigeration cycle driven by an electric motor. The refrigerant is compressed and releases heat in a condenser, and then it subsequently expands within the cycle, extracting more heat in the evaporator before being re-compressed. The key environmental aspect of heat-pump systems is that more useful heat is generated from the electricity used in the compressor pump, compared to the fuel consumed by traditional gas or electric heating systems. A typical system may produce 4 kW of useful heat for each 1 kW of electricity consumed.

The standard heat source is the ground, but alternatively exhaust air or groundwater flow may be used. Ground air and water in the UK are usually at around 12°C and exhaust air is typically at 21°C. For ground-source systems the size of the installation depends...
upon the anticipated load. The ground loop may be a horizontal array of pipes to a minimum depth of 0.6–1.0 m or a set of vertical bore heat exchangers. Soil conditions significantly affect the efficiency of ground-source systems. Specifically, dry sandy soils with low thermal conductivity are unsuitable for horizontal systems.

Small-scale heat pumps are usually used for space heating alone, as the maximum temperature attainable for domestic hot water from conventional refrigeration systems is 48–50°C. Many domestic installations are linked to underfloor heating systems. Large units for non-domestic applications can produce outputs up to 130 kW and water temperatures up to 70°C. The design of heat pump heating systems is described in BS EN 15450: 2007.

**Groundwater cooling systems**

Groundwater cooling systems offer an alternative to energy-intensive air-conditioning systems in larger buildings. Groundwater in the UK is typically at 12°C, although in London 14°C is common. Subject to official approvals, groundwater may be abstracted for use in cooling systems, provided that appropriate provision is made for its disposal. This may be to lakes, rivers and surface drainage systems, or returned to aquifers by injection into the ground. The system used for Portcullis House, Westminster, uses two 150 m boreholes in the chalk aquifers. These extract water at 14°C, use it within heat exchangers and then discharge it at 21°C. Some greywater is used for non-potable functions.

**Wind catchers**

Wind catchers have been standard architectural features on the roofs of buildings in hot-dry climates for centuries. However, in order to reduce energy costs associated with air-conditioning systems, this additional source of natural ventilation can now be designed into larger temperate-climate buildings to supplement other natural ventilation systems.

A wind catcher operates by capturing the air on the windward side of the shaft and deflecting it down one quadrant by a series of vanes. The force of the wind drives it into the space below. As the entering air is cooler and denser than that within the building, it displaces the warm vitiated air which rises by natural stack ventilation through the other quadrants of the shaft, leaving through the leeward side of the wind catcher. With a symmetrical system one quadrant will predominantly face the prevailing wind to act as the catcher and the opposite quadrant will provide the majority of the stack-ventilation effect. A glazed top to the wind catcher, which heats up further the vitiated air, can enhance the stack effect. Dampers can be used to reduce air flow during winter months and permit night-time cooling only in hot summer months. Wind catchers should be located near to the ridge on pitched roofs to maximise their efficiency. Solar-powered wind catchers incorporating a photovoltaic panel (Fig. 16.7) can automatically switch on a low-energy extractor fan to increase natural ventilation in sunny conditions.

For functioning in multi-storey buildings, wind catchers require appropriate ducting and damper systems, and may incorporate heat exchangers from the central heating system to admit tempered fresh air in winter operation.

**Wind turbine systems**

Micro-wind turbines can generate up to 1.5 kW of electrical power depending on the wind turbine diameter and local wind speed conditions. A 1.75 m diameter turbine operating at its rated wind speed of 12.5 m/s will generate 1 kW, with a minimum operating wind speed of approximately 4 m/s. The dc current generated is converted to standard voltage at 50 Hz ac for use before additional current is taken from the national supply. Micro-wind turbines are only intended to supplement the user’s standard mains electricity supply. Planning issues relate to visual impact and noise production.

The exact location of micro-turbines, for example on house roofs, is important as the power in the wind is proportional to the cube of wind speed. Urban wind speeds are significantly affected by the immediate topography. Recent research indicates that the best location for domestic micro-wind turbines is above the ridge at a gable end. Local wind speed data should be obtained from a local meteorological station or the NAOBL wind speed database. Certain environments in large urban conurbations are unsuitable for micro-wind turbines due to the low average wind speeds. This would mean that the environmental cost of the installation, its maintenance, ancillary equipment and decommissioning would never be returned during the active life of the micro-turbine.
Water management systems

RAINWATER HARVESTING SYSTEMS

Rainwater harvesting systems allow for the collection, filtering and storage of roof rainwater for use in non-potable functions. The storage capacity depends on the roof area and the anticipated demand. In certain systems, when the storage tank level is insufficient, the mains water supply automatically switches in.

