Engineering coatings

- design and application
Engineering coatings
- design and application

Second edition
Edited by
Stan Grainger and Jane Blunt
# Contents

*Preface to the first edition*  ix  
*Preface to the second edition*  xi  

1 **Introduction**  1  
Surfacing materials and processes  2  
Material/process selection  7  
Using the book  7  

2 **Mechanisms of wear and corrosion**  10  
Basic mechanisms of wear  10  
Corrosion  25  
Mixed attack mechanisms  32  
Testing for wear resistance  35  
Material selection for wear resistance  38  
Testing for corrosion resistance  40  
Material selection for corrosion resistance  41  

3 **Weld surfacing**  42  
Characteristics of weld surfacing  42  
Materials for weld surfacing and their selection  45  
Arc welding processes for surfacing  61  
Other fusion welding and forge welding processes  68  
Metallurgical and related effects of weld surfacing on  
substrate and coating  82  
Preparation and design for weld surfacing  92  
Practical considerations  111  

4 **Thermal spray processes**  119  
Thermal spray materials  122  
Characteristics of generic low energy thermal spray processes  132  
Characteristics of the high energy thermal spray processes  137  
Spray fused coatings  152  
Design for thermally sprayed coatings  157  
Coating production  161  
Conclusions  166
## Contents

5 Electrodeposited coatings  
Basic principles  167  
Selection  167  
Design  186  
Substrate materials  192  
Surface preparation  192  
Finishing  192  
Specifications, inspection and quality assurance  193

6 Physical and chemical vapour deposition techniques  
The processes  194  
Vacuum evaporation  194  
Evaporation sources  199  
Gas scattering deposition  200  
Sputter coating  203  
Ion implantation  209  
Chemical vapour deposition  214  
Comparison of processes  215  
Design for physical and chemical vapour deposition  220

7 Polymer coatings  
Materials  226  
Application processes  227  
Designing for polymer coatings  233  
Summary  234

8 Finishing of surface coatings applied by welding and thermal spraying  
The need for machining  235  
Spray fused and welded coatings  236  
Thermally sprayed coatings  242

9 Ensuring quality in surfacing  
The quality plan  246  
Testing and inspection of coated components  251  
Conclusions  258

10 Safe working in surfacing  
Surfacing by welding and thermal spraying  259  
Vacuum deposition  260  
Electrodeposition  271  
Bibliography  274

11 Industrial applications of engineering coatings  
The aircraft industry  279  
Chemical and petroleum industries  280
<table>
<thead>
<tr>
<th>Contents</th>
<th>vii</th>
</tr>
</thead>
<tbody>
<tr>
<td>Earthmoving, agricultural, quarrying and mining</td>
<td>284</td>
</tr>
<tr>
<td>Internal combustion (piston) engines</td>
<td>286</td>
</tr>
<tr>
<td>Food</td>
<td>288</td>
</tr>
<tr>
<td>Forging</td>
<td>289</td>
</tr>
<tr>
<td>Glass</td>
<td>292</td>
</tr>
<tr>
<td>Pulp and paper</td>
<td>292</td>
</tr>
<tr>
<td>Plastics and rubber</td>
<td>295</td>
</tr>
<tr>
<td>Power generation</td>
<td>297</td>
</tr>
<tr>
<td>Steelmaking/general metal forming</td>
<td>298</td>
</tr>
<tr>
<td>Textiles</td>
<td>301</td>
</tr>
<tr>
<td>Timber</td>
<td>301</td>
</tr>
<tr>
<td>Transport</td>
<td>303</td>
</tr>
</tbody>
</table>

**Glossary of terms used in surfacing**  309

**List of contributors**  315

**Index**  316
Preface to the first edition

This engineering coatings handbook deals with materials and application processes used for improving the surface durability of engineering components in service. Surfacing technology has developed rapidly in recent years and, perhaps because of its specialised nature, its scope and the benefits it provides are not widely recognised.

The purpose of the handbook is to acquaint readers with the subjects and to guide the choice of coatings and means of application to fit specific circumstances.

Production of this book was recommended by the Committee of the Surface Engineering Society which is affiliated to the Welding and Joining Society. The source material includes technical papers provided by committee members and illustrations and information supplied by many industrial organisations and individual authors. All their contributions, too numerous to mention separately, are gratefully acknowledged, as is the help and advice provided by individual staff at The Welding Institute.

Stan Grainger

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Preface to the second edition

Nine years have passed since the original work was published, and surfacing technology has advanced in several areas. New technologies have been included in this edition of the book – such as the newer thermal spray processes and laser surfacing. The book has also been revised to include more information about corrosion.

Major changes have taken place in health and safety legislation, and the chapter on health and safety has been revised significantly as a result.

Units have been rationalised to the SI system, wherever feasible. The presentation has been changed to index notation, i.e. 30km/h would be presented as 30kmh⁻¹. In tables and graphs the slash / may be read as ‘in units of’.

Thanks are due to colleagues in the committee of the Surface Engineering Society for the provision of up-to-date material and photographs. All sources are acknowledged in the captions, and a list of contributors is included at the end of the work.

Jane Blunt
Chapter 1

Introduction

The progressive deterioration of metallic surfaces in use, because of wear or corrosion, ultimately leads to loss of plant operating efficiency and at worst a breakdown. The cost to industry of wear and corrosion in equipment is high, and recognition of this fact lies behind the continuing development of the technology known as surface engineering, which includes the application of coatings to metal surfaces to improve their performance in specific working conditions. This is a subject of great importance to industry, which relies on long, trouble-free operation of plant to obtain uniform product quality and lowest possible product cost.

This book, produced under the auspices of the Surface Engineering Society, brings together a wealth of coating materials and processes of application currently available, together with examples of how the technology is being used successfully in industry.

An important feature of engineering plant is that individual components must be designed with three objectives in mind:

1. To present a series of surfaces to interact with other parts or with process media.
2. To support these surfaces with adequate strength to withstand service stresses.
3. To endow selected surfaces with resistance to wear and/or corrosion.

The terms wear and corrosion are used here to mean any of the destructive forms of attack that cause deterioration of metallic surfaces in use. These are described in greater detail in Chapter 2. Sometimes all surfaces of the part are exposed but, more frequently, only a proportion of the surface is required to display high resistance to a specific form of attack.

The compositional requirements of the material necessary to provide adequate strength in a component are usually different from those that provide wear or corrosion resistance, so a composite product consisting of a structural material with specially protected exposed surfaces naturally comes to mind.

Surfacing technology is used to cut the cost of component deterioration in service by providing:

- acceptable service life/reduced downtime costs;
- the minimum first cost of the part consistent with the above, including use of the cheapest constructional materials, and minimum use of
special materials that provide wear or corrosion resistance on selected surfaces;

- the opportunity, where possible, to repair the part after use by resurfacing.

While the bulk of surfacing is used to prevent wear or corrosion, there are several other uses of surfacing that are of economic importance. Surfacing can be used to introduce layers that are electrically resisting or electrically conducting, a layer that abrades in use to allow parts to conform, and lastly a layer that acts as a thermal barrier between the part and its working environment. Materials for all these purposes are included in this text.

**Surfacing materials and processes**

**Materials**

Most metals, alloys, ceramics and some intermetallic compounds can be applied as coatings either individually or as mixtures, but their characteristics often limit the processes that can be used for their application. Refractory oxides, for example, cannot be applied by welding processes, and carbides require a metal or alloy matrix.

The material/process relationship not only identifies where they can be used together but also determines properties that can be expected from the coating, such as density and adhesion to the substrate.

It is not possible to list all the individual surfacing consumables available for welding and thermal spraying, but Table 1.1 summarises basic types. Each type may be available in more than one variation of composition, each of which has been developed to possess specific properties to optimise life in given circumstances. Tables 1.2 and 1.3 cover coatings applied by electrodeposition and by physical and chemical vapour deposition.

**Processes**

**Welding**

Welding provides the highest bond strength between deposit and substrate, is capable of applying deposits of considerable thickness if required and, with few exceptions, can be operated manually or be mechanised and programmed. A particular application of a given coating can often be achieved satisfactorily by more than one welding process and this flexibility of choice is a useful characteristic of the group. There is a wide range of materials available for welded coatings.

**Thermal spraying**

Thermal spraying offers two significant advantages over weld surfacing. The first is that both non-weldable coating materials, such as ceramics, can be deposited as well as materials which are also weldable. The second is that
### Table 1.1 Summary of consumables for welding, thermal spraying and related application processes

<table>
<thead>
<tr>
<th>Group</th>
<th>Type</th>
<th>Weld deposition methods</th>
<th>Approximate deposit hardness/HV</th>
<th>Thermal spray methods</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>OA</td>
<td>MMA</td>
<td>MIG</td>
</tr>
<tr>
<td>A Ferrous alloys</td>
<td>1 Carbon steels</td>
<td>&lt;250</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>2 Low alloy steels</td>
<td>250–650</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>3 Martensitic Cr steels</td>
<td>350–650</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>4 High speed steels</td>
<td>600–700</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>5 Austenitic stainless steels</td>
<td>200/500*</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>6 Austenitic Mn steels</td>
<td>200/500*</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>7 Austenitic Cr–Mn steels</td>
<td>200/600*</td>
<td></td>
<td></td>
</tr>
<tr>
<td>B Irons</td>
<td>1 Austenitic irons</td>
<td>300–600</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>2 Martensitic irons</td>
<td>500–750</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>3 Hi-Cr austenitic irons</td>
<td>500–750</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>4 Hi-Cr martensitic irons</td>
<td>500–750</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>5 Hi-Cr complex irons</td>
<td>600–800</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>6 Fe-Cr-Co-Ni-Si</td>
<td>375–550</td>
<td></td>
<td></td>
</tr>
<tr>
<td>C Nickel alloys</td>
<td>1 Nickel</td>
<td>160</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>2 Ni–Cu, Ni–Cu–In</td>
<td>130</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>3 Ni–Fe</td>
<td>200</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>4 Ni–Mo–Cr–W</td>
<td>250/500*</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>5 Ni–Cr–Si–B (also with Cu + Mo)</td>
<td>200–750</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>6 Ni–Mo–Fe</td>
<td>200–300</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>7 Ni–Al bond coat</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td></td>
<td>8 Ni–Al–Cr bond coat</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>9 Ni–Cr B320</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>10 Ni–Cr–Fe</td>
<td></td>
<td></td>
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</tr>
<tr>
<td></td>
<td>11 Ni–Cr–Al–Y</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>12 Ni–Cr–Mo–Al–Ti</td>
<td></td>
<td></td>
<td></td>
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## Table 1.1 (cont.)

<table>
<thead>
<tr>
<th>Group</th>
<th>Type</th>
<th>Weld deposition methods</th>
<th>Approximate deposit hardness/HV</th>
<th>Thermal spray methods</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>OA</td>
<td>MMA</td>
<td>MIG</td>
</tr>
<tr>
<td>D Cobalt alloys</td>
<td>1 Co−Cr−W low alloy</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>2 Co−Cr−W medium</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>3 Co−Cr−W high alloy</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>4 Co−Cr−W−Ni</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>5 Co−Cr−Mo</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>6 Co−Cr−Mo−Si</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>7 Co−Cr−W−Ni−Si−B alloy</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>E Copper alloys</td>
<td>1 Brass</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>2 Silicon bronze</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>3 Aluminium bronze</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>4 Tin bronze</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F Composite materials</td>
<td>1 Ni−Cr−Si−B + Ni−Al or Mo</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>2 Ni−Cr−Si−B + carbid es</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>3 Co−Cr−W−Si−B + carbid es</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>4 C steel + carbid es</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>5 Ni + silicon carbide</td>
<td></td>
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<tr>
<td></td>
<td>6 WC + cobalt</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td></td>
<td>7 Cr−carbide + Ni−Cr</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>8 Cermets</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>9 Cr−B‡</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

- OA: Oxygas Arc
- MMA: MIG
- MIG: MIG
- FCAW: FCAW
- TIG: TIG
- SAW: SAW
- PTA: PTA
- Approximate deposit hardness/HV:
  - D Cobalt alloys: 380–430
  - E Copper alloys: 130
  - F Composite materials: 600–750
  - Thermal spray methods:
  - Oxygas: 380–430
  - Arc: 130
  - Plasma spray: 600–750
<table>
<thead>
<tr>
<th>Group</th>
<th>Type</th>
<th>Weld deposition methods</th>
<th>Approximate deposit hardness/HV</th>
<th>Thermal spray methods</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>OA</td>
<td>MMA</td>
<td>MIG</td>
</tr>
<tr>
<td>G Elements</td>
<td>1 Aluminium</td>
<td>OA</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>2 Zinc</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>3 Tungsten</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>4 Copper</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>5 Molybdenum (also as a bond coat)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>H Ceramics</td>
<td>1 Alumina</td>
<td>OA</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>2 Zirconia</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>3 Chrome oxide</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>4 Alumina/titania</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>5 Titanium dioxide</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>6 Magnesium zirconate</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>I Polymers</td>
<td>1 EVA</td>
<td>OA</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>2 Nylon</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td></td>
<td>3 Polyester</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>4 Polypropylene</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>5 Low density polyethylene</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Table 1.2 Electrodeposited coatings

<table>
<thead>
<tr>
<th>Type</th>
<th>Materials</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pure metals</td>
<td>Chromium, nickel, gold, silver, platinum, ruthenium, rhodium, lead, tin, iron</td>
</tr>
<tr>
<td>Alloys</td>
<td>Copper-tin, cobalt-tungsten, cobalt-molybdenum, tin-nickel</td>
</tr>
<tr>
<td>Composites</td>
<td>Cobalt-chromium carbide, nickel-silicon carbide, metal-poly tetrafluoroethene</td>
</tr>
</tbody>
</table>

Table 1.3 Physical and chemical vapour coatings for wear resistance

<table>
<thead>
<tr>
<th>Type</th>
<th>Materials</th>
</tr>
</thead>
<tbody>
<tr>
<td>Elements</td>
<td>Carbon, silver, chromium, molybdenum, tungsten, nickel, titanium</td>
</tr>
<tr>
<td>Compounds</td>
<td>Carbides, nitrides, oxides, sulphides. Borides of: titanium, aluminium, silicon, tantalum, tungsten, chromium, molybdenum</td>
</tr>
<tr>
<td>Alloys</td>
<td>M–Cr–Al–Y alloys, cobalt–chromium</td>
</tr>
</tbody>
</table>

all these materials can be deposited on substrate materials that are unsuited to welding either because of their composition or because their thin section would distort excessively when welded.

Compared with welded coatings, thermally sprayed deposits exhibit some porosity and a lower bond strength to the substrate. In general, deposit thickness is less than is possible by welding. Provided that these factors are taken into account at the design stage, thermal spraying can, and does, provide excellent and reliable service in applications as demanding as aircraft gas turbine engines.

**Electrodeposition**

A somewhat more limited range of coating materials is offered by electrodeposition but the low temperature of deposition provides advantages of low distortion, better access to internal surfaces and accurate control of deposit thickness.

**Vapour deposition**

Vapour deposition provides a limited range of coating material possibilities, but can be used with materials that are difficult or impossible to apply by other techniques, or to produce thin coatings of controlled thickness.

**Coating thickness**

The thickness of the coating material applied is normally consistent with the amount of material loss that can be permitted before the component is no longer fit for use, and takes into account an allowance for machining or
grinding the deposit to a specified dimension and finish. Table 1.4 gives a guide to the ranges of thickness that are possible using the various processes. In practice a different range may result from the characteristics of a particular coating material.

### Material/process selection

Faced with the wide range of possibilities indicated in the tables, selection of material and process may seem difficult, but is normally straightforward. Often there are constraints placed on the choice because of availability. In many cases there is a precedent, but when considering a new problem it helps to follow a checklist of the type shown in Fig. 1.1.

The sequence of decisions to be made covers several fundamental points. The first is the need to be clear about service conditions, based on experience or plant design data. This is the key to material selection. The second decision is the choice of application process for that material. This will involve questions of compatibility with the coating material. Reference to the tables in this chapter will show that not all the materials can be applied by all processes.

A further question of compatibility arises between both material and process with the substrate. All these issues are covered in subsequent chapters.

### Using the book

Chapter 2 is devoted to mechanisms of wear and corrosion and is intended to assist in the analysis of previous component failures and assessment of how parts of newly designed equipment are likely to suffer in service. In
Table 1.4 Comparison of surfacing processes and deposits

<table>
<thead>
<tr>
<th>Process</th>
<th>Thickness (mm)</th>
<th>Component geometry</th>
<th>Component size</th>
<th>Substrate material temperature (°C)</th>
<th>Pretreatment</th>
<th>Post-treatment</th>
<th>Coating porosity (%)</th>
<th>Bond strength (MPa)</th>
<th>Bond mechanism</th>
<th>Control of deposit thickness</th>
<th>Distortion of substrate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vapour deposition</td>
<td>0.001-0.2</td>
<td>Versatile</td>
<td>Limited by chamber size</td>
<td>PVD - ion bombardment</td>
<td>None</td>
<td>None/stress relief</td>
<td>Nil to small</td>
<td>100</td>
<td>High</td>
<td>Good</td>
<td>Low</td>
</tr>
<tr>
<td>Thermal spraying</td>
<td>0.1-1.0</td>
<td>Versatile</td>
<td>Almost limitless</td>
<td>CVD - various</td>
<td>Chemical cleaning and etching</td>
<td>100</td>
<td>200</td>
<td>High</td>
<td>1-15</td>
<td>Low</td>
<td>Strong forces</td>
</tr>
<tr>
<td>Spray fusing</td>
<td>0.05-1.5</td>
<td>Versatile</td>
<td>Almost limitless</td>
<td>PVD - ion bombardment</td>
<td>None/substrate stress relief as required</td>
<td>100</td>
<td>Nil</td>
<td>High</td>
<td>20-40</td>
<td>High</td>
<td>Mechanical</td>
</tr>
<tr>
<td>Welding</td>
<td>1-20 or more</td>
<td>Access to internal surfaces controlled by size of torch/gun</td>
<td>Limited by fusion facility</td>
<td>Substrate anneal/stress relief as required</td>
<td>Chemical cleaning and etching</td>
<td>100</td>
<td>1400</td>
<td>High</td>
<td>Nil</td>
<td>High</td>
<td>Mechanical</td>
</tr>
</tbody>
</table>

PVD = physical vapour deposition, CVD = chemical vapour deposition.
conjunction with later chapters, which give information on the ability of various coating materials to resist different forms of wear or corrosion, a search can be made for a suitable coating material.

Four chapters (3–6) deal with the main groups of coating processes outlined earlier. Those dealing with weld surfacing and thermal spraying are dealt with in some detail, recognising the widespread availability and use of these processes in industrial maintenance and production shops. Each chapter also describes consumable materials available for deposition by the processes. Application of polymeric materials by thermal spraying is discussed in Chapter 7.

Wear-resistant materials are more difficult to machine than steels and other alloys used for component construction, but precision finishing of most coatings is possible using appropriate techniques described in Chapter 8. Successful service of engineering coatings depends, as with other materials and processes, on quality assurance and control. Special features of coating materials and processes that are important in this regard are dealt with in Chapter 9. The practice of surface coating uses combinations of materials and processes that differ from those met frequently in industry. It has been considered important therefore to include a section on safe working practice (Chapter 10).

Much can be learned from successful uses of service coatings in industry. Chapter 11 illustrates the wide field of use to which the technology is put, not only to protect new parts, but also to repair and extend the life of worn parts. Much of the growing use of surfacing has resulted from the need to find ways to extend plant life.

Surface coating technology has developed its own jargon and a glossary of terms is included to minimise the risk of misunderstanding.
Mechanisms of wear and corrosion

The aim of this chapter is to describe the processes by which surfaces are damaged by mechanical or chemical removal of material, and to indicate properties needed in materials that are required to operate in such aggressive environments. The principal mechanisms of wear are discussed in the first part, and the principal mechanisms of corrosion in the second, followed by a few well-recognised combined attack mechanisms. A few wear and corrosion tests are described, which can give some valuable information about the probable performance of a coating or surface treatment.

Basic mechanisms of wear

Wear can be defined as the progressive loss of material from the operating surface of the body, occurring as a result of relative motion of the surface with respect to another body. The concept embraces metal to metal, metal to other solids and metal to fluid contact, and the definition clearly associates the process with the surfaces of materials.

Abrasion, adhesion and contact fatigue are generally regarded as the three basic wear mechanisms that result in material being removed from the surface of a component. An industrial survey has shown that abrasion and adhesion occur in 50% and 15% respectively of all wear situations. Other processes – such as corrosion – may intensify damage. The principal features are described below and summarised in tables.

However, in practical situations, more than one type of wear is generally encountered and, in addition, one type may predominate for part of the operating cycle and another for the remainder. Finally, the products of one type of wear may themselves cause secondary wear by another mechanism.

It is also true that wear is a systems phenomenon, the behaviour of the wearing surface being affected to a greater or lesser extent by the other parts of the system. These factors combine to make wear a complex subject and, while it is convenient to discuss the different processes separately, their interdependence should be borne in mind.

Abrasive wear

Abrasive wear arises from the penetration of one surface by a harder body or surface. Damage involves a cutting and ploughing action. It may involve
Mechanisms of wear and corrosion

2.1 Types of abrasive wear: (a) gouging; (b) low stress abrasion; (c) high stress abrasion (e.g. rod and ball mills); (d) three body abrasion.

particles moving over a surface (two body abrasion), hard particles moving between two moving surfaces (three body abrasion) or a roughened surface moving over and penetrating an opposing surface.

In practice, abrasion is often considered under separate headings, such as gouging, scoring, low stress and high stress (Fig. 2.1), but these identify the severity of the problem. In all, the basic abrasive action is similar. Where surfaces are subjected to abrasive conditions, the majority of particles cause little damage, simply rolling or sliding over the surface. It is only those whose attack angle lies between about 80° and 120° that give rise to significant wear damage.

Hardness and wear

It is possible to think that wear rate should be inversely related to hardness. However, practical results on abrasive wear tend not to confirm this. In materials of simple microstructure, there may be a simple relation between hardness and wear rate, as has been shown for example for commercially pure metals. However, with materials of more complex microstructure (typified by most engineering alloys), this is not so. In steels, the relation of wear to hardness is affected by the carbon content and by the microstructure of the matrix (Fig. 2.2).

The presence of secondary phases in the structure is also important. Carbides especially, but also borides, nitrides and phosphides, are widely used with success to provide resistance to abrasion. The degree of improvement depends on the composition, amount and morphology of the hard phases (as well as upon the operating environment).
### Table 2.1 Hardness of some abrasives and mineral phases

<table>
<thead>
<tr>
<th>Raw material, or mineral hardness/HV</th>
<th>Material phase hardness/HV</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coal</td>
<td>Ferrite</td>
</tr>
<tr>
<td>Gypsum</td>
<td>Pearlite</td>
</tr>
<tr>
<td>Lime</td>
<td>Austenite</td>
</tr>
<tr>
<td>Calcite</td>
<td>Martensite</td>
</tr>
<tr>
<td>Fluorspar</td>
<td>Basalt</td>
</tr>
<tr>
<td>Coke</td>
<td>Cementite</td>
</tr>
<tr>
<td>Iron ore</td>
<td>Chromium carbides</td>
</tr>
<tr>
<td>Glass</td>
<td>Alumina</td>
</tr>
<tr>
<td>Feldspar</td>
<td>Niobium carbide</td>
</tr>
<tr>
<td>Sinter</td>
<td>Tungsten carbide</td>
</tr>
<tr>
<td>Quartz</td>
<td>Silicon carbide</td>
</tr>
<tr>
<td>Corundum</td>
<td>Vanadium carbide</td>
</tr>
<tr>
<td></td>
<td>Boron carbide</td>
</tr>
<tr>
<td></td>
<td>Diamond</td>
</tr>
</tbody>
</table>

![Graph](image)

2.2 Relative wear resistance of steel against: (a) hardness; (b) carbon content; (c) carbides.
Mechanisms of wear and corrosion

With many materials, it is possible to reach a specified hardness by different means. Thus, improvement in resistance of steel to abrasion can be much greater if the matrix is hardened by alloying, than if either quenching and tempering or precipitation hardening is used to achieve the same result (Fig. 2.3). Again, in cast iron, if the matrix structure is kept constant and the hardness altered by changing graphite distribution, wear resistance can improve with reduction in bulk hardness of the alloy. Although these results refer only to the particular test conditions used, they do illustrate that not only hardness but also the method of achieving it are important.

In considering hardness, the difference between the abrading body and the other surface is important. Table 2.1 lists typical hardness data. Wear of a surface tends to be progressively reduced if the ratio of its hardness to that of the abrading body increases over the range 0.5–1.3. However, this is complicated by the fact that the wear rate/hardness curve generally shows discontinuities (Fig. 2.4), the transitions depending upon the hardness of the abrading phases.

Most metallic surfaces work-harden, and a few transformation-harden, during wear. Discussions of the role of hardness have generally centred on the hardness measured from unworn samples, whereas the relevant hardness is that after wear. These values can be quite different, the most significant example being austenitic manganese steel, which, from an initial hardness of 200HV, can be hardened to 600HV. Indeed, this material only gives good wear resistance to abrasive conditions if there is sufficient impact loading to ensure transformation of austenite to martensite during operation.

These observations should not be taken to indicate that hardness – even as usually measured on unworn components – is unimportant, but rather that it alone does not determine the ability of a material to resist abrasion; other properties also have an influence. Structure and composition play an
important part, and differences in manufacture can also influence wear resistance, through their effects on material structure.

**Erosion**

Erosive wear is a special form of abrasion in which contact stress arises from the kinetic energy of solid or liquid particles in a fluid stream encountering a surface. The damaged surface may show a fine granular appearance, rather similar to that observed in brittle fracture. Particle velocity and impact angle, together with abrasive size, give a measure of the kinetic energy of the erosive stream, and the wear volume is proportional to the cube of the stream velocity.

In general, there is a correlation between the erosion resistance of a material and its 'ultimate resilience' (defined as \(\frac{UTS}{2E}\) where UTS is ultimate tensile strength, and \(E\) is the modulus of elasticity).

This expression is a measure of the amount of energy that can be absorbed before cracking occurs. If the impact angles in erosion are small, cutting wear is the predominant mode of damage, and hardness of the surface is the main requirement. Hard materials, even if brittle, are generally satisfactory. Soft materials may be suitable and rubbers – because of their low moduli of elasticity – are often best (Fig. 2.5).

With larger impact angles, wear involves deformation, and a measure of toughness becomes necessary. Indeed, the mechanism of material loss is likely to become fatigue at large angles. This is discussed in the section on fatigue below.

**Practical diagnosis of abrasive wear**

The appearance of abrasive wear is influenced by many factors, including the material involved, pressure and speed of rubbing impact, turbulence of movement, etc. As a result the appearance of a metal surface exposed to
Mechanisms of wear and corrosion

2.5 Dependence of rate of erosion on angle of attack of impinging particles.

Wear by abrasive materials is usually characterised by regular or irregular scoring, scratching and pitting.

The presence of corrosion may not be apparent on an abraded surface because of the continual cleaning effect of the abrasive but there may be signs of corrosion products in any stagnant areas of movement. The metal may claim good resistance to the medium/media involved, but if this depends on the formation of protective surface films, the abrasive will remove these and continually expose clean areas to the combined effects of both, as indicated earlier in this chapter.

If three-body abrasion is involved, for example a metal to metal bearing that is exposed to abrasive material, there is a possibility that abrasive particles will be pressed into the surface of the softer of the two materials and act as a lap. After a period, the lap shows little or no signs of wear while the harder material is significantly worn. A good example of erosion is shown in Fig. 2.6, which is the internal surface of a pipe bend exposed to air carrying quartz particles up to 500\(\mu\)m at a velocity of 27.4\(\text{ms}^{-1}\) (90\(\text{ft} \text{s}^{-1}\)).

Erosion of surfaces exposed to high velocity fluids such as steam, which contain no abrasive particles, often exhibit a quite smooth surface either over an extensive area or in a series of eroded grooves.

Control of abrasive wear

For pure abrasion, the ideal structure tends to be a hard matrix (e.g. martensite) with evenly distributed hard particles (e.g. carbides). In seeking abrasion-resistant materials, it is necessary to ensure there is sufficient strength to resist imposed stresses from the wear mechanism and sufficient toughness to withstand impact. If there is a large amount of impact loading, then it may be necessary to revert to a tougher matrix (e.g. bainite) at the expense of some wear resistance, or possibly use an
Abrasion involves penetration of one surface by a harder body or surface.

Abrasion is often described by different terms – such as gouging, scoring, etc., but these identify the severity of the wear. In all of these, the basic mechanism of damage is similar.

In general, wear resistance is not related to hardness in a simple way. Other factors, particularly composition and structure, markedly influence resistance to abrasion.

In choosing wear-resistant materials, both strength and toughness are normally required and this involves some degree of compromise in selection.

2.6 The pattern of erosive wear in a steel pipe bend carrying quartz grains in air at a velocity of 27.4 ms\(^{-1}\) (courtesy CEGB).

Initially relatively soft matrix which transforms during service (e.g. unstable austenite). This dual requirement for strength and toughness demands a compromise. Where there is any doubt, the balance should be tilted in favour of extra toughness; a higher than optimum wear rate reduces component life but is less disastrous than component failure. In general, the hardness of a material chosen to resist abrasive wear should be at least 80\% of the hardness of the abradent, if this is feasible.

It cannot be overstressed that erosion is most easily (and economically) overcome by changing the angle of attack, or by reducing the velocity of flow, rather than simply by providing more erosion-resistant materials, whether bulk or coating.

The key points regarding abrasive wear are given in Table 2.2.
2.7 Contacting asperities.

**Adhesive wear**

The established theory of adhesive wear is based on the premise that surfaces of crystalline (metallic) solids, although apparently smooth, are rough on a microscopic scale and consist of a series of peaks and valleys. When two such surfaces are brought together, they make contact only at the opposing asperities (Fig. 2.7). The area of these asperity contacts may be as little as $10^{-4}$ times that of the nominal contact area; therefore even a modest load applied normally to the surface causes high local pressures (and temperatures). The elastic limit of one (or both) materials may be exceeded and the asperity contacts undergo plastic deformation, until the real contact area has increased sufficiently to support the applied load. Such plastic flow is likely to disrupt protective surface films and so allow clean metals to come into intimate contact. This may cause welding, and the junctions so formed tend to rupture upon relative movement of the surfaces. As sliding proceeds, this sequence of asperity welding and junction rupture is repeated. If rupture takes place on the original joint interface, minimal damage is caused. However, if the welds work-harden (as is common with metals), rupture is more likely to occur behind the interface and fragments will be torn out of one (or both) of the contacting surfaces, which thereby suffer damage. In addition, the loose debris produced may itself contribute to wear by an abrasion mechanism.

It has been claimed that friction and wear are directly related but, as Table 2.3 shows, this need not be so. High friction may be associated with low wear and vice versa. Neither friction nor wear is an intrinsic property of a material and their values are determined by the engineering system. For instance, there is evidence that metals that have good mutual solubility for one another will more readily form welds, and therefore contact between such pairs is best avoided. It is therefore unwise to attempt to extrapolate wear performance from friction results. At best, changes in coefficient of
Table 2.3 Wear and friction data for different materials

<table>
<thead>
<tr>
<th>Materials</th>
<th>Wear rate/cm³/cm⁻¹</th>
<th>Coefficient of friction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mild steel on mild steel</td>
<td>1.57 × 10⁻⁷</td>
<td>0.62</td>
</tr>
<tr>
<td>60/40 leaded brass</td>
<td>2.4 × 10⁻⁵</td>
<td>0.24</td>
</tr>
<tr>
<td>Polytetrafluoroethene (PTFE)</td>
<td>2.0 × 10⁻⁹</td>
<td>0.18</td>
</tr>
<tr>
<td>Stellite</td>
<td>3.2 × 10⁻¹⁰</td>
<td>0.60</td>
</tr>
<tr>
<td>Ferritic stainless steel</td>
<td>2.7 × 10⁻¹⁰</td>
<td>0.53</td>
</tr>
<tr>
<td>Polyethylene</td>
<td>3.0 × 10⁻¹¹</td>
<td>0.65</td>
</tr>
<tr>
<td>Tungsten carbide on tungsten</td>
<td>2.0 × 10⁻¹²</td>
<td>0.35</td>
</tr>
</tbody>
</table>

Load 400g; speed 1.80m/s⁻¹, materials run against hardened tool steel, except for mild steel and tungsten carbide.

friction within a single wear couple may be taken to indicate possible changes in the dominant wear mechanism.

Mild and severe wear

Adhesive wear is noted for marked transitions in wear rate resulting from changes in load or sliding speed; this is a consequence of the fact that there are two different regimes of damage, known as mild and severe wear. Figure 2.8 illustrates such transitions for steel. Below T1, the wear rate is low and the process conforms to mild wear. At T1, the wear rate increases greatly as severe wear begins and this high rate continues until it suddenly drops at T2 to another regime of mild wear.

Mild wear depends upon the presence of an oxidised surface, which keeps clean metals apart, and the wear process involves the production of fine, non-metallic debris from this surface layer. The protective oxide films are
mainly generated during rubbing or sliding, so that conditions that favour oxide formation are most likely to confine wear to the mild regime. Oxide particles that become detached can act on the system either as a lubricant or as an abrasive, depending upon their nature.

When the load on the surfaces becomes sufficiently high, fine cracks begin to form in the oxide layer and these may be followed by plastic flow of the supporting material. This causes the onset of the severe wear regime. There is significant material transfer from one surface to the other, metallic fragments are torn out and damage may be as much as $10^4$ times as great as in mild wear. There are at least two distinctive stages in the process: the removal of metal fragments from the wearing surfaces and the formation of wear particles which may be five to ten times as large. There may also be a third stage, in which the transferred material is oxidised.

During sliding under dry or imperfectly lubricated conditions, high temperatures can be generated by friction and these can lead to another transition in wear rate. High temperatures may cause intense surface hardening and possibly structural transformation in the sliding members. In addition, the elevated temperatures may affect the nature of the oxide formed on the surfaces, often to a more protective variety. The consequence of these changes is frequently a material more resistant to deformation and on which a protective surface layer can be re-established. If this happens, the wear rate drops again, to a level consistent with mild wear.

**Running-in wear**

During the initial stages of sliding, particularly with unused components, metal surfaces tend to be clean, with little surface film protection; under these circumstances, special care must be taken to avoid damage. This transient type of condition is known as running-in, and it has been observed for many materials under a variety of operating conditions. Running-in may produce few adverse effects, since an oxidised or lubricated surface layer is quickly established and this leads to an acceptable wear rate. However, under certain operating conditions, a form of severe wear known as scuffing can occur. This is characterised by roughening of the surface, beginning in narrow, localised bands and gradually spreading around the rubbing surface. Components seem to be most at risk from scuffing at loads just below the mild/severe transition (Fig. 2.9) and under conditions of poor lubrication.

**Fretting**

Fretting is the consequence of two surfaces having an oscillatory relative motion of small amplitude. The surfaces do not lose contact with each other and the products of wear are unable to escape, so that seizure may occur because of blockage of the free space and prevention of access for lubricant. Although it is convenient to treat it separately, fretting is not a distinct type of wear, but is a form of adhesive wear.

Fretting probably involves three main stages. Firstly, mechanical action disrupts the oxide films on the surfaces and exposes clean, reactive metals.
Although these may re-oxidise during the next half cycle, they are disrupted again on the return half cycles. The metal particles are then removed from the surface in a finely divided form by mechanical grinding or by the formation and breaking of welded asperities. Finally, oxide debris, either from the first process or by oxidation of the metallic particles released in the second, forms an abrasive powder which continues to damage the surface.

Fretting can be removed by eliminating the source of the vibration. Alternatively, surrounding the contact area with lubricant reduces friction and the adhesion propensity of the surfaces. With ferrous components in particular, elimination of oxygen is beneficial, as it militates against formation of oxides. Specific fretting difficulties can often be overcome by application of surface coatings, and treatments that reduce metallic adhesion, such as phosphiding, have been particularly successful in this respect.

**Control of adhesive wear**

If direct and intimate contact of opposing surfaces can be prevented, there will be no adhesive wear. Surface films, whether of oxides or absorbed atom layers, are extremely important and when these are deliberately removed, a high wear rate results. Material with good oxide-forming capacity may be expected to perform well but the situation is complicated by the fact that the sliding action tends to disrupt the films.

Materials of high elastic moduli have been claimed to offer advantages in resisting wear, as the area of asperity contact should be limited by the high modulus. However, as with hardness, it is difficult to equate modulus in any simple way with wear performance. Perhaps one of the most important properties is the ability of the material to work-harden, as this controls the position at which an asperity junction rupture takes place and so affects the degree of damage.

Materials options to control or minimise adhesive wear are, in practice, usually based upon one (or more) of three approaches:
Mechanisms of wear and corrosion

Lubrication using oil films to prevent direct metal/metal contact. However, it should be emphasised that it is not possible to maintain full hydrodynamic lubrication at all times and some contact of mating surfaces is inevitable.

Use of materials that are mutually insoluble. This may not always be possible, because of other constraints in choice of materials for opposing surfaces.

Application of surface coatings (or treatments). The success of such techniques depends mainly upon their ability to interpose a dissimilar and non-reactive barrier between otherwise reactive surfaces. The porosity inherent in some types of coating can provide a further benefit, by providing a reservoir for oil which can be helpful during periods of temporary lubricant starvation.

Practical diagnosis of adhesive wear

Adhesive wear is responsible for the well-known and characteristic condition of seizure, which can occur in closely fitting rubbing metal surfaces, but if clearance is sufficient to expel wear debris, wear can continue and results in significant loss of size to the surfaces involved. This is evident in Fig. 2.10 which shows the worn pin of a link chain. Corrosion and temperature can play an important role in increasing or decreasing adhesive wear, depending on the influence of the surface films that they produce.

Fretting is often characterised by the presence of metal oxide powder in a dismantled joint, and may be associated with significant amounts of wear. Figure 2.11 illustrates wear scars 500μm deep on a steel shaft caused by two 6mm wide piston-type seal rings. The adjacent three marks were caused by the lands on the ring carrier. In this example the assembly was exposed to considerable vibration in an axial flow compressor driven by a 5MW motor. There is no reciprocating or rotary movement between rings and shaft.

The key points regarding adhesive wear are given in Table 2.4.

Fatigue

Several types of wear condition can give rise to fatigue failures. Sometimes relatively large pieces of material are lost from the surface without prior warning, since the damage has been growing beneath the surface.

Contact fatigue

Contact fatigue (also known as surface fatigue) occurs in surfaces in rolling contact, such as ball and roller bearings and gears. During rolling, surfaces are exposed to cyclic stress conditions and, if the endurance limit is exceeded, fatigue failure eventually occurs. The stress distribution in the bearing surface underneath a ball or roller results in a maximum shear stress just beneath the surface of the material. Cracks may initiate from this point, and propagate through the material. They may join up to isolate pieces of
Adhesive wear damage to surfaces of pins in a double row link chain of 25.4 mm pitch, after 11 000 cycles of movement through 90° over a pulley at a frequency of 0.5 Hz. Note the difference in wear between the upper and lower pins, because of wear debris trapped between the lower pin and bush (courtesy CEGB).
Mechanisms of wear and corrosion

Table 2.4 Adhesive wear

- Adhesion of surfaces can take place and cause significant wear damage only if the asperities of the opposing surfaces make metal-to-metal contact.
- Adhesive wear is characterised by sharp transitions in behaviour, and wear may change from mild to severe (and back again) with only slight changes in load or sliding speed.
- For the majority of engineering materials, there is no evidence that hardness can be directly linked to adhesive wear resistance.
- Control of adhesive wear is usually achieved by one (or a combination) of three methods: lubrication, use of mutually incompatible materials in the wear couple and/or application of surface treatments/coatings that interpose a non-reactive layer between the metallic surfaces in relative motion.

Fracture, under these circumstances, takes place by a normal fatigue process.

Failure occurs suddenly, after an incubation period during which there is little (if any) loss of performance. Direct contact of the opposing surface is not necessary, as the stresses can be transmitted through an intervening material such as a lubricant film.

The location of the failure should be defined by the position of maximum shear stress (Fig. 2.12), but, in practice, it is greatly influenced by the presence of defects in the material. Wear caused by contact fatigue is assessed in terms of useful life, defined as the number of revolutions under a given stress that will be exceeded by 90% of the test components.

Contact fatigue can be classed under two headings, depending upon whether cracking has initiated at the surface or within the material. The former tends to occur with hard components in mutual contact in the presence of rolling and sliding. This type of failure is known as pitting and the particles of debris produced are usually triangular in shape. Failure
2.12 Variation of shear stress with distance below the surface.

originating below the surface generally stems from voids, inclusions or hard particles. Cracks propagate parallel to or beneath the surface and subsidiary cracks appear, which if they join up can give rise to loose particles that may fall out. With case-hardened material, if there is insufficient depth of hardening, it is not unusual for cracking to be located at the interface between case and core of the component.

**Erosion**

This was first discussed in the section on abrasion. When a stream of solid particles is in relative motion to a surface, and the angles of impact are much less than 90°, the wear mechanism is likely to be abrasion. However, if impact is at near-normal angles, the mechanism of wear can become fatigue, owing to the continual bombarding effect of the particles.

**Cavitation erosion**

Cavitation erosion is a particular form of contact fatigue, where a solid and a fluid are in relative motion. Bubbles formed in the fluid, which contain only the saturated vapour of the fluid, are unstable and implode against the surface of the solid, causing a strong compression wave. The stability of the bubble depends on the internal and external pressure differential, and on its surface energy, which is a measure of the potential damage on collapse. Cavitation erosion is commonly found on propellers, hydraulic machinery and turbines.

Another special form of erosion is produced when an electric spark occurs between two surfaces and causes permanent damage; this type of spark erosion produces problems for electrical contacts.

**Practical diagnosis**

Contact fatigue can be recognised by the loss of relatively large pieces from bearings and other surfaces in rolling contact. Failure of this type is
frequently catastrophic, as the loss of lubrication and normal alignment of parts can cause widespread mechanical damage to other working parts.

Cavitation can be characterised by an extremely rough surface, consisting of connected pits produced by fatigue fracture of small particles progressively from the surface.

**Control of contact fatigue**

Possibly the biggest single factor in contact fatigue wear is the load. Even a small reduction greatly increases the life of a part. In terms of material properties, an increase in hardness may increase pitting resistance. Further, particularly in respect of surface-initiated fatigue, surface finish is a determining factor. However, too great an increase in hardness can militate against the surface adjustment which is desirable when running-in. Compromise is therefore necessary and this commonly involves manufacturing one of the contacting surfaces slightly softer than the other. With subsurface fatigue, hard materials are again desirable, but they become less conformable as hardness increases. Yet again, a compromise is effected by making the surface only as hard as is required.

Surfacing materials and processes may offer a solution to this type of wear problem, but selection may be difficult if the failed part itself is a material with good fatigue properties. The best properties for this purpose, available from surface coatings, are associated with weld surfacing processes using ferrous alloys which rely on martensitic hardening, either during welding or from subsequent heat treatment.

Lubrication has little direct effect on contact fatigue, as the process depends on the stress versus cycles criterion, and the stress can be transmitted through the lubricant layer. However, it can help indirectly by keeping surface dirt-free and smooth, by eliminating adhesive wear and corrosion effects and by smoothing out local stress concentrations.