Three types of rainwater harvesting systems are described in BS 8515: 2009. Water from the storage tanks may be delivered by gravity or self-activating pump directly to the points of use; alternatively, the stored water is pumped to an elevated cistern which then gravity feeds the points of use. Filtration systems require routine annual maintenance and should retain particles of size $>1.25 \text{ mm}$. Rainwater is suitable for flushing toilets, laundry operations, car washing, garden watering and other non-potable functions. Rainwater outlets must be clearly marked as non-potable.

Guidance on calculating the required storage capacity is given in BS 8515: 2009. For the approximate...
calculation of the required storage capacity, a domestic consumption of 50 l/day/person is appropriate. To anticipate fluctuations in supply, storage capacity should equal 5% of the annual demand or 5% of the average annual rainwater yield, whichever is the lesser. Suitable provision is required for overflow from the storage tank.

**GREYWATER RECYCLING SYSTEMS**

Greywater from domestic showers and baths can be recycled to flush toilets. One electronically controlled system collects the greywater from showers and baths and, without the addition of chemical agents, skims off the floating material, allows the settlement of sand residues and retains up to 100 l of greywater. This is sufficient for approximately 20 flushes of the directly connected WCs. The system automatically purges itself as appropriate and is topped up with fresh water when necessary. The system conforms to the strict water regulations which prevent contamination of potable water supplies. The system can save up to 30% of normal domestic potable water use, and in new housing contributes to the Code for Sustainable Homes which requires reduced water consumption.

**Code for Sustainable Homes water specifications:**

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<tr>
<td>≤105</td>
<td>Levels 3 and 4</td>
</tr>
<tr>
<td>≤80</td>
<td>Levels 5 and 6</td>
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</table>

**Phase change systems**

A small unit filled with phase change material (PCM) can act as a large thermal mass by drawing in hot air and cooling it by absorption of the latent heat of fusion of the phase change material during excessively hot periods. The process is reversed as the ambient temperature falls when heat is released by the solidification of the phase change material. Passive cooling systems based on PCM technology can be located on outside walls to draw in either fresh or recycled air. Phase change materials are described in Chapter 12 Plaster (page 337).

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Introduction

Realisation of the finite nature of global resources and the greenhouse effect of ever increasing carbon dioxide emissions has promoted consideration of the potential for recycling into construction of many mass-produced waste products which are currently either burned or buried in landfill sites. Such materials include plastics, cardboard, straw, paper and tyres. Whilst some recycled products are only in the experimental stage, others are now becoming recognised as standard building materials. The re-use of building materials is well illustrated by the former Earth Centre, Doncaster, which is constructed using many recycled and reclaimed products including crushed concrete, telegraph poles, glass and radiators (Fig. 17.1).

Straw bales

Straw bales, a by-product of the mechanical harvesting and threshing of grain, are produced in large quantities in mechanised agricultural countries. The traditional rectangular bales, which are cheap and can be manhandled individually, are appropriate for building. The large cylindrical and very large rectangular bales, which require mechanical lifting, are less useful in construction and are not considered here. Standard bales (typically $330 \times 530 \times 1050$ mm) are produced within the baler by compressing quantities of straw into flakes about 100 mm thick. These layers are built up along the length of the bale, which is then automatically tied, usually with two polypropylene strings. There is inevitably some variation in length, and the ends are slightly rounded. For construction, the bales should be well compressed in manufacture, dry (maximum 20% moisture) to prevent the growth of moulds and fungi and with the minimum amount of remaining grain, which might attract rodents.

In building construction, bales are stacked, large faces down, making the orientation of the straw fibres predominantly horizontal. At ground level, straw bales must be protected from rising damp and from any risk of saturation from surface water. Additionally, steel mesh protection from rodents is necessary. Adjacent bales must be firmly packed together to ensure stability and to reduce settling under load both during and after construction. Bales are normally secured with metal spikes or hazel rods from coppiced timber and may be sprayed with insecticide for added protection. Externally, lime/clay or lime render on wire mesh is appropriate as it is flexible, self-healing, and will breathe to prevent the build-up of trapped moisture. Alternatively, a rainscreen, separated from the external face of the bales, may be used. Internally, straw bales are usually finished with gypsum plaster on wire mesh. Openings in straw bale construction may be formed with timber framing, but careful detailing is required to prevent water penetration at these locations. Roofs are normally set onto a timber wall plate fixed through the top bales for stability. A two-storey load-bearing wall of straw bale construction has been built within the UK with a $U$-value of 0.13 W/m² K.

An alternative approach to using load-bearing straw bales is timber or steel-frame construction with straw
bales as the insulating infill (Fig. 17.2). Whilst fire is a risk during straw bale construction, the non-combustible internal and external finishes and the compact nature of the straw make the completed construction resistant to fire. (The thermal conductivity of baled straw is approximately 0.050 W/m K.)

Cardboard

The cardboard classroom at Westborough School, Essex (Fig. 17.3), illustrates the potential of this largely recycled product as a useful construction material. A combination of flat composite panels and tubes forms the structure of this building which has an estimated life of 20 years.

In the recycling process, waste paper and cardboard are broken down and converted to pulp, which is a suspension of cellulose fibres in water. The pulp flows onto a conveyor belt, where it is drained of the excess water and compressed, causing the fibres to felt together, producing a long roll of paper. Flat cardboard sheets are formed by gluing together successive layers of paper. Tubes are manufactured from multiple layers of spirally wound paper plies, starting on a steel tube former of the appropriate size, the adhesive being starch or PVA glue. The first and last layers of paper can be of a different quality, for example, impregnated or coloured to create the required surface finish. For the Westborough School building the flat sections are composite panels consisting of multiple cardboard sheets and honeycomb cardboard interlayers, surrounded by a timber frame to facilitate ease of fixing between units. Adjacent roof and walls panels are articulated to ensure overall structural rigidity.

PROPERTIES

Cardboard, like timber, is combustible and can be treated to improve its fire performance, particularly in relation to the surface spread of flame test. However, some fire-retardant materials are environmentally unfriendly and should be avoided if the material is to be subsequently recycled.

The structural strength of cardboard is seriously affected by water. Cardboard, even if specially treated in its manufacture, is a hygroscopic material which will readily absorb moisture. It is therefore necessary to protect it from warm moist air within the building using an impermeable membrane, and externally from rain using a breathing membrane, the latter preventing the build-up of trapped interstitial moisture. In the Westborough School building, the inner plastic membrane is protected from physical damage with a further 1 mm cardboard layer, and the external membrane is covered by cement-bonded fibreboard for fire and rain protection.

Cardboard is potentially vulnerable to rot and insect attack. This could be prevented by treatment with boron products; however, this would adversely affect the potential for ultimate recycling of the cardboard. As cardboard is a recycled material it has a low embodied energy, and can legitimately be considered green.

Rammed-earth and cob construction

Earth construction is one of the oldest forms of building used by mankind. Rammed-earth buildings can be found in most countries, and many have survived...
Fig. 17.2 Straw bales in construction. *Architects and Photograph*: Sarah Wigglesworth Architects

Fig. 17.3 Cardboard classroom — Westborough School, Essex. *Engineers*: Buro Happold. *Architects*: Cottrell + Vermeulen Architecture. *Photograph*: Copyright Adam Wilson/Buro Happold
and many old cob buildings still exist in Devon. The construction was common in many parts of the UK achieved without the use of shuttering. This type of mum 450 mm stone plinth. Free-form designs may be tamped together, usually commencing from a mini-

duction process, the mixture of the straw and clay is cause cracking of the completed structure. In the con-

tion. In order to introduce sufficient light, at least one elevation is usually glazed, and in addition, interesting effects can be achieved with roof lights or light pipes. Ventilation may be mechanical but is normally provided through opening glazing. At the same time, unwanted cold air infiltration is eliminated by the earth enclosure.

The Hockerton Housing Project at Southwell, Not-

hundreds of years. The ideal material is a well-graded mixture of gravel, sand, silt and clay fines. The clay con-

tent should be sufficient to act as an efficient binder, but not in excess to cause large moisture movement or cracking of the finished construction. In modern rammed-earth construction Portland cement is fre-

Earth-sheltered buildings

Earth-sheltered buildings, including homes, are defined as those where the roof and some sides are covered by earth. Increased depth of earth cover improves the thermal performance, but this has to be balanced against consequential increased structural strength requirements. Typically, 400–450 mm of earth cover is appropriate, and the weight of this material is usually supported by concrete or masonry construction. Water exclusion is key to the design, requiring land drainage and the use of reinforced membranes.