**Control of erosion**

In addition to surface treatments, erosion may be reduced by several other measures. The damage done by impacting particles may be reduced by reducing their impact velocity. Reducing the angle of impact is also beneficial. It may be possible to reduce the size of the particles – for example, a probe may be placed in front of the radome of an aircraft to reduce the size of liquid droplets striking the surface.

The key points regarding contact fatigue are given in Table 2.5.

**Corrosion**

Corrosion of metals can be divided into two major categories: reactions with gases and reactions with liquids. While there is a strong chemical similarity between them, the transport mechanisms for the reacting species differ significantly.
Table 2.5 Contact fatigue

- Fatigue occurs suddenly, after an incubation period during which there is little (if any) loss of performance. Direct contact of the opposing surfaces is not necessary. In these respects, contact fatigue differs from abrasive and adhesive wear.
- Failure may take place either at the surface of the component or within the materials. The precise location of failure is greatly influenced by the presence of defects in the components.

Reaction with gases

Gases that can corrode surfaces include oxygen, carbon dioxide, sulphur dioxide, chlorine and hydrogen sulphide. This is not an exhaustive list, and clearly not all metals are susceptible to attack by all gases.

The underlying mechanism involves the metal forming positive ions, e.g.

\[ M \rightarrow M^{n+} + ne \]

where \( n \) is the oxidation state of the metal. Several metals can form more than one positive ion, for instance Fe can readily form Fe\(^{2+}\) and Fe\(^{3+}\). Ionic or covalently bonded solids are then formed, which in the case of oxides of metals ranging from an oxidation state of 1 to 4 are \( \text{M}_2\text{O}, \text{MO} \) and \( \text{M}_2\text{O}_3, \text{MO}_2 \) respectively.

The Gibbs free energy change (or Gibbs function) \( \Delta G_o \), generally quoted in terms of \( \text{kJ}\text{mol}^{-1}\text{(oxygen)} \), determines whether this chemical process is possible. As an example, Fig. 2.13 shows the Gibbs free energy change for a number of common oxides. At all temperatures where the free energy change is negative, oxides may form. Oxides with a large negative Gibbs free energy change are likely to form near-perfect crystal lattices, with few defects.

However, in most cases it is clearly not the Gibbs free energy that determines the extent or the rate of oxidation, since aluminium normally corrodes much slower than iron. The kinetics of the oxidation process usually exert a much stronger influence.

Growth of the oxide layer may depend upon the accessibility of the metal surface to the corroding gas, or on the diffusion of ions and/or electrons. If the compound formed by reaction at the surface is volatile, or continually falls off, it offers no protection for the surface and corrosion will proceed rapidly. However, if the compound at the surface adheres well to the surface, the growth may be limited by the diffusion processes. Ions may diffuse if there are defects in the lattice of the growing compound. Electrons flow by semiconductor mechanisms. This gives rise to three rate-determining mechanisms, which for simplicity use an oxide as an example.

In the first, Fig 2.14(a), metal ions and electrons diffuse faster than oxygen ions. This causes the oxide to grow at the outer surface, and is the mechanism seen in iron, copper, chromium and cobalt. In the second, Fig. 2.14(b), oxygen ions and electrons diffuse through the existing oxide layer faster than metal ions. This causes the oxide layer to grow at the metal/oxide
Mechanisms of wear and corrosion

2.13 Gibbs free energy of formation of some common oxides: (1) \( 2C + O_2 \rightarrow 2CO \); (2) \( 2Ni + O_2 \rightarrow 2NiO \); (3) \( \frac{4}{3}Fe + O_2 \rightarrow \frac{2}{3}Fe_2O_3 \); (4) \( \frac{4}{3}Cr + O_2 \rightarrow \frac{2}{3}Cr_2O_3 \); (5) \( Ti + O_2 \rightarrow TiO_2 \); (6) \( \frac{4}{3}Al + O_2 \rightarrow \frac{2}{3}Al_2O_3 \).

2.14 Mechanisms of oxidation at a surface.

interface, a situation seen in titanium and zirconium. Finally, electron conduction may be rate-determining, and if the oxide is a good insulator the oxide thickness is determined by the distance that electrons can tunnel through it. This is the case with aluminium, and it limits the natural oxide thickness on aluminium to approximately 3 nm.

Oxides formed on the surfaces of metals are frequently non-stoichiometric.
Table 2.6 Typical generic alloys developed for oxidation resistance

<table>
<thead>
<tr>
<th>Alloy</th>
<th>Chief constituents</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aluminium bronze</td>
<td>Copper, with 10% aluminium</td>
</tr>
<tr>
<td>Silicon iron</td>
<td>Iron, with 8% silicon</td>
</tr>
<tr>
<td>Stainless steel</td>
<td>Iron, with 18% chromium and 8% nickel</td>
</tr>
<tr>
<td>Nimonics</td>
<td>Nickel, with 20% chromium</td>
</tr>
</tbody>
</table>

- that is, the metal and oxygen may not be present in exact ratios. For example the oxide on copper is usually Cu$_2$O, instead of Cu$_4$O. Oxide layers also frequently contain the metal in more than one oxidation state, with the highest state nearest the free surface, where the oxygen supply is most plentiful.

As seen in Fig. 2.13, the Gibbs free energy for most oxidation reactions becomes less negative as the temperature is increased, so the thermodynamic driving force for change is reduced as the temperature rises. However, the mechanisms that allow oxidation to proceed generally entail diffusion of one or more species, and overcoming the activation energy of the oxidation reaction itself. Diffusion becomes more rapid as temperature rises, and generally the probability of a reaction taking place also increases. The net effect is that, in most cases, reactions increase in rate as the temperature is increased.

As already mentioned, once formed, the oxide layer may give a degree of protection to the metal surface. For best protection, the oxide should be in compression, have good adhesion to the metal surface, have good cohesion, and have poor conductivity for both electrons and ions. These conditions are well met by oxides formed on aluminium, chromium, silicon, zinc and titanium. This property is used when alloying, in the combinations shown in Table 2.6. However, oxides on copper and iron crack easily, continually exposing fresh surface underneath. The oxides on molybdenum and tungsten vaporise.

Alloying is a relatively expensive way of achieving corrosion resistance, and the large quantities of alloying elements that are required to confer this resistance have a significant effect on the mechanical properties of the material. Hence, these are often the very alloys that are used for surface coatings.

**Reaction with liquids**

Corrosion in liquids is electrochemical, and requires an anode, a cathode and a conducting circuit. The anode is the area that is corroded away, and the cathode receives the electrons from the conducting circuit, which includes the metal and the liquid surrounding it. Note that it is not necessary for the metal to be totally immersed, a single drop of conducting liquid is sufficient for a corrosion reaction.

Typical reactions at anodes are:
Iron Water

Fe²⁺ Fe(OH)₂

Cathodic area

2.15 Corrosion of iron in aerated water.

 Typical reactions at cathodes are:

\[
O_2 + 2H_2O + 4 e^- \rightarrow 4 OH^- \quad \text{(commonest reaction, in neutral solution)}
\]

\[
M'^+ + e^- \rightarrow M \quad \text{(electroplating reaction)}
\]

An extremely important cathodic reaction in steel is

\[
2H'^+ + 2e^- \rightarrow H_2 \quad \text{(in acid solution)}
\]

If this reaction proceeds excessively, hydrogen may dissolve in the steel, and it can cause hydrogen embrittlement.

For example, the corrosion of iron in aerated water proceeds as shown in Fig. 2.15. While some of the corrosion product forms on the surface, it can also form some distance from the surface. In a good electrolyte diffusion is rapid, and hence this corrosion mechanism proceeds rapidly.

Anodes and cathodes arise spontaneously within the material, either on a random basis, or as a result of stress or compositional differences. Common mechanisms that give rise to anodes and cathodes are described under the heading of galvanic cells below.

The products formed at the surface depend on the conditions of pH and potential, and some may have protective value. The equilibrium conditions for a large number of alloys are illustrated in Pourbaix diagrams. These are equilibrium diagrams, but they show where corrosion is feasible and can be a useful aid to the selection of materials. A typical diagram, for iron, is shown in Fig. 2.16.
Table 2.7  Standard electrode potentials measured against a hydrogen electrode

<table>
<thead>
<tr>
<th>Metal reaction</th>
<th>$E_e/V$</th>
<th>Noble and cathodic/Base and anodic</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\text{Fe}^{2+} + e \rightarrow \text{Fe}^{0}$</td>
<td>0.77</td>
<td></td>
</tr>
<tr>
<td>$\text{Cu}^{2+} + 2e \rightarrow \text{Cu}^{0}$</td>
<td>0.34</td>
<td></td>
</tr>
<tr>
<td>$\text{Sn}^{2+} + 2e \rightarrow \text{Sn}^{0}$</td>
<td>0.15</td>
<td></td>
</tr>
<tr>
<td>$2\text{H}^+ + 2e \rightarrow \text{H}_2$</td>
<td>0 (= arbitrary reference point)</td>
<td></td>
</tr>
<tr>
<td>$\text{Pb}^{2+} + 2e \rightarrow \text{Pb}^{0}$</td>
<td>-0.13</td>
<td></td>
</tr>
<tr>
<td>$\text{Sn} \rightarrow \text{Sn}^{2+} + 2e$</td>
<td>-0.14</td>
<td></td>
</tr>
<tr>
<td>$\text{Ni} \rightarrow \text{Ni}^{2+} + 2e$</td>
<td>-0.25</td>
<td></td>
</tr>
<tr>
<td>$\text{Co} \rightarrow \text{Co}^{2+} + 2e$</td>
<td>-0.28</td>
<td></td>
</tr>
<tr>
<td>$\text{Cd} \rightarrow \text{Cd}^{2+} + 2e$</td>
<td>-0.40</td>
<td></td>
</tr>
<tr>
<td>$\text{Fe} \rightarrow \text{Fe}^{3+} + 2e$</td>
<td>-0.44</td>
<td></td>
</tr>
<tr>
<td>$\text{Cr} \rightarrow \text{Cr}^{3+} + 3e$</td>
<td>-0.74</td>
<td></td>
</tr>
<tr>
<td>$\text{Zn} \rightarrow \text{Zn}^{2+} + 2e$</td>
<td>-0.76</td>
<td></td>
</tr>
<tr>
<td>$\text{Al} \rightarrow \text{Al}^{3+} + 3e$</td>
<td>-1.66</td>
<td></td>
</tr>
<tr>
<td>$\text{Ti} \rightarrow \text{Ti}^{4+} + 4e$</td>
<td>-1.63</td>
<td></td>
</tr>
<tr>
<td>$\text{Mg} \rightarrow \text{Mg}^{2+} + 2e$</td>
<td>-2.36</td>
<td></td>
</tr>
</tbody>
</table>

2.16 Pourbaix diagram for iron.

**Galvanic cells**

Anodes and cathodes are required for electrochemical corrosion, and there are many situations in which, by the presence of a combination of different materials, or microstructural features, a galvanic cell may be present.

As a rule, the more anodic a metal, the greater its tendency to corrode. In theory, this is determined by the electrode potential of the metal, and examples of the electrode potentials of common metals are shown in Table 2.7. While this series gives the theoretical basis for corrosion, it is complicated by the fact that some metals form passive films, that the electrode potential depends on the electrolyte, and that the actual corrosion rate will depend on the medium. For this reason, there are ‘galvanic series’ compiled for various media. As an example, Table 2.8 shows the galvanic series for seawater.

Dissimilar metals in electrical contact with one another in a conducting...
### Table 2.8 Galvanic series in sea water

<table>
<thead>
<tr>
<th>Galvanic series</th>
</tr>
</thead>
<tbody>
<tr>
<td>Increasingly cathodic (protected)</td>
</tr>
<tr>
<td>Graphite</td>
</tr>
<tr>
<td>Titanium</td>
</tr>
<tr>
<td>Silver</td>
</tr>
<tr>
<td>18-8 Stainless steel, in presence of plentiful oxygen supply</td>
</tr>
<tr>
<td>Monel/bronzes/copper/brasses</td>
</tr>
<tr>
<td>Nickel</td>
</tr>
<tr>
<td>Tin</td>
</tr>
<tr>
<td>Lead</td>
</tr>
<tr>
<td>18-8 Stainless steel, in absence of oxygen</td>
</tr>
<tr>
<td>Steel/cast iron</td>
</tr>
<tr>
<td>Decreasingly anodic (corroded)</td>
</tr>
<tr>
<td>2024 Aluminium</td>
</tr>
<tr>
<td>Cadmium</td>
</tr>
<tr>
<td>Aluminium</td>
</tr>
<tr>
<td>Zinc</td>
</tr>
<tr>
<td>Aluminium</td>
</tr>
<tr>
<td>Magnesium</td>
</tr>
</tbody>
</table>

\[
\text{Zn}^{2+} + 0.5 \text{O}_2 + \text{H}_2\text{O} + 2\text{e}^- = 2\text{OH}^- + \text{Zn}^{2+}
\]

\[
\text{tin} \\
\text{steel} \\
2\text{e}^- \leftrightarrow 2\text{e}^- \\
\]

\[
0.5 \text{O}_2 + \text{H}_2\text{O} + 2\text{e}^- = 2\text{OH}^- \\
0.5 \text{O}_2 + \text{H}_2\text{O} + 2\text{e}^- = 2\text{OH}^- \\
\text{tin} \\
\text{steel} \\
\text{Fe} = \text{Fe}^{2+} + 2\text{e}^-
\]

**2.17** Corrosion in (a) zinc and (b) tin-plated steel.

The solution will, because of their differing electrode potentials, set up a galvanic cell, in which one of the metals is likely to corrode. This is illustrated in Fig. 2.17, which shows the corrosion mechanisms on zinc- and tin-plated steel.

The same mechanism can operate on the microstructural scale, where different solid phases coexist. An example is illustrated in Fig. 2.18, which shows the mechanism by which brass loses zinc, pearlite loses ferrite, and iron is leached from cast iron since it is anodic with respect to carbon. Grain boundaries tend to be regions of higher energy than the bulk, and therefore tend to be more anodic than the bulk grains. This leads to etching away of the grain boundaries.
The following suggestions may prevent corrosion by galvanic cells:

1. Avoid electrical contact – provide washers.
2. Use coatings to build up electrical resistance.
3. Provide sacrificial protection, e.g. Al, Zn, Mg on iron.
4. Impress an electric potential to suppress the corrosion current – this is expensive to set up.
5. Provide inhibitors: anodic inhibitors, such as phosphates, chromates, nitrites; cathodic inhibitors, such as Mg²⁺, Zn²⁺, Ca²⁺ in solution, which form hydroxide precipitates on the cathode.

Concentration cells
Oxygen dissolved in the liquid may play a key role in corrosion, particularly if its concentration in the liquid varies significantly. Areas with a plentiful supply of oxygen tend to be cathodic. Many factors can lead to a situation where the oxygen concentration varies significantly, for example, crevices, gaskets and joints tend to be depleted of oxygen.

The following suggestions may prevent corrosion by concentration cells:

1. Weld joints (do not use rivets).
2. Use caulking compounds to seal joints.
3. Design to avoid stagnant regions.

Mixed attack mechanisms
In practical situations, both wear and corrosion may proceed together. The effects may be considerably worse than the sum of the two mechanisms of attack in isolation. The role of applied stress, or residual stress, is also significant.

Abrasion/corrosion
When sliding takes place in a corrosive environment, surface reactions occur and reaction products are formed on one or both surfaces. Various
types of reaction product can be formed. In dry air at elevated temperatures, the product is an oxide and in moist air an oxide or hydroxide. With some metals the product may be a carbonate, and chlorides, sulphides and nitrates are common in industrial atmospheres. In all of these, the reaction product tends to adhere poorly to the surface and further rubbing removes it mechanically. In this way, bare metal surfaces are continuously exposed and the medium may appear much more aggressive than it is under static conditions. Fretting and erosion can be particularly acute combined with corrosion.

The presence of a lubricant can have a beneficial effect on the system for two reasons. First, it may protect surfaces from the corrosive media and reduce the frictional heating; secondly the lubricant itself may react with the surface to produce a different form of reaction product. This latter effect is deliberately promoted with EP additives in oil, in which an active chlorine or sulphur compound is used to form anti-scuff and anti-seizure coatings.

**Stress corrosion**

In some materials and environments, cracks may grow at a stress level below the critical level predicted by simple fracture mechanics by a mechanism that is a combination of stress and local corrosion attack. Under such conditions, a structure that was safe when new becomes unsafe with time. It is observed in a wide range of alloys, and some of the most common combinations of material and environment to be aware of are shown in Table 2.9. Solutions to this problem include reducing the stress, or choosing an alternative material.

**Hydrogen-induced cracking**

Hydrogen tends to lock dislocations and make plastic flow more difficult, rendering the metal more brittle as a result. Hydrogen can diffuse through
metals, and if internal cavities exist it is possible for the atomic hydrogen to form molecular hydrogen within the cavity. This can cause high pressures within, and can lead to blisters. A metal embrittled by hydrogen can crack if it is subjected to tensile stresses. Occasionally it will take place without an external source of stress.

The source of the hydrogen may be corrosion of the metal itself, or of a baser material in direct contact with it – for example, cracking in zinc-galvanised high strength steel. Alternatively the hydrogen may have entered the steel during manufacturing, such as pickling, cathodic cleaning, electrodeposition or welding processes.

**Corrosion fatigue**

The normal fatigue process consists of the progressive growth of a crack. This is strongly promoted by corrosion, since the corrosion can attack the extrusions and intrusions on the metal surface and prevent work-hardening and strengthening of the material. It is seen in both pure metals and alloys, and is not linked to specific environments – it occurs sooner or later in all corrosive environments. This mechanism may reduce the fatigue life of a steel component in saltwater by a factor of 4.

**Elevated temperature**

The temperature of the rubbing surface is important in wear, and elevated temperatures can have a number of effects on wear performance. Strength and hardness of metallic materials decrease with increasing temperature, and the reduction can be rapid as temperatures become high. The relative strength of any adhesive joints formed become much greater under these conditions and this increases the likelihood of surface damage. Materials in the wear couple are often specified to have high hot hardness to counteract these effects; tool steels and alloys based on cobalt, with chromium and molybdenum or tungsten are commonly used. Plastics, and plastic based materials, suffer greater loss of strength properties than metals when heated. Ceramics, and some cermet, retain hardness at high temperature better than metals or plastics and, for highest temperature operation, ceramics (or possibly cermet) may be the most appropriate choice, provided the lack of ductility can be accepted.

Surface-contaminating films are affected by temperature. The formation of different types of oxide at different temperatures has already been discussed. Another effect is the removal, by heating, of adsorbed surface films – usually moisture or organics – whose presence tends to prevent severe metallic wear taking place.

Lubricants such as oil suffer degradation at quite moderate temperatures (<300°C) and even synthetic oils cannot be used for lengthy periods at much higher temperatures. Above about 500°C, some solid lubricants may be suitable but, again, their use is limited by chemical factors.

The effects of elevated temperature on wear are generally difficult to predict with accuracy but usually it would be expected to make the wear rate greater.
Testing for wear resistance

Reliable testing for wear resistance is notoriously difficult. Although many wear tests have been carried out, on a variety of machines, results cannot always be used to predict behaviour in service. The conclusion that seems to have been established is that test equipment should simulate the service conditions of the component as closely as possible. This is understandable if the wide variations in service conditions are considered. For example, excavator bucket teeth are subjected to non-metallic abrasion at ordinary temperatures, conditions totally different from those arising, say, in the shearing of hot steel billets. Generally, results obtained with a test machine, which carefully simulates certain specific service conditions, do bear a relationship to service results and the machine can be used to test different alloys to sort them out. For example, rubbing blocks on disks to test screw flight/barrel combinations for plastic processing equipment resulted in marked differences. Three barrel materials – martensitic white iron 58–64 RC iron; a corrosion-resistant Co–Ni–Cr–B alloy 48–54 RC; and a composite of tungsten carbide particles dispersed in a hard, corrosion-resistant Ni base matrix – were run against four surface-treated steels and two surfacing alloys. The results are shown in Table 2.10.

The tests clearly show that there is no direct relationship of wear resistance to material hardness and that the compatibility of any two materials together is not related to their relative hardness, but the data are useful if used judiciously as a guide to the better alternatives. Adhesive wear test results cannot provide the only criterion to material selection – corrosive conditions must be considered, and also material strength requirements. Thus although the case-hardened 4620 steel shows excellent compatibility with the first two barrels, it may not have sufficient strength for many applications. Past service experience with the Co base alloy (an intuitive choice) has shown results better than indicated, in the majority of cases. Where problems have occurred, the tests can suggest probable improvements.

Three tests are commonly used to assist in ranking materials: dry sand on rubber wheel, ASTM G65; erosion test, ASTM G76; and the pin-on-disc test, ASTM G99.

Dry sand/rubber wheel abrasion test, ASTM G65

This test involves the abrading of a standard test specimen with a grit of controlled size and composition. The abrasive is fed at a controlled flow rate into the gap between the test specimen and a wheel which has a rubber rim of a specified hardness. The test specimen is pressed against this wheel with a specified force, and the wheel rotates in the direction of the abrasive flow. The wear rate is measured as a loss of mass from the test specimen in a specified time period. The principle is illustrated in Fig. 2.19(a).

Erosion test, ASTM G76

This test delivers a stream of gas containing abrasive particles, which impact on the test specimen. The abrasive particles are introduced into a gas flow,
Table 2.10 Block on disc adhesive wear tests for various screw flight/barrel combinations of materials

<table>
<thead>
<tr>
<th>Barrel material (B)</th>
<th>Screw material (M)</th>
<th>Martensitic white iron 58–64 RC</th>
<th>Corrosion resistant Co–Ni–Cr–B alloy 48–54 RC</th>
<th>Composite WC in hard, Ni base matrix</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Weight loss (M/mg)</td>
<td>Weight loss (B/mg)</td>
<td>Scar width/mm</td>
<td>Weight loss (M/mg)</td>
</tr>
<tr>
<td>Co–Cr–W–C surfacing alloy (40 RC)</td>
<td>7.6</td>
<td>2.4</td>
<td>2.01</td>
<td>7.35</td>
</tr>
<tr>
<td>Ni–Cr–Si–B surfacing alloy (60 RC)</td>
<td>1.2</td>
<td>2.3</td>
<td>2.00</td>
<td>0.4</td>
</tr>
<tr>
<td>Carburised SAE 4620 steel</td>
<td>3.3</td>
<td>1.5</td>
<td>1.83</td>
<td>3.8</td>
</tr>
<tr>
<td>Hardened SAE 4140 steel</td>
<td>27.05</td>
<td>1.1</td>
<td>1.68</td>
<td>27.1</td>
</tr>
<tr>
<td>Nitrided SAE 4130 steel</td>
<td>7.35</td>
<td>1.4</td>
<td>1.79</td>
<td>8.0</td>
</tr>
<tr>
<td>Nitrided Nitralloy</td>
<td>7.0</td>
<td>2.0</td>
<td>2.04</td>
<td>4.05</td>
</tr>
</tbody>
</table>

Loads used in the tests represented extreme conditions. There was no interfacial film between barrel and screw materials, as in actual use.
Mechanisms of wear and corrosion

Rubber-lined wheel

(a)

2.19 Wear test geometries: (a) sand abrasion; (b) erosion.

and fed through a nozzle to strike the specimen. There must be a means of measuring and controlling particle velocity and particle flux. The abrading material must be uniform in such respects as particle size, moisture, chemical composition, etc. The principle is illustrated in Fig. 2.19(b).

Pin-on-disc test, ASTM G99

Two materials are required for this test, which tests the wear of one material against another, under nominally non-abrasive conditions. In this test a specimen (pin) with a radiused tip is pressed using a known force against the surface of a rotating disc of the second material. The pin describes a circular path on the disc surface. The wear rate is measured for both the pin and the disc as a function of sliding distance.
A variant of this test uses an abrasive wheel as the disc. This abrasive wheel must be of a specified composition and a known grit size, and its speed of rotation is controlled. It is usual to introduce a lubricant, such as water, to enable the wear debris to wash away from the surface under test. The wheel must also be maintained in a well-dressed condition. The wear rate may be measured as a mass loss, or a thickness loss, from the disc, depending upon the preference of the user. A photograph of a typical apparatus is shown in Fig. 2.20.

### Material selection for wear resistance

The harder the surface, the better the resistance against abrasion. This is true when dealing with single phase materials, but most alloys contain two or more phases and many of the surfacing alloys have complex structures with hard phases dispersed in a softer matrix. Hardness of constituent parts, their distribution and size have to be considered in relation to the operating regime. Generally, the macrohardness of the material to resist abrasion should be at least 80% of the hardness of the abradent.

In industrial practice greatest resistance to wear has been found in materials containing the maximum amount of the hardest carbides or borides which can be supported adequately in a suitable matrix. The
balance between the amount and size of the hard particles and the properties of the matrix must be related to the wear conditions, which may be metal-to-metal at high or low velocity, fretting, abrasion by lightly or heavily loaded particles of small or large size, impact or erosion by liquid or gas. Any of these factors may be complicated by a hot or corrosive environment.

For many applications involving abrasive wear, tungsten carbide is a preferred material. The greater the percentage of carbide, the better the wear resistance, so massive carbide with 6–25% Co as the binder (percentage adjusted to the degree of toughness needed) or applied as a coating, although expensive, may justify the cost. However, in an application in which the abrasive is fine, performance may be poor and far below a coating with only perhaps 30–50% tungsten carbide in a matrix of an Ni-based alloy. This is because the soft Co phase may be eroded, leaving the unsupported carbide to fall out, while the harder Ni alloy resists the erosion better.

Often trials on the plant must be made to determine the best surfacing solution. Results reported by an iron and steel company on protection of breaker tips in the sinter plant show the improvements achieved. The chosen method involved use of a chromium and tungsten carbide with a life about double that of the minimum tolerable interval between maintenance schedules. To obtain maximum wear resistance, tubular tungsten carbide rods were deposited by gas welding as this minimises solution of carbide in the steel matrix alloy compared with arc welding and this gives maximum wear life. With the introduction of a high percentage of foreign ore into the sinter mix, wear increased considerably. To reach tolerable maintenance levels extensive use of tungsten carbide became essential; even then results were not as good as with the less abrasive home ore (see Table 2.11).

Another method of deposition was used for tests on nozzle guide vanes which suffered wear in aeroengines flying around Iceland where the atmosphere contains much volcanic ash. The detonation gun-deposited tungsten carbide test areas showed excellent resistance so adequate surface protection is now designed into the sections vulnerable to wear. This emphasises the desirability of examining wear patterns as part of the design process.

<table>
<thead>
<tr>
<th>Sinter practice</th>
<th>Breaker tip type</th>
<th>Service life</th>
<th>Initial cost</th>
<th>Cost per tonne sinter/tonne</th>
<th>Sinter output/Mt</th>
</tr>
</thead>
<tbody>
<tr>
<td>Home ore</td>
<td>26–28% Cr tip</td>
<td>18</td>
<td>2500</td>
<td>0.20</td>
<td>1.25</td>
</tr>
<tr>
<td>Foreign ore</td>
<td>26–28% Cr tip</td>
<td>9</td>
<td>2500</td>
<td>0.42</td>
<td>0.6</td>
</tr>
<tr>
<td></td>
<td>Hardfaced with</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>high tungsten</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>carbide</td>
<td>12–14</td>
<td>4500</td>
<td>0.50</td>
<td>0.9</td>
</tr>
</tbody>
</table>

Table 2.11 Typical maintenance levels required by hard surfaced sinter plant
With the work referred to above, the more expensive tungsten carbide was applied only to the areas experiencing greatest wear.

**Testing for corrosion resistance**

As with wear testing, there are several tests available, and it is important to choose a test that reflects the conditions that the component will experience in service if the results are to be of greatest usefulness.

**Humidity cabinets**

In a humidity cabinet the humidity is raised to a value chosen as appropriate to the material under test. The temperature is generally cycled, so that the specimen is exposed to alternating humid air and condensation. The apparatus is automated to ensure that conditions are controlled within narrow limits. Other corrodent materials, such as sulphur dioxide, may also be introduced.

**Salt spray cabinets**

In salt spray chambers specimens are exposed to a mist of sodium chloride solution of a prescribed concentration and temperature. It is important to direct the salt spray so that it does not hit the specimens directly. Additions to the salt solution, such as ethanoic acid, or copper chloride, can increase the rate of corrosion of the specimens, so that a test for 16 hours can be equivalent to normal atmospheric exposure 100 times as long. A typical test is described in ASTM B3680-68, and BS 1224.

**Electrochemical methods**

Since corrosion is an electrochemical process, there are several corrosion tests that are based on electrical methods. These all involve the evaluation of the corrosion current, which gives a measure of the rate of loss of material from the specimen. One method is to measure the current in an electrochemical cell as a function of polarisation.

A potentiostatic or potentiodynamic method may be used to record the anodic polarisation curve; see Fig. 2.21. This is a particularly useful method
Mechanisms of wear and corrosion

for the development of new alloys. The current peak at the passivation potential defines the tendency of the alloy to become passive, and the passivity current indicates the corrosion rate in the passive state.

Material selection for corrosion resistance

As already noted, several metals and alloys are inherently corrosion-resistant – such as steel alloys containing sufficient chromium to maintain a self-healing chromium oxide layer on the surface, and nickel or copper alloys. These are good choices for corrosion-resistant coatings. However, if they are cathodic to the metal underneath, they will only maintain their corrosion protection when the coating is undamaged.

Zinc and aluminium are commonly used as coatings which offer sacrificial protection to iron and mild steel in aqueous environments, and these have the advantage that damage to the coating does not lead to the immediate corrosion to the material underneath.

Lead coatings are used in many applications, since lead resists attack by many chemical substances, including most acids.

Coatings derived from ceramics, such as alumina, titania and other oxides, silicides, aluminides or polymers, offer protection owing to their inherent corrosion resistance. Most are not only corrosion-resistant themselves, but also form a physical barrier preventing access of the corroding material to the metal surface.
Chapter 3

Weld surfacing

Characteristics of weld surfacing
The principal characteristics of welding processes which distinguish them from other methods of surface coating are as follows.

Coating thickness
Welded deposits of surfacing alloys can be applied in thicknesses greater than most other techniques, typically in the range of 3–10mm, with some restrictions for certain materials. This facility is essential when protection in depth is necessary. Conversely, there are a few welding processes that will produce very thin coatings where necessary.

Adhesion to substrate
Correctly selected materials and properly operated processes provide a metallurgical bond to the substrate which withstands thermal and mechanical shock without detachment. In the fusion welding processes this is achieved by melting a thin layer of the substrate material while applying the coating. In the other processes (friction and explosive cladding) it is achieved by a forging action.

Range of processes
Most conventional welding processes are used for application of surface coatings, bringing the operation within the scope of most engineering activities and enabling on-site work to be carried out under certain circumstances.

Processes have been divided into two broad sections, (a) arc processes, comprising

- manual metal arc,
- tungsten inert gas,
- plasma transferred arc,
- metal inert/active gas,
Table 3.1 Arc weld surfacing processes

<table>
<thead>
<tr>
<th>Process</th>
<th>Abbreviations</th>
<th>Approximate deposit thickness minimum/mm</th>
<th>Deposition rate/kg h⁻¹</th>
<th>Dilution single layer/%</th>
<th>Typical uses</th>
</tr>
</thead>
<tbody>
<tr>
<td>Manual metal arc</td>
<td>MMA</td>
<td>3</td>
<td>1–4</td>
<td>15–30</td>
<td>Multilayers on heavier sections</td>
</tr>
<tr>
<td>Tungsten inert gas</td>
<td>TIG, GTAW</td>
<td>1.5</td>
<td>≤2</td>
<td>5–10</td>
<td>High quality, low dilution work</td>
</tr>
<tr>
<td>Plasma transferred arc</td>
<td>PTA</td>
<td>2</td>
<td>≤10</td>
<td>2–10</td>
<td>High quality, lowest dilution work</td>
</tr>
<tr>
<td>Metal inert gas</td>
<td>MIG, GMAW</td>
<td>2</td>
<td>3–6</td>
<td>10–30</td>
<td>Faster than MMA, no stub end loss, positional work possible</td>
</tr>
<tr>
<td>Flux-cored arc</td>
<td>FCAW</td>
<td>2</td>
<td>3–6</td>
<td>15–30</td>
<td>Similar to MIG, Mainly for iron base alloys for high abrasion resistance</td>
</tr>
<tr>
<td>Submerged arc (wire)</td>
<td>SAW</td>
<td>3</td>
<td>10–30</td>
<td>15–30</td>
<td>Heavy section work; higher quality deposits than FCAW</td>
</tr>
<tr>
<td>Submerged arc (strip)</td>
<td>SAW</td>
<td>4</td>
<td>10–40</td>
<td>10–25</td>
<td>Corrosion-resistant cladding of large areas</td>
</tr>
<tr>
<td>Submerged arc (bulk welding)</td>
<td>SAW</td>
<td></td>
<td></td>
<td></td>
<td>Similar to SAW wire but other alloys possible</td>
</tr>
</tbody>
</table>

- flux-cored arc,
- submerged arc,

and (b) other processes including friction and forge processes, comprising

- oxyacetylene welding,
- powder welding,
- laser cladding,
- electroslag welding,
- resistance welding,
- friction surfacing,
- explosive cladding.

The key characteristics of the processes are summarised in Tables 3.1 and 3.2.
### Table 3.2 Other fusion welding processes and forge welding processes

<table>
<thead>
<tr>
<th>Process</th>
<th>Abbreviations</th>
<th>Approximate deposit thickness (mm)</th>
<th>Deposition rate (kg h⁻¹)</th>
<th>Dilution single layer (%)</th>
<th>Typical uses</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oxyacetylene OA</td>
<td>1.5</td>
<td>≤1</td>
<td>1–5</td>
<td></td>
<td>Small area deposits on light sections</td>
</tr>
<tr>
<td>Powder weld PW</td>
<td>0.1</td>
<td>0.2–1</td>
<td></td>
<td></td>
<td>Small area deposits on light sections</td>
</tr>
<tr>
<td>Laser cladding</td>
<td>0.25</td>
<td>&lt;3</td>
<td></td>
<td></td>
<td>High quality, low dilution coatings</td>
</tr>
<tr>
<td>Electroslag ESW (strip)</td>
<td>4</td>
<td>15–35</td>
<td>5–20</td>
<td></td>
<td>High quality deposits at higher deposit rates than SAW. Limited alloy range</td>
</tr>
<tr>
<td>Friction surfacing</td>
<td>None</td>
<td></td>
<td></td>
<td></td>
<td>Inlay cladding, knife blades</td>
</tr>
<tr>
<td>Resistance welding</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Repair of polymer, rubber and diecast moulds</td>
</tr>
<tr>
<td>Explosive cladding</td>
<td>0.011</td>
<td>None</td>
<td></td>
<td></td>
<td>Flat plate cladding, e.g. food trays, storage tanks</td>
</tr>
</tbody>
</table>

The spray fuse process is covered in Chapter 4.

### Resurfacing

The opportunity to carry out repairs on worn parts, whether previously weld surfaced or not, is a feature exploited in several industries. Welding processes involve application of some heat to the component being processed, and depending on the material from which it is made and its condition, certain precautions may need to be taken. These are described in detail in this chapter.

During fusion weld surfacing, the coating material is raised to its melting point, which means that metals and alloys used for the purpose must have a melting point similar to, or less than, that of the substrate materials. Exceptions to this general rule are found in friction surfacing and explosive cladding. Other coating materials with higher melting points, such as ceramics, may be applied by thermal spraying processes, which are described in the following chapter.
Materials for weld surfacing and their selection

Most materials used for weld surfacing have hardnesses greater than 200HV and for this reason they are often referred to as hardfacing or hardsurfacing alloys. Because of the large number of these materials it is convenient to classify them into groups. Such a system developed by British Steel is shown in Table 3.3, which gives details of the composition and typical uses of each type.

Table 1.1 is also based on the British Steel system, but is in simpler form and incorporates surfacing materials that can only be applied by other surfacing processes. This has been done to simplify the selection of a material to suit given service conditions and to identify the application processes for which it is suited.

Group 1 materials

Carbon steels

Plain carbon steel, containing up to 0.5% carbon, can be deposited by gas or arc welding processes to produce a weld deposit of about 240HV (20RC). The deposit can subsequently be heat treated to a higher hardness if required. The main application of this type of alloy is building up components for subsequent flame hardening, or for use as a buffer layer between a softer parent metal and a harder surfacing alloy. Few manufacturers of consumables supply this type of filler metal because its abrasion resistance is low and buffer layers can be made with hydrogen-controlled basic covered electrodes which are readily available.

Characteristics

High tensile and compressive strength, high impact strength, low abrasion resistance, low resistance to tempering. The maximum service temperature is 200°C.

Applications

Building up gear teeth, etc., for subsequent flame hardening and buffer layers.

Low alloy steels

Martensitic alloy steels are the most widely used hardfacing alloys and are characterised by low cost and a wide range of properties depending on composition. They contain, in addition to carbon, varying amounts of chromium, manganese, molybdenum and nickel, as well as smaller additions of tungsten and vanadium.
<table>
<thead>
<tr>
<th>Classification</th>
<th>Typical composition, %</th>
<th>Notes</th>
<th>Deposited Vickers hardness applicable to deposit</th>
<th>Characteristics applicable to deposit</th>
<th>Typical applications</th>
</tr>
</thead>
<tbody>
<tr>
<td>Group Type</td>
<td>Fe  C  Cr  Mn  Mo  V  W  Co  Ni  B  Nb  Cu</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 Steels</td>
<td>Bal 1.3 5</td>
<td>Up to 250, responds to heat treatment</td>
<td>Tough, crack free, machinable, low wear but good impact resistance, maximum build-up and buttressing layers, by gas or electric arc welding</td>
<td>Repair of steel components, build-up, alternate layers in laminated surfaces</td>
<td></td>
</tr>
<tr>
<td>2 Low alloy steels</td>
<td>Bal 0.1 0.5</td>
<td>Elements indicated up to 250, responds to heat treatment</td>
<td>Properties depend on composition: wide range mainly martensitic structure. Higher alloys: harder and more wear-resistant, but lower alloys give toughness and impact resistance – generally machinable – low crack susceptibility may require pre- and post-heat treatment</td>
<td>Punches, dies, earthmoving equipment, gear teeth, railway points</td>
<td></td>
</tr>
<tr>
<td>3 Martensitic chromium steels</td>
<td>Bal 0.1 0.7 10 15</td>
<td>Elements indicated up to 350-650, responds to heat treatment</td>
<td>Improved wear resistance over class 1/2 with increased oxidation and corrosion resistance. Medium impact resistance, decreasing with high C types.</td>
<td>Suitable for metal to metal wear up to 600°C. High C types suitable for hot work tool applications. Shear blades, mill rolls, roll necks, crane</td>
<td></td>
</tr>
</tbody>
</table>
May require wheels, hot preheat work dies and punches.

Higher carbide types provide good wear resistance, moderate impact resistance. Suitable for elevated temperatures. Lower carbide types for higher impact. Generally grindable; anneal to machine.

Used as a ductile buttressing layer, e.g. when depositing high Mn steels on to carbon steel base to avoid brittle bend zone. Furnace parts, chemical plants, heavy impact applications. Crusher and excavator equipment, railway points and crossing, crusher hammers.

### 4 High speed steels

<table>
<thead>
<tr>
<th>Bal</th>
<th>0.3</th>
<th>10</th>
<th>15</th>
<th>3</th>
<th>20</th>
<th>12</th>
<th>5</th>
</tr>
</thead>
</table>

600–750 hot hardness to 600 °C

### 5 Austenitic stainless steels

<table>
<thead>
<tr>
<th>Bal</th>
<th>0.0</th>
<th>17</th>
<th>32</th>
<th>7</th>
<th>22</th>
<th></th>
</tr>
</thead>
</table>

Mo and Mn total up to 10%

Tough, high corrosion and heat-resistant with low abrasive wear resistance. Impact resistance lower than class 1/6

As-deposited 200 up to 500 on work hardening

Used as a buttressing layer, e.g. when depositing high Mn steels on to carbon steel base to avoid brittle bend zone. Furnace parts, chemical plants, heavy impact applications. Crusher and excavator equipment, railway points and crossing, crusher hammers.

### 6 Austenitic manganese steels

<table>
<thead>
<tr>
<th>Bal</th>
<th>0.5</th>
<th>11</th>
<th>16</th>
<th></th>
</tr>
</thead>
</table>

Cr, Ni, Mo and V total up to 10%

Tough, impact-resistant, work hardens under heavy impact. Base metal cooling necessary during welding to reduce likely carbide embrittlement. Working temperatures not to exceed 200 °C. Buttering may be desirable on carbon steel base metals. Arc weld applications only.
<table>
<thead>
<tr>
<th>Classification</th>
<th>Type</th>
<th>Typical composition, %</th>
<th>Notes</th>
<th>Deposited characteristics applicable to deposit</th>
<th>Characteristics applicable to deposit</th>
<th>Typical applications</th>
</tr>
</thead>
<tbody>
<tr>
<td>Group</td>
<td>Type</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>Austenitic</td>
<td>Bal 0.3 12 12, 2 Bal 0.5 15 15</td>
<td>Mo, V and Ni total up to 4%</td>
<td>Similar to class 1/6 but high chromium inhibits carbide embrittlement, thus can be deposited on to carbon steels; not restricted to 200 °C working temperatures. More abrasion-resistant than class 1/6</td>
<td>Crusher and excavator equipment, railway points and hammers</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>1</td>
<td>Bal 4 12 20</td>
<td>300–600, responds to heat treatment</td>
<td>High abrasion resistance, moderate impact resistance. Grindable, pre- and post-heat to reduce cracking</td>
<td>Buttering layer to chrome irons. Crushing equipment, pump casings, impellers, excavator teeth</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>Martensitic</td>
<td>Bal 1 1 10</td>
<td>500–750, responds to heat treatment</td>
<td>High abrasive wear resistance, low to moderate impact resistance (Ni hard type). Nb improves wear resistance in hot conditions up to 400 °C</td>
<td>Suitable for conformal contacts in adhesive wear situations. Scrapers, bucket tips, forming rolls, cutting tools</td>
<td></td>
</tr>
<tr>
<td>Grade</td>
<td>Chromium</td>
<td>Carbon</td>
<td>Silicon</td>
<td>Maximum</td>
<td>Bal</td>
<td>Applications</td>
</tr>
<tr>
<td>-------</td>
<td>----------</td>
<td>--------</td>
<td>---------</td>
<td>---------</td>
<td>-----</td>
<td>--------------</td>
</tr>
<tr>
<td>5</td>
<td>Bal 2</td>
<td>20</td>
<td>5</td>
<td>40</td>
<td></td>
<td>Extremely high</td>
</tr>
</tbody>
</table>
Table 3.3 (cont.)