One method of construction uses fibrous plaster shells to create the internal organic form. These are sprayed with a layer of lightweight aggregate concrete insulation, followed by 100 mm of structural concrete. After the concrete is structurally sound, the building may be fitted out and covered externally with soil and grass. An alternative construction system uses extruded polystyrene insulation between the structural concrete and the soil backfill. Either approach uses the temperature stabilising effect of the mass concrete and soil cover to significantly reduce energy consumption. In order to introduce sufficient light, at least one elevation should be of lime plaster rather than a Portland cement render, which cracks or breaks away allowing rainwater to penetrate the wall. As with rammed-earth construction, the eaves should be deep enough to protect the walls from severe weathering.

The ‘House for Stories’ designed by Tono Mirai at the Bleddfa Centre for the Arts in Powys, Wales, is constructed from mud and straw. The building, which is intended to be a quiet space for contemplation and imagination, is partly subterranean, but spirals out of the ground into the delightful helical form shown in Fig. 17.4.
residents generate their own energy, harvest their own water, and recycle waste materials eliminating pollution and carbon dioxide emissions. Only the south elevation of the development overlooking the reed pond is visible, as grass covers the majority of the construction.

**Clay products**

The concern over the intrinsic energy in most manufactured building products has led to the further development of a range of clay-based products including clay boards and plasters. Unfired clay building products are hygroscopic and have the positive effect of controlling internal environments by absorbing odours and stabilising humidity and temperature.

**CLAY BOARDS**

Clay boards, which are an alternative to gypsum plasterboards, are manufactured from clay and layers of reed along and across the board length. Hessian on both faces acts as reinforcement and a key for a 2–3 mm finish of lime-based or earth plaster. Joints should be scrimmed before skimming, although clay boards if sealed can be painted directly. The 25 mm boards may be used for drywall construction and ceilings, where they should be screw-fixed at 600 or 400 mm centres, respectively.

**CLAY PLASTERS**

Clay plasters, also known as earth plasters, are available in a range of self colours, which do not require paint decoration. Clay plaster, manufactured from a blend of clay and fine aggregate, may be applied in two layers of 10 and 3 mm, respectively, or as a single 10 mm coat. If necessary, an initial 1 mm bonding coat may be applied to the substrate. Clay plasters harden only by drying out without any chemical processes. As with all clay products, these plasters absorb moisture and are responsive to environmental conditions helping to control the internal relative humidity.

**Recycled tyres**

Recycled tyres have been used to create the structural walls of new homes called earthships in Fife and near Brighton, UK. The tyres are laid in courses, filled with rammed earth and finished internally with plaster and externally with solar tiles. To date only a few small housing units have been built, but subject to Building Regulations, there is no theoretical limitation
The UK produces 40 million used tyres per year, enough to create 20,000 low-carbon autonomous earthships annually.

**Papercrete**

Papercrete is made from recycled paper and/or cardboard with sand and Portland cement. Pulverised waste glass from recycled bottles may be used instead of sand, and glossy magazines can be mixed in with standard newsprint. The material is made by dry mixing shredded paper with sand and Portland cement in the approximate ratio of 3:1:1. Water is added to make a paper mache slurry that can be cast into block units or into monolithic structures. Where papercrete blocks are used in construction, the same material can be used as the mortar. The material dries to a grey colour. It is, however, very water-absorbent, and must be protected from moisture and weather by appropriate detailing. Externally a stronger mix of 1:1 paper
and cement may be used as a stucco layer, and internally papercrete plaster may be used to give a textured or patterned finish. Papercrete is still an experimental material, but it has the potential to remove up to 20% of the waste material currently deposited in landfill sites. As a lightweight material it has good insulating properties and the cement content significantly increases its fire resistance.

Sandbags

In recently constructed office/home accommodation in Islington, London, sandbags were used for a sound-absorbing facade adjacent to a noisy main railway line (Fig. 17.6). The sandbags, filled with a sand–cement–lime mixture and exposed to the elements, will eventually set hard. Over a further period of time, possibly 30 years, the hessian bags will disintegrate leaving the undulating concrete exposed, imprinted with the texture of the hessian fabric. The wall is experimental in that the weathering effects cannot be predicted, as with other more standard forms of construction.