<table>
<thead>
<tr>
<th>Classification</th>
<th>Typical composition, %</th>
<th>Notes</th>
<th>Deposited hardness, Vickers HV</th>
<th>Characteristics applicable to applications</th>
</tr>
</thead>
<tbody>
<tr>
<td>Group</td>
<td>Type</td>
<td>Fe</td>
<td>C</td>
<td>Cr</td>
</tr>
<tr>
<td>3 Nickel alloys</td>
<td>1 Nickel</td>
<td>8</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>2 Nickel-copper</td>
<td>3</td>
<td>0.35</td>
<td>2.5</td>
<td>60</td>
</tr>
<tr>
<td>3 Nickel-iron</td>
<td>Bal</td>
<td>2</td>
<td>1</td>
<td>45</td>
</tr>
</tbody>
</table>
### Nickel-molybdenum-chromium-tungsten Alloys

<table>
<thead>
<tr>
<th>Alloy Description</th>
<th>Fe, C, Mo, W, Co, Nb</th>
<th>Mechanical Properties</th>
</tr>
</thead>
<tbody>
<tr>
<td>5 Nickel-chromium-boron</td>
<td>Fe, C, Mo, Si, total up to 10%</td>
<td>200-750</td>
</tr>
<tr>
<td>6 Nickel-molybdenum-iron</td>
<td>Fe, C, Mo, W, Co, Nb</td>
<td>200-300</td>
</tr>
</tbody>
</table>

### Cobalt-chromium-tungsten Alloys

<table>
<thead>
<tr>
<th>Alloy Description</th>
<th>Fe, C, Mo, W, Co, Nb</th>
<th>Mechanical Properties</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Cobalt-chromium-tungsten low alloy</td>
<td>1.4 32</td>
<td>Bal</td>
</tr>
<tr>
<td>2 Cobalt-chromium-tungsten medium alloy</td>
<td>1.7 32</td>
<td>Bal</td>
</tr>
<tr>
<td>3 Cobalt-chromium-tungsten high alloy</td>
<td>3.0 35</td>
<td>Bal</td>
</tr>
<tr>
<td>4 Cobalt-chromium-tungsten-nickel alloys</td>
<td>2.0 25</td>
<td>Bal</td>
</tr>
<tr>
<td>Classification</td>
<td>Typical composition, %</td>
<td>Notes</td>
</tr>
<tr>
<td>-------------------------</td>
<td>------------------------</td>
<td>-------</td>
</tr>
<tr>
<td>Group Type</td>
<td>Fe  C  Cr  Mn  Mo  V  W  Co  Ni  B  Nb  Cu</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>5 Copper alloys</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 Copper, zinc, brasses</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2 Copper, silicon, silicon bronzes</td>
<td>Up to 40% Zn, balance Cu</td>
<td></td>
</tr>
<tr>
<td>3 Copper, aluminium, aluminium bronzes</td>
<td>Up to 4% Si, balance Cu</td>
<td></td>
</tr>
<tr>
<td>4 Copper, tin, Phosphor bronzes</td>
<td>8-15% Al, balance Cu</td>
<td></td>
</tr>
<tr>
<td>6 Tungsten carbide</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 + 0 mesh (+1.7 mm)</td>
<td>Minimum of 40% tungsten carbide normally in an iron base matrix but can be in a copper or cobalt matrix</td>
<td></td>
</tr>
</tbody>
</table>

Classification is according to the tungsten carbide particle size.
The designation -10 + 20 mesh means material which will pass through a sieve of 10 mesh and does not pass through a sieve of 20 mesh.

Figures in parentheses are the corresponding nominal aperture size of sieve in accordance with BS410.

| Chromium boron paste | 1 | 80 | 20 | 750-850 | Fine mesh powder in suitable binder to form a paste. Applied to steel base material and fused by carbon arc, or gas weld techniques. Very hard wear-resistant surfaces, low impact resistance | Medium abrasive sliding wear situations. Chutes, selected blade applications |

Source: after British Steel.
Characteristics
Hardness of 250–800HV (20–62RC). A wide range of combinations of abrasion and impact resistance. Some alloys are heat-treatable, i.e. can be hardened for increased wear resistance or softened for machining.

Applications
Agricultural implements, earthmoving equipment, crushing machinery, gear teeth, railway points and crossings.

Martensitic chromium steels
Martensitic chromium steels containing about 12% chromium have increased heat and corrosion resistance compared with low alloy steels.

Characteristics
Resistance to metal wear at 300–600°C and moderate impact resistance.

Applications
Mill rolls, roll necks, crane wheels, hot work tools.

High speed steels
High speed steels are tool steels that are used to cut metals at high rates and withstand temperatures of up to 600°C without softening.

Characteristics
Higher carbon types have good wear resistance and moderate impact resistance. Lower carbon types have higher impact resistance. They are heat treatable, i.e. can be hardened for increased wear resistance or softened for machining.

Applications
Cutting tools, hot work, ingot tongs, punches, shear blades.

Austenitic stainless steels
Stainless steels have high corrosion and heat resistance, but an important use is 'buttering' carbon or alloy steels before manganese steel is deposited. Buttering avoids formation of brittle phases which occur at the interface between manganese steel and carbon or low alloy steels. Austenitic stainless steels withstand oxidation at temperatures of between 450 and 600°C, depending on composition. Room temperature hardness is low and these steels are unsuitable for abrasion resistance.
Weld surfacing

Characteristics
Tough, but lower impact resistance than austenitic manganese steel. Oxidation resistance up to 600°C, depending on composition. As-deposited hardness is around 200HV, which increases by work-hardening up to 500HV.

Applications
Chemical plant, furnace parts, buttering layer on carbon alloy steels before deposition of manganese steel.

Austenitic manganese steel
Weld metal containing 12–14% manganese has an austenitic structure which is soft (200HV) but which work-hardens at the surface to about 600HV (54 RC) under heavy impact. In the soft, as-deposited condition there is little resistance to low stress scratching abrasion. Thus, for example, the weld metal is steadily lapped away when digging in sandy soil. However, if impact is involved, as with rock crushing hammers, the weld metal deforms and work hardens at the surface, giving high resistance to further impact. This type of deposit also work-hardens if used for digging in soil that contains boulders, and the surface develops resistance to abrasion by soil particles.

Characteristics
High work-hardening capacity and extremely high impact resistance. Becomes embrittled if heated above 400°C. Not suitable for deposition by oxyacetylene welding because of embrittlement on slow cooling. Resistance to embrittlement by heat is increased by addition of 3–5% nickel or 0.5–1.5% molybdenum. Molybdenum-containing deposits have higher yield strength, which gives a greater resistance to deformation. Dilution by carbon or alloy steel parent metal produces a brittle interface, which can be avoided by a buttering layer of stainless steel.

Applications
Rock crushers, pulverising hammers, railway points and crossings, excavating equipment in rocky soil.

Austenitic chromium–manganese steel
Austenitic chromium–manganese steel is used for similar applications to austenitic manganese steel, but has certain advantages in spite of the higher cost. Because of the high alloy content these electrodes can be used to weld directly on to carbon steel parent metal without the formation of a brittle martensitic interface, and also to weld manganese steel inserts directly to carbon steel.
**Characteristics**

High work-hardening capacity, and extremely high impact resistance. Can be deposited directly on to carbon or low alloy steel without the need for a stainless steel buttering layer. Because of high chromium content, the weld metal cannot be cut or gouged with an oxygas flame.

**Applications**

Rock crushers, pulverising hammers, railway points and crossings, excavating equipment in rocky soil.

**Group 2 materials**

**Austenitic, martensitic, high chromium and complex irons**

High chromium, austenitic, martensitic and complex irons often referred to as 'chromium carbide' types, contain about 30% chromium and the microstructure of the deposit consists of chromium carbides in a matrix which can be austenite, martensite or a mixture of both, depending on the composition. These alloys are available in the form of cast rods or steel tubes containing chromium carbide particles, and can also be produced as a chromium steel core wire with alloying additions included in the flux covering. Coils of tubular flux-cored wires are also available for semi-automatic or fully mechanised welding.

**Austenitic irons: characteristics**

High abrasion resistance, moderate impact resistance and non-machinable.

**Austenitic irons: applications**

Crushing equipment, brick, cement and clay processing equipment, impellers, excavator buckets and teeth.

**Martensitic irons: characteristics**

Higher abrasion resistance than austenitic iron, low to moderate impact resistance and non-machinable.

**Martensitic irons: applications**

Crushing equipment, brick, cement and clay processing equipment, excavator buckets.

**High chromium austenitic irons: characteristics**

High abrasion resistance under low stresses, low to moderate impact resistance, oxidation-resistant and hot hardness up to 450°C.
High chromium austenitic irons: applications
Agricultural machinery in sandy soil, chutes.

High chromium martensitic irons: characteristics
High abrasion resistance under low or high stresses, good impact resistance if well supported by rigid parent material, oxidation-resistant, heat treatable and hot hardness up to 450°C.

High chromium martensitic irons: applications
Ball mill liners, impellers for gravel, dredging equipment, mine and quarry screens, sand blasting plant.

High chromium complex irons: characteristics
High abrasion resistance under low or high stresses, good impact resistance if well supported by rigid parent material, oxidation-resistant and maintains hot hardness up to 600°C.

High chromium complex irons: applications
Hot wear conditions, e.g. sintering plant, chutes, pulverisers, screens, steel mill guides.

Group 3 materials

Nickel alloys
Pure nickel, nickel copper and nickel–iron alloys listed are designed to weld cast irons.

Nickel: characteristics
Soft and machinable.

Nickel: applications
Surfacing cast iron, buttering layer on cast iron.

Nickel–copper: characteristics
Soft and machinable.

Nickel–copper: applications
Surfacing cast iron.
Nickel–iron: characteristics
Machinability lower than nickel or nickel–copper but improved by preheat of 150–300 °C.

Nickel–iron: applications
Surfacing cast iron, buttering layer on cast iron.

Nickel–molybdenum–chromium–tungsten
Nickel–molybdenum–chromium–tungsten alloys are used primarily for corrosion resistance but they also have good heat resistance.

Characteristics
Good corrosion and heat resistance, good resistance to erosion by hot furnace gases carrying abrasive particles, good impact resistance and hot hardness up to 500 °C.

Applications
Blast furnace bell and hopper seals. Hot work tools, e.g. die blocks.

Nickel–chromium–boron
The most common nickel-based hard-facing alloys contain chromium, boron and carbon, and their microstructure consists of wear-resistant chromium carbide and borides in a nickel–chromium matrix.

Characteristics
Oxidation resistance up to 850 °C, hot hardness up to 500 °C, good corrosion resistance against steam, saltwater and salt spray, and low impact resistance.

Applications
Valves, seating rings, screw conveyors, pump shafts, chemical plant.

Nickel–molybdenum–iron

Characteristics
Resistant to corrosion by hydrochloric acid, salt spray and alkalis.

Applications
Pumps, valves, chemical plant.
Group 4 materials

Cobalt alloys

The most widely used hardfacing alloys in the non-ferrous group are those based on the ternary system cobalt–chromium–tungsten developed under the trade name of Stellite. Their composition endows the alloys with hot hardness so that they can be used at service temperatures above 600°C with minimum softening or deformation. The presence of chromium forms a closely adherent oxide film which provides oxidation resistance, and this element, together with tungsten, brings about appreciable secondary hardening by the precipitation of carbides. The Stellite alloys have good resistance to erosion, cavitation and adhesive wear, and these properties, combined with good corrosion resistance, enable the alloys to resist the combined effects of different wear types.

Cobalt–chromium–tungsten low alloy: characteristics

Resistant to heat, corrosion and oxidation under impact stresses.

Cobalt–chromium–tungsten low alloy: applications

Exhaust valves for petrol and diesel engines. Fluid flow control valves, hot metal working tools.

Cobalt–chromium–tungsten medium alloy: characteristics

Higher abrasion resistance than low alloy type, but lower resistance to impact or thermal shock.

Cobalt–chromium–tungsten medium alloy: applications

Knives used in paper, carpet and chemical industries.

Cobalt–chromium–tungsten high alloy: characteristics

Highest abrasion resistance and lower shock resistance of this group.

Cobalt–chromium–tungsten high alloy: applications

Pumps, bearing surfaces, seals.

Cobalt–chromium–tungsten–nickel alloy: characteristics

High erosion and corrosion resistance at elevated temperatures, high thermal shock resistance.

Cobalt–chromium–tungsten–nickel alloy: applications

Exhaust valves for petrol engines.
Group 5 materials

Copper alloys
Copper–aluminium alloys are used for bearing, corrosion-resistant and wear-resistant surfaces. Tin bronzes are used for bearing surfaces and occasionally in corrosion-resistant applications. Silicon bronze is used for corrosion resistance only and brasses are used occasionally for bearing surfaces.

Brasses: characteristics
Low hardness, low abrasion resistance.

Brasses: applications
Limited use for bearing applications.

Silicon bronzes: characteristics
Good corrosion resistance.

Silicon bronzes: applications
Overlaying silicon brass.

Aluminium bronzes: characteristics
Good bearing properties, good corrosion resistance.

Aluminium bronzes: applications
Bearing surfaces deposited on steel, slideways, wear plates.

Tin bronzes: characteristics
Good bearing properties.

Tin bronzes: applications
Bearing surfaces deposited on steel slideways.

Group 6 materials

Tungsten carbide
Tungsten carbide has extremely good wear resistance. Hardfacing rods are actually steel tubes containing tungsten carbide particles (both WC and W,C) with the relative amount of tungsten carbide to steel at about 60:40. The carbide particles vary from 200 to 8 mesh (0.002 to 3 mm) but surfacing
rods generally contain different ranges of mesh sizes, e.g. 10/20, 20/30, depending on service conditions. Tubes may be bare for oxyacetylene deposition and either bare or covered for arc welding. Rods are also available in sintered form which contain up to 80% tungsten carbide. Coils of tubular flux-cored wires are also available for semi-automatic or fully mechanised welding.

**Characteristics**

Oxyacetylene or TIG deposits contain unmelted carbides giving extremely high abrasion resistance. Arc welding melts most of the carbide granules giving a homogeneous weld having slightly lower abrasion resistance.

**Applications**

Oxyacetylene or tungsten inert gas (TIG) deposits are used to surface rock cutting tools. Arc weld deposits are used for sand mixer blades, pug mill knives and coal cutter picks.

**Group 7 materials**

**Chromium boride paste**

Chromium boride paste is spread over steel parent material and is fused by carbon arc, manual metal arc (MMA), TIG, metal inert gas (MIG) or plasma welding.

**Characteristics**

Extremely high abrasion resistance.

**Applications**

Chutes.

**Arc welding processes for surfacing**

This section describes arc welding processes that are used for application of wear-resistant coatings. Each has specific characteristics that were summarised in Table 3.1. Particular types of work can only be carried out by certain processes. However, the variety available does mean that some work can be done by several processes, which permits flexibility of choice. This can be important when undertaking repairs or replacements after a breakdown and in the field. This flexibility and the wide availability of process equipment and welding skills make weld surfacing a popular technique.

While Table 3.1 shows typical figures for various process characteristics, it is important to bear in mind that results achieved depend to a large extent on the way the process is used, i.e. the welding procedure adopted. This
subject is dealt with later in this chapter, but one characteristic of particular interest to the production engineer is the rate of deposition. Ranges quoted are typical of those used in practice and represent performance with 100% 'arc on' time. Adjustment must be made for 'arc off' time in calculating an actual deposition rate. It is also important to recognise that the maximum deposition rate achievable on a given job depends on the extent to which dilution must be controlled, as this will increase as the component temperature increases; high deposition rates require high welding currents and these encourage a faster rise in temperature.

Manual metal arc (MMA)

In manual metal arc welding (Fig. 3.1), an electric arc is maintained between the electrode and the work piece. The electrode consists of a core wire (1.6–8mm diameter) with a flux covering.

The arc melts the parent metal and electrode to form a molten pool, which is protected from the atmosphere by liquid slag, and gas formed by melting and vaporising the flux covering. The slag formed from the molten flux adheres to the weld surface, protecting it as it cools, and must be chipped away after each weld pass. The flux may also be used as a vehicle for adding hard particles or alloying elements to the weld pool.

Advantages

1. It is adaptable to small and large complex parts.
2. Positional welding is possible, i.e. vertical.
3. It can be used in places with limited access.
4. A wide range of consumables is available.
5. Deposition rates up to 4kg h⁻¹ are possible.
6. It is ideal for one-off and small series work.
Weld surfacing

Disadvantages

1. A skilled operator is required for high quality deposits.
2. Removal of slag is necessary, reducing the duty cycle.
3. Dilution tends to be high.
4. Granular carbides in tubular electrodes are usually melted.

Tungsten inert gas (TIG)

In TIG welding an electric arc is maintained between the non-consumable tungsten electrode and the workpiece (Fig. 3.2). The filler material is usually in rod or wire form.

The arc melts the parent material to form the molten pool which is protected from the atmosphere by the inert gas shielding. Filler wire is fed into the pool manually to produce the surface coating.

Advantages

1. Low penetration is achieved, with dilution 5–10% depending on technique.
2. The process can be closely controlled by the welder, and small areas can be surfaced, e.g. small pores in hardfacing deposits.
3. The process can be used manually with a hand-held torch and hand-fed filler rod, or mechanised for special applications.
4. A deposit thickness of 2 mm upwards is achievable.
5. It provides a deposition rate of up to 2 kg h⁻¹.
6. It is capable of high quality deposits.
Disadvantages
The process is not suitable for site welding, as draughts tend to disrupt the gas shield. It is generally restricted to a workshop.

Plasma transferred arc (PTA)
The plasma transferred arc process (Fig. 3.3) uses an argon shielded tungsten arc as the source of energy. A direct current (DC) pilot arc is established between a central tungsten electrode within the torch and a water-cooled copper annulus which surrounds the electrode. The function of this pilot arc is to facilitate initiation of a heavier current transferred arc between the electrode and the workpiece when surfacing commences. A separate DC power source is usually connected between the tungsten electrode and the workpiece and controls the transferred arc, which is constricted by a narrow orifice in the copper annulus.

This is a mechanised process which is used to deposit alloy powder conveyed from a hopper to the torch by a carrier gas. It should not be confused with plasma spraying, which uses a non-transferred arc to generate the heating plasma for spraying alloy or ceramic powders (see Chapter 4).

Advantages
1. It produces low penetration and dilution.
2. It is mechanised and provides close control of surface profile with minimum finishing required.
3. The deposition thickness is in the range 2–5 mm.
4. The deposition rate is higher than with the TIG process, averaging 3.5 kg h⁻¹.
5. The range of torches available includes those for surfacing bores down to 35 mm diameter by 400 mm deep and internal valve seats.
Weld surfacing

The equipment is not readily portable.
1. Equipment costs are relatively high.

Disadvantages

Metal inert gas/metal active gas (MIG/MAG)

In the MIG/MAG process an electric arc is maintained between the electrode wire and the workpiece (Fig. 3.4). The parent metal and the consumable wire are melted by the arc to form a molten pool, which is protected from the atmosphere by gas fed coaxially with the wire through the welding gun nozzle.

Advantages

1. It is a continuous process that is used semi-automatically with a hand-held gun, or is wholly mechanised by traversing the gun and/or the workpiece.
2. Deposit thickness of 3 mm upwards is achievable.
3. Deposition rate is 3–6 kg h⁻¹.
4. No slag removal is required.
5. It provides a positional surfacing capability.
6. Controlled transfer pulse techniques produce less spatter and provide greater control of weld bead characteristics.
7. Guns are available for internal bore work.
Disadvantages
1. Use of gas shielding makes the process marginally less transportable than MMA and gas must be selected to suit the surfacing alloy.
2. Welding must generally be carried out within about 4 m of the wire feeder, but best results are obtained with the gun as close as possible to the feeder, especially when feeding stiff wires.
3. Dilution may be high unless appropriate procedures are used.
4. High levels of UV radiation are produced especially when using high peak current pulse welding.

Flux-cored arc (FCAW)
Flux-cored arc welding is similar in principle to the MIG/MAG process, but the core of the tubular electrode contains a flux which decomposes to provide a shield to protect the molten pool.

Advantages
1. It is a continuous process that is used semi-automatically with a hand-held gun or is wholly mechanised.
2. A wide range of consumables is available.
3. No shielding gas is required.
4. Thickness of deposit is 3 mm upwards.

Disadvantages
1. Dilution is 15–30% depending upon technique, so it is not suitable for use with non-ferrous surfacing alloys.
2. Regular maintenance of equipment is necessary.
3. Deposit quality may be lower than the MIG/MAG process.

Submerged-arc (SAW)
In submerged-arc welding an electric arc is maintained between the electrode and workpiece (Fig. 3.5). The electrode dips into the covering of flux dispensed from a hopper and an arc is struck to the workpiece. The arc remains submerged under the flux coating, some of which melts and protects the molten pool from oxidation. The electrode is fed at a controlled rate to maintain a stable arc of constant length, and flux which has not melted can be recovered and reused.

The process, which is mechanised, can use a wire or strip consumable. It is possible to give supplemental alloy additions to the molten pool from a separate feeder.

Advantages
1. It is a mechanised process.
2. Deposit thickness of 3 mm upwards is achieved.
3. Deposit rate is 10 kg h⁻¹ upwards on suitable workpieces.
4. A wide range of consumables is available.
Disadvantages

1. It is suited to large workpieces which can tolerate the high current and high deposit rates without overheating.
2. It is intended primarily for workshop use in a fixed installation.
3. Dilution may be high unless appropriate procedures are used.
4. Applications are generally limited to cylindrical or flat components; there is limited access to internal surfaces of larger bores.
5. Flux costs must be taken into account.

Wire consumables

A wide range of wires is available. Using careful overlapping of individual beads, large areas can be covered with smooth deposits which need little machining allowance.

Strip consumables

When covering large areas, the use of strip consumables offers advantages in deposition rate. The range of materials available in this form is more limited than for wire, but the process is popular for applying corrosion-resistant cladding to large vessels and process plant.

Alloy additions to the molten pool – bulkwelding

In bulkwelding powdered alloy additions are metered into the weld zone and covered with flux. The consumable can be solid or tubular (cored) wire; the layer of powdered metal 'cushions' the arc and reduces substrate penetration and dilution. Precise control of the mixture of powdered alloy metal and electrode metal produces the required composition of weld metal.

One application of this process is production of wear plates consisting of high chromium iron deposits on a mild steel backing.
Considerable development has taken place recently to improve the performance of power supplies, based on recognition of the need for accuracy and stable conditions to ensure repeatability of results to the desired quality standard. Conventional power supplies, based on transformer/rectifier systems operating at mains frequency, are now being replaced by units based on power electronics, capable of supplying waveforms suited to each welding process.

The new designs are compact and lighter than their predecessors and micro-electronic control enables output to be selected to suit each process so that the power supplies are truly multifunctional. The accurate control of pulse waveform necessary for controlled transfer pulse welding is provided by the new supplies, which also allow the waveform to be varied automatically to a preset programme when a change in current is required, providing synergic control.

Examples of two such units are shown in Fig. 3.6 and 3.7, both of which can be used for MIG/MAG, TIG, FCAW and MMA. The unit shown in Fig. 3.6 is also capable of supplying up to 30kW of HF power for induction pre- or post-heating.

These power supplies are suitable for welding operations that need to be mechanised and can follow a predetermined weld cycle without operator involvement once parameters are established.

Other fusion welding and forge welding processes

Other processes are summarised in Table 3.2.
**Oxyacetylene**

In this process flame adjustment is important and is described by the length of the outer cone – the feather containing excess acetylene – as a ratio to the inner cone; see Fig. 3.8. The acetylene feather contains particles of carbon which tend to carburise the surface of the base metal and also reduce oxides on it.

Flame adjustments for the different types of surfacing alloys are as follows:
Neutral: nickel base self-fluxing alloys.

Slightly reducing: carbon or alloy steels containing a high percentage of iron.

1x–2x feather: deposition of Co-based alloys onto austenitic stainless steels, to prevent carburising the base material.

3x feather: Co base alloys on other substrates.

4x feather: surfacing rods comprising tungsten carbides in a ferrous tube.

Advantages

1 Minimum melting of the parent metal is possible with low dilution of the surfacing alloy. This is advantageous when using highly alloyed consumables and is also important if a thin coating only is desired.

2 There is minimum solution of carbide granules from tubular rods.

3 The process is under close control by the operator.

4 Small areas can be surfaced.

5 Thin, smooth coatings can be deposited.

6 Grooves and other recesses can be filled accurately.

7 The operator can contour the deposit to minimise final machining.

Disadvantages

1 The process is slow and not suitable for surfacing large areas.

2 The build-up of heat may overheat the component and lead to distortion.

3 The range of iron base consumables is limited.

4 As it is usually a manual process, results are dependent on operator skill, fitness and degree of fatigue.

5 Good technique is essential to ensure that the bond at the interface is sound, especially when fusion with the substrate is not involved. There is a lack of non-destructive testing (NDT) methods to check adhesion between coating and base material.

Powder welding

Powder welding uses a modified oxyacetylene torch fitted with an integral hopper to contain a self-fluxing surfacing alloy powder. The powder flow, under finger lever control, is entrained in the flow of combustion gases and emerges with the torch flame. The nickel base alloys normally used with this process can be deposited in thin layers and the technique lends itself to building up worn corners. Grit blasting to provide a mechanical key to the substrate is unnecessary as the deposit has a metallurgical bond to the substrate, but it is often the easier method of ensuring that the surface is clean. Preheating the part speeds up deposition as the interface has to be brought up to the bonding temperature of 1000°C; without this the surfacing alloy is just cast onto the surface. There must be enough heat and time for the self-fluxing action of the alloy to clean the surface of interfering oxides to ensure a full metallurgical bond between it and the coating. Once the initial coating has been established, the deposi-
Laser cladding; geometry of the process (courtesy Praxair Surface Technologies Ltd).

The deposition of subsequent layers is fast and easy. By making sure that the initial layer is not too thick, deposits can be made on small areas of large components.

Self-fluxing alloys possess complex structures and have a wide temperature range between liquidus and solidus. In between these temperatures the alloy has a pasty consistency and by careful control of the heat input the deposit can be built up into thin sharp edges, a good example being protection of the edges of cast iron moulds used in the manufacture of glass containers. Preheating the base metal to about 600°C, after applying a thin protective coating of the surfacing alloy on the cold metal, enables high deposition rates to be maintained and produces a deposit that needs minimum finish machining.

**Advantages**

This surfacing method requires less skill than a gas weld deposit and should be considered for surfacing small parts, small areas on large components or for generation of irregular deposits.

**Disadvantages**

The method shares the same disadvantages as oxyacetylene welding, and in addition there is a limited range of consumables suited to the process.

**Laser cladding**

Laser cladding uses a collimated laser beam, directed through a system of beam ducts, focused to a spot on the workpiece, see Fig. 3.9. The power of the laser and the focal spot diameter can be precisely controlled, and
the beam is scanned and translated over the workpiece to melt and solidify the pre-placed cladding material.

The cladding material solidifies rapidly, giving microstructures characterised by fine grain size, fine dendrite arm spacing and a more uniform dispersion of microconstituents such as carbides, nitrides, Laves phases, etc. Dilution is routinely controlled to less than 3%, giving coatings that are metallurgically superior to those produced by arc welding processes. Owing to the low heat input of this process, distortion of the component is negligible.

The width of the deposited coating can range from 0.15 mm to 50 mm, and almost any size of component can be treated. Beads can be laid side by side, and layers can be built up to any desired thickness. The maximum thickness in a single pass is approximately 2 mm. Figure 3.10 shows a large waterwall panel being surfaced with a corrosion-resistant material.

Materials available for laser cladding include several nickel alloys, cobalt alloys and iron chromium alloys, prepared as gas-atomised spherical powders.

Electroslag

Electroslag welding uses equipment similar to the submerged arc process for strip cladding. Deposition takes place under a blanket of flux, whose composition is formulated to produce a molten layer of slag by passage of the
Weld surfacing

welding current. No arc is used; the electric current is maintained by conduction through the molten slag. This, as well as protecting the metal transfer from the electrode to the substrate, provides sufficient heat to melt the electrode and to raise the surface of the substrate to a temperature at which bonding with the coating can take place. Materials available for use with this process are limited to those available in strip form, such as some stainless steels and nickel base alloys.

Compared with submerged arc strip cladding, advantages claimed for electroslag include low defect levels, higher deposition rates and better control of dilution. The more popular use of this process is cladding large areas with corrosion-resistant alloys.

Resistance welding

Small units have been designed to repair moulds for plastics, die casting and rubber. The power supply creates a series of high energy discharges through a capacitance circuit. It can be used to weld a consumable in the form of a powder, paste or strip. The application is via a hand-held electrode, similar in size to a soldering iron, for accurate placement of material. The heat affected zone (HAZ) is very small. Consumables available include nickel alloys, iron–chromium–aluminium alloy and stainless steel.

Explosive weld cladding

It was well known during the First World War that bullets or shrapnel could adhere to the metal surfaces they impacted. This scientific phenomenon was not appreciated as a welding process at that time and it was only in 1944 that Carl recognised it as a solid phase welding process. This chance observation has been developed since then to what is today a well-established industrial process. Bond strengths in explosive cladding approach those of conventional fusion welding processes.

Although explosive welding (like any other process) suffers some limitations, it is one of the most effective techniques for welding dissimilar metals. It is generally used for the production of composite metal sheets and for lining tubular geometries with thin layers of alloy materials for corrosion protection.

Theory of the process

Explosive welding is achieved when one metal plate is accelerated by an explosive charge to a high velocity oblique collision with another metal plate. Figure 3.11 demonstrates the basic configuration of explosive welding, where a flyer plate is accelerated under the influence of a detonation to impinge obliquely on to a base plate. In so doing, a jet of metal (from both plates) in the form of a spray is ejected ahead of the collision point, cleaning the weld surfaces of any oxide films or surface impurities. Pressure at the collision point is estimated at 250 kbar, which is well in excess of the yield strength of any metal. Thus, under these conditions, the interfaces of the two metals are subjected to an interatomic contact where the cohesive energy, or
more simply the balance of the interatomic forces between the metal atoms, results in a solid phase weld.

Stainless steel, nickel, titanium and other metals are frequently chosen for their corrosion resistance, but when used in the solid state, most of their thickness is required for physical strength. As only the exposed surface, plus a few millimetres (depending upon the corrosion/erosion rate) is required to give protection, considerable savings can be made by using the expensive metal where it is needed and a less expensive one to provide strength.

Explosive welding can clad a wide range of metals to almost any backing. The process offers several important advantages:

1. Up to 100% of the area is bonded.
2. A metallurgical bond is formed which is stronger than the weaker of the two metals.
3. There is no HAZ for many combinations.
4. Butt welding of the finished product is possible.
5. The backing metal can be double clad with a different material on either side.

**Practical applications**

The main applications of explosive welding are in flat plate cladding and welding or cladding of cylindrical geometries. Explosively clad plates are widely used in the following sectors of industry:
Stainless steel/aluminium composites are used in production of freezer trays for the fast freeze food industry.

Thick steel plates are clad with a thin layer of corrosion protection materials such as titanium, Monel and nickel alloys. These plates are used for fabrication of storage tanks for process plants.

Explosive cladding of cylindrical surfaces was an early development in explosive welding. Initially it was applied in particular to nozzles in steam power plants for corrosion protection. Recent developments on new explosive compositions and production clamp designs have produced commercially viable components, such as long weld neck flanges clad with nickel alloy materials (Fig. 3.12), spool pieces and flowlines clad with nickel alloy 625 (Fig. 3.13) for subsea production systems. A major contribution to the pipeline welding industry is the explosive welding of stainless steel duplex pipe, with both practical and commercial advantages in laying of pipelines for the offshore oil industry.

Further, explosive welding can be controlled so that it can be applied to production of microwelds between metal foils as thin as 11 μm. An example of this is the coating of titanium sheets with platinum.

Explosive welding has been used in other areas of industry, but the practicality and the commercial viability of each system must be assessed on its own merits.

**Friction surfacing**

In 1941 Klopstock and Neelands filed a UK patent application for ‘an improved method of joining or welding metals’. The information contained in the patent introduced the concept of depositing metals by friction surfacing. More recently, however, there has been renewed interest in this method of deposition which shows promise for a wide range of applications. Essen-
Friction surfacing is a solid phase deposition technique whereby a rotating consumable bar, usually in the range 10–40mm in diameter, is pressed on to a laterally moving substrate (Fig. 3.14).

Friction surfacing has the advantage of allowing materials to be used that would otherwise be metallurgically incompatible with the substrate, yet still providing a layer of deposited material with a high degree of adherence. Special combinations of material properties that cannot be realised in monolithic materials can be provided, which conserve more expensive and strategic materials.

Metal combinations and properties

The deposit, a product of a hot forging action, is inherently homogeneous and of good mechanical strength; see Table 3.4. The solid phase nature of the process ensures that negligible dilution is achieved yet good basic adhesion properties are maintained. The interface region usually remains intact, even after resisting loads at the ultimate tensile strength of the weaker material.

With respect to composite clad layers, corrosion-resistant and hardfacing materials usually have superior mechanical properties compared with the substrate material. It is not unexpected, therefore, that for optimised deposits, subjected to through-thickness tensile testing, failure occurs at loads equalling the ultimate tensile strength of the substrate. Lateral restraint provided by a stronger deposit material to the interface region usually means that failure occurs well away from the joint. With like to like deposit/substrate materials failure sometimes occurs in cohesion within the bulk deposit. With relatively weaker deposit materials failure usually occurs in cohesion of the bulk deposit.

Comparative corrosion-resistance properties established from potentiodynamic polarisation curves for stainless steel deposits in chloride and acidic solutions, between machined deposits and consumable specimens
<table>
<thead>
<tr>
<th>Deposit</th>
<th>Substrate</th>
<th>Welding machine settings</th>
<th>Mechanical tests TTT* Tensile strength/N mm⁻²</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Consumable diameter/mm</td>
<td>Rotation speed/rpm</td>
</tr>
<tr>
<td>Mild steel</td>
<td>Mild steel</td>
<td>25</td>
<td>975</td>
</tr>
<tr>
<td>Mild steel</td>
<td>Low alloy steel</td>
<td>25</td>
<td>975</td>
</tr>
<tr>
<td>Austenitic stainless steel</td>
<td>Mild steel</td>
<td>25</td>
<td>550</td>
</tr>
<tr>
<td>Austenitic stainless steel</td>
<td>Austenitic stainless steel</td>
<td>25</td>
<td>550</td>
</tr>
<tr>
<td>Alloy 625</td>
<td>Mild steel</td>
<td>20</td>
<td>410</td>
</tr>
<tr>
<td>Hastelloy CW-12M-1</td>
<td>Austenitic stainless steel</td>
<td>20</td>
<td>330</td>
</tr>
<tr>
<td>Stellite grade 6</td>
<td>Austenitic stainless steel</td>
<td>20</td>
<td>330</td>
</tr>
<tr>
<td>Aluminium alloy (2011)</td>
<td>Aluminium alloy (20, 4A)</td>
<td>25</td>
<td>778</td>
</tr>
</tbody>
</table>

* TTT = through-thickness tensile.
Engineering coatings

indicate that corrosion resistance properties are maintained. Pin-on-disc comparative wear tests between cobalt base alloys (Stellite grade 6) have shown that the deposit wear properties at least equal those of the consumable.

Physical mechanisms

During initial dynamic contact between the rotating consumable and the substrate, the resultant relative motion under an axial load produces a scouring action which removes the oxide layer from both contact faces. After the initial contact and after traverse has been initiated, the scouring action necessary to disperse the intrusion of the substrate oxide barrier is continued, not by the contact face of the consumable directly, but by the plasticised layer produced from the consumable. While it is recommended that the plasticised material has lower mechanical strength, it is expected that, provided suitable axial force is maintained, oxide dispersal continues, resulting in sound bonds.

The unequal temperature distribution between the comparatively small consumable bar and the bulk substrate leads to preferential consumption of the bar. This asymmetrical temperature distribution is further enhanced during friction surfacing by continually introducing new substrate at ambient temperature.

Although regions of the rotating consumable are repeatedly exposed to the atmosphere, sound deposits of good integrity can be produced. More reactive material, however, may benefit from a suitable shielding gas.

Lack of symmetry characterises the relative motion and material transfer of the process. One observable characteristic, more noticeable at lower rotation speeds, is the slight difference in appearance between the two side edges. Figure 3.15 shows that relative movement between the bar and the substrate is enhanced on the forward or advancing side and conversely is diminished on the retreating side. The introduction of lateral motion of the substrate to the angular rotation of the consumable produces asymmetric effects during deposition. Annular deposits produced with the consumable and substrate revolving in the same angular direction (Fig. 3.16), nominally minimise this asymmetric effect, especially with increasingly larger diameter consumables.

The uniform surface ripple, Fig. 3.17(a), results from the interaction of rotation and substrate traverse. Any point towards, and including, the periphery of the bar, moves in a cycloidal path, known as a compact superior trochoid. The almost semicircular ripples on the deposit surface point towards the start of the run and are produced by the final sweep of the trailing edge of the rotating consumable. Investigation has shown that at the substrate surface the scouring marks point towards the stop end of the deposit. Thus a complete reversal of relative motion must take place (Fig. 3.18) within the central zone of the deposit thickness. However, the extreme edges of the deposit only experience one direction with respect to the rotating consumable, i.e. one side retreating and the other side advancing with respect to the traverse.
Rotation of consumable. Inward (retreating) moves on to existing deposit and produces turbulence and back wash (extreme conditions)

Rotation of consumable. Outward (advancing) moves away from existing deposit and maintains straight uniform edge

3.15 Deposit characteristics.

3.16 Friction surfacing annular deposits: (a) consumable and substrate, offset, with same angular direction; (b) consumable and substrate, offset, with opposing angular direction.
3.17 (a) 500mm long friction surfaced deposit of stainless steel on mild steel produced with a 25 mm diameter consumable; (b) transverse macrosection of deposit, consumable speed 650 rpm, force 62 kN, traverse rate 7.5 mm s⁻¹.

3.18 Motion on the upper and lower surfaces of the deposit.

**Unequal heat generation**

The effect of the relative heat generation can be seen in any transverse macrosection. Figure 3.17(b) shows a shallow crescent-shaped HAZ of maximum depth in the centre. Insufficient heating and lack of bond at the
outer edges of the deposit are also shown. Tubular consumables, in addition to shaping of the substrate, modify this effect.

Lack of bonding at the outer edges of the deposit is inherent to operating on flat substrates. The outer edges of the deposit only receive action at tangential points corresponding to the outer diameter of the bar. On the other hand in the centre zone the full bar acts upon the plasticised layer and deposited material for a longer period. In addition there is not the same degree of pressure at the outer edges of the consumable bar, and this contributes to the unequal distribution of frictional effort.

Applications
Friction surfacing has already been used for reconditioning of worn shafts and good quality deposits have been reported. Successful hardfacing of cutting edges and agricultural tools has also been carried out by Russian workers.

Interest in the process continues and as a result applications, mainly in localised cladding are being investigated. Various substrate geometries are shown in Fig. 3.19. Typical examples include the anticorrosion overlay of slide valve plates, surfacing of the annular contact face of composite pipe flanges, disc brakes, etc. (Fig. 3.20a), inlay cladding of strategic material positioned to suit bearings or ‘seal’ contact areas of shafts (Fig. 3.20b), and cladding of the exposed regions of shafts that see service in pernicious environments. Further suggestions include manufacture of specialised wear tiles for sinter plant, wire rubbing pads (Fig. 3.21), quarrying equipment, bi-metal linear bearings and also in situ reclamation of worn railway points. Turbine and guillotine blades are also considered examples of suitable applications for friction surfacing.

The process will become increasingly concomitant with that of other solid phase large overlay techniques. Repair and reclamation of materials already in clad form can also be carried out. The Welding Institute considers that artillery shell banding may be another application area, where suitably soft material can be friction surfaced to shell bodies.

Advantages
1. The solid phase friction surfacing process minimises distortion.
2. Friction surfacing allows deposits to be made in all positions.
3. Thin layers of negligible dilution but considerable bond strength can be obtained for a range of corrosion- and wear-resistant materials.

Disadvantages
1. Unless the consumable is wider than the substrate component or deposition is to be made into a recessed groove, lack of bonding at the outer edges of the deposit is noticeable. This distracting feature is an inherent effect which reduces the effective bond width to just less than the diameter of the consumable.
2. The substrate needs to be able to react to high compressive loads; inter-
Suggested geometric arrangements for friction surfacing.

**Metallurgical and related effects of weld surfacing on substrate and coating**

This section deals with changes that may take place in the component material, and alterations that can occur in the coating material as a result of the welding operation. The principal factor causing these changes is the
3.20 Friction surfaces components produced at The Welding Institute: (a) bimetal brake disc inlayed with wear-resisting material; (b) tubular mild steel shaft inlayed with stainless steel and tool steel clad layers on to flat and tubular components.

3.21 A wire rubbing pad inlayed with wear-resisting materials which has greatly improved the surface life of the component (courtesy Friction Technology Ltd).

temperature reached in the component and coating during the process. These remarks are particularly applicable to the fusion welding processes, and are less applicable to the forging processes.

Most steels may have their properties changed by a controlled thermal treatment; temperatures used are similar to those reached during welding. Lack of control of the temperature cycle during welding may result in hardening/embrittlement of the steel, with effects that may become evident during the process, or later in service. The problem can usually be overcome by controlling the temperature cycle so that the part remains in an unhardened and tough condition.

There are two main types of coating material. One is affected in its main properties by the temperature cycle like a hardenable steel, so the cycle must
be controlled accordingly. The second type is not dependent on thermal
treatment to acquire its properties. Both may be affected by what is called
‘dilution’, i.e. intermixing of the deposit with the substrate material, which
can alter the material’s response to the thermal cycle.

A further feature of weld surfacing is development of residual stresses in
the part. This arises from differences in thermal expansion characteristics
between coatings and substrate and from the thermal gradients present
during the coating operation. These stresses may result in distortion of the
part or, in the case of the harder coating alloys, in contraction cracks in the
coating.

A knowledge of the characteristics of the chosen substrate and coating
materials is used to design thermal cycles for surface coating operations that
produce acceptable finished parts.

The temperature cycle

Heat treatment before surfacing

Substrate materials should be specified in the soft condition for weld surfac-
ing. The temperatures involved produce some tempering of the structure
of a hardened steel. Parts made by casting or forging, or from bar stock that
has been extensively machined, should be stress relieved before the final
preparation of the part, especially when distortion of the part during the
welding process must be kept to a minimum.

Selection of thermal treatments for surfacing steel components must take
into account the various factors mentioned below, and in addition compo-
nent size, shape and practical problems of handling the parts during and
after welding, either through size or production volume.

The need for preheat

It is well known that molten metals shrink when they solidify. For cast-
ings, the pattern is typically made around 2% larger than the casting size
needed.

Weld-deposited metals shrink similarly unless – as is usual – they are
restrained from doing so by attachment to the substrate material. This leads
to residual stress in the component. By applying preheat to the part, it
expands, and contraction on cooling after welding partially offsets weld
shrinkage. Residual stress is thereby reduced and this helps to reduce any
tendency to contraction cracking in a deposit of surfacing alloys near the top
of the hardness range. It also helps to control distortion of the part by
reducing temperature gradients.

In steel substrates that are capable of hardening if cooled in air, a thin
layer of the HAZ under the weld deposit may harden. This transformation
involves a volume expansion which increases any residual welding stresses
and can cause cracks both in the deposit and in the HAZ. The latter will
cause detachment of the coating.