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**BRE Reports**


Introduction

In 2008, the UK Government published a strategy for sustainable construction which reflected the industry's commitment to reduce its carbon footprint and consumption of natural resources, whilst maintaining a strong construction sector. The initiative was within the wider context of concern for global warming and climate change, which has led to UK national targets for reduction in carbon dioxide emissions by 34% from 1990 levels by 2020 and by 80% by 2050. An indicator of the problem is that the middle-range emissions scenario for London predicts by 2020 a $1.5^\circ$C increase in mean summer temperature ($+1.3^\circ$C in winter), with 6% more rainfall in the winter and 6% less rainfall in the summer.

As buildings account either directly or indirectly for approximately 44% of the UK’s carbon emissions (27% from homes and 17% from non-domestic buildings), it is appropriate that the targets placed on the construction industry reflect this significant factor. The aim is therefore to move to zero carbon for homes by 2016 and for other buildings by 2019. In addition, the production of materials and transportation adds another 10% to the UK’s carbon emissions.

Whilst technological solutions can ameliorate carbon emissions from domestic, commercial and industrial buildings, the cornerstone for sustainable construction is holistic design taking into consideration site, local climate, orientation, external form, building envelope and openings.

Zero carbon targets

CODE FOR SUSTAINABLE HOMES

The Code for Sustainable Homes was launched by the UK Government in 2007 as part of the process for moving towards zero carbon new homes by 2016. All new homes are now required to have a Code rating, prepared by a licensed assessor, which should be included in the Home Information Pack (HIP) when the property is transferred. Code assessment procedure is normally a two-stage process which includes a Design Stage Assessment (DSA) and a Post-Construction Review (PCR). Only on completion of both stages is the property awarded its Code level certificate. The Code features a rating system from one to six stars, with one star the entry level, just above the minimum required by the 2006 Building Regulations Part L (Conservation of Fuel and Power).

The Code relates to nine identified design issues each of which is assessed on the one to six star basis.

**Code for Sustainable Homes Design categories:**

- Energy and carbon dioxide emissions
- Water
- Materials
- Surface water run-off
- Waste
- Pollution
- Health and well-being
- Management
- Ecology
By 2010, new homes should be at Code level 3, requiring a 25% energy efficiency improvement over the 2006 Building Regulations base. By 2014, a reduction of 44% (Code level 4) is required, and by 2016, zero carbon (Code level 6) is mandatory. In Wales, all new dwellings must be at Code level 5 by 2011.

In addition to energy considerations, the other ecological and social factors are star rated, each corresponding to point scores. The overall star rating is calculated by a summation of the point scores in each of the nine categories, giving final totals which range from level 1 (36 points) and level 3 (57 points) to level 6 (90 points). The property will receive a Final Certificate with an overall one to six star rating, and a breakdown into the nine design categories indicating the percentage score achieved in each. Initially, the main focus of the Code is on the measurable issues including energy, carbon, water and waste.

To achieve level 1, a house must be 10% more energy efficient than the thermal requirements of the 2006 Building Regulations. In addition, it must be built of materials rated ‘D’ or better in the ‘BRE Green Guide to Specification’, and designed to use no more than 120 l of water per day. In order to achieve a level 3 rating, the house must be 25% more efficient than the 2006 baseline and be designed to use only 105 l of water per day.

A level 6 house must be completely zero carbon with respect to space and water heating, cooking and use of all appliances. The anticipated water use should be only 80 l per day, which means that approximately 30% of the water requirement will be taken from rainwater harvesting or recycling. To achieve level 6, high levels of insulation and airtightness are required. In addition, the design must provide good daylighting levels, whilst incorporating appropriate solar control to prevent summertime overheating. A range of renewable energy technologies will include solar water heating and combinations of other systems, such as photovoltaics, heat recovery, passive cooling and ventilation, a biomass boiler and highly efficient water and waste systems.

**Prototypical zero carbon homes**

A number of experimental homes have been built to demonstrate the alternative approaches to achieving various Code level homes. These include several houses on the BRE Innovation Park at Watford, although only the Sheppard Robson ‘Lighthouse’ (Fig. 18.1) and the Barratt ‘Green House’ (Fig. 18.2), both of which featured some off-site construction, achieved a Code level 6 rating. The University of Nottingham ‘Tarmac Homes Project’, built to Code levels 4 and 6 using traditional masonry building techniques, is illustrated in Fig. 18.3. Bill Dunster has completed six Code level 6 homes at Upton in Northamptonshire.

The ‘Lighthouse’ built with high-performance structural insulated panels (SIPs), features a 40°
pitched roof with a photovoltaic array and incorporates an integrated wind catcher for passive cooling and ventilation with a sun pipe for bringing additional daylight in the core of the home. West and east elevations windows are shaded to prevent excessive solar gain. Building services include a biomass boiler, low water consumption and recycling systems, solar water heating, mechanical ventilation with heat recovery, also water-saving and rainwater harvesting systems. External shading is operated by automatic shutters to optimise useful solar gain and prevent summer overheating. The Barratt Green House achieved 92 points, just two over the requirement for an overall Code level 6 rating. The key U-values are listed in Table 18.1.

The University of Nottingham ‘Tarmac Homes Project’ reflects the different requirements of Code level 4 and Code level 6 housing within a pair of semi-detached homes. Both halves of the pair are built in masonry, one (Code 6) with solid walls and the other (Code 4) with cavity construction, but these forms could, in principle, be interchanged with appropriate adjustments to their insulation levels. Both properties incorporate 3 m² solar thermal panels, rainwater harvesting and sun pipes, and they share an individually monitored biomass boiler. The key differences are in the insulation levels associated with the external walls and glazing.

The Code level 4 house of brick and block construction has a 150 mm cavity, partially filled with 100 mm of low-emissivity foil-coated polyisocyanurate (PIR) insulation. The inner leaf is constructed with 100 mm lightweight blockwork and a 12.5 mm dry plasterboard lining, giving an overall wall U-value of 0.20 W/m² K.

The Code level 6 house is constructed from 215 mm large format aircrète blocks using thin joint

---

**Table 18.1  Barratt ‘Green House’ U-values**

<table>
<thead>
<tr>
<th>Building element</th>
<th>U-values (W/m² K)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Roof</td>
<td>0.09</td>
</tr>
<tr>
<td>Walls</td>
<td>0.11</td>
</tr>
<tr>
<td>Ground floor</td>
<td>0.09</td>
</tr>
<tr>
<td>Exposed floor to car port</td>
<td>0.11</td>
</tr>
<tr>
<td>Glazing (triple and gas filled)</td>
<td>0.70</td>
</tr>
<tr>
<td>Doors</td>
<td>0.68</td>
</tr>
</tbody>
</table>

**Notes:**
- Airtightness: 1.0 m³/(hour m²) @ 50 Pa
- Heat-loss parameter (HLP): 0.72 W/m² K, cf. Code 6 HLP 0.8 W/m² K
construction. It is externally insulated with 150 mm of expanded polystyrene and thin-coat render. With an internal finish of thin-coat projection plaster, the overall wall $U$-value is 0.15 W/m² K.

The Code level 4 house has PVC-U windows with 16 mm argon fill and low-emissivity hard coat, giving an overall $U$-value of 1.90 W/m² K. The Code level 6 house has timber windows with 16 mm argon fill and low-emissivity soft coat, giving an overall $U$-value of 1.50 W/m² K and a BFRC rating of band A. The Code 6 south elevation has a sun space to prevent summer overheating but is designed to admit the required solar gain during the cooler months.

The Code level 4 house incorporates a mechanical ventilation system with heat recovery, whilst the Code level 6 house uses a passive heat recovery wind cowl ventilation system.

The 72 photovoltaic tiles on the Code level 6 house have a peak generating power of 3.74 kW, which equates to the anticipated peak domestic load of 3.52 kW.

The six heavyweight timber-frame system-built homes at Upton in Northamptonshire incorporate eco-concrete ceilings and wall linings to give thermal mass. Building services include water harvesting, photovoltaic and solar panels with a back-up biomass boiler.

### ZERO CARBON NON-DOMESTIC BUILDINGS

The challenge to meet the target of zero carbon non-domestic buildings by 2019 is complex due to the wide variety and forms of commercial and industrial buildings and their associated business activities. The UK Government has set a zero carbon ambition of 2018 for all new public sector buildings, with a focus on schools and colleges to achieve the goal by 2016. The general interim reductions of 25% by 2010 and 44% by 2013 are the same as for new dwellings. Only the regulated energy consumption, that is the energy defined by Part L 2006 of the Building Regulations and predicted by the Simplified Building Energy Model (SBEM), applies. Energy used for industrial processes is classified in this context as unregulated, but it is subject to a range of Government incentives aimed at encouraging reductions.

### EXISTING BUILDING STOCK

Whilst the current focus is on new building, it can be assumed that at least 70% of the current building stock will still be in use by 2050. The issue of addressing retro-fit energy conservation technologies will therefore increase in relative importance as higher standards become the norm for all new construction. The Building Regulations specify appropriate requirements when building work is carried out on existing buildings.