The use of preheat, maintained at a specified minimum temperature
during the welding cycle, can ensure that no hardened HAZ is created during this period, by reducing the speed at which the HAZ cools and by limiting its lower temperature while welding proceeds. Depending on the transformation characteristics of the steel, this may be sufficient to allow the HAZ to transform into a soft structure, but a safe precaution is to arrange an immediate post-welding heat treatment which ensures that only a soft structure exists throughout when cooled at room temperature. In oxyacetylene welding, somewhat higher temperatures may be desirable than the minimum required to avoid a hardened HAZ. This is because the flame is a low intensity heat source, and by using a higher preheat, faster deposition is possible without risk of increased substrate melting and deposit dilution.

The upper limit of possible preheat is governed by several factors. The first is that the upper surface of the component may oxidise ahead of the weld, incurring the risk of defective deposits – porosity caused by oxide inclusions and/or improper bonding to the substrate. A limit is around 400°C for low alloy steels and 600°C for oxidation-resistant materials.

The second factor is that with increased temperature, the strength of the substrate may be significantly less than the coating alloy, and this may increase rather than reduce the distortion risk. A further risk is that of increased intermixing of the deposit with the substrate, as the latter is more rapidly raised to its melting point by the heat source. The consequences of this effect – dilution – are discussed elsewhere in this chapter. High temperatures during welding can also lead to grain growth in some steels.

**Interpass temperature control**

Having stipulated the minimum and maximum temperatures at which weld surfacing can take place safely, as described above, the question of interpass temperature control is settled. In practical terms this may mean that deposition must be stopped periodically to avoid overshooting the upper limit, or welding switched to a cooler portion of the workpiece, if this is possible.

At the other extreme, it means that at no time may any portion of the workpiece fall below the minimum set temperature, as the HAZ in that region may then transform and harden, with possible risks of cracking. This means that, in practice, local preheat may be needed to provide adequate control on a large component. When depositing alloys designed to develop their hardness by a martensitic transformation, hardening may be prevented by the use of preheat or a rise in component temperature during welding. Specialist advice should be sought from the consumable manufacturers.

**Post-weld heat treatment**

For components made from hardenable steels, this can take two forms:

1. Temperature equalisation on the welding set-up, followed by slow cooling in an insulating powder such as vermiculite.
2 Temperature equalisation followed by immediate transfer to a heat treatment furnace preheated to the minimum interpass temperature.

The subsequent temperature cycle can then follow various paths:

1. Cool to just below the martensite transformation temperature, hold, and then raise to temper the martensite.
2. Raise to the austenitising temperature and then cool at a rate to produce a pearlitic (soft) structure.
3. Raise to the temperature for most rapid transformation of any austenite to pearlite; hold for completion and then slow cool.

The selection of appropriate treatment times and temperatures requires a study of the available isothermal or continuous cooling transformation diagrams for the steel.

Subsequent heat treatment

Components that have been cooled in an insulating powder after welding are likely to contain more residual stress than those heat treated in a furnace. Heat treatment at temperatures recommended for stress relief of steels at around 650°C is effective in reducing movement when parts are subsequently machined and also for dimensional stability in use. Slow heating and slow cooling is needed for best results; cooling at 100°C per hour is often used.

Substrate materials – special precautions

Some steels may be embrittled by exposure to elevated temperatures; for example 11–14% manganese steel must be kept cold by only intermittent welding or water cooling. Some Ni–Cr steels can be susceptible to temper brittleness, or a reduction in the ductile–brittle transformation temperature, while some high alloy stainless steels can form the brittle sigma phase when welded. Specialist advice should be sought.

Other substrate materials

There is a wide range of engineering metals and alloys in use. When considering the possibility of applying a welded coating for improved service life, the first question to be asked is whether a cheaper substrate material of much superior welding suitability can be substituted – often at greatly reduced cost.

If the existing material must be considered, the following lists characteristics that may make weld surfacing difficult or even impossible:

- melting point lower than the surfacing alloy;
- free cutting properties;
- contains easily oxidised elements, e.g. Al, Cu, high Si;
- is hard and/or brittle;
- a stainless steel not stabilised for welding;
- a steel containing nitrogen as a deliberate addition.
Grey cast irons
The melting point of cast iron is lower than that of steel and this limits the coating alloys that can be deposited without melting the substrate and diluting the deposit. Other problems include the risk of substrate cracking caused by welding stresses – it has a low tensile strength and, depending on composition, may develop hardened HAZs of even greater brittleness. Problems are increased by the presence of any casting defects. The oxyacetylene process offers the best possibility of controlling dilution, but use of a cast iron welding flux may be necessary.

White cast irons
White irons are extremely brittle and weld surfacing is not recommended.

Ni-hard cast irons
These irons are also difficult to weld surface, although some success has been reported by use of high levels of preheat, a buttering layer of a ductile material such as nickel under the surfacing alloy and a very slow rate of cooling.

Influence of substrate defects
The presence of defects in the materials from which a component has been made is likely to affect the quality of the welded deposit. The importance of such defects depends on the quality standards demanded for the finished job.

Much surfacing work is carried out to improve the service life of parts used in critical items of equipment, including chemical and petroleum plant, power stations and heat engines. Standards for these applications are high and after finish, machining/grinding parts are examined to ensure freedom from defects in the coatings.

Two important sources of defects in welded coatings arising from the substrate are:

- discontinuities in the substrate, e.g. porosity or cracks;
- inclusions, e.g. mould material in castings, scale in forgings, slag particles/stringers, sulphur segregation.

Careful inspection of the substrate is therefore indicated, at a level appropriate to the quality of the finished part. Discontinuities are identified on the machined surface of the part before welding by dye penetrant inspection. Defects should be excavated to verify that they are not an outcrop of a larger cavity. Inclusions should also be excavated to remove them entirely and the resultant holes filled by welding, using a consumable to match the substrate, and then dressed flush with the surrounding surface.

When defects become apparent during the welding process by local collapse of the surface of the component, attempts to 'weld them out' usually result in failure, either because of the appearance of porosity in the welded
surface after machining, or even the presence of cracks caused by local increase in thickness of the coating material or to porosity acting as a stress raiser.

Repair of defects in welded deposits is dealt with later in this chapter.

**Dilution**

A feature of weld-deposited coatings is the strong bond with the substrate, and temperatures required to achieve this always result in some melting of the substrate. Intermixing of the welded deposit and the substrate is inevitable, but the extent of the intermixing is determined by the materials and the welding parameters.

The consequence of this intermixing is that properties of thesurfacing material may be altered and its performance changed. Excessive dilution during surfacing is always to be avoided and the question arises as to how much can be tolerated and how it is to be controlled.

Many ferrous surfacing alloys, supplied as electrodes or wires for arc surfacing processes, that are to be deposited on steel substrates are identified by their typical weld metal composition when applied by specified processes and recommended welding parameters. As both substrate and coating are iron-based alloys the composition of the consumable can take into account some dilution and provide the desired weld composition and properties.

If the surfacing alloy is not iron-based, but for example an alloy in group 3 (nickel) or group 4 (cobalt) base, and the introduction of iron into the composition is known to affect wear-resistant properties, the alloy designer must recommend that dilution is kept as low as possible for best results.

There are several ways in which this can be achieved:

1. Use of a low energy heat source such as oxyacetylene flame. By using the braze welding technique mentioned earlier with cobalt base alloys, it is possible to keep dilution below 5% on much work.
2. Self-fluxing alloys containing silicon and boron are also deposited with similar levels of dilution with oxyacetylene.
3. If a higher deposition rate is required, the PTAW (plasma transferred arc welding) process can be used and dilution levels similar to oxyacetylene welding are achievable on single layer deposits.

On components of variable cross-section, or those whose temperature may rise significantly during welding (which encourages increased dilution), it is normal to regard figures of less than 10% dilution as realistic for many applications. This is more difficult to achieve with direct arc processes such as GMAW and submerged arc welding (SAW) on single beads, but by use of overlapping beads, negative polarity and suitable parameters where appropriate, dilution of 15% or less can be achieved. If lower figures than this are required, a two-layer deposit should be considered.

Apart from the influence of dilution of a surfacing alloy with iron from the substrate, other elements may be introduced from the same source. For example, an austenitic steel surfacing material dependent on a low carbon content for maximum corrosion resistance could be affected adversely by pick-up of carbon from the substrate.
Dilution between the coating alloy and the substrate can sometimes produce undesirable properties. This is well illustrated in surfacing austenitic manganese steel with a material giving a hard surface as deposited. Interdiffusion causes the surfacing alloy to mix with the Mn steel, reducing the Mn content so the substrate becomes brittle near the surface. A similar effect is produced when surfacing carbon and low alloy steels with an Mn steel. In this example a brittle layer is formed in the overlay. In either, the interdiffusion can be prevented by using a buffer or buttering layer of an austenitic stainless steel. Alternatively, hard-facing rods of a work-hardening nature, but containing 12% Cr and only 3% Mn, can be deposited without formation of a brittle layer if reasonable precautions are taken. Other modifications can be used, the requirement being to preserve the austenitic structure even with some change in composition.

The figures for dilution given in Table 3.2 show a range because of the process variations mentioned above. In direct arc processes such as MIG and SAW, dilution is influenced inter alia by welding current, which also governs the deposition rate. Hence these factors must be considered together when determining the process and parameters to be used.

Coating properties and structure

It is important that the designer appreciates the variation in methods of deposition associated with each alloy group and the effect that this can have on the ultimate properties of the coating. Apart from dilution with the substrate material, both the micro- and macrostructure of the deposit can be changed by alteration in the surfacing method.

Surfacing rods containing tungsten carbide

Surfacing rods containing tungsten carbide produce different microstructures between arc weld and gas weld deposits. Higher heat input in arc weld depositing takes much more of the carbide into solution, hardening the matrix and reducing the amount and the size of the carbide particles. The structure is also greatly influenced by the initial particle size of the carbide grains (Fig. 3.22).

Coarse particles give a better cutting action on rock, finer grades give better and more uniform resistance to wear. The filler metals comprise steel tubes filled with tungsten carbide particles and are deposited with an oxy-acetylene torch or the usual arc processes. The melting steel takes up the tungsten and carbon from the carbide to form a matrix anchoring the remainder of the carbide particles. The amount of carbide that dissolves in the steel depends on the temperature and the length of time the weld pool is molten. The extreme is reached when surfacing with a very fine carbide using a high amperage electric arc. In this case all the carbide may dissolve, giving a very hard brittle tungsten steel liable to weld cracking and containing few, if any, carbide particles. Much less solution occurs with gas welding, and the carbide distribution remains more uniform.

With arc welding the surface of the substrate is melted and interdiffusion with the molten surfacing alloy provides the strong metallurgical bond. Fluxes may or may not be used, depending on the nature and amount of the
oxide on the surface of the blank. Application temperatures are high, and there can be much glare and fume from the flux, adding to the difficulty of putting down an accurate deposit. The designer must therefore provide as much help as possible. Part of the surface melts, so it is not feasible to design sections so thin that they will melt or taper to a feather edge.

### Cobalt base alloys

Cobalt base alloys have lower melting points than steel and a different technique is used for oxyacetylene deposition on ferrous base materials. The flame is kept in a reducing state and the carbon from it alloys with the iron base substrate, forming a skin of gradually reducing melting point until this reaches that of the surfacing alloy, around 1300°C. At this temperature the surfacing alloy can be melted on to the ‘sweating’ surface with which it forms a strong bond. Oxides created ahead of the pool float to the surface. The high chromium content of the surfacing material provides extremely good high temperature oxidation resistance, so the deposit remains clearly visible and, provided the blank design embodies good reference points, the operator can build up the surface accurately. The alloy will develop reasonably smooth surfaces, so the machining allowance is less than is required for arc weld deposition. On substrates that cannot produce a molten surface using a carburising flame, a flux is necessary to clean off surface oxides to assist bonding to the surfacing alloy melted on to it.

A hypoeutectic cobalt base alloy (30% Cr–8% W–1.6% C), when deposited by gas welding, shows dendrites of a Co–Cr solid solution, infilled with a eutectic of chromium carbides and the Co–Cr matrix (Fig. 3.23). Deposited by arc welding, the microstructure is finer, and because of the dilution by
the base metal, there is very little carbon to form carbides so there is little eutectic. The effect of a lower carbon content is seen more clearly in a Co base alloy (28% Cr–4.5% W–1% C), the percentage of the dendritic solid solution being much higher.

**Nickel base surfacing alloys**

Nickel base alloys vary greatly in composition. Ni–Mo–Fe alloys are used for corrosion resistance, Ni–Mo–Cr–W alloys for corrosion and heat resistance. Both need high application temperatures to melt the alloy, whether gas or arc welding is used. The most widely used Ni base surfacing alloys contain boron and silicon, which provide in-built self-fluxing properties. Melting at about 1050°C the molten alloy cleans off all but the most tenacious surface oxides from the substrate, to which it bonds by a brazing mechanism, requiring negligible intersolution with the surface of the base metal. Flow of the deposit is controlled easily, so the operator can cover complicated contours, and produce thin smooth coatings on large or small areas. At the working temperature the deposition is easily seen; however, it is still essential that the blank design provides visual guides for position and depth of deposit. Formulation, as well as deposition method, may produce markedly different microstructures in Ni base alloys having the same chemical composition.
**Buffer layers**

The term ‘buffer layer’ is used to describe the presence of an intermediate deposit between the base metal and the hardfacing weld material. The use of more than one type of hardfacing alloy may be necessary in some circumstances to reduce stress, to prevent cracking or to improve wear life of heavy deposits. There are a number of applications where this practice occurs:

1. When the ‘harder’ surfacing alloys are used on a soft base material, e.g. mild steel, there is a tendency for the hardfacing layer to sink under high load conditions. Under extreme conditions this may result in the surfacing material spalling off. To overcome this, a layer of strong, tough material is deposited on the workpiece before the hardfacing.

2. With gas welding techniques, if differences in thermal expansion are significant and the surface hardness exceeds 50RC, it may be useful to apply an alloy of compatible composition, hardness 25–30RC, underneath to prevent cracking of the hard overlayer. Where the design calls for a heavy build-up, full thickness may be achieved using alternate layers of hardfacing alloy and buffer material. In arc welding processes, natural dilution from the base metal usually provides the necessary gradation of properties.

3. If a component is subject to heavy impact or flexing, there is a risk that deposits that do not relief-check during welding will develop fine transverse cracks. These are not detrimental to the hardfacing but there is a danger that in service the cracks will act as stress concentrators and progress through into the base material. This tendency is most pronounced when the base metal is a high strength steel. Use of a buffer layer prevents such crack propagation.

**Preparation and design for weld surfacing**

Successful hardfacing depends on adequate surface preparation. The surfaces to be coated must be free from rust, grease, dirt or other material likely to be detrimental to the final deposit. Grinding, machining, filing, chipping, grit blasting, etc., are all used for preparation.

A sound base is required, and in reclamation work this may necessitate removing fatigued or rolled over material, high ridges or other major surface irregularities. Cracked or defective base material should be repaired. It is usually desirable to remove severely work-hardened or case-hardened surfaces before hardfacing.

For work such as excavator digger teeth, bulldozer blades, etc., which are hardfaced by arc welding, extensive preparation is not necessary, as occasional blowholes in the deposit are not important. On precision components such as engine valve seats, however, it is essential for the surfaces of the blank to be clean, and the job should be smooth machined before welding. Rough machining marks are an indication of torn metal which may oxidise and cause pinholes in the deposit.

To prevent contamination of the deposit, sharp corners must be removed from the area of the blank to be hardfaced, as they are apt to melt if the
arc or flame dwells on them. This is particularly important if parts have to withstand corrosive conditions in service and are being surfaced with iron-free stainless alloys, or Ni or Co base alloys. Any serious contamination of the deposit with iron may lead to premature breakdown in service through corrosion, and is likely to result in reduced hardness in the deposit as discussed earlier.

When building up edges of cutting tools, dies, etc., a recess is required to provide adequate support for the hardfacing material.

Wherever possible, the workpiece should be positioned so that hardfacing can be performed in the downhand position (workpiece horizontal). An uphill inclination of about 10° can sometimes be of assistance in laying down heavier weld passes. If work must be done out of position, detailed attention must be given to selecting suitable consumables and surfacing processes.

If the facilities are available, blanks should be degreased immediately before hardfacing. Where surfacing is being applied as a reclamation or repair process, all cracks should be removed by machining, rough grinding or gouging, filled in using compatible electrodes, and machined smooth before final surfacing.

**Deposit thickness**

Weld deposits are generally at least 3mm thick and can be designed to any thickness required, although the harder, more brittle alloys should be restricted to not more than two layers, with a total deposit of about 8mm maximum to reduce the risk of contraction cracking. Oxyacetylene and TIG weld deposits can be thinner, down to about 1mm thickness. The advantage of thick coatings is that they provide longer service life before resurfacing is necessary, assuming that the change in dimensions as the part wears can be tolerated. It is bad economics to design a deposit thicker than the allowable wear. Although components such as shear blades, gear wheels, conveyor screws and extrusion dies show a reduction in efficiency as wear takes place, other components such as gyratory crushers can tolerate wear of many centimetres before resurfacing becomes necessary.

If the deposit is to be ground, it should be at least 1mm thickness and in most cases preferably not thinner than 1.5mm. During deposition some distortion of the blank probably occurs and also scaling of the centreholes or chucking or locating faces. Consequently, it may be difficult to ensure that the deposit cleans up to an even thickness all over. It could be ground off entirely in some areas unless it is more than 1mm thick. On larger jobs, such as dies 100mm diameter, even a 1.5mm deposit (after grinding) is not sufficient to take care of possible variation because of distortion or contraction, and 2.5 or 3mm of finished deposit is usual.

In addition to the need to design a deposit thick enough to allow distortion to be corrected by machining and still leave the desired thickness for the working life of the part, it must be remembered that welding generally produces an uneven surface profile which may measure up to 2mm peak to trough in manual weld and less with mechanised surfacing.

If the deposit is heated in service, such as deposits on hot shear blades, it
must be designed sufficiently thick (usually 3mm minimum) to ensure that the base metal is not so heated that it becomes soft enough to allow collapse of the deposit. Again, a thicker deposit is needed on areas where high pressures are exerted, such as the line of entry or throat of a drawing die.

Examples already quoted show how design calls for variation in the thickness of deposit depending on the amount of wear predicted in the various sections of a part or assembly. For reasons of economy, the thickness of the deposit, after any necessary finishing, should be the minimum required for adequate component performance. The optimum thickness varies greatly from one application to another, since the ‘wear tolerance’ for the part may vary from as little as 0.05 mm for a precision component to as much as 50 mm for earthmoving equipment.

Process considerations can limit the minimum thickness that can be deposited. Guidelines for the designer are given in Tables 3.1 and 3.2.

Finally, it should be borne in mind that when designing coatings for wear resistance, the finished coating thickness should be greater than the permitted wear tolerance. The coating should never be allowed to wear through so that the base metal becomes exposed, or further wear will be very rapid. In detailing the extra thickness the designer must make allowance for the variation in deposit thickness which may occur across the part because of distortion in processing, as already discussed.

**Blank preparation and deposit design**

**General**

Reference has been made to the effects of welding temperatures on steel components, i.e. some oxidation of adjacent surfaces and possibly some distortion, depending on the job. So for components that require finishing to accurate limits with a clean machined/ground surface, a machining allowance is necessary on all unwelded surfaces.

If a deposit is to be machined to a sharp corner, difficult corner build-ups by welding can be avoided by making the part locally oversize, later machining back to size and producing the corner. If a weld-deposited surface is to be precision finished flush with adjacent surfaces, then a recess should be provided for the deposit. This has the added advantage of providing a ready guide – especially to a manual welder – to the exact location and thickness of deposit required. Recesses should never have square corners, since protruding corners heat more rapidly than surrounding surfaces and tend to melt, diluting the deposit and reducing hardness, wear resistance and corrosion resistance of many surfacing alloys. Recessed sharp corners are slow to reach welding temperature even when the adjacent surfaces have done so, and can be the site of defects such as porosity and an imperfect bond to the substrate, which may be exposed on machining.

When designing blanks, always provide a reference for the machinist to pick up, such as centres, shoulders or faces. Once the part has been welded, the marking or recess provided for the welder is obscured, and with no
reference, the deposit may easily be machined so far that it is unable to do the job; indeed some deposits have been totally removed because of this lack of foresight. Remember, the accuracy of the finished job depends on a correctly designed blank (Fig. 3.24).

**Design details**

For welded coatings on flat or circular sections, the recess (Fig. 3.25) should not be less than 13 mm wide, even for a shallow gas-welded deposit on a thin section blank. The rim of metal at the side should be at least 3 mm even when using a small flame. The depth should be the finished thickness plus
grinding allowance. Any surface to be coated should preferably be smooth machined.

To prevent contamination of the deposit, sharp corners must be removed from the area of the blank to be hardfaced, as they are apt to melt if the flame dwells on them. This is of particular importance on jobs that have to withstand corrosion and that are being surfaced with iron-free stainless alloys such as cobalt-based materials. Serious contamination with iron could lead to premature breakdown caused by corrosion.

Figure 3.25 indicates minimum recommendations for various thicknesses of deposit. A 45° angle should be used for the side of the recess so that there is no corner to melt easily. If a radius is used it should be at least 3 mm. The rim of metal must not be less than 1.5 mm even on a small job needing a small flame, and with a large flame 6 mm may be required to prevent melting.

If a large component is being manually surfaced by welding it is locally red hot, and the operator will have difficulty in deciding the exact location and extent of the area to be surfaced. A recess provides desirable assistance helping to control the thickness of the deposit, preventing waste of alloy and reducing subsequent machining time and cost. A typical example for gas welding is the steam throttle valve (Fig. 3.26). This illustrates several design features:
The recess for the deposit is radiused smoothly. For oxyacetylene welding the recess is wide enough to take the large welding flame necessary to hardface the mass of the metal. With a narrow recess the two top corners would probably be melted before the correct temperature was attained in the base of the groove. The outside wall, which is machined off after facing, is thick enough to prevent its melting during deposition. The top of the recess indicates the correct height of the deposit. A smooth, level surface can be put down keeping excess surfacing material and finish machining to a minimum.

Small diameter components (Fig. 3.27) can have an oversize deposit of metal laid on the end (a) if the sharp corner is chamfered to prevent its melting. Alternatively if the protection does not need to cover all the end a shallow depression can be formed (b). A groove with thin walls should not be designed into the part (c). On larger diameters a machined recess can be used (d) but there must be adequate metal left on either side. An alternative is to use a shallow groove (e) which avoids the side wall and so can use a less massive blank with savings in steel and finish machining.

Rock crusher hammers (Fig. 3.28) are hardfaced to improve wear life. Wear on these starts at the leading corner, requiring the thickest coating there, tapering off down the front face. It should be borne in mind that the life of the hammer will be increased a number of times by corner protection so the extent of surface protection should be designed accordingly. The end face of the hammer need not be hardfaced: it does little work, and when the front corner wears through it would probably be stripped off. A thin deposit is, however, sometimes of advantage down the side faces if they wear rapidly. Surfacing, as in Fig. 3.28(a), could cause premature failure of the coating on the other corner.

**Distortion control**

Contraction of the weld deposit during solidification and cooling can cause distortion of the component. This problem is more common in weld surfacing than in fabrication welding because there is little chance of balancing welding stresses by welding both sides of the neutral axis. There are three main factors:

1. Weld metal contracts during solidification and cooling, from a temperature that may be considerably above that of the substrate. Shrinkage is approximately 2% on cooling to room temperature and this tends to pull the workpiece in an arc along the direction of the weld run (Fig. 3.29).
2. Metal adjacent to and at a distance from the weld run suffers different degrees of expansion and contraction. This problem is accentuated most on thin base materials. The metal close to the weld zone becomes hot and starts to expand. Being restrained by the colder stronger metal further from the weld zone it can only move by buckling around the weld area. This movement will not be fully reversed during the cooling cycle and the workpiece may remain permanently distorted. Alternatively the hotter metal may undergo plastic deformation and on
3.27 Design of blanks for surfacing (a)–(c) smaller diameters – (a) and (b) correct, (c) incorrect; (d) larger diameter, machined recess; (e) larger diameter, shallow groove.
3.28 Hardfacing of rock crusher hammers as in (a) will break and fail rapidly, as in (b) where the top has been stripped off; (c) for economy the hammer is protected on both edges so that when the first deposit wears the hammer can be reversed.

3.29 Shrinkage of weld metal during solidification and cooling tends to pull the workpiece in an arc along the direction of the weld run.

3.30 Small 'relief check' cracks which form across the weld bead reduce the amount of stress distorting the base metal – often such cracking is not detrimental to the performance of the component surface.

cooling will not stretch again and so cause distortion, such as dishing of a valve seat.

3 Apart from temperature differences there will be differences in the coefficient of expansion and contraction of the surfacing alloy and the substrate. With many combinations this is relatively slight, but it can be an important factor with Ni and Co base alloys, cast irons and materials containing an appreciable percentage of tungsten carbides.

Often distortion, especially of the first type considered above, can be overcome by restraining the part so that it is not free to move. This can be done by clamping or tack welding the part to a firmer support. For flat items, such as crusher jaws, two parts can be clamped or tack welded back to back and hardfacing applied to each alternately.

Surfacing using hard alloys may lead to cracking in the deposit if such restraint is used (Fig. 3.30). If such cracking cannot be tolerated, a high preheat may prevent cracking in heavily restrained parts. Temperatures may have to be up to 400°C for arc welding and 650°C for gas weld surfacing. The use of a soft buttering layer may reduce stresses between the overlay and the substrate. Multilayer deposits of hard alloys are sometimes
To combat distortion a workpiece can sometimes be preset or pre-bent in the opposite direction.

For surfacing grader blades or bulldozer cutting edges place two blades back to back and pre-bend by clamping at each end with a spacer bar in the centre. Skip weld, working on each blade alternately applying a deposit 37 mm wide in one run.

made with alternate layers of soft ductile metal to prevent propagation of cracks. A corrosion-resistant interlayer may also eliminate corrosion of the substrate should this occur through the cracks in a hard wear-resistant top coating.

By slowing down the cooling rate, and thus the rate at which hot metal contracts, preheating can sometimes be used to prevent, or certainly reduce, distortion. This allows more time for stresses to become equalised in areas immediately adjacent to the weld rather than distorting the workpiece.

Presetting or prebending the part in the opposite direction to that in which it would distort can be used effectively on metal up to about 12 mm thick (Fig. 3.31). Contraction stresses pull the surface straight. The amount and actual nature of the presetting required for a given job must be established by experience. An alternative to presetting the part, illustrated by the surfacing of bulldozer cutting edges (Fig. 3.32), and knife blades is to clamp the two components back to back with a spacer bar in the centre. Contraction of the surfacing alloy on release pulls the blade straight and puts the deposit into compression, a desirable state. A single blade could be clamped to a strongback with a spacer under its centre to make it convex during hardfacing.

The use of intermittent welding techniques, such as backstepping, may greatly reduce distortion when hardfacing a long blade (Fig. 3.33). Several centimetres at the centre should be hardfaced first and similar lengths on each side of the deposit should be laid down. The direction of deposition should always be towards the centre. Each run should be the maximum width and thickness possible or necessary, so as to reduce to a minimum the number of times the blank is heated and cooled. Again, if in surfacing a
3.33 Backstepping can reduce distortion when hardfacing a long blade.

3.34 To prevent warping when hardfacing shafts, a short run should be made parallel to the axis and then a similar run made on the opposite side to equalise stresses.

pair of components they are fixed back to back, distortion is minimised if hardfacing is performed alternately on each.

Similar considerations apply to manual hardfacing of shafts (Fig. 3.34). To prevent warping, a short run should be made on one side, parallel to the axis, then the shaft should be turned over and a similar run made on the opposite side to equalise the stresses. If a long run were made on one side only, this would cause a permanent set that would not be removed by the next deposit on the opposite side. The sequence of runs should be as shown in Fig. 3.34. If the length of the area to be hardfaced is over 200mm long, it should be deposited in 100 or 150mm sections in the above manner. The job must be well supported at close intervals to prevent sagging, particularly if it is preheated to a red heat.

The problem of distortion of shafts is greatly reduced by mechanised deposition. The shaft is rotated about a horizontal axis and a continuous overlapping bead of weld applied in a controlled spiral. Mechanisation, when possible, usually helps to reduce distortion as a result of more uniform heat input and deposit thickness than is possible manually; economies in machining allowance are therefore possible.

The allowance to be made for the effects of distortion, especially if the part has to be machined to accurate dimensions, depends on many factors, and must rely greatly on experience. If a plate is preset it is better to overdo the
convexity so that on flattening the deposit is compressed. It must be remembered that if straightening is carried out hot, perhaps at red heat, further distortion occurs on cooling to room temperature so the process should leave the blank still slightly convex, of the order of 0.25 mm per 100 mm. Shrinkage, as well as producing bending in a plate, causes some shrinkage in the base material. The designer should make the blank overlong so that, where necessary, it can be machined back to size. As a guide there should be a minimum of 1% added to the length.

When hardfacing the bore of a die or the periphery of a sleeve, radial shrinkage takes place. The amount of shrinkage depends on the thickness and the extent of the deposit. Other factors are the compositions of the blank and surfacing alloy, the amount of preheat used and the temperature to which the blank rises. All materials lose strength as they are heated, some to a much greater extent than others, and the size of the blank has a great effect on the resistance it shows to deformation. Thus with gas welding where the heat input is greater and more widespread than with arc welding, distortion caused by plastic deformation of the blank is likely to be considerably greater. An austenitic steel is much stronger at elevated temperatures than a carbon steel, so will distort far less. A thicker blank shows less contraction especially if the temperature is kept low during deposition, so that the colder, stronger sections can support the hotter, weaker areas. A narrow band of surfacing on a long sleeve provides less contraction both on diameter and length than a much wider deposit. A figure of 1–2% is a reasonable first estimate.

Many components are so shaped that correction of weld shrinkage distortion is not possible by application of force, as mentioned in the example of a plate or shear blade. In such cases it is necessary to envisage the effect that welding stresses will have on the shape of the part, and to premachine it to a shape that will be largely self-correcting. One such example is shown in Fig. 3.35 which depicts a ring that requires a deposit on one face to form a valve seat or sealing ring. Weld shrinkage will cause the ring to contract in diameter and, at the same time, shrinkage on the deposited face will be greater, leading to the shape shown. By premachining an angle on the top face before welding, the deposit–steel interface becomes nearly flat and the risk of a locally thin deposit is avoided. An element of trial and error is involved in deciding the actual shape of a given component before welding, depending as it does in the finished size, shape and material used, as well as the process and weld parameters employed. A second example (Fig. 3.36), is that of a hollow cylinder to be faced on the inside diameter. Shrinkage will tend to reduce the diameter of the part but this will not be uniform along the length. Typically the ends will shrink more than the middle, so that a taper premachined in the ends of the bore provides compensation and the finished deposit is substantially uniform in thickness.

In all these examples, the result will be more uniform and predictable when the job is mechanised; machining allowances in both the component and the welded deposit can then be reliably reduced with a saving in both materials and time.
Valve seat ring or sealing ring face

Blank design

As-welded

Finish machined

Note: proportions exaggerated for emphasis

3.35 Blank design to accommodate welding distortion on rings.

Internal deposit in bore

Blank design

As-welded

Finish M/C

3.36 Blank design to accommodate welding distortion in bushes.
For protection against fine abrasion weld runs should be at right angles to the direction of travel.

3.37 For protection against fire abrasion weld runs should be at right angles to the direction of travel.

Correct

Incorrect

3.38 There should be sufficient overlapping of weld runs if protection is required against fine abrasion.

Weld patterns

The actual pattern used in the welding operation can greatly affect the efficiency with which the surfaced component resists wear. This is an aspect of design that requires considerable experience and it cannot be reduced easily to design rules. A few examples, however, may give an indication of some of the factors involved.

Hardfacing deposits are generally applied in one of three patterns: continuous cover, stringer beads or individual dots. Selection of the pattern to use depends on a number of factors including function of the component and service conditions.

Continuous coverage is used for hardfacing parts that have a critical size or shape, such as rolls, shaft, tracks, crusher jaws and cones. It is often required on parts subjected to much fine abrasion or erosion. Typical examples would be pump and fan impellers, sand chutes, valve seats, dredge bucket lips and pug mill augers.

For protection against fine abrasion or erosion, weld runs should be at right angles to the direction of travel of the abrasive material (Fig. 3.37), and care should be taken that sufficient overlapping of the weld runs is given to ensure adequate coverage of the area being treated (Fig. 3.38).

This design feature does not apply if a very hard deposit is being laid down and relief cracking is intended to eliminate excessive, harmful locked-in stress in the deposit. Electrode movement should be parallel to the direction of flow of the fine abradent as the cracking will be at right angles to this and across the direction of flow.
Choosing weld bead patterns for maximum life in service: (a) single surface, unsatisfactory; (b) both surfaces, also unsatisfactory; (c) optimum solution.

Stringer beads are often used when it is not necessary to cover the base material completely. Typical examples are dragline buckets and teeth, ripper teeth and rock chutes.

There are a number of fundamental guidelines for the designer. If surfacing of shovel teeth (Fig. 3.39) is considered, hardfacing one face will not be effective, since, as the unprotected edge wears away, the hardfacing will tend to chip off because of lack of support (Fig. 3.39a). However, coating both faces (Fig. 3.39b) will not give optimum results either, because as the hard surface wears, cavitation of the substrate is probable with the likelihood of chipping of both deposits. The best results are found with a surfacing pattern shown in Fig. 3.39(c), where additional protection of the sides greatly assists resistance to wear, and hardfacing the top, which is more highly stressed by the abradent, reduces the rate of abrasion of the base material, and is a self-sharpening action with the gradual wear along the bottom.

An example of the economical use of deposit patterns is illustrated in Fig. 3.40. This shows a feed screw used in fertiliser manufacture, handling granules containing limestone. The screw is 2.1 m (7 ft) long and 0.8 m (2.5 ft) in diameter and the face of the flights are deposited in a lattice pattern with a chromium carbide applied by manual metal arc welding.
3.40 The life of an unprotected carbon steel feed screw, used in fertiliser manufacture, is about one month. A chrome carbide deposit of tungsten carbide on the crest of the flights increases life to 9–12 months (courtesy ICI plc).

For teeth working in coarse rocky conditions, it is desirable (Fig. 3.41), for the stringer beads to run parallel to the path of the material being handled. This allows the large lumps of rock, etc., to ride along the top of the hardfacing beads without coming into contact with the base metal. For teeth working in fine sandy conditions the stringer beads are better placed at right angles to the direction of travel. This allows the fine material to compact in the intermediate spaces and provide protection to the base metal. Usually, however, a mixture of coarse and fine material is encountered. For these conditions a combination pattern is particularly effective where the material being handled is damp or stiff, e.g. clay, but it can be ineffective where the material is dry or free flowing. Performance achieved depends greatly on the spacing and height of the weld beads and should be determined experimentally for each application.
For areas where wear is less severe, such as along the rear of buckets and shovels, a dot pattern (Fig. 3.42) is often used. This reduces the cost of material and application time. While not as effective as a waffle pattern it may allow material to compact between the dots, and the high spots offer protection against abrasion from large lumps of rock.

Generally the dots are 15–20 mm in diameter and 10 mm high at about 50 mm centres for earthmoving applications but there can be great variation. One specification for quarrying, also for damp material using material containing tungsten carbide, is for dots 20 mm high with the centres spaced more closely, 6–30 mm apart. Hard and fast rules cannot be formulated. A dot pattern is useful on manganese steel castings to keep down the heat input, and also to restrict excessive heating of the substrate when surfacing quenched and tempered steel.

In the example of the crusher roller shown in Fig. 3.43, transverse beads 12.5 mm wide are applied. Skip welding is used to avoid overheating. The use of a positioner permits welding to be carried out much more easily. Longitudinal beads assist in gripping the material, but such a pattern would be most undesirable in, say, a pump where turbulence would result in decreased performance. For this, complete coverage should be used with the beads deposited parallel to the direction of flow.
For the most severe wear conditions, complete hardfacing is needed and this may be supplemented by superimposed dots of alloys containing chromium carbide or tungsten carbide. This pattern is useful when hardfacing is subject to high tensile stresses, e.g. on rolls, as it can avoid notches which can introduce the risk of fracture.

Where the edge and face of a component is built up with a multiplicity of weld beads (Fig. 3.44), it is essential to specify an adequate deposit on the corner, otherwise rapid wear can take place at the edge.

**Machining allowances**

Part of the design process involves decisions on machining allowances and these depend on two factors:

1. Any change in shape of the component itself caused by welding stresses.
2. Variations in deposit thickness, normally greater on manual deposits than those that are mechanised.

The tendency for distortion and shrinkage of the substrate material, and the importance of providing datum faces for subsequent machining operations,
have been dealt with earlier in this chapter. Because of the variety of circumstances that are met in practice, no design figures can be quoted that will apply to actual jobs, and experience with a given type of work has a key role in blank design.

For work that is likely to be repeated, the following approach is suggested:

1. Apply reasonably tight dimensional limits to the blank dimensions, i.e. minimise variations in size between blanks.
2. Make the surfacing operation as reproducible as possible, e.g. by mechanisation and use of stipulated and controllable welding parameters.
3. Ensure that post-weld heat treatments are always reproduced accurately.
4. Measure and record changes in shape and size of the job, and use them to make controlled adjustments to the blank design of subsequent parts. The object is to economise on substrate and surfacing material, welding and machining time.

When parts having a low chance of repeatability are involved it is likely to be cheaper to err on the generous side of allowances, as the additional cost is likely to be less than the cost of rectification if the part fails to meet drawing requirements after the first attempt.

It usually pays to take a proof cut over the deposit before machining unsurfaced areas. If a need arises to resurface locally, because of a low spot or area, there is then a finishing allowance still on the unsurfaced areas to cope with any small additional distortion from the second welding operation.

When choosing datum faces for the operator to use when machining the coating, choose one as close as possible to the deposit, to minimise errors that may result from shrinkage or distortion of the welded part.

Coating area

It is important to define clearly the area of the component that is to be treated. It must not be too small, as serious wear may occur outside the surfaced portion, but it should not be unduly generous as this adds unnecessary cost. A detailed knowledge of the operation of the component is often essential to obtain optimum results, as the actual positioning of the coating can greatly affect the efficiency with which the part performs. This aspect of design cannot be reduced to simple rules, as it depends on knowledge of the individual application or examination of the worn part.

When a coating is designed for a wear-vulnerable area and successfully reduces the wear at this location, it may be found that other areas of the component – which before had not given wear problems – may start to wear to such an extent that their life determines the life of the whole. This was illustrated dramatically with a sand muller. The machine incorporated a carbon steel shaft mounted vertically in upper and lower bushes. Installed without any hardfacing it was found that after 280 hours' operation the shaft was running eccentrically as the upper bushing and rubbing areas were badly worn. The lower bushing and shaft, however, had worn very little because they were enclosed and well protected from sand, grit and other
abrasive materials. The upper bushing had considerably less protection. Because wear was slight in the lower section it was decided to hardsurface the worn area only. This involved a hard nickel base alloy applied by spraying and fusing.

After nearly 2000 hours the shaft was again found to be running eccentrically. This time it was found that the lower bushing and shaft had worn. Little wear had taken place on the upper bearing area. Surfacing the lower section of the shaft ensured many thousands of hours of satisfactory operation.

Conveyor screws, auger screws, extrusion screws, etc., provide varying wear patterns in different sections along the length. Pressures rise at the exit end requiring harder coating materials, thicker deposits and more extensive coverage of the face of the flight in many circumstances. The central shaft may also need protection, possibly from abrasive wear or deterioration which is caused primarily by corrosion. The extent to which different areas need protection is often determined by experience, but the designer needs to be aware of the probability of wear, and take as many precautions as possible.

**Physical and mechanical properties of coatings**

The manufacturers of surface coating materials for application by weld surfacing publish data about their products. These figures may represent the undiluted weld metal or the deposited metal applied according to the maker's instructions. Generally, products are designed to provide resistance to various forms of wear and/or corrosion, possibly over a wide temperature range. To achieve these properties, hardness, structure and corrosion resistance take precedence over tensile strength, elongation and other mechanical characteristics. It is normally assumed that the latter properties are provided by the material from which the body of the part is made and that this will provide adequate support for the coating.

Coating materials designed for resistance to wear usually feature hardness as an important characteristic, and sometimes the strength and toughness of these diminish as the hardness increases. However, this seldom gives a problem in service, although a part exposed to a significant degree of fatigue loading might develop cracks in the coating during use. A softer, tougher coating material may be a compromise solution.

The question of dilution of deposit with substrate material has already been referred to, and this is a variable effect depending on welding process and parameters used. It can be expected to modify the behaviour of the coating alloy from properties quoted for undiluted deposits. With cobalt and nickel base alloys deposited on low alloy steels, the pick-up of iron causes a fall in hardness, an increase in toughness, a drop in erosion resistance and some change in the corrosion resistance, depending on the corrosive medium involved. These changes are relatively small at dilution levels in the region of 5%, a level commonly achieved in good quality deposits. For many purposes, up to 8–10% dilution can prove acceptable in service.

Compared with cobalt and nickel base alloys, which contain small
amounts of iron, many iron-based surfacing consumables have compositions that allow the dilution associated with the welding process. For example, when using MMA electrodes with coating alloys such as austenitic steels, dilution can affect corrosion resistance because of a reduction in the effective chromium content, or an increase in carbon content through pick-up of carbon from the steel substrate.

Reference has been made earlier to the influence of welding conditions and process on the metallographic structure of deposits. Coating alloys that show large primary carbide development when applied by gas welding (a slow cooling rate) may have this suppressed by the faster cooling rates of some arc processes. This could lead to an increase in the toughness of the alloy deposit and a reduced risk of contraction cracking.

Practical considerations

Manual versus mechanised surfacing

Weld surfacing originated from a need by plant users to restore worn parts to a usable condition and if possible to extend life between breakdowns. For this purpose, manual processes may be appropriate to the infrequent nature of the work. However, some variation in quality of the deposits may be a consequence of this approach. The demand for parts bought as replacements to have deposits of wear-resistant alloys incorporated in their manufacture naturally led to the need for mechanised surfacing processes to deal with the larger batch quantities as efficiently as possible.

Important benefits of mechanised processes are as follows:

- higher rates of deposition, greater productivity;
- greater precision of weld bead shape and position;
- less surfacing material used;
- smaller machining allowances needed, and possible savings on machining time;
- improved product uniformity and performance;
- fewer rejects and rework;
- less operator skill needed;
- reduced operator fatigue and discomfort.

Certain weld surfacing processes, such as submerged arc, are unsuited to manual operation. Mechanisation of some manual processes can be carried out but it is those processes that use a continuous form of surfacing consumable which lend themselves best to mechanisation. The MMA process is not a candidate for mechanisation and the oxyacetylene process, although capable of mechanisation is being displaced by electric arc processes.

Apart from submerged arc referred to above, those most suitable for mechanisation are TIG/GTAW (Fig. 3.45), PTA, MIG/GMAW and FCAW (see Table 3.1). The first two are non-consumable electrode processes and require separate adjustment of welding current and consumable feed rate. The other two regulate consumable feed rate in accordance with welding current. A consequence of this difference between the two groups is that the
latter will automatically compensate for small changes in gun to workpiece distances while the former may require a separate arc length control mechanism.

The principal items of additional equipment required to mechanise a process are those that provide relative movement between torch and workpiece. A positioner is used to rotate a circular or cylindrical workpiece at the required surface speed. The example shown in Fig. 3.46 is fitted with
Plasma transferred arc system for mechanised surfacing of extenders screws up to 2.4m long and 680kg weight (courtesy Deloro Stellite Group Ltd).

A tilting table and variable speed drive. Such a unit is not required if, for example, weld deposits are to be applied only to flat surfaces.

A manipulator is used to control the position of the welding head and to provide movement to this during the welding cycle. One such unit is the column and boom shown alongside the positioner in Fig. 3.46. The boom is adjustable for height on the column and is provided with a variable speed drive laterally. Thus a cylindrical workpiece secured in the chuck attached to the faceplate of the positioner can be surfaced on its diameter with a continuous spiral weld bead as the manipulator traverses the welding head along the workpiece.

Sometimes, it may be desirable to weave the weld bead, and to achieve this an oscillating mechanism is attached to the end of the boom. An alternative form of manipulator is shown in Fig. 3.47. This is a side beam carriage and provides linear movement only. In this example the side beam is teamed with a lathe-type manipulator designed to rotate long shaft-type workpieces between chuck and back centre. To cater for variations in the shaft diameter, an arc length control system is fitted between the torch and the carriage.

The equipment illustrated in Fig. 3.46 and 3.47 could be used for TIG, PTA or MIG/MAG and, of the two, the column and boom set-up would provide the more flexible arrangement for a shop handling a variety of work.

Figure 3.48 shows a purpose-built unit for incorporation in a production line. It is designed to weld surface small circular components such as valves and seats or seal rings. Built to use the PTA process, it incorporates an enclosed welding cabinet which contains a small tilting and rotating table to carry the workpieces. The PTA torch is carried on the end of an arm
extending through the cabinet wall to the oscillating mechanism fitted to the left side of the machine, which provides weld bead weaving where needed. The surfacing consumable is an alloy powder held in the container on top of the cabinet and fed through a metering device to the torch. On top of the welding power supply is a system that controls the whole welding cycle to preset parameters. This can include mechanised loading and unloading of the workpieces through the rear of the cabinet.

The advantage of such a system is that once weld surfacing parameters have been set, the unit operates with minimum attention provided that supplies of workpieces and surfacing consumables are maintained. Provision can be made to store a series of programmes to suit a range of workpieces.

**Accessibility**

The problem of access arises in weld surfacing when the region to be deposited is located inside a component, in such a way that it is difficult to get the welding torch and consumable feed inside, and present them at the required angle to the work surface. Even if this is possible, it may be difficult to see what is happening and to make suitable adjustments to the welding parameters while deposition is taking place.

In such circumstances manual deposition is usually not possible because
of poor visibility and consequent loss of control. The difficulty can be avoided by fabricating the piece after deposition of the temporarily accessible surface, welding or brazing castings of the surfacing alloy in position, or melting precast shaped inserts in position by means of a modified TIG torch of slender proportions.

Two developments have, however, made this type of operation easier for designers. The first is the introduction of small PTA torches which work with a powder consumable and are capable of working inside a bore of 35mm (Fig. 3.49) with a nozzle set at 90° or 45° to the torch axis. Current models have a ‘reach’ of about 450mm. A longer reach is possible using specially designed MIG/MAG guns, but in this case minimum diameter is about 75mm. Both types of torch/gun are designed for mechanised use only and it is normal to secure area coverage of the deposit by arranging a slow spiral deposit with suitable bead overlap. This requires a suitable speed relationship between the positioner and the manipulator. To obtain satisfactory results, the component must be true and concentric to the axis of rotation. By turning end to end, it is possible to cover a bore double the length of the torch reach, assuming access from both ends.

**Weld defects**

Weld-deposited coatings are extensively used in industry and in widely differing circumstances, so quality requirements also differ. The highest quality demands a trained, qualified and experienced workforce, but much weld surfacing is carried out successfully under different circumstances when fitness for purpose is the requirement. This section deals with imperfections that may arise and, if considered to be prejudicial to the service conditions that will be met, how to deal with them.

The specification under which work is carried out should not only lay down the surfacing procedure to be used and precautions to be observed but should define tests and acceptance standards to be met. If the finished parts, on inspection, fail to meet these standards, it is essential to identify faults which may be rectified and the rectification procedures which are permissible. The inspection schedule may include in-process inspections at various stages of the work – some are obvious, such as the cleanli-
ness of the blank, adherence to dimensional tolerances, thickness of deposit, deslagging between successive deposit layers and so on. This section is confined to comments on inspection after surfacing has been completed. The designer can use these in development of a specification suited to each job.

Inspection is facilitated if the work, after weld surfacing, is grit blasted to remove scale and oxides. Grit blasting after rough machining is a valuable test for soundness in the deposit as it can reveal porous areas. Proof machining of components that have to be finally ground or machined to precise dimensions is essential, as an inspection at this stage reveals undersize areas that can be built up further while there is sufficient material in the rest of the part available to cover scaling or distortion as noted earlier. The rectification allowable depends on the process and alloys used, and can range from spotting and more extensive additional surfacing, observing the necessary heating requirements, to complete removal of the deposit and re-initiation of the surfacing cycle. A repair procedure should be laid down, rather than be left to the operator.

The deposit must be viewed carefully for cracks, porosity, pinholes and inclusions which may have to be rectified. Visual inspection may be aided by dye penetrant inspection. When used on an as-welded surface care must be taken in interpretation of dye penetrant tests. Such tests should be repeated after proof machining. At this stage additional tests may be specified, such as hardness, radiographic or ultrasonic examinations. As surface cracking may be caused by incorrect grinding of harder deposits after final machining, dye penetrant testing should be repeated, as well as dimensional checks.

The results of a testing procedure can vary according to the skill and training of the operator and the techniques that are used. It is therefore desirable that techniques be laid down in detail, and audited from time to time. An acceptance standard should be stated for each job.

Cracks

Contraction cracking

Contraction cracking is normally visible to the naked eye and may not necessarily affect the performance of the component adversely. In the harder surfacing materials, providing maximum wear resistance, such cracking (relief cracking) may be encouraged to release locked-in tensile stresses. Such cracking rarely involves the risk of the coating breaking away from the base metal, provided there is no hardening of the HAZ, and a satisfactory bonding to the substrate has been achieved. However, cracks cannot normally be tolerated in applications such as:

- sealing surfaces of valves, mechanical seat rings, printing rolls, etc.;
- surfaces subject to erosion such as flow control valves;
- surfaces designed to provide both wear and corrosion resistance;
- surfaces subject to severe fatigue stresses in service;
- surfaces that must not pick up any process material that could contaminate subsequent batches, such as in plastic extrusion.
Grinding cracks

Grinding cracks are usually extremely shallow, but as they may extend under mechanical or thermal stress they must be treated as contraction cracking.

Subsurface cracks

If the deposit lifts from the base metal, areas may flake off or spall even under slight mechanical pressure, or relatively small surface temperature change and therefore such parts must be rejected, not repaired. If the condition arises from faulty processing, and the reason can be identified, the deposit should be removed by machining or grinding before reprocessing, taking steps to avoid a recurrence of the problem. Avoid repair situations that result in excessive deposit thickness, for example by applying an initial buttering layer (for example, mild steel or austenitic stainless steel – see the section on group 1 materials earlier in this chapter and that on treatment of defects below).

Porosity

The degree of porosity, if any, and the size of the pores can vary. There can be a few specks barely visible to the unaided eye or large pores and visibly spongy areas if poor preparation or technique had been used. Pictorial standards are desirable from which a grading can be specified for each job. A numerical approach is an alternative, e.g. a maximum of $X$ pores per square $Y$, the pore size not to exceed $Z$ micrometres. The designer should not call for a higher quality than is required for the efficient performance of the surface, and this may differ from one area to another. Thus with a seal or valve seat the hardfacing frequently extends beyond the actual sealing zone and some porosity outside this zone may not be detrimental. The component drawing should show the different zones and indicate the acceptable standard for each.

Inclusions

Inclusions, like porosity, can range from visibly obvious defects to very fine dispersions. Removal on machining or during the working life of the part produces the same problems as pores. Their identification and classification can be treated similarly.

Treatment of defects

Cracks

For applications where cracking of any sort is unacceptable, it is sometimes possible to remove the entire deposit and to start again, provided that:

- the cause of the cracking is known and recurrence can be prevented;
- there is adequate machining allowance on uncoated areas of the component:
the part, after removal of the faulty deposit, is heat treated to remove welding stresses, and to correct any unacceptable metallurgical structures such as a hard HAZ;

- the increased thickness of the deposit resulting from machining to remove the faulty deposit and to get down to clean, uncracked substrate material, is acceptable from the design and use point of view, and unlikely to cause recurrence of the cracking problem;

- the newly prepared blank is checked by dye penetrant testing and found to be free from all defects.

Once formed, grinding cracks in some materials, although initially shallow, propagate more deeply if further stock removal takes place, so total removal may sometimes be necessary before rewelding.

Any defects penetrating locally into the blank should be excavated by local grinding to give an open recess, not a drilled hole, so that the welding flame or arc can access the bottom of the hole. The defect can then be rewelded, stress relieved, remachined and tested before resurfacing. Never use the surfacing alloy to fill excavations that have penetrated the substrate, as the local change in deposit thickness can cause further cracks. Instead, repair the substrate with a compatible, ductile, non-hardening filler material, and then follow with the surfacing alloy.

Do not attempt to repair cracks in the deposit or the substrate without excavation to remove the defect entirely; the heat of welding is likely to propagate the defect even more deeply.

Porosity and inclusions

The distribution of porosity, if unacceptable, determines the method of rectification. A localised defect, possibly in the final weld crater, may be repaired by local excavation, dye penetrant testing to verify complete removal of the defect, and rewelding. As with cracks, attempts to repair defects by welding without their prior mechanical removal invariably fail.

More widely distributed defects suggest a fundamental problem with the materials used, or the procedures, and this will require investigation. If the cause is identified and can be avoided in future, it is probably best to remove the whole deposit and reweld.

Inclusions should be treated in the same way as porosity.

The case for and against repair of defects, assuming that it is considered to be a practical proposition, depends on factors such as the cost compared with using a new blank, whether sufficient machining allowance is still available on uncoated areas, and on metallurgical considerations of effects of further exposure to welding temperatures on the substrate.
Chapter 4

Thermal spray processes

Thermal spray processing is a very rapidly expanding field of surface engineering. The driving forces for development include the economics of aero-engine and oil and gas industries. However, the technologies that are being developed to suit their needs may easily be transferred to a myriad other applications.

The principal characteristics of thermal spraying processes that distinguish them from weld-deposited coatings are as follows:

1. The strength of the bond between coating and substrate covers a wide range, depending on the materials and process used. It can vary from a relatively low strength, to figures approaching those of welded bonds if the process involves high temperature diffusion between coating particles and substrate.

2. Thermal spraying can apply coating materials to substrates that are unsuited to welding because of their composition or tendency to distort. This feature offers the designer scope to use materials with desirable properties that would not be possible by other means.

3. Sprayed deposits can be applied in thinner layers than welded coatings, but thick coatings are possible under certain circumstances.

4. Almost all material compositions may be deposited (provided at least one constituent has a stable liquid phase), metals, ceramics, carbides, polymers or any combination. This chapter deals with coatings other than polymers, which are covered in Chapter 7.

5. Most processes are 'cold' compared with welding and there is no dilution or metallurgical degradation of the substrate.

6. Thermal spraying processes are all-positional and most can be operated in air, so they possess great flexibility.

In all thermal spraying processes, the consumable coating material fed to the spray gun is raised in temperature and projected in particulate form to strike the workpiece. On arrival, the hot particles form splats which interlock and gradually build up a coating of the desired thickness (Fig. 4.1). The particles must be at least partially molten in transit for splats to form. The density and cohesion of the coating depends on the material, its temperature when it strikes the workpiece, and its impact energy.

Adhesion of the coating to the substrate depends on the same factors plus the surface condition of the substrate, which must be clean and suitably
roughened. In most cases adhesion is believed to be largely due to mechanical keying. Owing to the nature of the bond between coating and substrate, it is generally not recommended that a sprayed coating should be loaded in shear.

Processes available for thermal spraying have been developed specifically for the purpose and fall into two categories – high and low energy processes. The key processes and their energy sources are summarised in Table 4.1.

The lower energy processes, often referred to as metallising, are arc and flame spraying. These are extensively used for spraying metals for corrosion resistance such as zinc and aluminium, for service at or near ambient temperature, on large structures and in circumstances where thermal and mechanical shock or abrasive wear are small.

Some porosity will always be present in these coatings. This may be beneficial, as residual stresses in porous coatings tend to be lower, and thicker coatings can be applied without the risk of lifting from the substrate. Porosity can also act as a reservoir for lubricant for a sprayed shaft running in a bearing. Porosity can be sealed by a material such as a lacquer or resin in circumstances where corrosion could otherwise penetrate to the substrate.

After preparation machining and degreasing, the substrate is prepared for spraying by shot blasting with angular iron grit or occasionally by rough machining (e.g. threading).

The higher energy processes, which include plasma, D-gun and high velocity combustion spraying, were developed to provide coatings of much lower porosity and improved adhesion to the substrate. They can also handle materials of higher melting points, thus widening the range of applications, to include coatings having resistance to higher temperatures and to thermal and mechanical shock.

With higher energy processes, the porosity is much lower and bond strengths are higher, both within the coating itself, and between the coating...
### Table 4.1 Definition of spraying processes

<table>
<thead>
<tr>
<th>Process</th>
<th>Energy source</th>
<th>Other names for the process</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Low energy processes</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Flame spraying</td>
<td>Oxyfuel combustion</td>
<td>Oxyfuel gas – powder spraying</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Oxyfuel gas – wire spraying</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Metallising</td>
</tr>
<tr>
<td>Arc spraying</td>
<td>Electrical</td>
<td>Electric arc spraying</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Twin-wire arc spraying</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Metallising</td>
</tr>
<tr>
<td><strong>High energy processes</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Plasma spraying</td>
<td>Electrical</td>
<td>Air plasma spraying</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Vacuum plasma spraying</td>
</tr>
<tr>
<td>Detonation flame spraying</td>
<td>Oxyfuel combustion</td>
<td>D-gun</td>
</tr>
<tr>
<td>High velocity oxyfuel spraying</td>
<td>Oxyfuel combustion</td>
<td>HVOF spraying</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High velocity oxygen fuel spraying</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High velocity flame spraying</td>
</tr>
<tr>
<td></td>
<td></td>
<td>High velocity air fuel</td>
</tr>
</tbody>
</table>

and substrate. This is because of the disruption of any oxide films present on the particles or the workpiece surface, and there is some evidence that some diffusion bonding takes place. However, cleaning and roughening of the workpiece is just as important as for the lower energy processes if good results are to be obtained consistently. Alumina or silicon carbide grits are often used in place of angular iron grit, as a finer particle size is preferred.

Other differences distinguish the two groups of processes mentioned above. The low energy ‘metallising’ group has lower capital cost and is relatively portable, while the higher energy group has a higher capital cost and tends to be used in fixed installations. However, there are some industrial applications where a process from either group could be used successfully.

Consumables used for thermal spraying are available in two forms. Solid drawn wire is used for metallising processes. In addition some materials are produced as a fine dispersion of particles in an organic binder, which is also in wire form and is known as a ‘cord’. For higher energy processes a powder consumable is used. This has advantages for non-metallic materials and can also be transported more easily to the point of spraying, especially for torches designed to be used in small bores. The optimum specification of powder depends on the process, and includes the particle size distribution, morphology and manufacturing route.

Included in this chapter is the spray fuse process. This technique involves spraying to apply a coating of special self-fluxing alloys based on cobalt or nickel. The coating is then raised in temperature to between the solidus and the liquidus of the alloy, when the self-fluxing action allows diffusion bonding between the sprayed particles and the substrate.
Thermal spray materials

Material form

Powder morphology

Powders for thermal spraying vary a great deal, depending on the route by which they have been manufactured. The morphology can be spheroidal, as shown in Fig. 4.2(a), or may be more angular, as in Fig. 4.2(b). This can affect the way in which the powder feeds through the delivery system. Some compositions are supplied as a mechanical blend, and it is important to make sure that the constituents are thoroughly mixed before spraying begins. Others are pre-alloyed and are therefore homogeneous. In all cases it is important to ensure that the powder is thoroughly dry before spraying, by storing it in a warmed cabinet.

Wires for spraying

Wires for spraying are used in two processes – oxygas spraying, and arc spraying. For the former process, they may be pure metals, alloys or composites. This enables a wide range of materials to be used.

Wires for spraying by the arc process must possess certain characteristics. They must be good conductors of electricity, which limits the degree to which the wire material may be alloyed. They must also be robust enough to be fed through the wire delivery system. Extra alloying elements, or second phases such as carbides may be introduced by making the wire cored, but the range of compositions is more limited than the range possible in pow-
Thermal spray processes

dered materials. Nevertheless, there is a useful range of cored wires available for arc spraying designed to give composite or highly alloyed coatings.

**Materials with good bonding characteristics**

Experience has shown that certain materials provide an excellent bond to the substrate. This fact has led to their use as an interlayer and also in the form of a blend with some surfacing alloys, which would otherwise exhibit rather poor adhesion. Common bond coat materials are listed in Table 4.2. Several are also used for surfacing in their own right, and these appear in other tables also.

**Ferrous alloys**

Mild low alloy and carbon steels are most frequently used to build up worn areas of components where corrosion resistance is not required (Table 4.3). Stainless steels can also be sprayed and are frequently used for reclamation where corrosion resistance is needed. Iron–chromium alloys are also sprayed for corrosion resistance.

**Nickel base alloys**

All nickel base alloys are corrosion-resistant, and those containing chromium have good resistance to oxidation at elevated temperatures (Table 4.4). Nickel–chromium–boron–silicon alloys can be fused after spraying to give coatings that are free from porosity and are metallurgically bonded to the substrate.

A newer development is to attempt to form intermetallic compounds, such as nickel aluminide either during spraying, or by suitable heat treatment afterwards. These coatings are expected to show superior corrosion resistance.

NiCrAlY coatings are used for resistance to high temperature oxidation and sulphidation. They are particularly useful in gas turbines.

**Cobalt base alloys**

Alloys based on cobalt have high resistance to softening at elevated temperatures, together with corrosion and wear resistance (Table 4.5). Alloys modified by inclusion of boron and silicon can be fused after spraying to give pore-free metallurgically bonded overlays (see spray fusing later in the chapter). Co–Cr–Al–Y coatings are used for similar applications to the nickel-based alternative.

**Other non-ferrous metals and alloys**

Several other metals and alloys are thermally sprayed; see Table 4.6. Aluminium and zinc find specific application where properties such
**Table 4.2 Bond coats**

<table>
<thead>
<tr>
<th>Materials</th>
<th>Characteristics and applications</th>
<th>Suitable for (processes)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Flame (wire)</td>
</tr>
<tr>
<td>Ni-Al</td>
<td>95-5 and 80-20 composite and pre-alloyed materials. Bond coat and for building up dimensions</td>
<td>\</td>
</tr>
<tr>
<td>Ni-Cr–Al</td>
<td>As for nickel-aluminium but giving higher temperature resistance</td>
<td>\</td>
</tr>
<tr>
<td>Mo</td>
<td>Useful also as a wear resistance coating, see Table 4.6</td>
<td>\</td>
</tr>
<tr>
<td>Ni–Cr</td>
<td>Bond coat for ceramics, see also Table 4.4</td>
<td>\</td>
</tr>
<tr>
<td>Ni–Cr–B–Si/Ni–Al</td>
<td>Typical of a number of blended powder materials</td>
<td>\</td>
</tr>
<tr>
<td>Ni–Cr–B–Si/Mo</td>
<td></td>
<td>\</td>
</tr>
<tr>
<td>M–Cr–AlY (M = Co, Ni or Fe)</td>
<td>Used as bond coats for yttria-stabilised zirconia. See also Tables 4.4 and 4.5</td>
<td>\</td>
</tr>
</tbody>
</table>
as resistance to atmospheric corrosion are of prime importance. Bronzes are used for bearing surfaces, as are Babbitt alloys. Copper is an excellent electrical conductor and can be used for electromagnetic shielding.

Molybdenum spray parameters are chosen to minimise oxidation in the deposit when it is used as a bond coat. However, the hardness of the deposit rises with an increase in oxide content, giving a range of 400–800HV, producing useful wear-resistant properties. Deposits are applied to the synchronising rings in gear boxes, a typical hardness for these being 650HV. Harder deposits are resistant to galling and fretting and are used, for example, on engine piston rings. The maximum service temperature is 300°C.
<table>
<thead>
<tr>
<th>Alloy type</th>
<th>Characteristics and applications</th>
<th>Suitable for (processes)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nickel</td>
<td>Dense, easily machined coatings, used for reclamation</td>
<td>Flame (wire)</td>
</tr>
<tr>
<td>Ni-Cu</td>
<td>Gives high density machinable coatings with good corrosion resistance, especially under marine conditions</td>
<td>Flame (powder)</td>
</tr>
<tr>
<td>Ni-Cr (80/20)</td>
<td>Resistance to corrosion and high temperature oxidation. Used as a precoat for application of ceramics</td>
<td>Spray-fused coatings</td>
</tr>
<tr>
<td>Ni-Cr-Fe</td>
<td>Corrosion and high temperature-resistant coatings</td>
<td>Electric arc spray</td>
</tr>
<tr>
<td>Ni-Cr-Mo-W</td>
<td>Resistant to corrosion and wear; work hardens under impact</td>
<td>Plasma spray</td>
</tr>
<tr>
<td>Ni-Cr-Mo-Fe (alloy 625)</td>
<td>Corrosion-resistant, suitable for marine environment</td>
<td>HVOF/detonation spraying</td>
</tr>
<tr>
<td>Ni-Cr-B-Si</td>
<td>A range of alloys with hardnesses from 200 to 700HV. High wear and corrosion resistance and good retention of hardness at temperature</td>
<td></td>
</tr>
<tr>
<td>Ni-Cr-Al-Y</td>
<td>Resistant to high temperature oxidation and sulphidation</td>
<td></td>
</tr>
<tr>
<td>Ni-Cr-Fe-Si-B</td>
<td>Corrosion resistance</td>
<td>(low pressure)</td>
</tr>
<tr>
<td>Ni-Cr-B-Si-Cu-Mo</td>
<td>High corrosion, wear and heat resistance with wide fusing range enabling it to be used on complex shaped components</td>
<td></td>
</tr>
<tr>
<td>Alloy type</td>
<td>Characteristics and applications</td>
<td>Suitable for (processes)</td>
</tr>
<tr>
<td>--------------------</td>
<td>--------------------------------------------------------------------------------------------------</td>
<td>--------------------------</td>
</tr>
<tr>
<td>Co-Cr-W</td>
<td>A range of alloys with hardinesses from 300 to 600HV. High resistance to wear, corrosion and heat</td>
<td>Flame (wire)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Flame (powder)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Spray-fused coatings</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Electric arc spray</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Plasma spray</td>
</tr>
<tr>
<td></td>
<td></td>
<td>HVOF/detonation spraying</td>
</tr>
<tr>
<td>Co-Cr-Ni-W</td>
<td>Used for coatings where heat resistance combined with resistance to fretting wear is required</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Co-Cr-Mo</td>
<td>Hardness 300HV. This alloy has higher corrosion resistance and ductility than the above. Resistant to fretting</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Co-Cr-Al-Y</td>
<td>Resistant to high temperature oxidation and sulphidation</td>
<td>(low pressure)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Co-Cr-Mo-Si intermetallic</td>
<td>Good corrosion resistance and wear resistance</td>
<td></td>
</tr>
<tr>
<td>Co-Cr-W-B-Si</td>
<td>A range of very hard alloys, from 400 to 700HV. Used for production of spray-fused coatings</td>
<td></td>
</tr>
<tr>
<td>Co-Cr-W-Ni-Fe-C-S</td>
<td>Abrasive wear, corrosion</td>
<td></td>
</tr>
<tr>
<td>Materials</td>
<td>Characteristics and applications</td>
<td>Suitable for (processes)</td>
</tr>
<tr>
<td>-------------------</td>
<td>-------------------------------------------------------------------------------------------------</td>
<td>-------------------------------------------------------------</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Flame (wire)</td>
</tr>
<tr>
<td>Al</td>
<td>Used principally as a corrosion-resistant coating on steels, for neutral and acidic environment. When heat treated provides a high temperature oxidation-resistant alloy coating on steels. Repair of Al substrates</td>
<td></td>
</tr>
<tr>
<td>Al-Si</td>
<td>Repair, tooling</td>
<td></td>
</tr>
<tr>
<td>Zn</td>
<td>Used for corrosion-resistant coatings on steels, neutral and alkaline environment</td>
<td></td>
</tr>
<tr>
<td>Mo</td>
<td>Low friction, wear-resistant to 400 °C, metal-to-metal wear</td>
<td></td>
</tr>
<tr>
<td>Bronze</td>
<td>For bearing surfaces</td>
<td></td>
</tr>
<tr>
<td>Aluminium bronze</td>
<td>Corrosion-resistant coatings with good bearing properties. Also used as a bond coat with arc spray process</td>
<td></td>
</tr>
<tr>
<td>Cu</td>
<td>Used for electrical conductivity and screening from electromagnetic interference, reclamation of mis-machined parts</td>
<td></td>
</tr>
<tr>
<td>Cu-Ni-In</td>
<td>Good resistance to galling and fretting</td>
<td></td>
</tr>
<tr>
<td>Cu-Zn brasses</td>
<td>For decorative work</td>
<td></td>
</tr>
<tr>
<td>Sn</td>
<td>Food processing vessels, EMC protection</td>
<td></td>
</tr>
<tr>
<td>Sn-Zn</td>
<td>Capacitor end connections</td>
<td></td>
</tr>
<tr>
<td>Babbitt metal</td>
<td>Heavy duty bearings</td>
<td></td>
</tr>
</tbody>
</table>
**Tungsten carbide and chromium carbide**

As a sintered composite with cobalt, tungsten carbide is used for oxyfuel, plasma and detonation coatings (Table 4.7). It blends with nickel or cobalt base self-fluxing alloys and can be used to provide spray-fuse coatings. Chromium carbide is used, generally in conjunction with Ni–Cr, to confer resistance to high temperature wear. Research is currently searching for alternative carbides, to reduce the cost of these coatings. Candidate materials include titanium carbide and vanadium carbide.

**Ceramics**

Ceramics have high temperature resistance and are resistant to wear and erosion (Table 4.8). Coatings sprayed by high energy processes, such as oxyfuel gas spraying from rods or cords, or from powders applied by plasma or detonation, have high density. Lower energy flame spray coatings made using powders may be porous and may sometimes need to be sealed. Porosity is sometimes deliberately sought to give higher thermal insulating properties. Many proprietary blends of ceramic powders are used, but those given in Table 4.8 are considered the most important.

Zirconia and magnesium zirconate are prized for their thermal insulating properties, and are used in equipment operating at high temperatures to shield the underlying metal. This allows the operating temperature of the engine to be raised, thus improving its thermodynamic efficiency. Examples are shown in Fig. 4.3 and 4.4.

**Abraudable coatings**

A special class of materials called abraudable coatings is included in Table 4.9. These materials are specifically formulated to wear away wherever parts are required to have minimal operating clearances. As may be imagined, care has been taken to ensure that the material that abrades does not then produce wear debris which will cause damage elsewhere. Applications include the compressor section of jet engines.

**Graded coatings**

Because of wide differences in the expansion characteristics of ceramics and steels it is sometimes necessary to use a cermet or graded coating system to reduce the tendency to cracking and spalling under thermal cycling. In these situations it is often possible to use a first coating with 70–80% metal component, a second layer of 20–30% metal and a final layer wholly of the required ceramic.

Cermets are useful in other circumstances, for instance to combine wear resistance of a ceramic material with the higher thermal conductivity of a metal.
<table>
<thead>
<tr>
<th>Materials</th>
<th>Characteristics and applications</th>
<th>Suitable for (processes)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Flame (wire)</td>
</tr>
<tr>
<td>Tungsten carbide/Co</td>
<td>Composite powders with 10–20% cobalt. Used for coatings subject to wear plus impact or erosion at temperatures up to 450°C</td>
<td></td>
</tr>
<tr>
<td>Tungsten carbide/Ni</td>
<td>For abrasive wear</td>
<td></td>
</tr>
<tr>
<td>Tungsten carbide/Co–Cr</td>
<td>Abrasion in water-based solutions. Co–Cr matrix gives high abrasion/corrosion resistance, up to 600°C</td>
<td></td>
</tr>
<tr>
<td>Tungsten carbide/Ni–Cr–B–Si alloy blended powders</td>
<td>Fused coatings having very high wear resistance</td>
<td></td>
</tr>
<tr>
<td>Tungsten carbide/Co–Cr–W–B–Si alloy blended powders</td>
<td>Fused coatings having very high wear resistance</td>
<td></td>
</tr>
<tr>
<td>Chromium carbide/Ni–Cr</td>
<td>Composite powders for coatings subject to wear at temperatures up to 800°C</td>
<td></td>
</tr>
<tr>
<td>Materials</td>
<td>Characteristics and applications</td>
<td>Suitable for (processes)</td>
</tr>
<tr>
<td>---------------------------------------</td>
<td>--------------------------------------------------------------------------------------------------</td>
<td>-------------------------</td>
</tr>
<tr>
<td>Alumina</td>
<td>Wear-resistant coating with high electrical resistance</td>
<td>C</td>
</tr>
<tr>
<td>Zirconia (yttria-stabilised)</td>
<td>Thermal barrier coatings having good resistance to thermal shock</td>
<td>C</td>
</tr>
<tr>
<td>Zirconia (calcia-stabilised)</td>
<td>Severe thermal barrier applications</td>
<td></td>
</tr>
<tr>
<td>Chromium oxide</td>
<td>Very high wear resistance, corrosion-, erosion- and oxidation-resistant</td>
<td>C</td>
</tr>
<tr>
<td>Alumina–titania</td>
<td>Compositions available from 2 to 40% titania; coatings give higher wear resistance than 'pure' alumina</td>
<td>C</td>
</tr>
<tr>
<td>Titania</td>
<td>Resistant to wear, high temperatures and corrosion</td>
<td>C</td>
</tr>
<tr>
<td>Magnesium zirconate</td>
<td>Thermal barrier coating with high resistance to erosion by fine particles and molten metals</td>
<td></td>
</tr>
<tr>
<td>Hydroxyapatite</td>
<td>Biocompatible coatings for medical prostheses and dentistry</td>
<td></td>
</tr>
</tbody>
</table>

C = available as a plastic covered cord.
### Table 4.9 Abradable coatings (all for air plasma spraying)

<table>
<thead>
<tr>
<th>Base material</th>
<th>Other constituents</th>
</tr>
</thead>
<tbody>
<tr>
<td>Al</td>
<td>Graphite</td>
</tr>
<tr>
<td>Al</td>
<td>Si, polyester</td>
</tr>
<tr>
<td>Ni</td>
<td>Graphite</td>
</tr>
<tr>
<td>Ni</td>
<td>Cr, Al, polyester</td>
</tr>
<tr>
<td>Ni</td>
<td>Cr, Al, bentonite</td>
</tr>
<tr>
<td>Ni</td>
<td>Cr, Fe, boron nitride</td>
</tr>
<tr>
<td>Cu</td>
<td>Al, polyester</td>
</tr>
</tbody>
</table>

### Characteristics of generic low energy thermal spray processes

**Oxyfuel gas wire spraying**

In oxyfuel gas wire spraying the coating material, in the form of a rod, wire or cord, is fed by a variable speed motor into the centre of the multijet flame which melts the tip of the wire. An annular gas jet then propels the molten material on to the substrate (Fig. 4.5 and 4.6).

**Principal applications**

Wire spraying is used extensively for deposition of zinc and aluminium coatings for corrosion protection. Worn or mismachined parts can be built up using steels, copper and copper alloys, nickel and its alloys or with ceramics.

**Advantages**

1. Relatively low cost equipment.
2. High spray rates.
3. Wide range of surfacing materials can be sprayed.
4. Adaptable for differing wire diameters.
5. Spraying can be mechanised.

**Disadvantages**

1. Limited to consumable in wire form.
2. Lower density deposits and lower adhesion strength than arc spray.

**Oxyfuel gas powder spraying**

The surfacing material, in the form of a powder, is contained in a hopper from which it is conveyed into the spray flame, either by reduced pressure generated by a venturi in the gas stream or in a carrier gas (usually air); see
4.3 Future diesel engines may be insulated for improved efficiency. Components of a diesel engine coated with a 2mm thickness thermal barrier applied by plasma spraying (courtesy of Materials Engineering Centre, AERE).
4.4 Microstructure of a typical plasma sprayed three layer thermal barrier system - metallic bond (NiCrAl + ZrO₂), zirconia (A = substrate, B = NiCrAl; C = cermet; D = zirconia) (courtesy of Materials Engineering Centre AERE).

4.5 Wire combustion spray method.

4.6 Typical oxyfuel gas wire spray gun for low melting point metals including zinc, aluminium, tin and Babbitt (courtesy Metco Ltd).
Fig. 4.7. Some designs of powder torch use additional air to propel the heated powder particles on to the substrate, to confine the spray stream, and/or to cool the substrate.

**Principal applications**

Powder spraying is used extensively for application of wear-resistant coatings of nickel or cobalt base alloys, which are subsequently fused to give fully dense, metallurgically bonded alloys.

Metal powders such as steels, copper alloys, nickel alloys and others are used to restore worn surfaces.

**Advantages**

1. The most suitable flame spray process for high alloy and self-fluxing surfacing materials.
2. Gives a continuous supply of surfacing material.
3. Can be used to provide coatings from materials which cannot be produced as rods and wires.

**Disadvantages**

1. Requires consumable in powder form of suitable mesh range.
2. Gives lower density deposits, of lower adhesion strength than arc spraying (except in self-fluxing alloys that are subsequently fused).

**Arc wire spraying**

Electric arc wire spraying uses twin wires which are fed together into a spray head in which an arc is struck between the wire ends and the resulting molten material is projected from the arc as a series of molten droplets by means of a high pressure gas jet (usually air); see Fig. 4.8 and 4.9.
A heavy duty arc spray pistol for mechanised and automated plant rated at up to 600A (courtesy Metallisation Ltd).

The gas jet used for propulsion of the coating material is normally air, but for specialised applications, for example where a copper coating of very low oxide content is needed, nitrogen can be used for the gas jet.

**Principal applications**

The arc spray process is most suitable for surfacing large components such as rollers for the steel, paper, paint and plastics industries, and to surface and reclaim hydraulic rams and pistons, shafts, bearings, etc.
Advantages

1. Either solid or tubular wires can be used, giving a wide range of coating alloys.
2. Very high spray rates are possible.
3. By using two different wires, composite or 'pseudo-alloy' coatings can be produced.

Disadvantages

1. Limited to consumables available in wire form.
2. Lower density deposits, of lower adhesion than the high velocity processes.

Characteristics of the high energy thermal spray processes

Plasma spraying in air (APS)

A plasma arc spray torch contains a tubular copper anode, in the rear of which is a tungsten cathode. Both electrodes are water cooled, and are surrounded by an insulating body, which holds them in correct relation to each other and serves as an arc chamber. A high current arc is generated within the torch, and a gas injected into the arc chamber where it is heated and, on passing through a constriction in the anode bore, is converted into a high temperature plasma. Powdered surfacing material is injected into this plasma jet and is thus heated and accelerated onto the substrate (Fig. 4.10 and 4.11).

There are significant variations in torch design. The injection of the spraying material axially into the centre of the plasma jet is claimed to give the best results, with high deposition rates and efficiencies. A water-stabilised plasma torch is claimed to give up to ten times the deposition...
Plasma spray torches have been developed in a variety of configurations to satisfy the requirements of particular work, involving manual and mechanised installations and for applications on external and internal surfaces (courtesy Plasma-Technik AG).

Water-stabilised plasma spray torch (courtesy Institute of Plasma Physics, Academy of Sciences of the Czech Republic).

Rates of the more conventional torches, with similar particle speeds. The torch is illustrated in Fig. 4.12.

Robots may be used to hold the torch, enabling the spraying of complex geometries. One such system is shown in Fig. 4.13.
4.13 Robot-assisted plasma spraying for complex component geometries (courtesy Euromat GmbH).

**Principal applications**

APS is used mostly for deposition of refractory high melting point materials, ceramics, carbides and high temperature alloys. Coatings are deposited on aircraft engine components, textile machine parts; wear-resistant coatings are deposited on pump and valve spindles; and APS is used for the application of electrical and thermal insulating coatings.

The water-stabilised torch is used for the production of free-standing forms, such as ceramic tubes. These are deposited on reusable mandrels, which are removed soon after spraying. Tubes are from 20 to 400mm diameter, 2 to 10mm thick and up to 1.8m long.

**Advantages**

1. The high temperature in the plasma enables almost all materials to be sprayed.
Deposits are of high density, and strongly bonded to the substrate. Very low heat input to the substrate.

Disadvantages
1. Higher capital cost than gas and arc spraying, bulky equipment.
2. Deposits of lower density and adhesion than vacuum plasma spraying (VPS).

Vacuum plasma spraying
This process, known as vacuum plasma spraying or low pressure plasma spraying (VPS/LPPS; Fig. 4.14) involves spraying in a chamber initially evacuated to a pressure of $10^{-2}$ mbar and then backfilled with argon to 40 mbar.

By excluding oxygen from the environment, neither the sprayed particles nor the substrate surface becomes oxidised during spraying, and deposits are dense and well bonded. The plasma jet is longer than in air and the workpiece reaches higher temperatures.

Principal applications
Figure 4.15 shows an installation incorporating a robot manipulator with a rotary table positioner used for coating developments. The high quality coatings produced by this process have led to applications on aero-engine parts which are coated with materials such as the M–Cr–Al–Y series, to resist corrosion and wear at high temperatures.

Figures 4.16 and 4.17 show the microstructures of 50–50 Ni–Cr alloy deposited by VPS/LPPS compared with APS, and illustrate the significant increase in density and reduction in oxidation obtained by low pressure operation.
4.15 Developments in plasma spraying include use of robotics and low pressure plasma – the interior of the Materials Engineering Centre’s chamber facility at Harwell (courtesy of AERE Harwell).
Advantages

1. High density, high adhesion, low oxide coatings.
2. Coating properties are reproducible to close limits.

Disadvantages

1. High capital cost, bulky, fixed installation.
2. High heat input to workpiece may cause distortion or metallurgical changes in the substrate.
Detonation coating

Detonation coating is a proprietary process. A device similar to a rifle barrel has powdered coating material and an oxygen/acetylene gas mixture metered into it, which is then ignited by a spark discharge (Fig. 4.18 and 4.19). The mixture is detonated and the powder is propelled from the barrel at high temperatures and velocities up to $760 \text{ m s}^{-1}$. The operating cycle is repeated four to eight times a second to provide a continuous deposit. Because of the noise created by the process it is carried out in a soundproof chamber and is fully automated.

A recent development is a gun that operates at approximately double the frequency of the original, and gives higher particle velocities, see Fig. 4.20. The coatings produced by this process have high density and bond strength, and they form the standard of excellence for thermal spray coatings. Applications include protection of gas turbine engine parts against various forms of wear at high temperature, textile machine parts, rollers for
steel, paper and plastics industries, bearings and other parts for the nuclear industry, and cutting edges.

**Advantages**

1. Very high density deposits.
2. High deposit adhesion.
3. Heavy grit blast surface preparation is not needed.
4. Low heat input to workpiece.
5. Fully mechanised and controlled process.

**High velocity oxyfuel processes**

This group of processes, often abbreviated to ‘HVOF’, is designed to give high levels of coating density and adhesion to the substrate. The density and adhesion of the coating depend on a number of factors, which include the temperature and velocity of the sprayed particles when they strike the workpiece. Higher values of these factors are achievable from the specially designed spray guns burning oxygen and a fuel, than those obtained from the conventional gas and arc spraying processes already described. In HVOF systems the combustion process is internal to the gun, and the gas flow rates are much higher than in conventional flame spraying. The combination of these two factors leads to supersonic flame speeds, up to approximately 2000 m s\(^{-1}\). The particle velocities may reach 800 m s\(^{-1}\).

**Fuel gas**

Fuel gases include hydrogen, propane and propylene, which all produce supersonic speeds. Hydrogen is capable of producing coatings that are very dense and have excellent adhesion, owing to a greater degree of melting of the particles in transit to the substrate. Several systems operate with acetylene fuel, but this does not produce such high velocities in the flame. While
this may appear to be a disadvantage, since the adhesion and porosity of the coating is lower than for hydrogen, for materials such as tungsten carbide/cobalt more of the hard phase, tungsten carbide, is retained and the coatings produced have better wear resistance. Figure 4.21 shows the change in porosity of an alumina coating, sprayed with hydrogen and acetylene.

Propylene and propane are popular fuels for HVOF, but require a well-designed delivery system in order to ensure the high flow rates required for the process. Relying on the vapour pressure of the fuel alone is not satisfactory, particularly in cold weather. Nitrogen is used to pressurise the head space above the liquid in the fuel cylinder, which is then forced into a dip tube and into the manifold. From here it is piped to the thermostatically controlled immersion heater. Application of trace heating and thermal insulation is essential to prevent condensation. The control system ensures that liquid is prevented from entering the system, and the pressure is controlled at the outlet of the vaporiser to be 790–900 kbar to ensure adequate pressure at the console. A liquid withdrawal installation for propylene is illustrated in Fig. 4.22.

A range of equipment is now available to use liquid fuels, such as kerosene. These systems are capable of much higher deposition rates than conventional gas-fuelled units.

**Gun design**

There are significant differences between the guns available. These include the powder feed position, the gas flow rates, the fuel gases that can be used, the internal geometry and the oxygen to fuel ratios available.

Powder is fed directly into the combustion chamber in the CDS, HV2000 and Top Gun systems (Fig. 4.23). It is fed into the exhaust barrel in the Jet Kote and Diamond Jet systems (Fig. 4.24a and b). Diamond Jet has been developed to a second generation, the Hybrid Diamond Jet, which incorporates a longer air cap with a converging/diverging geometry. This enables higher gas velocities to be achieved, 2700 m s⁻¹ compared with the original 1250 m s⁻¹. This latter design is immune to backfiring, and gives coatings with lower porosity, lower oxide content, lower loss of alloying elements and better adhesion.

The JP5000 system (Fig. 4.25) also has a converging/diverging exit nozzle, which allows a lower powder feed pressure. The powder is fed through a radial injection system into the barrel. Powder velocity and temperature are evenly distributed across the spray.

Figure 4.26 shows Hybrid Diamond Jet being used to spray a roll surface. Figure 4.27 shows Jet Kote equipment in use manually, while Fig. 4.28 shows the high quality of deposit attainable, in this case 83/17 tungsten carbide/cobalt on a titanium substrate.