Energy Performance Certificates (EPCs) are required for all buildings when they are sold, built or rented. Certificates are rated A to G. Currently, less than half of the existing housing stock within England and Wales achieves the average D rating. Furthermore, many government buildings fall into the lower categories with respect to energy efficiency.

The Energy Saving Trust has suggested a range of best practice standards (Table 18.2) for the refurbishment of the existing building stock. Clearly, a 60% reduction in carbon is easier to obtain on the poorer quality properties, than those already at a higher standard. The best practice represents the optimum balance between environmental performance and practicality.

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**Table 18.2** Energy Saving Trust best practice targets for the refurbishment of social and private housing

<table>
<thead>
<tr>
<th>Building element or function</th>
<th>Improvements and target $U$-values</th>
</tr>
</thead>
<tbody>
<tr>
<td>Walls</td>
<td>Insulation to a maximum $U$-value of 0.30 W/m² K</td>
</tr>
<tr>
<td>Pitched roofs</td>
<td>Insulation between rafters to a $U$-value of 0.16 W/m² K</td>
</tr>
<tr>
<td>Flat roofs</td>
<td>Insulation to a $U$-value of 0.25 W/m² K or better</td>
</tr>
<tr>
<td>Windows</td>
<td>BFRC rating of C or above. Any retained windows to be draught-proofed</td>
</tr>
<tr>
<td>Doors</td>
<td>Solid doors to a $U$-value of 1.0 W/m² K, half-glazed doors to a $U$-value of 1.5 W/m² K</td>
</tr>
<tr>
<td>Space heating</td>
<td>Ideally, a domestic wet central heating and hot water system except where electricity is the only option.</td>
</tr>
<tr>
<td>Airtightness</td>
<td>A target permeability of 5 m³/(m²)@50 Pa</td>
</tr>
<tr>
<td>Ventilation</td>
<td>A ventilation system to be installed</td>
</tr>
<tr>
<td>Light and appliances</td>
<td>When re-wiring make 75% of the units for low energy lamp fittings only. Specify low-energy appliances</td>
</tr>
<tr>
<td>Renewable and low-carbon technologies</td>
<td>Encourage the provision of solar hot water heating and other renewable technologies such as photovoltaics</td>
</tr>
</tbody>
</table>
### Specification of materials

As energy considerations have risen up the construction agenda, building material manufacturers have pressed their ecological credentials, often with claims and counterclaims between competing materials and systems. The spin-off from this debate has been a rush to reduce embodied energy in the production processes (Table 18.3), to reduce waste both within the factory and on site and also to emphasise recyclability.

Metals, such as steel, aluminium and copper, have long had good records of recycling from within the construction industry, but now, for example, significant volumes of PVC-U from the replacement windows industry are recycled. Some plastics products can be co-extruded with a core of recycled material and external layers of virgin material. Plastic bottles may be recycled as loft insulation or into composites as plastic lumber. Low-grade timber waste can be used in non-traditional products such as permanent insulating concrete formwork. These changes significantly affect the balance of ecological ratings traditionally placed on construction materials. Current research on cement is investigating alternative products which have much lower embodied energy and may even be carbon neutral. Excess bottle glass, not required for direct recycling, can be converted into a foamed aggregate for lightweight concrete.

### Responsible sourcing of materials

Responsible sourcing of materials requires a well-managed supply chain, from the point where the material is mined or harvested through processing and manufacturing to use, reuse, recycling and waste disposal. The BRE Global Certification Scheme BES 6001 provides a framework for responsible and sustainable resourcing of construction products. It also offers a route to gaining credits within the materials section of the Code for Sustainable Homes and BREEAM certification schemes.

### Life-Cycle Assessment

A full life-cycle assessment includes extraction, processing, manufacturing, transportation, maintenance, reuse, recycling and final disposal. This clearly gives a more realistic environmental impact than consideration of embodied energy alone (Table 18.3). For example, steel has an embodied energy of approximately 24 MJ/kg, but it is recycled many times over without any loss of performance. By comparison concrete, which can normally only be recycled as aggregate, has an embodied energy of approximately 1 MJ/kg. Also, the quantities used for equivalent structural purposes differ significantly and the energies involved in their deconstruction and recycling are different.

### Green Guide to Specification

In order to give guidance to designers and specifiers on the environmental impact of various building materials and construction systems, the BRE has produced The Green Guide to Specification, which lists a wide range of construction systems against a standardised set of environmental criteria, giving each criterion a letter grading (A+ to E) together with an overall summary rating (A+ to E). The Green Guide is intended for use with whole building assessment tools such as the BRE Environmental Assessment Method (BREEAM) and the Code for Sustainable Homes.

The Green Guide records the relative environmental impacts of the construction materials used in the six generic building types:

- Commercial buildings – including offices
- Educational buildings – schools, universities, colleges
- Healthcare buildings – including hospitals
- Retail outlets

### Table 18.3 Embodied energy in construction materials

<table>
<thead>
<tr>
<th>Material</th>
<th>Embodied energy (MJ/kg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bricks</td>
<td>4.6</td>
</tr>
<tr>
<td>Portland cement (wet process)</td>
<td>5.9</td>
</tr>
<tr>
<td>Portland cement (dry process)</td>
<td>3.3</td>
</tr>
<tr>
<td>Concrete (1 : 2 : 4 mix)</td>
<td>0.95</td>
</tr>
<tr>
<td>Autoclaved aerated blocks</td>
<td>3.5</td>
</tr>
<tr>
<td>Steel (42% recycled)</td>
<td>24.4</td>
</tr>
<tr>
<td>Stainless steel</td>
<td>56.7</td>
</tr>
<tr>
<td>Softwood—sawn</td>
<td>7.8</td>
</tr>
<tr>
<td>Glued laminated timber</td>
<td>12</td>
</tr>
<tr>
<td>Hardboard</td>
<td>16</td>
</tr>
<tr>
<td>MDF (medium-density fibreboard)</td>
<td>16</td>
</tr>
<tr>
<td>Plywood</td>
<td>15</td>
</tr>
<tr>
<td>Glass</td>
<td>15</td>
</tr>
<tr>
<td>Glass—toughened</td>
<td>23.5</td>
</tr>
</tbody>
</table>

Notes:
Data from Institution of Civil Engineers (Ref.: Hammond and Jones: 2008) derived from published information.
Residential buildings
Industrial buildings

Materials and components are arranged within the key building elements:

- Ground floors
- Upper floors
- Roofs
- External walls
- Windows
- Internal walls and partitions
- Insulation
- Landscaping

Within each category the materials are compared on a like-for-like basis, so that equivalent forms of construction, such as a steel or concrete column, can be directly compared. Where appropriate, materials are compared on a common $U$-value equating to equivalent heat loss.

The environmental ratings are based on a life-cycle assessment which includes greenhouse gas emissions, mineral and water extraction, toxicity of waste and pollution, landfill or incineration, fossil fuel depletion, atmospheric pollution and acid rain. Specifications are based on a 60-year period, with factors included for shorter anticipated periods to replacement. Recycled content is taken into consideration, but this is constantly improving with greater realisation of its environmental importance.

Recycling and deconstruction

Despite the high and increasing landfill tax, approximately 30 million tonnes of construction waste is sent to landfill each year within the UK. The issue of waste on building sites (Fig. 18.4) tends to be disregarded by much of the industry, despite the financial and ecological costs involved. However, some producers such as plasterboard, block and insulation manufacturers will take back clean excess and off-cuts for reprocessing.

With the increasing need to recycle materials, it can be anticipated that in the future more attention will be given to design for deconstruction rather than demolition, as this can lead to greater reuse of resources. Deconstruction accounts for typically 6–8% of the whole life-cycle energy used within building construction.

Fig. 18.4 Building site waste

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BR 502: 2009 Sustainability in the built environment. An introduction to its definition and measurement.

**GOVERNMENT PUBLICATIONS**


**ADVISORY ORGANISATIONS**

Association for the Conservation of Energy, Westgate House, 2a Prebend Street, London N1 8PT, UK (020 7359 8000).
BREEM Centre, BRE, Garston, Watford, Hertfordshire WD25 9XX, UK (01923 664940).
Centre for Energy and the Environment, University of Exeter, Stocker Road, Exeter, Devon EX4 4QL, UK (01392 264144).
Energy Saving Trust, 21 Dartmouth Street, London SW1H 9BP, UK (0845 120 7799).
Sustainability Centre, East Meon, Hampshire GU32 1HR, UK (01703 823166).
Sustainable Homes Ltd., Marina House, 17 Marina Place, Hampton Wick, Kingston-upon-Thames, Surrey KT1 4BH, UK (020 8973 0429).
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