Owing to the high temperatures generated within the guns, water cooling is generally used. In the case of Diamond Jet air cooling was used on the original, but a combination of air and water cooling is used on the Hybrid version.
4.2.1 Comparison of alumina coatings produced using different fuel gases: (a) hydrogen; (b) acetylene (courtesy TWI).
4.22 Liquid withdrawal propylene installation for HVOF (courtesy of BOC Gases).

4.23 Diagram of Top Gun HVOF system.
Substrate cooling

Although high velocity oxyfuel systems are designed to be ‘cold’ processes, the substrate can become hot owing to the close proximity of the flame, since the optimum ‘stand-off’ between the gun and the substrate is of the order 100–200mm only.
Keeping the substrate cool reduces the residual stresses in the coatings, which arise through a mismatch in thermal expansion coefficient between substrate and coating. This is especially important with ceramic coatings on metals, where lowering the substrate temperature during deposition decreases residual compressive stresses and generally also improves thermal shock and thermal fatigue resistance. It has also been demonstrated that a low deposition temperature can lead to a decrease in the stress gradient in the coating, thereby improving cohesive strength and avoiding cracking within the coating.

Significant benefits can be gained through using liquid carbon dioxide for
cooling, owing to its high latent heat of sublimation. Figure 4.29 shows an attachment for plasma or HVOF torch which enables carbon dioxide to be sprayed on the substrate simultaneously with the spray process. The cooling system has controls allowing a choice of cooling profile for the substrate.

**Principal applications**

The high quality of HVOF sprayed coatings has led to their rapid acceptance in a wide range of applications. These include

- tungsten carbide cobalt coatings for fretting wear resistance in aeroengine turbine components, and on gate valves;
- wear-resistant cobalt alloys on to fluid control valve seating areas;
- coatings for printing rolls, including copper, alumina and chromia;
- NiCrBSi coatings (unfused) for glass plungers;
- NiCr coatings for high temperature oxidation/corrosion resistance;
- alumina and alumina–titania dielectric coatings;
- biocompatible hydroxypatite coatings for prostheses.
Jet Kote Nova gun in use applying a coating to a pump liner (courtesy Deloro Stellite Group).

Deposit of 83/17 WC/Co on titanium (courtesy of Praxair Surface Technologies Ltd).

**Advantages**

1. Deposits of high density and adhesion to the workpiece.
2. Low heat input to the component.
3. Tight spray pattern allows accurate placement of the deposit.
4. The gun to workpiece distance is relatively insensitive.
5. Manual or mechanised capability.
Spray fused coatings

The two stage spray fuse process was developed to improve the strength of bond between coating and substrate which is obtained with a normal thermal spraying process, and at the same time take advantage of the typical smoothness and uniformity of the coating (which can be quite thin, e.g. less than 1 mm).

The coating is typically sprayed using an oxyacetylene torch, followed by a fusing operation that involves heating the coating and the underlying substrate to a temperature between the solidus and liquidus of the coating alloy, usually about 1000°C. This develops a metallurgical structure in the coating and a bond with the substrate, which in mild steel has a strength of 380-550 MPa, depending on the coating alloy. To be successful the coating alloy requires self-fluxing characteristics and these are possessed by a range of cobalt and nickel base alloys which contain a proportion of boron and silicon. Self-fluxing alloys are also applied by a process known as powder welding which was dealt with in Chapter 3.
The spray fuse process lends itself particularly well to coating of cylindrical parts such as pump shafts and sleeves, for which spraying can easily be mechanised to give a uniform smooth coating.

Fusing of the coating can be carried out in various ways. Manual fusing using an oxyacetylene torch fitted with a large multi-jet nozzle (Fig. 4.30) is widely practised. The large flame introduces heat gently into the component and this is important to avoid the coating heating too rapidly and expanding away from the substrate before it is fused. This would prevent attainment of a metallurgical bond when fusing temperature is reached.

A cylindrical component is rotated in, for example, a lathe chuck and the flame is directed at one end of the coating. At a temperature between the solidus and the liquidus a band on the surface of the coating develops a glossy, light-reflecting finish which indicates that the correct temperature has been reached. The torch is then traversed slowly across the coated area to produce this effect all over and the component is then allowed to cool, or is heat treated if required.

4.30 Manually fusing the thermally sprayed deposit of a Cr–Si–B alloy on the surface of a pump rotor (courtesy Deloro Stellite Group).
As the temperature in the component is relatively high, precautions similar to those used in weld surfacing must be taken to suit the substrate material, remembering that the HAZ is likely to be deeper than that associated with welding.

An alternative method of fusing involves the transfer of the sprayed component to a vacuum furnace. With the heating, fusing and cooling cycle carried out in a vacuum the oxides formed during spraying are reduced, and no further oxidation occurs. Sufficient time at fusion temperature is allowed for diffusion processes to take place within the coating and at the coating substrate interface, and this provides cohesion and adhesion strengths much superior to normal thermal spraying and close to those of welded deposits. With this process the cooling stage can often be adjusted to suit the desired metallurgical structure in the substrate.

Vacuum fusing offers advantages over manual fusing shared with the spray fuse process. Vacuum fusing

- is ideal for quantity production using batch fusing techniques;
- is easily controlled and repeatable, giving consistent results;
- provides uniform heating of the whole component, imposing less stress on the bond between the coating and the substrate as the temperature rises;
- avoids the problem of temperature control experienced when fusing coatings manually on parts whose cross-section suffers sudden local changes;
- enables components of thicker cross-section to be coated than is practicable with manual torch fusing.

Figure 4.31 shows taper plug valves coated with a NiCrSiB alloy with a hardness of about 60RC. The coin of 22mm diameter gives a guide to
the size of the plugs and the smooth, even, as-fused surface is clearly seen. On a component such as a ball valve, manual fusing would be difficult, and incur the risk of the sprayed coating lifting from the substrate while other areas are being fused. Uniform heating in a vacuum furnace, Fig. 4.32, avoids the thermal stresses that cause this problem and also minimises any tendency for the part itself to distort. These advantages of the spray and vacuum fuse process mean that less material, time and machining are required than if a weld surfacing process had been used.

The temperature needed to fuse the coating can also be developed by induction heating, provided that the rate of heating and the uniformity of temperature required for satisfactory results can be attained; a further possibility is to use a laser beam. This was described in Chapter 3.

Substrate materials

The thermal cycle

It has already been noted that temperatures required to fuse sprayed deposits of self-fluxing alloys are in the region of 1000°C, and with manual fusing the whole part may exceed 800°C. With furnace fusing the whole part will be at fusing temperature. Information given in Chapter 3 on welded coatings therefore applies to a large extent to fusing of a sprayed coating. Differences in treatment are discussed below.

After fusing has commenced, if a hardenable substrate steel is in use, no area of the fused component should be allowed to cool below the temperature at which martensite forms until the fusion operation has been completed and the part subjected to a suitable heat treatment – as for welded deposits.

With large areas of coating and deposits of hard alloys, or on components of large mass, it is helpful to introduce extra heat into the component during spraying once the whole area has received an initial thin coating of spray. On conclusion of spraying and with a substrate temperature of about 500–
700°C there is less thermal shock and differential expansion stress on the coating when raising it to fusing temperature, so the risk of the bond to the substrate failing is minimised. Cooling stresses are also lowered when fusing is carried out with less steep temperature gradients in the component. The first sprayed coating protects the substrate from oxidising as the temperature rises later in the spraying operation.

**Unsuitable substrate materials**

The component material must have a solidus temperature greater than the fusing temperature of the coating alloys. The presence of, for example, a high phosphorus content in a cast iron can result in melting of low melting point constituents at the grain boundaries and collapse of the part when fusing the coating.

Carburised and nitrided surfaces should be removed before surfacing and free-cutting materials should be avoided. Materials capable of forming tenacious oxide films, caused, for example, by aluminium content can resist the self-fluxing action of the coating alloys and inhibit the formation of a strong bond. This can result in detachment of the coating.

**Coating alloys**

The alloys used for the spray fuse process are those of a nickel base and cobalt base; see Table 1.1. Examples are to be found in Tables 4.4, 4.5 and 4.7. A wide range of hardnesses is available from about 20 to 60RC. Those of medium to high chromium content have resistance to oxidation and corrosion at elevated temperature and their abrasive wear resistance increases with hardness. Cobalt base alloys have good high temperature hardness, and resistance to adhesive wear and erosion.

All the alloys are supplied in powder form for spray fusing and are frequently mixed with tungsten carbide in proportions ranging up to about 75% to provide even greater resistance to abrasive wear in use.

**Manual fusing**

As indicated earlier, to ensure the creation of a metallurgical bond between the sprayed particles and between the coating and substrate, the fusing temperature for the coating alloy must be reached throughout its thickness. This is achieved when temperature gradients are as low as possible, a condition readily achieved by furnace fusing but requiring some care when torch fusing.

The most important point to observe is that heat is introduced gradually, as it has to be transmitted through the thickness of the coating into the substrate. Too great a rate of heat input can result in the coating expanding away from the substrate; a condition that requires the damaged coating to be removed and the job restarted.

Once a thin coating has been applied by spraying and the substrate is
Thermal spray processes

4.33 Temperature differences in manual fusing of a nickel base alloy coating.

Thereby protected against oxidation, it is good practice when torch fusing is intended to use extra heat as necessary to ensure that the whole of the sprayed area of the component rises in temperature as spraying proceeds. When this stage is reached and the component is at perhaps 600°C, extra heat can be applied with the fusing torch generally before it is concentrated on one end of the deposit and fusing is commenced. This technique ensures that temperature gradients are kept low and satisfactory coatings are properly bonded to the substrate.

Figure 4.33 illustrates a nickel base alloy coating which requires a temperature of 1000°C minimum at the interface. With too low a temperature in the substrate the coating may be fused but not bonded to the substrate. In fusing by induction heating the substrate actually exceeds the coating in temperature, even when the core temperature is low.

Design for thermally sprayed coatings

As with weld-deposited coatings there are many factors to be considered when drawing up a design for thermally sprayed coatings, which are discussed below.

Component shape

Surfaces to be coated must be accessible to the spray guns available for the chosen process, and the component must provide adequate routes for the dispersion of the hot spray gases and overspray. It must be possible to direct the spray at right angles to the surface, although sometimes this may be reduced to 45° at the expense of increased overspray and some loss of adhesion to the substrate.

Thus re-entrant areas of any type may produce a problem, which increases with the depth/width ratio (unless a gun designed to work inside the recess is available). This is especially so if there is no through passage, providing adequate ventilation for gas and overspray.

If the recess performs no practical function it may be easier to seal it off with a welded-in plug, which after grinding flush leaves a smooth clean external surface to be sprayed.
Surface profile details

Fabrications

Figure 4.34 shows joint configurations that should be used and draws attention to the problems of unsealed surfaces and discontinuous fabrication welds. Unsealed joints can harbour dirt and moisture which can give rise to local defects in the sprayed coating that may become sites for premature failure.

Precision components

If the whole surface is not to be covered, it is normal to arrange for the edges of sprayed deposits to finish flush with the parent material of the workpiece. This involves creation of a recess to accommodate the required finish thickness of the deposit and the provision of a finishing allowance on adjacent areas. Bearing in mind the low heat input of most thermal spraying processes and the low risk of distortion, this finishing allowance can normally be quite small, typically less than 0.5 mm per face. The important feature of the recess is that it should be provided with the edges chamfered full depth at an included angle with the base of the recess not more than 45°; 35° would be better (Fig. 4.35).

When deposits must reach the end of a shaft or bore, the corner should be provided with a radius or chamfer as shown in Fig. 4.36, but a heavy local build-up of deposit (in an attempt to machine to a square corner) must be avoided. This would encourage additional stress and a possible risk of the deposit lifting. Corners of deposits produced in the manner described are always weaker than a deposit contained in a recess and the latter is to be preferred, especially if the part is likely to be exposed to shock on the corners when handling or in use.

In reclamation work there is always the possibility that holes or keyways will be present in the surface to be coated. The usual way to deal with these is to plug them with graphite, making a tight fit in the recess and protruding slightly higher than the finished thickness of the deposit. The plug will be revealed on machining to finished size and being soft is easily removed. However, this leaves a sharp and fragile edge on the deposit which should be radius chamfered or tapered to remove the sharp corner and to minimise damage in use.

Deposit thickness

General considerations

Factors to be considered concerning the appropriate deposit thickness vary for new and for reclamation work. For new work, the best guide is to apply the minimum necessary to do the job, taking into account any opportunity for regrinding before recoating is necessary. For repair work thicker deposits may be necessary. Excessive wear can often be made good by spraying a first coating of a softer material and then finishing with the desired top coating.
AVOID

Angles back-to-back
(inside cannot be protected)

AVOID

Lap joints with discontinuous welds

AVOID

Sharp corners and discontinuous welds

PREFER

Continuous welds

PREFER

Lap joints fully sealed by continuous welds. Smooth and free from slag and weld spatter

PREFER

Rounded corners and continuous butt welds

Butt joint fully sealed by continuous weld. Smooth and free from slag and weld spatter

Use similar section T bars

4.34 Crevices, lap joints and corners.
This is because soft materials applied by low energy processes can be applied in thicker coatings having lower residual stress than hard materials, particularly those applied by high energy techniques. An indication of typical thickness design figures is given in Table 4.10.

In reclamation work, irregular wear patterns should be machined to avoid a varying thickness deposit, but local increases in thickness can be accommodated using a 30° or 45° chamfer at each change in depth.

Coating thickness for corrosion protection on iron and steel structures sprayed with zinc or aluminium ranges from about 100 to 400μm and 100 to 250μm respectively, the latter coatings being used for longer life structures.
### Table 4.10 Typical design figures for thermal spraying

<table>
<thead>
<tr>
<th>Gas/arc</th>
<th>Temperature/°C</th>
<th>Max. particle velocity/ m s⁻¹</th>
<th>Adhesion/ MPa</th>
<th>Oxide content/ %</th>
<th>Porosity/ %</th>
<th>Spray rate/ kg h⁻¹</th>
<th>Relative cost/ low = 1</th>
<th>Typical deposit thickness/ mm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Flame</td>
<td>2600–3100</td>
<td>40 to 8</td>
<td>10–15</td>
<td>10–15</td>
<td>2–6</td>
<td>1</td>
<td>0.1–15</td>
<td></td>
</tr>
<tr>
<td>Electric arc</td>
<td>6000</td>
<td>100 10–30</td>
<td>2–20</td>
<td>2–10</td>
<td>5–20</td>
<td>2</td>
<td>0.1 to &gt; 50</td>
<td></td>
</tr>
<tr>
<td>HVOF</td>
<td>2600–3100</td>
<td>600–800 &gt;70</td>
<td>0.1–5</td>
<td>0.1–2</td>
<td>2–5</td>
<td>3</td>
<td>0.1–2</td>
<td></td>
</tr>
<tr>
<td>D-gun</td>
<td>4000</td>
<td>800 &gt;70</td>
<td>1–5</td>
<td>1–2</td>
<td>0.5</td>
<td>N/A</td>
<td>0.05–0.3</td>
<td></td>
</tr>
<tr>
<td>APS</td>
<td>12 000–16 000</td>
<td>200–400 to &gt;70</td>
<td>1–3</td>
<td>1–8</td>
<td>3–8</td>
<td>4</td>
<td>0.1–1</td>
<td></td>
</tr>
<tr>
<td>VPS</td>
<td>12 000–16 000</td>
<td>400–600 &gt;70 ppm</td>
<td>&lt;0.5</td>
<td>0.5–2</td>
<td>5</td>
<td>0.1–1</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Special considerations for spray-fuse coatings**

To the finished thickness of the deposit must be added allowances to cover the possible distortion arising from the coating or subsequent heat treatment operation, and a small machining allowance if the part is to be precision machined. The size of these allowances depends on the design and size of the part, particularly matters such as length/diameter ratio, which influence distortion. On short, stubby shafts which have been accurately centred at the ends and can therefore be accurately set up for machining, a combined allowance of 0.1–0.5 mm may be sufficient if spraying is effected mechanically to ensure uniformity.

An extra allowance must be added to arrive at the size of the part in the as-sprayed condition before fusing. This is to compensate for two factors. The first is that the sprayed coating shrinks some 12–20% on fusing, depending on the alloy and spraying conditions. The second is that, when measured, the sprayed coating and the component will be hot and therefore will have expanded.

**Coating production**

### Surface preparation

The importance of correct surface preparation for all thermally sprayed coatings has already been stressed as one of the main factors in achieving a high and consistent strength of bond between coating and substrate. The first requirement is removal of any grease or dirt, if possible by a degreasing operation. The second is roughening of the surface to promote an effective key with the sprayed particles. This can be achieved most readily by abrasive blasting, although for metallising processes rough machining (e.g. by thread cutting) has been used effectively.

It is important that a blast medium is kept clean and free from fines. For metallising an angular chilled iron grit is used on most substrates, and this must be replaced when it becomes blunt. Grit sizes range from 14 to 100, see Table 4.11.
Table 4.1 Grit sizes used for surface preparation

<table>
<thead>
<tr>
<th>Grit size</th>
<th>BS2451 equivalent</th>
<th>Nominal sieve opening/mm</th>
<th>Size distribution/mm</th>
</tr>
</thead>
<tbody>
<tr>
<td>14</td>
<td>G55</td>
<td>1.41</td>
<td>2.0–0.84</td>
</tr>
<tr>
<td>18</td>
<td>G39</td>
<td>1.0</td>
<td>1.68–0.71</td>
</tr>
<tr>
<td>25</td>
<td></td>
<td>0.71</td>
<td>1.0–0.42</td>
</tr>
<tr>
<td>40</td>
<td>G17</td>
<td>0.42</td>
<td>0.59–0.25</td>
</tr>
<tr>
<td>60</td>
<td></td>
<td>0.25</td>
<td>0.35–0.15</td>
</tr>
<tr>
<td>100</td>
<td></td>
<td>0.15</td>
<td>0.18–0.06</td>
</tr>
</tbody>
</table>

Smaller size particles permit blasting of a larger area per hour. Larger particles provide more rapid removal of material and give a rougher surface finish. Grit sizes 16 and 60 are used for metal substrates and 60–100 for most plastics. For thin coatings, particularly when used on thin substrates, fine grit should be specified. Coarser grit (14–25) providing rougher finishes is used for thick coatings (greater than 0.25mm) and optimum coating adhesion.

On cylindrical components the operation to machine a recess for the deposit is sometimes followed by cutting a V thread in the bottom of the recess at, say, 0.5mm pitch, as this gives a good key to the sprayed deposit. This may be followed by abrasive blasting. All traces of cutting fluid must be removed. For coatings applied by high energy processes, aluminous or silicon carbide grits in the 25–100 range are used.

The action of abrasive blasting increases the surface area of the workpiece significantly and in its clean and abraded state it is very active and oxidises readily. It should therefore be sprayed as soon as possible after the blasting operation and never handled directly by the fingers, which can transfer dirt and grease to the surface. Any surface contamination can impair adhesion of the coating. The abraded area should always stretch beyond the area to be coated to prevent risk of the overspray lifting and the bond failure propagating under the coated area, even for a short distance.

On high quality work, such as aircraft turbine engine parts, automatic grit blasting machines have been developed, which control air pressure, blast angles, nozzle stand-off and time of blasting. Further quality assurance standards are used to control grit size range, nozzle wear and grit damage with a particular substrate.

Abrasive blasting can lead to distortion of thin components and a reduction in grit size or impact velocity may not always overcome this. Some materials to be coated may be too hard to roughen adequately by grit blasting and to overcome the risk that a coating may not adhere properly under these circumstances, a preliminary layer of a bond coat is applied.

**Bond coats**

Some sprayed materials adhere strongly to clean smooth surfaces. They are adherent over a wide range of conditions and a thin layer can serve as an
'anchor' for materials sprayed on top. They are used in both the circumstances mentioned in the paragraph above. Coating thickness for a bond coat is normally 30–80 μm. Note that bond coats are not applicable to spray-fused coatings.

Commercially available bonding materials include nickel–aluminium and molybdenum. Nickel–aluminium is used for a large number of applications because of its good high temperature properties and low dependence on operator technique. It is used on a wide range of substrates including all common steels, stainless steel, hardened alloy steel, nitriding steel, cast steel (although nitrided and carburised surfaces are best removed before spraying), nickel, chrome nickel alloys, cast steels, magnesium, most Al steels, Monel, nickel and cast iron. It does not bond properly to copper, brass, bronze, chrome plate, nitrided surfaces or electroless nickel.

**Preheating**

Preheating the prepared surface to prevent condensation of water vapour is normal good shop practice. A temperature of 90 °C is adequate, but further heating to 175–260 °C may improve the bond. This operation should follow as quickly as possible after shot blasting to avoid risk of the surface becoming contaminated with rust or dirt.

On cylindrical parts preheating should be carried out while held in a rotating chuck so that heat intake is as uniform as possible.

**Masking**

With thermal spraying processes the use of a mask or shield prevents deposition on undesired surfaces. The mask should be positioned above the surface to be sprayed, since if placed directly on the surface, the coating will link the job and the mask. Subsequent removal is then likely to damage the coating. Masks are conveniently made from sheet metal, cut or bent to the required shape and supported rigidly so that the force of the spray does not dislodge them.

**Spraying**

Spraying should be as near normal to the surface as possible and never at an angle of less than 45°, so adequate access is essential. Special spray guns are available that enable bores of 150 mm diameter to be sprayed to a depth of 500–600 mm from an open end. The spraying operation should be mechanised where possible to ensure a uniform coating thickness and economy on time and material.

As already discussed, areas that are to be kept free from deposit can be masked with suitable tape to prevent damage during blasting and a metallic shield can be used as a shadow mask to limit the edges of the deposit. This must stand clear of the surface so that its removal does no damage to the deposit.

Thermal spraying processes generate overspray which can constitute a health and safety risk. This subject is dealt with in detail in Chapter 10.

Emphasis has been placed on the importance of correct design and surface
A programmable six-axis robot teamed with a five-axis positioner can accurately reproduce preset spray patterns. A memory holds up to 10 pre-programmed spray patterns (courtesy Sulzer Metco Ltd).

preparation of the component, if satisfactory results are to be obtained. Equally important to the result is control of spraying conditions. The ability of a correctly designed and selected deposit to give satisfactory service depends on such factors as coating density, oxide content and bond strength to the substrate. These are properties that are not easily verified by non-destructive testing, so when developing new applications it is usual to carry out destructive tests on prototype or simulated components. When necessary, adjustments can then be made to surfacing parameters until acceptable results are obtained.

The important next step is to ensure that conditions pertaining to the acceptable testpieces are accurately repeated on production work, and this means use of properly calibrated devices to measure each parameter. A mechanised process, incorporating microprocessor control of parameters and, when appropriate, robotisation (Fig. 4.37), will give improved
4.38 (a) Deposit of flame sprayed 13% chromium steel. (b) Arc sprayed deposit of the same steel using correct conditions. (c) As for (b), but poor interface because of excessively fine atomisation and use of inadequate extraction (courtesy Metallisation Ltd).

repeatability, compared with manual work, and will always be the preferred choice for critical work.

The photomicrographs in Fig. 4.38 illustrate structures of deposits of 13% chromium steel sprayed by flame and electric arc processes. The laminar structure characteristic of thermally sprayed deposits is clearly seen, with the arc deposit having lower porosity. By contrast, Fig. 4.38(c) shows a deposit applied with unsuitable conditions; this would not necessarily be obvious on visual examination of the surface.

Sealing

It is advisable to seal all deposits used in machined parts except those used in lubricated conditions under light load. This prevents ingress of fluids which may accelerate corrosion of the coating or penetrate to the substrate, causing corrosion at the interface and causing the coating to lift. Surface porosity may be impregnated by materials being processed which could contaminate subsequent batches.

One of the most important reasons for sealing the coating on shafts to be used for journals, pump plungers, etc., is to ensure a better, cleaner ground finish. Grit from the grinding operation is prevented from entering the pores in the coating.

Sealers are low viscosity fluids containing an inert resin binder in a solvent. This impregnates the pores in the coating, filling them as the solvent evaporates. When used on coatings such as zinc, use of a sealer does not
impair the cathodic protection of the coating. Silicone resins are available for service up to 550°C.

**Conclusions**

Thermal spray processes provide a wide range of possibilities for application of coatings that are capable of giving high resistance to wear, corrosion, oxidation and heat as well as those that can be used for thermal or electrical insulation.

Production of reliable coatings depends on:

- selection of the correct coating material;
- use of the correct coating process;
- proper surface preparation;
- adherence to correct operating parameters;
- process automation;
- precise quality control of powder consumables.
Electrodeposition is a well-established process for applying metallic coatings to improve surface properties of materials used in engineering practice. Although the principles are similar to those involved in the application of relatively thin coatings for decorative and corrosion protection purposes, engineering electrodeposition differs in several important respects and constitutes a specialised process in its own right. For the sake of completeness this chapter has been extended to describe the application of certain metal-based coatings by chemical reduction, by anodic oxidation and by chemical conversion. These processes operate in aqueous environments and are often used in conjunction with electroplating processes.

In theory, there is no limit to the thickness to which many metals and alloys can be electrodeposited, but the thickness needed to perform the required function is usually the basic criterion. Process economics are always important. Therefore it should be borne in mind that electrodeposition can be slow and costly, so where thick coatings are desired some other surfacing methods may be more cost effective. For thinner engineering coatings, however, electrodeposition is essential to the successful operation of innumerable components and it offers considerable scope and flexibility to the designer.

**Basic principles**

Electrodeposition or electroplating involves making the component to be coated the negative electrode or cathode in a cell containing a liquid or electrolyte which must allow the passage of electric current (Fig. 5.1). This electrolyte is usually a solution in water of a salt of the metal to be deposited, and is maintained at a controlled temperature which can be up to about 60°C. The electrical circuit is completed by a positive electrode or anode which is generally made out of the metal to be deposited and is located a short distance away from the cathode. Under the action of a direct current applied at a low voltage, positively charged metal ions in the electrolyte move towards the cathode, where they undergo conversion to metal atoms and deposit on the cathode, i.e. the component surface.

The structure and properties of the deposited metal depend on the chemical composition of the electrolyte as well as its temperature and degree of acidity or alkalinity, i.e. its pH value. These factors, and in
5.1 Plating bath.

particular the density of the electric current per unit area of the cathode surface, determine the rate of deposition. Current flow to projections on the cathode surface is greater than to recesses, and the consequent variation in current density influences metal distribution, since the thickness deposited is proportional to the current density. Uniformity of thickness, which is a function of the throwing power of the electrolyte, can be improved considerably by attention to the design of the component and to the conditions of deposition. As electrodeposited coatings are seldom of constant thickness over the entire surface it is usual to define that portion of the surface that is essential to the serviceability of the component as the ‘significant surface’ and to quote the minimum rather than the average thickness over this surface.

Electrodeposition is used extensively not only to apply coatings to new components to confer the required surface properties, but also to restore the dimensions of parts that have either worn excessively in service or been so over-machined as to be outside required tolerances. Another application is production of free-standing bodies by deposition on to shaped mandrels which are capable of subsequent removal. This process, known as electroforming, enables the shape and surface finish of the mandrel to be faithfully reproduced and in this way constitutes a simple method of fabricating parts of intricate shape in a single operation.

Two techniques are available for electrodeposition of coatings for engineering purposes, following the principles outlined earlier: vat and selective plating. The characteristics of both processes and the complementary processes of electroless or autocatalytic deposition and hard anodising are described below.
Vat plating

Principles
Virtually all electrodeposition for engineering applications is undertaken in tanks or vats which may have capacities up to several thousand litres. For some specialised applications the tank may be built round the workpiece, or a large cylindrical component requiring electroplating internally may function as its own tank. The electrolyte is usually mildly agitated either by air jets or by mechanical movement of the work, and is maintained at the working temperature by electric immersion heaters or steam coils; alternatively water cooling may be necessary. The anodes, suspended some centimetres from the workpiece, are sometimes inert in that they carry the current but do not dissolve in the electrolytic process, so the coating material is derived wholly from the solution. The workpiece itself is mounted on a rack or jig and suspended in the electrolyte. A transformer-rectifier set normally supplies the plating current at a voltage in the range 4–8 V. The current applied is read on an ammeter and the time to deposit the required thickness is estimated from this current and the known efficiency of deposition.

Deposition efficiency is taken as the ratio of the weight of metal actually deposited against the weight that should have been deposited by the electrical energy used. Some of the current passed is usually wasted in the unavoidable evolution of hydrogen at the cathode as described later in this chapter, and in resistive heating. Deposition efficiency varies with the electrolyte system in use, the plating conditions employed and the current density.

Non-metallic particles may be incorporated into the metal deposit, e.g. silicon carbide in nickel, by maintaining the particles in suspension in the electrolyte. The process is controlled by measuring the density and acidity of the electrolyte and, in the longer term, by chemical analysis.

A variation in vat plating enables small components to be plated in bulk. For example, components may be held in a perforated cylindrical barrel constructed in a polymeric material which is immersed in the plating solution and rotated continuously. The work in the barrel forms the negative electrode, and in tumbling over each other the components maintain electrical contact and simultaneously present fresh surfaces to the action of the electrolyte. Barrel plating is restricted to components weighing less than about 500 g which are of simple shape and capable of tumbling without locking together. Coatings so applied are thin and the main applications are for decoration, but engineering applications include deposition of gold and platinum group metals for corrosion and wear resistance.

Characteristics
1. As operating temperatures never exceed 100 °C the work should not undergo distortion or undesirable metallurgical changes.
2. Plating conditions may be adjusted to modify hardness, internal stress and metallurgical characteristics of the deposit.
Coatings are dense and adherent to the substrate. Bonding, which is molecular in nature, may be as strong as 1000 N mm⁻².

The thickness of the deposit is proportional to the current density and the length of the time of deposition.

As the current density over the workpiece surface is seldom uniform, coatings tend to be thicker at edges and corners and thinner in recesses and at the centre of large flat areas.

The rate of deposition seldom exceeds 75 μm h⁻¹, but it can be accelerated by forced circulation of the electrolyte.

There is no technical limit to the thickness of the deposits. Metals such as nickel may be 13 mm or more in electroforming and reclamation work, but most surfacing applications require much thinner coatings.

Application of coatings is not confined to the line of sight. Although the throwing power (i.e., the ability to plate round corners) may be limited, there is comparative freedom for the location of anodes, for example in coating the bores of narrow tubing.

Areas not requiring deposition may be masked.

The size of the vat limits the dimensions of the work.

The process is suitable for automation.

**Brush or selective plating**

Electrochemical reactions in brush or selective plating are the same as in vat plating and the essential features are shown in Fig. 5.2. The main distinction is that coatings can be applied manually to components such as moulds, dies, shafts and bearings in selected areas.

**Principles**

This process complements vat plating in that electrodeposits can be produced on localised areas with minimum masking or stopping off and without immersion of components to be coated. Principles are the same as for vat plating, but the anode is mounted in an insulated handle and enveloped by an absorbent pad which is soaked in the electrolyte. The
work is connected to the negative side of a DC source and upon bringing the pad into contact with the work the circuit is completed and electrodeposition takes place. Generally the operator traverses the pad manually over the areas to be coated, but this procedure requires much skill, as the rate of movement is critical in ensuring a sound and even deposit (see later).

The anode is a high grade carbon rod, so does not dissolve. Long fibre cotton wool may form the absorbent pad, but a resin-bonded plastics felt is preferable. The electrolyte may be renewed on the pad by dipping into a beaker, by drip feed or by pumping. Cleaning of the work before plating involves reversing the polarity to make the work anodic, in the presence of special solutions.

**Characteristics**

The first five characteristics of vat plating apply to selective or brush plating. Additional characteristics include:

1. The equipment is portable and can be taken to the work, so that moulds and dies for example can be plated in situ with minimum loss of production time.
2. Masking is generally unnecessary as only the area to be coated needs to be in contact with the electrolyte.
3. The electrolyte must be more robust than in vat plating to withstand the inevitable sharp changes in operating temperature and current density.
4. The power source which operates in the range 8–30 V incorporates cut-outs to minimise damage by short circuits and also a current time meter to monitor the progress of plating.
5. Deposition rates generally exceed those of vat plating and may reach 200–400 μm h⁻¹.
6. The electrolytes are costly, but small volumes only are required.
7. Manual operation is labour intensive and demands considerable skill.
8. The process can be mechanised or automated, for example in plating cylindrical bores.
9. Brush plating is not suited to high volume production.

**Electroless or autocatalytic deposition**

**Principles**

Electrodeposition involves the reduction of metal ions, arriving from the electrolyte, to produce metal atoms which are deposited at the cathode or workpiece surface. The current required for this reduction need not be supplied externally. By providing a reducing agent in the solution, the electrons for the deposition reaction can be furnished from within the solution. Metals such as nickel, copper, cobalt, gold, silver and palladium can be deposited from aqueous solutions of their salts by chemical reduction, and the initial layer catalyses the subsequent deposition. Hard nickel alloys deposited from solutions containing either phosphorus
or boron compounds as reducing agents are widely used in engineering. The process is generally operated in polypropylene or PTFE-coated stainless steel tanks containing the solution maintained at about 90°C, and fitted with facilities for accurate temperature control, agitation and solution filtration. The work must be cleaned just as efficiently as in electrodeposition.

Electrochemical displacement can also occur when one metal is immersed in a salt solution of another, less reactive, metal. Copper, for example, can be plated on to iron by displacement. The coatings are very thin and often so poorly adherent as to be generally of limited industrial use. These displacement reactions are not true electroless deposition reactions.

**Characteristics of electroless deposition**

1. The equipment is simple and economical as neither anodes nor DC electrical sources are required.
2. Deposits are uniform in thickness provided that the solution can be circulated over all the surfaces to be coated (Fig. 5.3).
3. The rate of deposition, which is dependent on temperature, is approximately 20µm/h.
4. Thicknesses of up to 125–200µm can be applied.
5. Areas not requiring deposition may be masked.
6. The size of the tank limits the dimension of the work.
7. Although the chemical materials are expensive, costs may be competi-
The equipment is more involved than that in electrodeposition as a refrigeration unit is often required to maintain the required temperature.

2 Hard anodising is a conversion process, i.e. the coating grows from the metal and is integral with it.

3 Throwing power, i.e. uniformity of coating thickness, is excellent.

4 Hard anodic coatings can be applied to aluminium and its alloys by selective or brush techniques.

5 Inserts of foreign metals, e.g. steel, must be masked to avoid dissolution.

6 The process can be automated.

**Phosphating**

Phosphating, like anodising, involves the chemical conversion of the substrate surface. The conversion process involves complex chemical reactions in the presence of phosphoric acid and other metals, so that, for example, a layer of zinc or manganese phosphate may be formed on steel.

Metals that can be treated in this way include ferrous metals, aluminium and zinc. The coating may be used as an intermediate layer, or as an engineering layer in its own right. A fully formed phosphate coating is tightly bonded to the substrate surface, will cover the surface completely, is mineral in nature and may be electrically insulating.
Table 5.1 Electrode potentials of engineering metals

<table>
<thead>
<tr>
<th>Metal</th>
<th>Standard electrode potential at 25°C/V (versus a hydrogen electrode)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aluminium</td>
<td>-1.66</td>
</tr>
<tr>
<td>Titanium</td>
<td>-1.63</td>
</tr>
<tr>
<td>Manganese</td>
<td>-1.18</td>
</tr>
<tr>
<td>Chromium</td>
<td>-0.74</td>
</tr>
<tr>
<td>Iron</td>
<td>-0.44</td>
</tr>
<tr>
<td>Cobalt</td>
<td>-0.28</td>
</tr>
<tr>
<td>Nickel</td>
<td>-0.25</td>
</tr>
<tr>
<td>Tin</td>
<td>-0.14</td>
</tr>
<tr>
<td>Lead</td>
<td>-0.13</td>
</tr>
<tr>
<td>Hydrogen</td>
<td>(0)</td>
</tr>
<tr>
<td>Copper</td>
<td>+0.34</td>
</tr>
<tr>
<td>Silver</td>
<td>+0.80</td>
</tr>
<tr>
<td>Gold</td>
<td>+1.68</td>
</tr>
</tbody>
</table>

Coating materials

Electrochemical principles govern the mechanism of electrodeposition. Standard electrode potentials, recorded in Table 5.1, represent the electrical potential established between the metal and a solution of one of its salts under standard conditions, and they indicate the relative ease of deposition of individual metals from aqueous solutions. In practice, the actual potential of the electrodes depends on the ambient conditions, in particular the solution concentration. Binary and ternary alloys may be electrodeposited provided that the plating conditions are chosen to equalise the deposition potentials of the constituent metals. Metals with higher negative potential than -1.2V (Table 5.1) generally cannot be deposited from aqueous solutions, thus aluminium and titanium can only be plated either from solutions in organic solvents or from molten salt electrolytes, both of which necessitate specialised equipment and techniques. Nevertheless these processes are becoming available commercially.

Table 5.2 summarises the mechanical and physical properties of the metals and alloys more usually electrodeposited for engineering purposes.

Chromium

Although its hardness when electrodeposited can be matched by other surface treatments, chromium possesses a unique combination of properties of value in engineering practice. The aggressive nature of the electrolytes makes the deposition of chromium alloys difficult, but claims for a 1% molybdenum alloy include improved resistance to both mechanical wear and corrosion in acid environments.

UK specifications for engineering chromium plating include BS4641, DEF STAN 03-14/1 and DEF GUIDE DG13.
Table 5.2 Mechanical and physical properties of some electroplated metals deposited for engineering purposes

<table>
<thead>
<tr>
<th>Coating</th>
<th>Usual finished thickness/mm</th>
<th>Melting point/°C</th>
<th>Density/kg m$^3$</th>
<th>Hardness/HV</th>
<th>Tensile strength/ N/mm$^2$</th>
<th>Elongation/%</th>
<th>Resistivity at 20°C/μΩ cm</th>
<th>Thermal conductivity/ W m K$^{-1}$</th>
<th>Linear coefficient of expansion/ x 10$^{-6}$/°C$^{-1}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chromium</td>
<td>Up to 0.13 sometimes to 0.50</td>
<td>1878</td>
<td>6930</td>
<td>800–1000</td>
<td>100–550</td>
<td>&lt;0.1</td>
<td>14–67</td>
<td>67</td>
<td>7.4</td>
</tr>
<tr>
<td>Electrodeposited nickel</td>
<td>Up to 0.50 and even 13+</td>
<td>1455</td>
<td>8907</td>
<td>150–450</td>
<td>340–1050</td>
<td>3–30</td>
<td>7.4–11.5</td>
<td>94</td>
<td>13.5</td>
</tr>
<tr>
<td>Electroless nickel – 9% phosphorus</td>
<td>0.013 to 0.05 (approx.)</td>
<td>890</td>
<td>8000</td>
<td>480 (1030*)</td>
<td>758 (69*)</td>
<td>1–1.5 (0.2*)</td>
<td>60–80 (20*)</td>
<td>†</td>
<td>13.0–14.5</td>
</tr>
<tr>
<td>Copper</td>
<td>Up to 0.50 and even 13+</td>
<td>1084</td>
<td>8933</td>
<td>60–150</td>
<td>180–650</td>
<td>Up to 40+</td>
<td>1.7–4.6</td>
<td>403</td>
<td>16.5</td>
</tr>
<tr>
<td>65/35 tin-nickel</td>
<td>0.01–0.02</td>
<td>800+</td>
<td>8400</td>
<td>650–700</td>
<td>†</td>
<td>†</td>
<td>†</td>
<td>†</td>
<td>†</td>
</tr>
<tr>
<td>Silver</td>
<td>Up to 1.3</td>
<td>961</td>
<td>10,500</td>
<td>60–120</td>
<td>240–340</td>
<td>12–19</td>
<td>1.6–1.9</td>
<td>427 (approx.)</td>
<td>21.0 (approx.)</td>
</tr>
</tbody>
</table>

* After heat treatment, otherwise data refer to as-deposited metals.
† Data either not available, or not known accurately.
Engineering coatings

Characteristics

1. High hardness, i.e. 800–1000 HV, conferring resistance to abrasion.
2. Low frictional coefficient and resistance to sticking, thus combating adhesive wear.
3. Resistant to corrosion, also to oxidation up to 800°C.
4. Retains room temperature strength up to about 300°C.
5. Deposits thicker than about 50 μm require finishing by grinding.
6. Thickness generally limited to about 0.5 mm, but thicker deposits for reclamation work are usually built up on an undercoat of nickel.
7. Brittle, not resistant to shock loading.
8. Tensile stresses are sufficiently high to induce cracking and so coating thickness should not be less than 50 μm for corrosion protection.
9. The crack pattern may be developed to produce an open porous structure for lubricant retention.
10. Deposition efficiency of chromium is low and so the process is energy intensive.
11. Hydrogen discharged simultaneously may dissolve into the workpiece.

Applications

1. Plastics moulds.
2. Metal forming and drawing dies.
3. Cutting tools.
5. Cylinder liners.
6. Piston rings.
7. Crankshafts.
8. Hydraulic rams.
9. Reclamation of worn parts.

Nickel

Nickel is used extensively in engineering to combat mechanical wear, corrosion, fretting and heat scaling. The range of plating solutions is wider than for chromium, and nickel is more economical to deposit. BS4758 and DTD 905A are relevant UK specifications. Brass is often electroplated as an undercoat to nickel to improve adhesion and corrosion protection.

The ease of application and the properties of the deposits enable nickel to be used extensively in electroforming (Fig. 5.4), for instance in making tools such as moulds and dies (which are themselves the source of other engineering products) or foil and mesh products.

Characteristics

1. Moderately hard, i.e. 150–450 HV.
2. Softer and more ductile than chromium and so more resistant to fretting corrosion.
Electrodeposited coatings

5.4 Miniature nickel bellows made by electroforming.

3 Resistant to corrosion, in that 125 μm thickness ensures protection in most chemical environments.
4 Resistant to corrosion fatigue.
5 Resists scaling up to 600°C.
6 No technical limit to deposit thickness.
7 Deposits can be finished by turning.
8 Internal stresses in deposits generally controllable.
9 Tends to gall when running against some metals, even when lubricated.

Applications

1 Hydraulic equipment, especially in marine service.
2 Processing of sensitive products such as food and textiles to avoid contamination, e.g. from rusting or staining.
3 Reclamation of worn parts.
4 Electroforming.

Electroless nickel

The deposits produced are essentially nickel alloys containing either 6–12% phosphorus or about 2–7% boron, depending on the reducing agent used. It is possible to produce ternary alloys such as nickel–phosphorus–copper and to make composite coatings by suspending non-metallic particles in the deposition solution DEF STAN 03-5/1 covers the requirements of nickel–phosphorus deposition (Fig. 5.5).
Hydraulic components electrolessly plated with a nickel-phosphorus alloy (courtesy W Canning Materials Ltd).

**Nickel-phosphorus (Ni-P): Characteristics**

1. Uniform thickness.
2. Components can be plated to size without any finishing operation.
3. Hardness 400–500HV as deposited can be raised to 1000HV by heat treatment at 400°C. (Properties are then generally comparable with hard chromium.)
4. Resistance to both corrosion and wear is superior to electrodeposited nickel.
5. Low frictional coefficient.
6. Retains room temperature strength up to 350°C and regains some strength on cooling from higher temperatures.
7. Difficult to plate high alloy steels, which may require an initial coating of electrodeposited nickel.
8. Materials more expensive than for electrodeposition.
9. Pre-treatment and solution control more critical than in electrodeposition.

**Nickel-phosphorus (Ni-P): Applications**

1. Components for water pumps.
2. Hydraulic and pneumatic valves.
3. Polymer moulds.
4. Reclamation of worn parts.
5. Equipment for handling food – screw conveyors.
6. Reaction vessels, heat exchangers.
7. Turbine blades, pitot tubes.

**Nickel-boron (Ni-B): Characteristics**

1. Generally similar to Ni–P, but harder, i.e. 500–750HV.
2. Can be heat treated, if required, to increase hardness.
3 Less ductile and higher internal stresses than Ni-P deposits.
4 Resists wear better than Ni-P deposits.
5 Operating costs greater than for Ni-P deposition.

**Nickel-boron (Ni-B): Applications**

As for Ni-P deposits but Ni-B preferred when hot hardness is important, e.g. in coating dies for glass moulding.

**Copper**

Copper electrodeposits, which are generally softer than nickel, can be applied easily from several well-established electrolytes. The metal may also be deposited electrolessly and so is used extensively in the electronics industry. Copper can be electroformed to produce components such as spark erosion tools and wave guides for radar technology.

Other applications include provision of lubrication during drawing processes, 'stop-off' layers for selective case hardening, and printing.

**Characteristics**

1 Soft and ductile, hardness 60–150HV.
2 Good conductor of electricity.
3 Resistant to fretting corrosion.

**Applications**

1 Stop-off for selective case hardening of steels.
2 Printing rollers.
3 Surface lubricant in metal working.
4 Electroforming.
5 Reclamation of worn parts.

**Copper–tin alloys**

Alloying of copper with 10–15% tin produces compositions that are harder than either constituent metal.

**Characteristics**

1 Hardness 300–400HV.
2 Relatively ductile.
3 Resistant to corrosion and wear.
4 Resistant to shock loading.

**Applications**

1 Anti-sparking coating for underground pit props.
2 Undercoat for hard chromium.
3 Stop-off for selective nitriding of steels.
Iron

Electrodeposited iron has been used from the earliest days for the reclamation of worn parts, etc., but its use appears to be considered seriously only when nickel is in short supply. Electroforming may be applied to make iron foil and also hollow roller bearing elements in iron–nickel alloys.

Characteristics

1. Inexpensive and strong.
3. Not resistant to corrosion.
4. Not easy to deposit because of generally high temperature required and consequent oxidation of electrolyte constituents.

Applications

1. Soldering iron tips.
2. Anti-scuff coatings in internal combustion engines.
3. Reclamation of worn parts.
4. Electroforming.

Cobalt

Unalloyed cobalt is used in computer technology because of its magnetic properties. However, alloys with up to 20% of either molybdenum or tungsten have important engineering applications, notably in improving the wear resistance of metal forging and cold pressing tools (Fig. 5.6). As these components are generally massive, and downtime must be minimal, the brush method can be used to plate in situ.

Characteristics

1. Hardness about 600HV.
2. Low frictional coefficient.
3. Resistant to corrosion.
4. Deposits as thin as 12μm are resistant to erosion and metal pick-up.

Applications

1. Hot forging dies.
2. Cold pressing tools.

Lead

The importance of lead in engineering applications lies in the range of alloys which, when applied as overlay coatings, improve the performance of plain bearings. The alloys may contain 10% tin, 8% indium, or 10% tin plus 2% copper.
Electrodeposited coatings

5.6 Electrodeposition of cobalt–molybdenum alloy on to hot forging die by brush plating (courtesy Aston University).

**Characteristics**

1. Soft and ductile, hardness 8–15 HV.
2. Low frictional coefficient, resistant to scuffing.
3. Resistant to corrosion by degraded lubricant oils.
4. Thickness about 25 μm.

**Applications**

Overlay coatings for copper and aluminium-based plain bearings, to resist the severe conditions associated with starting and stopping machine movements, and to entrap small pieces of dirt, so avoiding damage to the bearing or shaft.

**Tin**

Like lead, electrodeposited tin is soft and its engineering applications are generally concerned with adhesive wear problems. BS1872 and DEF STAN 03-8/1 cover the deposition of tin.

**Characteristics**

1. Soft and ductile, hardness about 12 HV.
2. Low melting point.
3. Low frictional coefficient, resistant to scuffing.
4 Can be soldered.
5 Resistant to corrosion.

Applications
1 Stop-off in the nitriding of steel.
2 Corrosion protection in absence of mechanical wear.
3 Flash coatings to facilitate running-in of moving parts.

63/35 tin–nickel alloy (covered by BS3597): Characteristics
1 High hardness, i.e. 700 HV.
2 Low frictional coefficient.
3 Resistant to corrosion.
4 Retains oil films.
5 Brittle, limited resistance to impact.
6 Upper limit about 360 °C.
7 Solderable.

63/35 tin–nickel alloy: Applications
Automotive braking systems.

Tin-based diffusion alloys
Some tin-rich alloys are electrodeposited on to steel and then heat treated to form diffusion alloys containing iron–tin and other hard intermetallic compounds, interspersed in a softer supporting matrix. Electrodeposited bronze diffusion coatings may also be applied to steel and cast iron.

Characteristics
1 Hardness 600–950 HV.
2 Low frictional coefficient.
3 Resistant to corrosion.
4 Resistant to wear when lubricating conditions poor.
5 Minimal reduction in fatigue strength.

Applications
1 Pump plungers and bearings.
2 Textile thread guides.
3 Punches and dies for stamping and cutting stainless steel.

Silver
Electrodeposited silver has mechanical, electrical and chemical properties that are useful in many industries. Alloys with lead, palladium or copper
Electrodeposited coatings extend the upper temperature limit. BS2816, DEF STAN 03-9/1 and DTD 939 cover the requirements of silver deposition.

**Characteristics**

1. Hardness 60–120HV.
2. Thickness ranges from 0.001 to 1.0mm.
3. Resistant to fretting corrosion.
4. Good frictional properties.
5. Good conductor of electricity.

**Applications**

1. Electrical contacts.
2. Special bearings, e.g. in aircraft fuel systems.
3. Combating high temperature seizure.

**Gold**

Because of its unique combination of properties, electrodeposited gold finds extensive use in electronics, control and communications equipment. The pure metal is relatively soft, i.e. about 70HV, and is free of surface oxide films so that galling may occur between mating surfaces, e.g. in sliding contacts. However, the incorporation of a small proportion of base metals, such as nickel and cobalt, as alloying constituents increases hardness to about 450HV. This factor, combined with the co-deposition of an organic polymer generated in the plating solution, leads to an improvement in wear resistance and so extends the range of applications of alloy gold coatings for electrical contact purposes.

**Characteristics**

1. Hardness of pure gold, i.e. about 70HV, can be increased by alloying with base metals, such as nickel and cobalt.
2. Low electrical contact resistance.
3. Withstands corrosion and oxidation over a wide temperature range.
4. Good solderability, which is retained even after long storage.

**Applications**

1. Edge connectors in printed circuit boards.
2. Transistor header components.

**Platinum group metals**

Electrodeposited coatings of palladium, platinum, rhodium and ruthenium are of practical engineering interest because of their hardness and resistance.
to corrosion. Thus thin coatings are applied to protect components against wear and corrosion in electrical and electronics equipment. The main characteristics and applications are summarised briefly below.

**Palladium**

Palladium may be deposited at relatively low cost and its hardness of about 300 HV commends it for use in heavy duty electrical contacts. Palladium coatings are also used to protect refractory metals, such as molybdenum, against oxidation.

**Platinum**

Platinum is especially resistant to corrosion. Very thin coatings applied to titanium produce electrodes which are widely used for cathodic protection purposes, and in industrial electrochemical processes.

**Rhodium**

Rhodium is easily deposited and has a hardness of about 800 HV. It is used in manufacture of rotary contacts operating in conditions of extreme wear, and is also employed to protect silver against tarnishing and against wear in contacts carrying heavy currents.

**Ruthenium**

Ruthenium is becoming of increasing interest and may replace rhodium in certain applications.

**Composite materials**

The ability to co-deposit hard or lubricant particles considerably extends the range of coatings applicable by electrodeposition. Hard particles, such as ceramics, can be incorporated into a coating to improve wear resistance. Polymer particles can be incorporated to impart lubricity.

Commercially available processes are based on cobalt and nickel matrices containing about 30% by volume of particulate materials (around 3 μm size), such as chromium carbide and silicon carbide. Co-deposited lubricants include PTFE.

Electroless deposition methods may be modified to produce composite coatings with an advantage of uniform thickness. DTD 943 covers deposition of cobalt–chromium carbide composite coatings (Fig. 5.7).

**Cobalt–chromium carbide: Characteristics**

1. Hardness 350–500 HV.
2. Resistant to oxidation and fretting up to 800°C.
Electrodeposited coatings

5.7 Typical microstructure of a cobalt–chromium carbide electrodeposited composite coating (courtesy Praxair Surface Technologies Ltd).

Cobalt–chromium carbide: Applications
1. Aircraft jet engine components.
2. Textile machinery components.

Nickel–silicon carbide: Characteristics
1. Hardness 350–500 HV.
2. Wear resistance not generally as good as cobalt–chromium carbide.

Nickel–silicon carbide: Applications
1. Rotary internal combustion engines.
2. Seals for operation in fresh- and seawater.

Hard anodising
Aluminium alloys can be easily coated with, for example, chromium to improve wear resistance, but hard anodising constitutes a useful alternative method. Although applicable to most alloys, those containing more than about 3% copper or 7% silicon generally require modified treatment. BS5599 and DEF STAN 03-26/1 are the relevant UK specifications. PTFE can also be incorporated into the anodised coating by co-deposition, and this can improve the wear lifetime of a component by up to a factor of four, compared with a component that has only been anodised.

Characteristics
1. Hardness 400–500 HV.
2. Wear resistance comparable with hard chromium.
3. Thickness 25–75μm, sometimes up to 175μm.
4. Porosity may be sealed with lubricants.
5. Brittle, not resistant to shock loading.
6 Can be sealed to improve corrosion resistance, but wear resistance is reduced.

Applications
1 Transport industries especially aerospace.
2 Hydraulic valves, especially when PTFE is incorporated into the coat.
3 High speed machinery.
4 Injection moulds as an economic alternative to steel.
5 Self-lubricating surfaces.

Titanium can also be anodised (see British Standard BS M 58, from the Aero series). This is done to prevent galling of titanium fasteners, and for protection against galvanic corrosion when in contact with less noble metals.

Selection
Wear is a complex phenomenon and in practice more than one form is generally operating (see Chapter 2). For example, the surface of components may be subjected to relative movement in a corrosive fluid containing an abrasive powder. In this case, a designer would choose a hard coating material, such as chromium, to resist both abrasion arising from scoring or erosion and corrosive attack. A low coefficient of friction is desirable to combat adhesive wear associated with scuffing or galling, and chromium or electroless nickel would be natural choices. Softer coatings, however, such as electrodeposited nickel or copper, are preferred to combat fretting corrosion.

The development of wear-resistant coatings in which hard particles are co-deposited with a matrix metal, generally nickel or cobalt, adds to the design potential of this surfacing process. For wear resistance, the hard particles are often carbides of chromium or silicon but softer materials such as molybdenum disulphide for lubrication can also be used. Electroless deposition methods may also be modified to produce composite coatings of uniform thickness. In selecting a coating the designer may also have to consider factors such as ease of machining and resistance to shock loading.

Table 5.3 indicates an empirical order of suitability of the more usual electrodeposited engineering coatings. The ratings quoted are generalised and subjective, and serve only as an approximate guide. It is important to consult a specialised industrial electroplater before reaching a decision on a suitable coating.

Design
Electrodeposition must be regarded as an integral part of the production process and the designer should consult the electroplater at an early stage to achieve the best results.

The characteristics of an electrodeposited coating are primarily determined by the following:
### Table 5.3 Guide to selection of an electrodeposited engineering coating

<table>
<thead>
<tr>
<th>Coating</th>
<th>Relative cost* (A = least costly)</th>
<th>Ease of deposition</th>
<th>Ease of machining</th>
<th>Abrasion resistance at normal temperatures</th>
<th>Protection against corrosion</th>
<th>Resistance to shock loading</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chromium</td>
<td>D</td>
<td>3</td>
<td>1</td>
<td>6</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>Electrodeposited nickel</td>
<td>A</td>
<td>6</td>
<td>6</td>
<td>1</td>
<td>5</td>
<td>6</td>
</tr>
<tr>
<td>Electroless nickel – 9% phosphorus</td>
<td>F</td>
<td>5</td>
<td>5</td>
<td>4 (heat treated)</td>
<td>6</td>
<td>4</td>
</tr>
<tr>
<td>65/35 tin-nickel</td>
<td>B</td>
<td>4</td>
<td>3</td>
<td>3</td>
<td>4</td>
<td>1</td>
</tr>
<tr>
<td>Cobalt–chromium carbide composite</td>
<td>E</td>
<td>1</td>
<td>4</td>
<td>2</td>
<td>3</td>
<td>5</td>
</tr>
<tr>
<td>Hard anodised aluminium</td>
<td>C</td>
<td>2</td>
<td>2</td>
<td>5</td>
<td>2</td>
<td>3</td>
</tr>
</tbody>
</table>

1 = lowest or worst, 6 = highest or best.

*Costs depend on many factors, such as the shape of the work and the volume of production, especially for electroless nickel coatings.

1. The electrolyte system in use.
2. The lack of local uniformity of current density on the workpiece.
3. The dependence of deposition rate on current density, deposition time and deposition efficiency.

Other general features include the inability of electroplating to fill holes, level rough material and cover defects satisfactorily. While it may be possible to deposit some plating over such defects, the deposit will be poorly bonded to the substrate. Furthermore cracks or surface depressions tend to trap air or gas bubbles which prevent effective solution control and result in poor plating.

The electrolyte system will generally not be under the control of the designer but he/she can do much to alleviate the problems arising from the other factors, and can be helped by good electroplating practice. These two aspects are considered separately.

### Component design

Figure 5.8 shows examples of component design features that avoid or minimise many of the undesirable effects generated by electrodeposition. The underlying principles are to avoid sharp edges, sharp changes of section, close spacing of members, protuberances, and traps for the solution or gas bubbles.

When plated parts have to be fitted together, allowance must be made for coating thickness and variations in this thickness. Screw threads, for example, present a problem, and specification of an electroless process, if possible, improves the uniformity of the deposit.

Since the part being plated must be connected electrically into the circuit, the component must be suitably held. Sometimes this is best achieved by designing lugs away from the working surface which can be removed...
5.8 Design guides and requirements for components to be electroplated.
**Avoid**

- Sharp edges on holes give rise to excessive plating on edges
- Protuberances draw current preferentially
- Ribs should be smooth in section and spaced as widely as possible
- This produces a solution trap and poor throwing power is achieved
- Eliminate sharp edges and corners of slots
- Large overlap on jointed sheets

**Prefer**

- Rounded or countersunk holes give more uniform plating on edges of holes
- Inside curved surfaces should have a minimum radius of 12.5mm
- Smaller overlap reduces effect of solution entrapment
- As above (spot welded)
- Reduced joint area lessens danger of solution entrapment

5.8 (cont.)
easily after plating. The lugs may also facilitate holding for surface preparation and polishing (Fig. 5.9). Insoluble particles in the solution may settle on the uppermost surfaces of the component, creating roughness and possibly reducing corrosion resistance, so the designer should arrange either for non-significant surfaces or for areas of the significant surface that are of less importance to be uppermost when the article is in the plating solution.

Although barrel plating avoids many of the constraints of vat plating, it introduces some special requirements. For example design considerations should take account of the non-uniform distribution of finish in a barrel, caused by masking effects between adjacent parts. Parts must not be prone to interlocking. Flat parts such as washers can be dimpled to prevent sticking. Hollow parts may not immerse easily so they should be provided with drainage holes.

**Process design for vat plating**

Deposit thickness is dependent on current density, and this is greater at edges, corners and projections, but less at the centres of large flat areas, and in surface recesses. Thus even a flat sheet will not have a uniform current distribution and hence will not receive a uniform deposit. Figure 5.10 shows the current distribution and the resulting coating thickness, which results from a simple uncompensated arrangement of anodes.

Good plating practice can overcome the problem of deposit uniformity to a large extent by making the current distribution as uniform as possible, e.g. by

- correct positioning and size of anodes;
- using anodes that approximate to the shape of the surface or part to be treated;
- positioning ‘burners’ or ‘robbers’ (electrically connected to the cathode) near to high current density areas and so reducing the amount deposited on those areas;
- using non-conductive shields to reduce current flow at sharp edges.

The diagrams in Fig. 5.11 illustrate these principles.
5.10 Distribution of current and electrodeposits on a cathode of rectangular cross-section.

5.11 Plating practices to aid deposit uniformity: (a) positioning and size of anodes; (b) conformal anodes; (c) robbers; (d) non-conductive shields.
Substrate materials

Electroplated coatings can be applied to all the metals and most of the polymers used in engineering but care is needed with certain materials. The constraints on polymeric substrates are concerned with process temperatures and compatibilities. Some steels are susceptible to hydrogen that is released as a competing process to the electrodeposition of metals. Hydrogen ions in the electrolyte are discharged as atomic hydrogen which may rapidly diffuse into steel workpieces. If the steel has a tensile strength exceeding 1000 N mm⁻² or if it has been heavily cold worked before plating it may consequently suffer from hydrogen embrittlement. Heat treatment at 200–250°C after plating should release the hydrogen. It is preferable, however, to stress relieve components before electroplating, if possible.

Surface preparation

Substrate surfaces should be sound and without significant pores or defects such as cracks. The surface should be clean, smooth and free from burrs and machining or polishing debris. Traps for debris are best avoided by design. A polished or fine mechanical finish is most suitable.

Precleaning involves degreasing by solvent and/or alkaline electrolytic bath. The components should be compatible with these processes but the major responsibility for precleaning lies with the electroplater.

To avoid unnecessary plating, those areas not requiring treatment may be suitably stopped-off by the use of lacquer, tape, a well-fitting mask, or by screen printing. Again, the work should be compatible with the chosen method.

As in all aspects of coating, consultation between the customer and electroplater is essential if problems are to be avoided.

Finishing

Reshaping of plated parts must be avoided or the coating will be damaged, but some finishing, such as polishing, can be done if necessary. Ideally, however, finishing on the coating closely resembles that on the original workpiece. For close tolerance work, however, machining is necessary.

Electrodeposits are usually in a state of stress which can reduce fatigue strength if the deposit is in tension. In stressed parts, such as aircraft components, coating tensile stresses may be countered by introducing compressive stresses in the substrate by shot peening before plating. Alternatively, stresses can be reduced by heat treatment provided that the mechanical properties of the coating and substrate are not unduly influenced. Coatings on polymeric substrates cannot be stress relieved by the above means, and so the electrodeposition process must be chosen to minimise residual stresses.
Specifications, inspection and quality assurance

The designer should specify significant surfaces on components, masked areas, the material to be deposited, thickness distribution and tolerances, coating hardness and other mechanical properties, surface finish to be achieved, appearance of the coating and other specific requirements. The electroplater will advise on product shape and finish and on requirements for cleaning, jigging, plating, etc.

A procedure for sampling and inspection to meet the above should be agreed with the electroplater, and a procedure for handling items that fail inspection should also be agreed. Many of the standards quoted in this chapter give detailed guidance on specification and inspection but it must be emphasised that non-destructive testing of thin coatings (of whatever nature) is limited, and so quality assurance must play a major role in production of any component that is to be used as a critical part.
Chapter 6

Physical and chemical vapour deposition techniques

The processes

This chapter covers methods for producing overlay inorganic coatings which are formed on the surface of a substrate by condensation or reaction from the vapour phase. It does not cover thermal or thermochemical diffusion treatments, but ion implantation (although not strictly a coating process) is included.

Physical vapour deposition (PVD) coating techniques are almost wholly confined to making relatively thin films (ranging from $10^{-7}$ to $10^{-4}$m) whereas chemical vapour deposition (CVD) is used both for thin films and for coatings in excess of 1 mm. Such coatings have an important industrial role because of the anti-wear and anti-corrosion properties that they confer on engineering components. In addition to these uses there are large markets in the fields of electronic and optical devices, product decoration and architecture. As a result, development is taking place in all these directions and as each application emphasises its own special set of requirements, hybrid techniques are continually emerging; in fact the present extensive overlap in PVD and CVD methods means that hard and fast distinctions are difficult to make. The commercial success of many of the variants is open to doubt because their characteristics are not yet fully known and they have not yet found their niche in the market. The processes and their characteristics are summarised in Table 6.1.

All vapour processes involve treatment in a chamber—either in a vacuum chamber or in one that can withstand the high temperature and corrosive gases used in CVD. This limits somewhat the size of object to be coated but this limitation is largely imposed by the capital expenditure involved rather than by any fundamental characteristics of the process. Providing that the substrate can be manipulated to face the coating source, the size and shape of objects to be coated is again only limited by capital expenditure on plant. Small objects are held in large numbers in baskets and barrel-coated as in electroplating; at the other end of the scale large objects such as aircraft undercarriage units are coated with aluminium in chambers of 2 m diameter by 3 m long, and window glass sheets up to 6 m by 8 m are solar coated in plant up to 60 m long (Fig. 6.1).

Most processes are operated on a batch basis (Fig. 6.2), but vacuum evaporation on to paper or plastic sheet may be run with a continuous air-to-air feed through differentially pumped inlet and outlet ports.
Table 6.1 Comparison of process characteristics

<table>
<thead>
<tr>
<th>Processing</th>
<th>Throwing power</th>
<th>Coating materials</th>
<th>Coating applications and special features</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vacuum evaporation</td>
<td>Room temperature (RT) → 700, usually &lt;200</td>
<td>Line of sight</td>
<td>Chiefly metal, especially Al (a few simple alloys and a few simple compounds)</td>
</tr>
<tr>
<td>Ion implantation</td>
<td>200–400 best &lt;250 for N</td>
<td>Line of sight</td>
<td>Usually N (B, C)</td>
</tr>
<tr>
<td>Ion plating ARE*</td>
<td>RT → 0.7 $T_m$ of coating¹, Best at elevated temperatures</td>
<td>Moderate to good</td>
<td>Ion plating: Al, other metals (few alloys) ARE: TiN and other compounds</td>
</tr>
<tr>
<td>Sputtering</td>
<td>RT → 0.7 $T_m$ of metal coatings, Best &gt; 200 for non-metals.</td>
<td>Line of sight</td>
<td>Metals, alloys, glasses, oxides. TiN and other compounds²</td>
</tr>
<tr>
<td>CVD</td>
<td>300–2000 usually 600–1200</td>
<td>Very good</td>
<td>Metals, especially refractory TiN and other compounds (see below) pyrolytic C, pyrolytic BN</td>
</tr>
</tbody>
</table>

¹Activated reactive evaporation (includes sputter ion plating and arc evaporation).
²$T_m$ = melting temperature (Kelvin).
³Compounds: oxides, nitrides, carbides, silicides, borides of: Al, B, Cr, Hf, Mo, Nb, Ni, Re, Si, Ta, Ti, V, W, Zr.

The PVD processes described in this chapter are those that illustrate the major factors governing the PVD characteristics and the nature of the coatings that they produce; they are also the most widely used. PVD coating vapours are generated either by evaporation from a (usually) molten source, or by ejection of atoms from a solid source that is undergoing bombardment by an ionised gas (i.e. sputtering). The vapour may then be left as a stream of neutral atoms in a vacuum (vacuum evaporation) or it may be ionised to a greater or lesser extent. A partially (say 0.1%) ionised stream is usually
6.1 (a) Continuous sputtering line for solar coatings on architectural glass. (b) The Motley No. 2 Building, Texas, USA (courtesy Airco-Temescal, BOC Technologies).
6.2 Plant for vacuum evaporation of aluminium on to plastic foil (courtesy Leybold-Heraeus).

mixed with an ionised gas and deposits on an earthed or biased substrate (ion plating and sputter coating), but a highly ionised stream that forms a plasma is attracted to a biased substrate (arc plasma evaporation). Alternatively a 100% ionised beam may be focused and accelerated to sufficiently high energies to penetrate the substrate (ion implantation). This chapter concentrates only on those processes that are in commercial use (Fig. 6.3). The section on CVD coatings is subdivided on the basis of the chemical reactions involved.

Surface preparation

No vapour deposition method gives an adhesion acceptable for engineering purposes unless the substrate is truly clean. The standard of cleanliness required is far beyond that needed for weld surfacing or for spraying and it involves removal of contaminant layers only a few tens of molecules thick. Such preparation cannot be done in air; the most appropriate technique is ion bombardment and it is assumed that many of the PVD processes to be described will have been preceded by such a cleaning cycle. Ion bombardment cleaning is performed by a low pressure gas discharge; the work chamber contains argon at a pressure of some $10^{-2}$ torr and the workpiece is made negative at 2–5 kV with respect to earth. Positive argon ions generated in the discharge are accelerated to the workpiece at high energies and eject surface atoms when they arrive. The substrate is thus cleaned by an
6.3 Methods of physical vapour deposition: (a) vacuum deposition processes; (b) gas scattering processes.
erosion process which ‘sputters’ contaminant and substrate atoms into the chamber.

CVD processes, on the other hand, normally require only that the substrate be properly degreased and is free from obvious oxide films; a cleaning cycle involving a reducing gas at elevated temperatures is often all that is required before coating begins.

**Vacuum evaporation**

Vacuum evaporation is probably the most widely used of PVD techniques and it accounts for the major proportion of both the equipment in use and the area coated. It is also the oldest of the PVD methods and is, in principle, the simplest, but is rather limited in its applications. Some of these limitations can be reduced by using coating material sources that ionise their output, but some methods are much more complex and cannot yet be said to represent a significant industrial process.

Vacuum evaporation is usually conducted in a hard vacuum of 0.1–10 mPa, at which pressures the mean free path of a gas atom (i.e. the average distance travelled before it suffers collision with another atom) is about 1–100 m. As this is very much greater than the chamber dimensions, an atom evaporating from a source will travel in a straight line and so the process is essentially one of line of sight; coating around corners or into re-entrants is not possible without physical movement of the substrate (Fig. 6.4).

Coating material vapour is usually produced by thermal evaporation. The vapour consists largely of single atoms or clusters of atoms, although in
some variations of the process there may be an appreciable degree of ionisation of the vapour stream. Except where ionisation is present, the energy of the vapour particles is relatively low and is merely that necessary for evaporation (around 0.1–1.0 eV).

The substrate to be coated is usually unbiased (i.e. earthed) and may be heated or cooled. Deposition rates in vacuum evaporation are amongst the highest in all PVD processes and can range from 75 μm min⁻¹ but a more usual figure is around 2 μm min⁻¹. Coatings can be deposited to preserve the finish of the underlying surface, but adhesion of thermally evaporated coatings is wholly dependent on the cleanliness of the substrate immediately before deposition starts. For many less demanding industrial applications, the ion bombardment cleaning cycle is omitted and adhesion consequently suffers.

The properties of the coatings are strongly dependent on the ratio of substrate temperature to the melting point of the coating material. At low temperature ratios coatings tend to have a high dislocation density, similar to a cold worked material, and mobility of the adherend species is poor, leading to columnar voids if the material does not arrive normal to the substrate. At high temperature ratios (above approximately 0.5), recrystallisation of the coating can occur, leading to a coating with properties resembling those of fully annealed bulk materials.

**Evaporation sources**

Vacuum evaporation at sensible rates implies a vapour pressure of the source material around 1 Pa which, for the majority of materials, requires a temperature well above the melting point. Thus some form of crucible must be used that can contain the melt without reaction and this can be a major problem. Heating the source is commonly accomplished by direct resistance heating of, for example, tungsten or molybdenum boats (Fig. 6.5), by induction heating, or by electron beam heating of the material in a water cooled copper crucible (Fig. 6.6). A major exception is aluminium which is evaporated from an electrically conducting refractory boat of boron nitride/titanium dioxide mixture, because the molten metal reacts with materials such as tungsten.

Within limits, alloy deposition is possible if account is taken of the different vapour pressures exerted by the alloy components. Although the problem can sometimes be avoided by dropping a fine powder of the alloy on to a superheated boat to cause total vapourisation without fractionation (flash evaporation), limitations are imposed by reactions between the powders and the boat.

Electron beam heating is a versatile process and avoids the problem of reaction between the molten source and the crucible. It can also be used for making alloy coatings by oscillating the beam between crucibles containing the individual constituents and correspondingly adjusting beam power to produce the correct vapour pressure above each. However, the beam generator must be shielded from the metal vapour and the systems are complex (Fig. 6.6).
Materials and applications

The method is easiest to apply to elemental materials that have low melting points and high vapour pressures. The major applications of vacuum evaporation are confined to coating of single metals such as aluminium (which constitutes 90% of all materials evaporated), chromium, silver, etc. Alloys whose constituents have similar vapour pressures can be passed into the vapour phase with little change in composition. Some compounds, such as SiO and MgF₂, which are widely used in optics, can also be evaporated without change of composition.

Engineering uses of vacuum coatings are limited – general aerospace applications use Al and Ni–Cr (for corrosion protection) and Al and Ag as solid lubricants. Hard coatings of Cr or Al₂O₃ are sometimes deposited on
steel or tungsten carbide tools but this application has now largely been taken over by sputter coating. MCrAlY coatings, where M may be Co, Ni or Fe, are used to coat nickel-based alloy turbine blades, and TiC and TiN coatings are used for high speed cutting tools. Probably the largest area coated is for decorative purposes, e.g. Al and Cr for automotive trim, Al for packaging and costume jewellery, and fluorides for iridescent effects. Evaporated coatings for electronic applications include Al and Ta on polymer films for capacitors, Al–Si and Ni–Cr by flash evaporation for thin film resistors and Si, GaAs and CdS for semiconductors and solar cells respectively. Optical coatings include Al and Ag for mirrors and reflectors and also
for solar coatings on windows; in addition evaporated MgF₂ is well known as an anti-reflection coating on lenses.

**Gas scattering deposition**

Vacuum evaporation, being a line of sight process, normally has poor 'throwing power'. However, the presence of gas molecules in the coating chamber reduces the mean free path of the evaporant atoms and, if there are sufficient collisions, the coating atoms will no longer impinge on the substrate in a straight line from the source. At 1 Pa, the mean free path is about 5 mm; thus in a system with a typical source to substrate distance of some 200 mm, evaporant atoms will suffer over 50 collisions en route and will arrive at the substrate from almost any direction; their arrival route will now be partially controlled by a gas diffusion process. The pressure range for this regime is from 0.1 to 5 Pa at which there is a substantial gain in throwing power without appreciable reduction in evaporation rates; furthermore, the introduction of a gas into the coating chamber offers a number of new variables which allow an improvement in coating quality and an extension of the number of materials which can be deposited as coatings. The major variables are: electrically neutral or ionised gas, and chemically inert or reactive gas – the combinations are illustrated in Fig. 6.7.

**Ion plating**

**General principles**

Evaporation into an electrically neutral gas has been virtually superseded for industrial purposes by use of an ionised gas as the scattering medium, i.e.
ion plating. A dynamic pressure of some 1 Pa of a gas such as argon is maintained by a continuous gas bleed and throttled pumping (a dynamic pressure is preferred to prevent the build-up of outgassed impurities in the chamber), and the substrate is negatively biased at 1–2 kV with respect to the grounded evaporation source (Fig. 6.8).

Evaporant atoms gain energy by collision with the ionised argon but contrary to the earlier belief, from which the name ion plating originated, the degree of ionisation of the vapour stream is probably no more than 0.1%; the term ion plating is therefore largely a misnomer, but is widely used. The combination of high energy (approximately 10 eV) evaporant atoms plus continuous substrate bombardment by gas ions results in excellent film adhesion and, by correct choice of deposition parameters, in a dense coating. At the same time the presence of the gas ensures a good throwing power (Fig. 6.9).

The evaporant charge is melted as in vacuum evaporation and gas ionisation may be advantageously increased by a device such as a heated filament electron emitter. There are several variants of the technique and all claim high film adhesion and density but all require the substrate to be held at a high DC potential or to be connected to an RF (radiofrequency) supply.

Materials and applications
Deposition rates are somewhat lower than in vacuum evaporation (some coating material is scattered away from the substrate) but they are nevertheless among the highest of the PVD techniques. Dense deposits are best achieved by deposition at elevated temperatures up to $0.7 T_m$ (melting temperature) and the grain structure of the deposit (columnar, equiaxed, etc.) is controlled by a suitable choice of deposition rate, gas pressure, substrate bias and substrate temperature.
Ion plating has excited considerable interest over the past 20 years but it is now used only for a number of important industrial applications. As with vacuum evaporation, the method is limited to those metals which can be evaporated, and the same limitation with regard to crucibles, etc., apply. Furthermore, because the substrate holder – which may be required to oscillate and rotate the parts to be coated – is maintained at a high potential, the complexity of the equipment is greater than that for simple vacuum evaporation; nevertheless the improvement in coating quality is such as to warrant that complexity.

It is not practical to electroplate aluminium for corrosion protection, and thermally sprayed coatings may not be suitable for some components, but ion plated films of aluminium are routinely applied to complex aircraft parts (Fig. 6.9). Ion plating rivals or has even displaced electroplating for some materials such as cadmium where there is a danger of hydrogen embrittlement of high strength steel substrates and where the poisonous nature of the solutions has caused them to be banned in some countries; chromium is increasingly being deposited on to polymeric materials for car trim for the same reason. Solid lubricant films of, e.g. silver, for aerospace applications are ion plated on to bearings, etc., that are to be exposed to heat, radiation and hard vacuum.

In general any material that can be vacuum evaporated can be ion plated with superior adhesion and density.
Activated reactive evaporation (ARE) techniques

Reactive ion plating

Vacuum evaporation and ion plating are both restricted to coating materials that can be evaporated without change of composition. There is a considerable demand, however, for wear-resistant coatings of oxides, nitrides, carbides, etc., and it was for this reason that glow discharge ARE (also known as reactive ion plating) was developed. The principle is similar to that of ion plating except that a reactive, instead of an inert, gas is used to generate the discharge. The metallic constituent of the compound is evaporated into the reactive gas whose reactivity is considerably enhanced by the glow discharge and reaction takes place either in the gas phase or, more commonly, on the substrate surface. Thus a thin, hard, wear-resistant coating of titanium nitride (TiN), for example, can be produced by evaporating titanium into a low pressure glow discharge of a mixture of argon and nitrogen; one variant of this technique is shown in Fig. 6.10.

The diagram shows what is basically an electron beam heated vacuum evaporator into which the reactive gas is introduced close to an auxiliary ionising electrode. The resultant plasma activates the reaction by ionisation of the reactive gas and by creation of metastable species. Further complexity is introduced by the necessity to control the auxiliary electrode and the balance of the reactive and inert gases, but ARE is one of the two evaporative methods used for making coatings of many compounds.
Arc evaporation ion plating

The reactive methods described earlier use some form of indirect heating to melt the coating material which is evaporated with little ionisation into a generalised glow discharge. Arc evaporation, on the other hand, uses a solid metal target which is locally and momentarily struck between the source (cathode) and either the chamber (extended anode) or a localised anode, although unlike atmospheric arcs no luminous discharge path is present. The high power density at the root of the arc (>10^9 W m^-2) causes an intense evaporation of metal vapour which (in contrast to ion plating and other forms of ARE) is extensively ionised and is attracted to the negatively biased substrate (Fig. 6.11). The substrate undergoes a vigorous bombardment by metal ions which clean and heat the surface without causing a build-up of coating material. A reactive gas is then admitted and the parameters are adjusted to give ion plating conditions.

The use of an arc to cause evaporation from an effectively solid target overcomes many of the limitations of fully molten sources. Thus, high melting and reactive metals are melted without crucible problems and multiple sources can be arranged vertically, horizontally downwards or at any angle to improve the speed and uniformity of coating. The combination of elevated substrate temperature and highly energetic coating ions ensures excellent adhesion and dense coatings, but droplet ejection from the target may cause a higher degree of surface roughness on the coating than is found in other reactive techniques.
Sputter ion plating (SIP)

Another commercially important technique for overcoming the limitations of evaporative vapour production (and one that does not involve any melting whatsoever of the coating source) is ion bombardment sputtering, which is more fully described later in this chapter. Sputtering does not require the coating source to be melted and it therefore adds considerably to the range of materials that can be deposited. The SIP process can be run with both inert and reactive gases, but it is much slower than the evaporative techniques. On the other hand, the characteristics of SIP include high purity and control of the constituents and components in the system. Furthermore, it is amenable not only to large chamber operation, but also to treating mixed batches of components; it is these factors, particularly, which give the technique a certain commercial advantage over its competitors.

Materials and applications

Oxides, carbides and nitrides are the principal compounds described by SIP and the various ARE techniques by matching the generation rate of the metallic constituent with the partial pressure of oxygen, ethane or nitrogen respectively. Molybdenum disulphide films are also produced, and other compounds such as borides or silicides have been applied as coatings, but there is, as yet, no major industrial use for these latter.

The compound that dominates this field is titanium nitride. Films of TiN some 5μm thick are now applied to bearings and to a wide range of tools, such as simple twist bits, complex gear hobs, punches, dies, taps, milling cutters and forming tools (Fig. 6.12). Tool life is improved three to ten times by the coating, and cutting speed can often be increased; moreover, the quality of the work is also improved.
This process is now an enabling technology for the expansion in use of titanium and titanium alloys. There is a growing use for nitrides and carbides to give the best surface characteristics in components made from high performance alloys in the automotive and aerospace sectors.

The wear-resistant films are golden coloured and are also used to provide decorative scratch resistance to, for example, stainless steel or gold watch cases. Substrate temperatures are generally held above 400°C for the more demanding tribological applications but films for decorative purposes may be deposited at much lower temperatures.

Although TiN is currently by far the most widely used of the wear-resistant thin film compounds, others such as titanium carbide, titanium carbonitride, titanium aluminium nitride, and carbides or nitrides of, for example, hafnium, chromium or zirconium are used on a much smaller scale. They may find specialised applications at, for example, temperatures above 500°C where TiN is unsuitable. Research into thin tribological coatings is presently very active, and undoubtedly new coatings, and multilayer coating systems, will emerge in the next few years.

In the optical/electronic fields, ARE and its associated techniques have allowed manufacture of scratch-resistant anti-flare coatings of compounds that could be evaporated, and there is also an important market for thin (<0.1μm), transparent and electrically conducting films of indium–tin oxide deposited as de-icing layers on, for example, aircraft windows.

There is presently much emphasis on thin tribological films, but metal and – to a much lesser extent – alloy coatings are also made by these ionisation-assisted techniques. Sputter ion plating produces dense adherent and pure coatings of refractory metals, and aluminium and cadmium coatings for corrosion protection have been reported. Aluminium conductor tracks are also ion plated on some electronic devices, as are copper tracks. However, the major area of operation of all of these techniques is to deposit thin coatings of materials – metals, alloys, compounds – that cannot be deposited by electroplating.

The various activated reaction techniques are in competition and there is much overlap between them, particularly for production of titanium nitride coatings, but it is too soon to tell which, if any, will eventually command the field. The different characteristics of the techniques have, in the past, led to different characteristics in coatings of nominally the same material, but these differences seem to be decreasing. The eventual dominance of one method may be influenced more by its commercial than by its technical characteristics.

Sputter coating

General principles

In the ion bombardment cleaning cycle that precedes evaporation coating, material is ejected from the face of the negatively biased substrate by argon ion bombardment. The ejected material is in the form of high energy atoms
which condense on the surrounding surfaces (the darkening that occurs at the ends of fluorescent light tubes as they age is material that has been sputtered from the electrodes onto the glass). Because the high energy of the bombarding ions is sufficient to overcome the binding energy of any substrate lattice, and because the source generates its own vapour not by heating and evaporation but by ion bombardment, the technique can be used for depositing a wide range of coating materials (the source is called the target: Fig. 6.13). It is this feature, above all others, that makes sputter coating the most versatile and probably the most widely used process for engineering, optical and electronic thin films.

The target is a solid and can be of an alloy or compound (or even a pseudo-alloy consisting of a mixture of the powdered constituents); the gas ionising potential is applied to it as DC or RF. The gas itself may be inert or reactive and the substrate may be earthed or biased; the combinations are shown in Fig. 6.14. The temperature reached by the substrate is governed by many factors, but unless higher values are required, it is generally below 200°C.

There are snags of course. The necessary gas pressures (around 0.5 Pa) reduce the mean free path to 30 mm or so, therefore the substrate must be approximately at this distance if the high energy of the coating species (with all the advantages that this confers) is not to be lost. This in turn means that throwing power is compromised and that the target must be similar in size and shape to the substrate. There are other snags of which the most unfortu-
nate is the low rate of deposition; the simple diode system shown in Fig. 6.13 cannot exceed coating rates of 0.1 μm min⁻¹ at best and with some coating materials it operates at one or two orders of magnitude lower. More complex systems involving auxiliary ionising sources may achieve 0.4 μm min⁻¹, but a slow deposition rate is not only unfortunate from a production aspect, it can also lead to a high gas content in the deposit. A film deposition rate of 0.1 μm min⁻¹ is equivalent to about three monolayers per second and this is about the same figure as the arrival rate of molecular oxygen at a partial pressure of 1 mPa. Good vacuum practice is therefore needed to maintain oxygen-free films; furthermore, subsequent heat treatment may be required to desorb occluded discharge gas. An important development, however, is the magnetron source, which can give very high rates of deposition. This is described later.

Despite its problems, the sputtering technique can deposit excellent (albeit thin) coatings of materials that cannot be deposited by any other method.

**Substrate biasing**

The usual sputtering technique is to apply an ionising potential to the target only, leaving the substrate earthed, but there is a variant of the technique known as bias sputtering in which the substrate is also biased, but at a lower potential than the target. The effect of substrate biasing is to divert some of the ion bombardment to the substrate and so to gain the advantages that apply to ion plating. Possibly this variant should be regarded as a form of ion plating in which sputtering (rather than evaporation) is used to generate
the coating material vapour. Substrate bias may be positive or negative, according to the ionised gas used and the composition of the film to be deposited.

**Gas ionisation**

Sputter coating has been in use since the 1920s but in its early form, as illustrated in Fig. 6.13, it had two major disadvantages, namely the necessity to confine its operation to electrically conducting targets (insulators became charged during bombardment and repelled gas ions) and a low coating rate (caused by a comparatively inefficient argon ionisation rate).

The problems of insulating targets becoming charged under DC bombardment are overcome by the use of RF excitation. If an insulated or non-conducting target is capacitively coupled to an RF generator the space charge that would otherwise build up is effectively abolished and the surface adopts negative charge. Frequencies above 10 MHz are used and the RF bias is typically 2.5 kV peak-to-peak. The target can therefore be of an insulating material, and oxides, for example, can be directly sputtered onto a substrate although the erosion rate of an oxide is generally only about 10% of that of the corresponding metal. The ability to produce coatings directly from non-metallic targets is a valuable property of RF sputtering.

Early attempts to increase the ionisation efficiency of the simple DC diode system were successful to some degree and sputtering rates were increased by a factor of three or so, but the introduction of magnetically assisted (magnetron) sputtering led to deposition rates that are comparable with those from vacuum evaporation. Magnetron sputtering has undoubtedly been responsible for the widespread adoption of sputter coating by many branches of modern technology.

Magnets are arranged behind the target such that the magnetic flux lines lie parallel to the target along part of their path and so lie at right angles to the electric field (Fig. 6.15). Electrons caught in this crossed field configuration are forced to spiral parallel to the target and so, by increasing electron path length, the chance of ionising collisions with gas atoms is also increased. A region of intense ionisation is created immediately above the target surface which is heavily bombarded and generates a high flux of coating atoms.

The method can only be used to make magnetic coatings when the target is held above the Curie temperature, otherwise ferromagnetic targets confine the magnetic field and so shield the discharge electrons from its effect. There are numerous variations on this basic theme, differing with respect to biasing, target configuration and magnet; Fig. 6.16 shows a rectangular planar source, but cylindrical sources are also available. Magnetron sources coupled with various forms of RF biasing have enormously increased the scope of sputter coating which is nowadays used on a large scale.
Materials and applications

Materials

Sputter coating, like ion plating and ARE, can use elemental sources running in an inert gas plasma to deposit elemental films, or elemental sources
in a reactive gas to deposit the corresponding compound. Sources that are themselves chemical compounds may also be used, but some change of composition is often found in the coating. The choice between reactive sputter coating from a metal target or coating from a compound is determined by many factors, but both methods are in widespread use as is shown by the vast range of materials available as sputtering targets. One catalogue lists more than 36 metals, 32 intermetallics and alloy types, 44 oxides, 19 borides, 14 carbides, 17 fluorides, 9 nitrides, 14 silicides, 9 selenides, 10 sulphides and 7 tellurides. It is also possible to make coatings of PTFE and some cross-linked polymers. In size and shape the targets range from metal plate several metres long to shaped or planar discs, rings, bars, cylinders, buttons, etc. The limits are generally financial rather than technical.

Applications
The following lists are only a sample of the many uses to which sputtered thin films (some only 5nm thickness) are being put.

Engineering
- Corrosion and oxidation resistance, e.g. Ni–Cr, M–Cr–Al–Y, polymers.
- Wear resistance, e.g. TiN, other nitrides, W, Mo, carbides, borides, diamond type carbon.
- Lubricants, e.g. Ag, In, MoS₂, PTFE, selenides, silicides, tellurides.

Electronic
- Metallisation, e.g. Al, Cr, Al–Si.
- Resistors, e.g. Cr–SiO, Ni–Cr.
- Insulators, e.g. SiO, SiO₂, TiO₂, silicides.
- Diffusion barriers, e.g. Si(ON), silicides, TiN.
- Semiconductors, e.g. GaAs, GaP, InAs, InP.

Optical, electro-optic
- Lens coatings, e.g. MgO, MgF₂.
- Transparent conducting films, e.g. In–Sn–Ox.
- Antiglare coatings, e.g. TiO₂–Ag, Cd–Sn–Ox, SnO₂–Cr(N).
- Interference filters, e.g. Ge, metal oxides, fluorides, Zn Se–S.
- Luminescents and light-emitting diodes (LEDs), e.g. MoO₃, ZnSe.
- Photoconductors, e.g. selenides, sulphides, tellurides.

Ion implantation

General principles
Ion implantation does not produce a coating; the process generates an intense beam of very high energy ions (much higher than in ARE or sputter-
ing) which penetrate the substrate surface and modify, rather than coat, it (cf. carburising). Extensive work in this field has been carried out by Dearnley and colleagues at Harwell who have concentrated on beams of ionised nitrogen (boron and carbon have also been explored). Nitrogen ion beams with energies in the range 80–150 keV are directed on to the surface of the workpiece and penetrate to depths of 0.1 µm. The process operates in a vacuum (and is therefore a line of sight technique) and the beam is typically of square cross-section which expands at an angle of about 20° to cover some 150 × 150 mm on the workpiece which may be masked or rotated. Treatment times vary from 1 to 10 hours according to the area to be covered and the temperature rise may be no more than a few tens of degrees Celsius, so that fully finished and heat treated parts can be implanted with virtually no change of dimensions or distortion. The only visible change is a slight polishing of the surface.

As the implanted ions are scattered beneath the surface, some of them become trapped in dislocations and other defects, thereby effectively pinning them; others form metastable nitrides in some alloys which decompose to give martensitic hardening (unlike gas nitriding of steels which forms stable nitrides). The result of these processes is an increase in hardness and wear resistance, a decrease in friction coefficient and galling tendency, and generation of compressive stresses in the surface which improve both the fatigue and corrosion resistance.

A unique and most useful feature of the process is the retention of many of these surface properties after wear to depths which may be 100 times greater than the original implanted depth. It is believed that some of the implanted nitrogen diffuses ahead of the wear front and maintains the original effects.

**Materials and applications**

Ion implantation is commercially applied to various steels, to tungsten carbide/cobalt materials, to alloys of titanium, nickel, cobalt, aluminium, copper, to chromium plate and even to diamond. Extrusion and moulding components for the plastics industry are extensively treated, as are slitter blades for rubber, paper and textiles.

Applications are limited to service temperatures below, say, 250°C for steels and 450°C for carbides, but within these restrictions there are many areas, particularly in metal forming and general engineering, where improvements in service life of a factor of ten are regularly reported for steels, and factors of three or four for carbides and non-ferrous alloys. The marked exception to this latter case is titanium where life improvement factors for prosthetic devices of several hundred are claimed. The cost of treatment is generally in the range of 10–30% of the original component cost.

**Chemical vapour deposition**

The techniques of chemical vapour deposition (CVD) have been in use for the best part of 100 years, but not specifically for coating purposes; for example the Mond process for purifying nickel via thermal decomposition
of the volatile nickel carbonyl is a CVD process, as are pack aluminising and chromising. As a coating process, however, CVD began to play a specialised, but important role only when the low melting halides of the refractory metals were systematically investigated. Nowadays, the term chemical vapour deposition covers a broad range of processes, all of which use gaseous reagents undergoing chemical reactions near or on the heated surface that is to be coated. The major reactions involved in coating processes are shown in Fig. 6.17.

In general, there are three steps in any CVD reaction: firstly, the production of a volatile carrier compound, e.g. nickel carbonyl; secondly, the transport of that gas—without decomposition—to the deposition site; thirdly, the chemical reaction necessary to produce the coating on the substrate. These steps may be quite distinct in both time and space, or they may all occur at the same time within the same reaction chamber—as in pack aluminising or, indeed, as in the quartz iodide lamp (in this latter case, the source—the tungsten filament—is being continually recoated by its own evaporated vapour).

The requirements of these three steps impose many limitations on the process; nevertheless, CVD usefully fills a gap where the desired element or compound cannot be satisfactorily deposited by other means to the required thickness. CVD is probably unique in its ability to make pyrolitic carbon and pyrolitic boron nitride.

**Process characteristics**

The high temperature and general nature of the reaction involved tend to require less stringent cleaning of substrates before coating them for PVD processes; equally the high temperatures ensure high density and good adhesion. The coatings are generally rougher than the underlying substrate, but this is partially a consequence of the thicker coatings which range from, say, 10μm to 1mm. Because the process uses gas at or near atmosphere pressure, CVD has excellent throwing power and coats all exposed surfaces to a high degree of uniformity. This also means that mixed loads of components can be treated (i.e. mixed materials as well as mixed sizes and shapes). The process is particularly well suited to coatings on sintered carbides (see below).
Reaction types

Thermal decomposition

Elemental coatings, chiefly of refractory metals, are produced by simple thermal decomposition of volatile compounds. The most commonly used compounds in this class are: carbonyls, e.g. Ni(CO); halides, e.g. WF₆; hydrides, e.g. B₂H₆; and certain organo-metallics, e.g. copper acetyl-acetonate.

The carbonyls usually decompose to the metal plus carbon monoxide at temperatures below 200°C but the substrate must often be much hotter than this if coherent and adherent coating is to be produced; generally, however, substrate temperatures can be kept below 600°C. One severe disadvantage of the carbonyls is their high toxicity.

Halides and hydrides tend to require much higher decomposition temperatures, say above 500°C, but coating requirements again dominate and temperatures above 1000°C are not uncommon. There are many more of these compounds suitable for CVD than there are in the metal carbonyl group.

Organic compounds are not widely used on their own for coating purposes because of the side reactions between the products of decomposition, but they decompose at lower temperatures than the halides and are essential in the production of pyrolytic carbon either as a coating or, more importantly, as free-standing shapes. Simple hydrocarbons such as the low paraffins are used to make pyrolytic carbon, but substrate temperatures above 1500°C are required.

Reduction to metal

Reduction — usually of halides — by hydrogen is possibly the most important route for the formation of refractory metal coatings; the chlorides tend to dominate this group. Although the presence of hydrogen reduces the reaction temperatures, higher levels are often demanded for dense coatings, e.g. 1100°C for titanium, but excellent deposits of molybdenum or tungsten are now routinely produced by hydrogen reduction of the gaseous chlorides at a rate of about 250μm·h⁻¹ at 600–700°C. The process is also used to form free-standing bodies, such as rocket nozzles or crucibles of tungsten and pyrolitic boron nitride (Fig. 6.18).

Reduction to compounds

A major commercial application of CVD is in the coating of cemented carbide tips on cutting tools, e.g. TiC which can be produced from the reaction between titanium tetrachloride, methane and hydrogen. Provided that pressures well below atmospheric are used, the reaction temperature can be reduced to 700°C and a TiC layer some 5μm in thickness will give a ten-fold increase in tool life. Titanium carbonitride is produced at somewhat higher temperatures but gives a rather longer life. Titanium nitride (from the chloride plus nitrogen and hydrogen) has excellent wear resistance and an attractive golden colour; it is now more widely used than the
carbide and is applied to a variety of components ranging from tool tips to watch cases. It is reported that more than 60% of all indexable carbide cutting tools now made in the USA are CVD coated with TiC or TiN, and to a lesser extent HfN, HfC or Al₂O₃ at a cost ranging between 5 and 50% of the original tool cost.

In addition to the above, many nitrides, carbides, oxides, silicides and borides are produced as wear-resistant coatings by CVD, but the reactants are frequently poisonous and temperatures above 1000°C are usually needed. One exception to this last is of W₆C which can be produced well below 700°C and may thus have considerable potential for coating heat-treated dies, etc., without the need for further heat treatment.
Disproportioning reactions

This group comprises a very small number of reactions, possibly the most important being the sequence involved in pack aluminising. Other metals treated in this way (e.g. germanium, gold) have little interest in engineering coatings in the present context.

**Plasma activated CVD (PACVD)**

The great improvements in PVD processes brought about by ionising the reactive species prompted a similar approach in CVD where the introduction of a glow discharge further lowers reaction temperatures by many hundreds of degrees Celsius. Whereas CVD of SiO₂ from tetraethoxysilane requires temperatures of some 700 °C, the use of RF PACVD reduces the reaction temperatures to 200–300 °C and therefore allows quartz films to be deposited on semiconductor materials. Reaction pressures must, however, be reduced below 100 Pa to avoid arcing and therefore coating rates are reduced compared with the non-activated reactions, and may be as low as 0.3 μm h⁻¹.

Both DC and RF plasmas (with frequencies into the microwave region) have been used, and it has been demonstrated that the quartz reaction above is efficiently activated by UV light. Microwave plasmas are currently of considerable interest for the activated manufacture of diamond films. The films are deposited from a mixture of hydrogen and organic gases containing methyl radicals at growth rates of some 10 μm h⁻¹, but it is reported that Russian workers are producing coatings more than 1 mm thickness. Diamond films have a huge potential as wear-resistant coatings and in various semiconductor applications, but the first commercial product was in the hi-fi field. The Japanese company Sony marketed a diamond film high frequency loudspeaker (tweeter) based on the piezoelectric properties of diamond; the film, being light and stiff, responds particularly well to the high audio frequencies.

**Materials and applications**

Metal coatings produced by CVD may be pure and ductile and can be deposited at virtually 100% density. Coatings of, for example, Cr or Ni can be made, but CVD is a more expensive process than electroplating and so is usually applied only to those metals that cannot be electrodeposited. The refractory metals, W, Mo, Re, Nb, Ta, Zr, Hf, etc. – and some of their alloys – are the major candidates – have coating rates that are dependent upon reaction temperature and range from 3 μm to 3 mm h⁻¹. The major applications for metals take advantage of the excellent throwing power of the process to produce coatings and free-standing shapes at temperatures well below the usual processing levels. Typical products are crucibles, rocket nozzles and other high temperature components, linings for chemical vessels and coatings for electronic components.

Pyrolytic carbon is remarkably inert and has an excellent thermal conductivity. It is used in chemical vessels, high temperature crucibles, as a
coating on nuclear fuels and, in one rather less exotic application, as a bowl liner in tobacco pipes. Pyrolytic boron nitride is chiefly used for crucibles in the manufacture of certain electro-optic materials, and as a coating for cutting tools.

A substantial field for CVD exists as yet another process for producing thin films of TiN, TiC and other compounds on metal working and forming tools and on components such as impellers, valves, nozzles, etc., that are subject to abrasion and corrosion. The widespread application of CVD coatings where very high abrasion resistance is required (e.g. thread rolling dies) is largely confined to TiC, but where high lubricity and galling resistance is paramount, TiN is used on, for example, carbide tips, taps, and punches. To a lesser extent, alumina, which acts both as a thermal barrier and as a protection for the cobalt binder, is applied to carbide tools that are being run at unusually high feed rates.

Although other processes have been applied to the deposition of diamond-like films (i-carbon), the CVD techniques are probably the only ones that can deposit coatings to thicknesses sensibly above a few micrometres. Diamond coatings for wear resistance and for electronic and possibly some optical applications are probably on the brink of a big expansion.

Other compounds that are deposited – particularly by PACVD – for electronic applications include quartz, silicon, silicon nitride and titanium nitride which are used as thin film substrates, dielectrics and insulating layers.

In spite of high coating rates, the generally high temperatures tend to make batch processing times comparable with the slower coating processes. Furthermore, the high treatment temperatures often require that alloy steels must undergo further heat treatment after coating. Lower temperatures are possible when the reactions are plasma activated, but PACVD coating rates are generally lower as a consequence of the lower gas pressures that are employed. These and other factors militate against simple comparisons with other processes and indeed against a clear-cut choice for a specific job, but the CVD processes do have a few outstanding characteristics that make them indispensable for some applications.

Comparison of processes

Although there is a lot of overlap between the characteristics of the various processes, it is difficult to make direct comparisons because so much depends upon the applications involved. Moreover, many of the processes are in an active stage of their evolution and so the sustainability of a particular technique for any given application may change dramatically with the advent of new development. However, it may be said, for instance, that ion implantation is limited almost entirely to wear-resistant applications; that CVD is restricted to substrates that can safely be heated to (preferably) 300°C or more, and that thicknesses sufficient for free-standing bodies and thick layers of pyrolytic materials can be made only by CVD. These statements apart, Fig. 6.19 (range of thicknesses), Fig. 6.20 (deposition rates) and Table 6.1 demonstrate the wide overlap that exists between
6.19 Comparison of thicknesses.

6.20 Comparison of deposition rates at typical working distances (e.g. sputtering at 50 mm, vacuum evaporation at 250 mm).
the processes in respect of coating thickness, coating rate, coating materials and the areas of application.

**Coating thickness**

The range of thicknesses (Fig. 6.19) does not necessarily apply to all materials available to each process, nor to all variants of that process, nor indeed do the values shown necessarily imply that these are the absolute limits, but the charts do show the most commonly encountered ranges for applications reported. Coating thickness is often determined by the application and not by the process capability.

The above remarks do not apply to ion implantation. Note that Fig. 6.19 does not show the thickness of ion implantation but the initial implantation depth — the concentration/depth curve being near Gaussian — however, a unique feature of nitrogen implantation into steel is the inward migration of nitrogen as the surface wears away. Thus, although the major concentration of the implant is initially confined to a thin subsurface layer of only a fraction of 1 μm, the wear resistance persists to a much greater depth.

**Coating rate**

Comparison of coating rates for a given material is complicated by the fact that different processes use different source to substrate distances. With the exception of ion implantation, the coating vapour stream issues as a cone, or even totally fills the reaction chamber, thus coating rates vary in a complex manner with coating distance; Fig. 6.20 shows the range of rates usually found in practice. Virtually all process rates can be increased by having a multiplication of sources and, to a lesser extent, by an increase in the power input, but the coating quality may suffer if too high a rate is attempted.

The magnetron sputter source is expected to continue to increase in efficiency as a result of the intense developments that are taking place. There has been such an increase in magnetron sputtering rates that the process has become commercially competitive with evaporation techniques for many materials and there is as yet no indication that the limit has been reached particularly for area coverage.

Ion implantation rates cannot be quoted in the same way as coating rates. The total dose of implant is the important factor (usually $3 \times 10^{17}$ nitrogen ions per square centimetre) and therefore the treatment rate is the time taken to implant unit area with the required dose. Treatment rates of up to $5 \text{cm}^2\text{min}^{-1}$ of nitrogen implant are currently possible and again there is no indication that the limit has been reached.

**Cost**

Direct comparisons of coating cost are not generally possible, but broad generalisations can be made. Vacuum evaporation by resistance heating
is probably the cheapest in all respects, but the introduction of electron bombardment heating or RF power can significantly affect the capital and running costs. Ion plating and its associated techniques require fairly expensive equipment in comparison with vacuum evaporation because there are so many more factors to control, and because coating efficiencies are lower.

The capital cost of sputtering equipment is high and its complexity is increasing but it is undergoing continuous development (thanks to the electronic industry) and basic unit costs are not increasing too rapidly in real terms. It is too early to make cost comparisons between ion implantation and the other methods because the process has only recently been launched on a commercial scale; area treatment direct costs of ion implantation are modest but capital costs are high. However, here again development, principally for machine tools, is continuing and is bearing fruit.

CVD is a remarkably inexpensive method of forming some of the refractory metals when compared with the traditional techniques, and even its unique ability to produce pyrolytic materials is not an inherently costly process, but the indirect costs caused by the toxicity of many of the compounds used can add significantly to the capital and running costs.

Design for physical and chemical vapour deposition

General considerations

Physical vapour deposition techniques are many and varied and include warm or hot treatment in vacuum whereas CVD employs at least two highly reactive gases which generate the coating by high temperature reaction either in the gas phase or on the substrate. The various processes have been considered in detail but some basic considerations are important for the designer.

The whole component whose surface (or part thereof) is to form the substrate must be compatible with the process. Thus, the substrate temperature for PVD will nominally be at ambient temperature or somewhat above, but temperatures in excess of about 500°C are rarely experienced by the part. CVD processes, on the other hand, tend to operate at higher temperatures, generally in the range 500–1200°C, but possibly much higher. The vacuum requirements of PVD, and the high operating temperatures of CVD, demand that the component being treated must not contain materials that have high vapour pressure under the deposition conditions, e.g. brazes containing Cd or Zn. Plastics, however, can be coated under certain conditions. Both CVD and reactive PVD techniques use reactive gases, therefore precautions must be taken against leakage of these gases onto uncoated areas. Articles to be coated must allow free access both for evacuation and for deposition, e.g. recesses may not be easily evacuated, or they may present difficulties during glow discharge cleaning, or they may not be sufficiently accessible to the coating vapours. In general, the design require-
ments that apply to, for example, electroplating, apply equally to PVD and CVD coating.

**Design requirements**

1. The design should not contain deep, blind holes.
2. Small bores are difficult to coat. The component preferably should be designed so that it can be split.
3. Holes or recesses with a width to depth ratio of 1:1 can be coated but a broader ratio is preferable.
4. There should be no surface voids and the component should not be porous or contain fissures, etc.
5. There must be no burrs on the edges of the surface to be coated.
6. Parts must be assembled, e.g. screwed together or shrink fitted.
7. Large variations in cross-section can give rise to temperature variations during the process and should be avoided where possible.
8. Masking must be mechanical and because fit-ups must be tight it entails careful design and probably consultation with the subcontractor.
9. Some means of fixturing may be necessary and should be discussed with the subcontractor.

**Surface condition**

1. The surface of the parts must be bright and untreated.
2. Suitable surface finishes are, for example, ground, polished, fine finish, spark eroded, or spray lapped. Blunt or worn grinding discs must not be used because cold laps must be avoided.
3. Polishing agents should be removed using an appropriate solvent.
4. Parts should be lightly oiled or covered with a sealant as soon as possible after machining or cleaning to protect against corrosion.
5. Parts must be free from machining chips and foreign particles, especially in blind holes.
6. The surface to be coated must be free from any coating or surface treatment such as electroplating, carburising, nitriding, chromating or aluminising.
7. The surface must not be burnished.
8. There must be no rust, paint or colour identification markings.

**Data to be supplied to the processor**

1. Type of material and specification if available.
2. Tempering temperature and any heat treatment to which the part has been subjected.
3. Drawing showing dimensions and indicating clearly the surfaces to be coated, which surfaces may be coated and any surfaces where coating is to be avoided.
4 If polished, details of the polishing process.
5 For machine parts, type of tool or description of its use.
6 Hardness, indicating the point where measured.
7 Identification marks/numbers to be retained.
Chapter 7

**Polymer coatings**

The use of polymeric materials for engineering coatings developed from the adaptation of thermal spraying torches, originally designed for spraying metal powders and patented from the 1920s, to handle polymers which started to become available in powder form in the early 1940s. The useful properties of many polymers have led to the development of thermal spraying equipment designed specifically for spraying them in powder form. In addition, other techniques, including electrostatic spraying and the use of fluidised beds, have been adapted to apply coatings. The various possible combinations of material and application process that have emerged have led to a wide range of uses for engineering and decorative purposes.

The majority of coatings are applied for protection to the surface of metallic substrates, often allowing use of cheaper construction materials. In addition, polymer coatings have been applied to substrates such as concrete for waterproofing and decorative gypsum tiles for improved appearance and waterproofing.

While such coatings have an upper service temperature limit of about 250°C or less, depending upon the polymer used, all have good resistance to corrosion by a wide range of media, are tough and have insulation properties. Hardness and wear resistance depend on the material type.

Polymer coatings are widely used for their corrosion resistance, either as a substitute for cathodic protection by zinc or aluminium, as an overlayer to these to extend life, or to provide a durable and decorative finish by using one of the many colours in which most polymers are available. To give some idea of the range of uses to which a polyethylene-based co-polymer may be used, the manufacturer recommends it for the following:

- Chemical industry – acid tank stands, pickling tanks.
- Paper and pulp handling industry – piping, tanks and mixer paddles.
- Marine environment – exposed steel above the water line.
- Waste water treatment plant – exposed piping, inlet water screens.

When used for protection against corrosion, polymer coatings offer significant benefits over paints in terms of ease of application and durability in service. In addition, polymers can provide superior resistance to abrasion and a wider usable temperature range. Coatings are available which are suitable for use in bearing applications, and to meet the hygiene requirements of handling food stuffs.
Application of polymeric powders to a component requires the presence of sufficient heat to fuse the powder. As this is only a few hundred degrees Celsius, there is little or no problem with metallurgical changes in the substrate and, provided that the metal surface is correctly prepared, adhesion of the coating is excellent.

**Materials**

A selection of materials used in industry is listed in Table 7.1. The majority are thermoplastics and offer a range of properties with an upper service temperature of about 250 °C for polyether ether ketone. For higher temperatures, a thermosetting epoxy resin is used. This type of material requires a post-coating curing treatment unless a curing agent is added to the powder.

The particle size of the polymer powder is important. The size for electrostatic spraying (30–60 μm) is too fine for thermal spraying as the particles would overheat in the flame; fluidised beds and flame spraying use a coarser powder of 80–200 μm.

A common material is polyamide (nylon) which has been in industrial use for about 40 years. Nylon 11 is the most widely used polyamide for coating purposes. In addition to characteristics listed in Table 7.1 it has a well-established use with foodstuffs. The powder is available in a variety of colours.

The ethylene/vinyl acetate co-polymer (EVA) is also widely used for coatings. In addition to the properties listed it has a wide processing temperature range of 180–350 °C and a relatively low cost.

Thermoplastic coatings are suitable for local repair without removal of the original coating.

Use of other polymers for coatings is under development and specific applications are reported from time to time in the technical press. Experimentation needs to be undertaken with caution, as the temperatures involved in applying the coating may lead to the emission of toxic fumes for some polymers.

In addition to being used as straight polymers an improvement in resistance in abrasive wear has been obtained by addition of mineral substances such as glass flake to the polymer. Blends of metallic alloys and polymers are also available – such as Al–Si with polyester, or Al–bronze with polyester.

**Application processes**

**Flame spraying**

Spraying polymers developed in the 1950s using modified equipment originally developed for spraying metals but, as the requirements of the materials are different, guns now in use have been designed specifically to handle polymers.

A build-up of hot polymer inside the gun caused by the flame temperature must be avoided by surrounding the powder jet with cooling air (Fig. 7.1 and 7.2).
<table>
<thead>
<tr>
<th>Material</th>
<th>Melting temperature/°C</th>
<th>Application techniques</th>
<th>Properties</th>
<th>Workpiece preheat temperature/°C</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Thermal spray</td>
<td>Electrostatic spray</td>
<td>Fluidised bed</td>
</tr>
<tr>
<td><strong>Thermoplastics</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ethylene/vinyl acetate copolymer (EVA)</td>
<td>108</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Low density polyethylene</td>
<td>111</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>High density polyethylene</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Polypropylene</td>
<td>120</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Polyamide/polyether copolymer</td>
<td>128</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Macromolecular thermoplastic polyester</td>
<td>174</td>
<td></td>
<td></td>
<td>X</td>
</tr>
<tr>
<td>Polyamide (nylon 11)</td>
<td>186</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Polyether ether ketone</td>
<td>334</td>
<td>X</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Polynvinyl chloride (PVC)</td>
<td>350</td>
<td>X</td>
<td></td>
<td>X</td>
</tr>
<tr>
<td><strong>Thermosets</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Epoxy resins</td>
<td>–</td>
<td>X</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
7.1 Magnum torch for plastics coating, designed for touch-up work on damaged components and for site work. The powder consumable is held in the torch-mounted hopper (courtesy Croboride Ltd).

7.2 Naval exhaust muffler being thermally sprayed with nylon (courtesy Croboride Ltd).

The flame is produced by combustion of a fuel gas such as acetylene, hydrogen or propane in oxygen or air. It is necessary to ensure that the particles of polymer are raised to their melting point but do not suffer any degradation in properties caused by overheating.

Powder is fed to the gun either from a hopper mounted directly on the gun itself or conveyed by air in a hose from a separate powder feeder. The latter
method has advantages for mechanised work, or manual work with long spraying times.

**Advantages**

1. A portable and versatile process, suitable for workshop or site work.
2. Thick coatings over 1 mm are possible.
3. Low capital equipment cost.
4. Rapid changes of powder are possible.
5. Can be mechanised or manually operated.

**Disadvantages**

1. Coating uniformity and quality is dependent on manual operator's skill.
2. Overspray losses may be high on small or narrow workpieces.
3. Surface finish may be inferior to other processes.
4. Line of sight process; access limitations.

Selection of correct spraying parameters and their control is essential to the production of deposits of the correct and repeatable quality.

**Fluidised bed coating**

An early technique consisted of preheating the component to be coated and then rolling or dipping it in polymer powder, causing the powder in contact with the workpiece to melt, building up a coating which, with further heating, developed a smooth, well-bonded, continuous film. However in the 1950s the technique was considerably enhanced by adapting the fluidised bed technique which provided improved coverage especially of irregular surfaces, and improved productivity. The principle is illustrated in Fig. 7.3.

It is suitable for polymers such as nylon, PVC, acrylics, polyethylene, polypropylene, silicones, EVA and polystyrene. Powders used are in the particle size range 1–200 μm, although 20–200 μm is preferred.

**Advantages**

1. Improved uniformity of coverage and coverage of irregular and internal surfaces.
Polymer coatings

2 Little or no waste of powder material.
3 Simultaneous coating of external and internal surfaces.
4 Better temperature control of workpiece is possible as coating is instantaneous all over.
5 Thick coatings possible in a short time.
6 Process is easily mechanised.
7 Environmentally sound, low energy process.

Disadvantages

1 Essentially a workshop process, not portable.
2 Higher capital cost of equipment than thermal spraying.
3 Large volume of powder held in the fluidised bed.
4 Quick colour changes not possible.
5 Masking difficult because of preheat temperature required.
6 Size of workpiece is controlled by size of fluidised bed and preheating oven.

Electrostatic spraying

In this process, the powder delivered from the spray gun is electrostatically charged and propelled at low velocity by air or a revolving spray head to the workpiece which is ‘earthed’. Care is necessary at this stage as too much air pressure causes the powder to bounce off the substrate and makes recesses difficult to coat.

The process produces a coating thickness which is self-limiting as the increasing thickness of the coating increases the electrical insulation of the surface of the substrate. This effect can be overcome by preheating the substrate and applying several coats. After spraying the component is transferred to an oven to cure the coating and to develop a continuous film. For this reason materials with low melt viscosity are best, which flow easily to form a continuous film. Coatings 50–75μm thick are produced on cold objects, and up to 250μm thick on heated objects.

A major advantage of electrostatic spraying is the wrap-around achieved. Spray particles follow the lines of the electric field, even to the reverse side of the object being sprayed. For example, the external surface of a 25mm diameter tube would be completely covered during the spraying process even if the tube were not rotated during spraying and the gun was only traversed along its length. Manufacturers of quality tubular articles such as furniture, bicycles, etc., are major users of the process.

Advantages

1 Good wrap around without workpiece positioning is possible.
2 Process can be easily mechanised.
3 Thin films are possible (50μm).
4 Good edge cover.
Temperature control of oven ensures correct fusing of coating.

Masking is easier than with other processes.

Disadvantages

1. High capital cost of plant.
2. Equipment not portable – a workshop process.
3. Size of workpiece is controlled by the capacity of the equipment.
4. The workpiece must be an electrical conductor.

Other techniques

Small parts

Parts that are too small to be handled individually are heated and dropped into a vibrating or tumbling bed of powder. The parts are later post-heated to produce a pore-free film. Some systems allow for coating without suspension points or touch marks. Up to 50,000 parts per hour are possible using unskilled labour.

Vessel interiors

For components such as fire extinguishers and hot water cylinders, a system exists for introducing a measured quantity of powder into the preheated part, and then moving the part in a planetary fashion to distribute the powder evenly on the inner surface, while heat applied from the outside fuses it in position.

Pipe interiors

To coat interior surfaces of complex parts, low vacuum is applied to pull the powder through the preheated component. Extremely complex pieces are easily coated by this system which has been used for lengths up to 10 m and diameters of 1 m. Heat exchangers have also been coated in a similar manner.

Continuous wire coating

In continuous wire coating, wire is drawn through a fluidised bed of powder and heated in it by an induction coil, thereby picking up a fused coating. A second coil above the powder flows out the coating.

Continuous tube coating

Continuous tube coating is similar to the wire coating process, but the powder is applied in a ‘cloud chamber’ and the hot tube is cooled in water before further processing. Continuous production is possible at high speeds with a closely controlled thickness of coating.
Flock spraying

Components that are too large or heavy to handle in a fluidised bed are heated in an oven and then sprayed by a simple venturi type gun, with or without electrostatic charge. Powder fuses to form coatings up to 1250 μm (0.05 inch) thickness if required, which are then restored if necessary to complete the fusion.

Designing for polymer coatings

Surface preparation

Coatings of polymers on metallic substrates must be applied under carefully controlled conditions. It is essential that the substrate is perfectly clean before any coating operation commences. This means that all traces of scale, rust, grease, paint or other contamination must be removed. Failure to do so will result in loss of adhesion between the metal and the coating.

The cleaned surface is normally blasted using angular iron grit. In addition to the cleaning effect this roughens the surface and provides a mechanical key for the coating. The surface after blasting is extremely active and will oxidise rapidly, especially in damp air, so coating must commence immediately after blasting. Grit used is to SA2½ standard, G17. For thin film coatings where the roughening effect of G17 grit would be too great, use of G7 grit is recommended.

As an alternative to grit blasting, chemical cleaning followed by phosphating or chromating is used, especially for thin coatings obtained by electrostatic deposition. Phosphate coatings are unsuited to the high temperatures used for fluidised bed deposition. Specialist advice is recommended when choosing such a treatment.

Although grit blasting is the best preparation for all thick coatings, use of a primer system before applying the coating is advisable in service applications. These improve the adhesion of the coating to the substrate and are usually recommended for electrostatic applications of nylon. Primers based on epoxy phenolic formulations are applied by conventional spray guns to a thickness of about 7–12 μm as a dry film. Spraying can be carried out with electrostatic assistance if required and will dry fairly quickly. The temperatures involved in thermal spraying may damage the primer and this may preclude their use with this process.

Thin components can easily be distorted by abrasive blasting – chemical cleaning and use of primers can be the answer.

Component design

Surfaces to be coated must be accessible not only for the coating process chosen but also for the surface preparation. Because of the heating required to fuse coatings of polymers it is necessary to provide vents in any hollow sections. Cleaning fluids will enter through these and they should therefore
be sited so that the fluids can be completely drained from the cavities. Subsequent sealing of these vents may be necessary to prevent internal corrosion of the unprotected surface during service.

It is good practice to avoid sharp corners on parts to be sprayed. The coating processes vary in their ability to coat these as uniformly as other surfaces, but the coating may be thinner on sharp corners and will be more vulnerable to damage in handling, especially if the part is large or heavy.

Summary

Given the wide range of industrial wear and corrosion problems, polymeric materials occupy a useful niche in selection of cost-effective coatings available to the design engineer. As with materials and processes covered in other chapters, coatings may be applied in the factory as part of the original design of the component, or in the field. The latter may be to repair a part that was polymer-coated when new, or to extend the life of an uncoated part. Depending on circumstances, manual or mechanised versions of the processes can be adopted. However, strict adherence to proven operating parameters is essential to satisfactory and predictable service life of the part. Where manual application is envisaged, the use of properly trained operators is important.

Points requiring special care are correct design and preparation of the part, control of pre- and post-heating temperatures and avoidance of moisture condensation on parts heated by direct flame.
It will be clear from previous chapters that surface coatings vary widely in their characteristics. Some are used in applications where they can be put into service in the as-coated condition, but many are applied to engineering components that demand a dimensional accuracy and surface finish that cannot be met without a precision finishing operation.

Coatings designed to resist wear are, by definition, likely to be more difficult to machine than ordinary construction materials and therefore call for different techniques. This chapter is devoted to the various finishing processes used for achieving dimensional accuracy and surface finish on coatings applied by weld deposition and thermal spraying. Finishing of electrodeposited coatings was dealt with in Chapter 5, and coatings applied by PVD and CVD in Chapter 6.

The need for machining

Inevitably machining operations performed on hardfacing materials are expensive because of the wear-resistant nature of the alloys, so consideration should be given to the reasons why machining is necessary. The most common requirements for machining involve the need to produce:

- closer dimensional tolerances than the deposited finish can achieve;
- better surface finishes than may be expected from deposition;
- geometric shapes which deposition methods cannot achieve.

The above need to be considered not only when selecting the method of machining but also in the original design of the component. The design engineer will need to know which alloys may be selected to enable dimensional tolerances and surface finishes to be achieved and must be aware of the relative costs of the procedures to be used in achieving those requirements, if cost is a major factor to be considered within the design parameters. As the required degree of precision increases, so does the cost. This is even more apparent when dealing with wear-resistant coatings. The relationship is not a linear one as it costs little, if any, more to make a part within 0.3 mm of its nominal size than to within 0.5 mm. However, it costs
much more to produce a part to within 0.003 mm than to within 0.03 mm of nominal size.

When considering the production economy of machining wear-resistant deposits it is necessary to examine the complete system. Achieving dimensional tolerances and surface finishes depends on a combination of

- the machine tool and its condition;
- the degree of difficulty in machining the material;
- the type of cutting tool required.

It is for these reasons that the cutting parameters suggested should be regarded only as ‘bench marks’ rather than the ultimate achievable.

Before considering machining parameters in detail, time should be given to understanding what may be achieved in terms of dimensional tolerances and surface finishes. These are set out in Fig. 8.1. The relationship between surface finish and relative production times for machining these materials is shown in Fig. 8.2.

**Spray fused and welded coatings**

By their nature, fused and welded hardfacings are of an irregular form and generally need to be machine finished. The purpose of this section is to discuss the various methods used in machining cobalt- and nickel-based alloy deposits, and to set some reference points from which to work, but it should be emphasised that the parameters given are only intended as a guide.
Turning

Most hardfacing alloys below 60RC can be turned. However, the set-up should be rigid so that the tool is presented to the work without any deflection or flutter. Slides should be tight, tool projection at a minimum and the tool section as heavy as practicable. For most applications a negative rake is preferred.

Knowing the condition and capability of each machine tool is essential if standards are to be maintained. Regular monitoring of machining operations by comparison against set standards plays a useful role in avoiding excess costs, while achieving the demanded quality standards. Deterioration of the machine tool will be detected, indicating that attention is required or even that the machine tool is no longer economically viable for a particular operation or standard. In the selection of feeds and speeds the three main considerations are:

- the amount of stock removal;
- the surface finish requirements;
- the dimensional tolerance requirements.

If the turning operation is to finish the diameter in question to size, it is usually necessary to take a roughing cut, followed by a finishing cut to size, taking surface finish requirements into account. The roughing operation would normally be carried out using the combination of feeds, speeds and depths of cut to give the best economic stock removal, but for the finishing cut other considerations must be made. Because these materials are resistant to wear it follows that there is a natural resistance to the turning tool. This has the effect of ‘tool push off’, making it almost impossible to pick up a cut within less than 0.1 mm where a tool may have failed. It is therefore essential, in making the finishing cut, to select a range of speeds and feeds that will guarantee a tool life at least as long as the cut will take, and that at the
end of the cut the tool will still be performing well enough to give the required surface finish. It is this feature of push off that makes final size prediction uncertain and close tolerance work difficult. Where fine feeds are required it normally follows that increases in surface speeds may be used. Similarly where coarser feed rates are advantageous in cutting some of the work-hardening materials then surface speeds should be reduced.

Three types of cutting tip are recommended:

- tungsten carbide in the ISO K10 range;
- ceramic – mixed;
- polycrystalline CBN.

Selection depends upon the grade of material and the soundness or consistency of that material. Typical speeds and feeds that may be used are shown in Table 8.1. Hardfacings with a hardness value of less than 60RC can be easily machined using tungsten carbide tooling as described. However, variations in both chemical composition and hardness values make machining with ceramics or polycrystalline CBN less predictable. Interrupted cuts, such as may occur in uneven coatings or with imperfections in the overlayed material, can cause catastrophic failure of the ceramic cutting tool. Other variations as yet more difficult to understand, but possibly an effect of base material dilution, tend to affect the machining parameters when using polycrystalline CBN. Where good results have been obtained, attention to detail in the deposition and heat treatment operations to give repeatability at the subsequent machining operations is essential. Many examples are available to indicate that most nickel- and cobalt-based deposits can be machined using polycrystalline CBN or ceramic tips but it is advisable to record successes for future reference.

**Milling**

Milling is restricted to the softer hardfacing alloys, and the cutting tools normally used are tungsten carbide. The nature of the milling operation is one of a series of interrupted cuts and ceramic tips cannot usually withstand
these forces generated in the milling of wear-resistant coatings, while polycrystalline CBN tools are not in contact with the workpiece consistently enough to achieve the desired cutting parameters. There will be, however, instances where both ceramics and polycrystalline CBN tools are effective. For the purpose of this chapter, only tungsten carbide tools are considered.

The three areas of machining where milling is an effective operation are:

- large flat areas;
- machining angles;
- machining grooves (i.e. keyways and grease grooves).

A vertical milling machine is considered to be advisable because of the rigid set-ups required to achieve the best results. Milling cutters should be selected which use indexable inserts of the negative geometric form (SNUN, TNUN, RNUN, etc) as this ensures maximum strength at the cutting edge. The size of the cutter selected should be such that the maximum length of cutting stroke is achieved, the best results being obtained in cutting a groove in the material (180° arc of contact). Surface speed at the cutting tip should be approximately 60% of that used for turning, with feed rates of around 0.05 mm per tip per revolution. Depth of cut should be no more than 3 mm, although this may be increased for some of the softer alloys. The wear pattern, flank or crater on the insert will indicate where the rpm or feed rates should be adjusted. When machining grooves or angles, roughing and finishing operations should normally be used if stepping or breakout is to be avoided on the sides of the grooves or along the angles being machined.

For example, for face milling a block 180 mm wide overlaid with cobalt-based deposit, hardness 36RC, use a milling cutter 200 mm diameter negative geometry with 14 inserts, cutter speed 20 rpm, depth of cut 2 mm, feed rate 14 mm min⁻¹.

**Drilling**

Production drilling of holes in most hardfacing alloys presents few problems. If close dimensional tolerances are required then the set-up must be rigid. Breakout can occur if drilling through holes, but this may be overcome by using progressively larger drills, and finally drilling into the deposit from either side.

The least expensive way to drill holes of up to 20 mm diameter in hardfacing materials is by using standard masonry drills. These usually cut well oversize and the clearance helps to avoid overheating. Speeds of around 400 rpm with a feed rate of 0.01 mm rev⁻¹ are usual when drilling holes of up to 7 mm diameter, while for holes between 8 and 20 mm diameter, speeds of around 170 rpm are usual.

It must be remembered that usually it is only necessary to drill through the deposit into the base material using the techniques described above. From the point where the drill meets the base material, conventional drilling techniques with regard to that material should be used. Coolant should always be used as it helps to avoid cracking and gives greater life to the cutting edge.
Grinding

All fused and welded hardface deposits can be ground to give a surface finish of 0.4 µm Ra. For small batch production, wheels normally used for grinding high speed steel can be employed, provided they can be kept open and free cutting and that a copious supply of coolant is used.

Cylindrical grinding

Before finish grinding it is advisable to turn the deposit to within 0.4–0.6 mm of finished size if possible. For most grinding operations aluminium oxide grit wheels in the range A60J with vitrified bonds can be used. An approximate peripheral speed of around 25 m s⁻¹ with a workspeed of around 15 m min⁻¹ is a good starting point, but with some of the harder deposits it may be necessary to reduce the wheel speed if possible, or, if this is not possible, the work speed to minimise the possibility of cracking.

Wheels using CBN normally give greater stock removal rates when grinding harder deposits but precise bands of feeds and speed will need to be selected to give the best surface finishes. For internal cylindrical grinding the same parameters are generally used but the rigidity of the grinding quill has an important part to play in achieving surface finish and good stock removal.

Surface grinding

Most surface grinding of hardfaced deposits is carried out on vertical spindle, rotary table grinding machines, using ring wheels or segmental wheels if economic stock removal is to be achieved.

Selection of wheel types depends on

- the surface area of deposit to be ground;
- the type of hardfacing alloy;
- if steel and deposit are being ground together.

Selection guidelines are as seen in Table 8.2. As with other types of grinding copious amounts of coolant are required.

Using conventional surface grinding techniques (horizontal spindle reciprocating table machines) wheels in the A60J to A120K range are recommended but advantages in stock removal rates may be found on the harder alloys by using CBN or diamond grinding wheels.

Honing

Honing is recommended as a reasonably fast and accurate alternative to grinding when finishing deposited bores, where either dimensional tolerances are close, or where a fine surface finish is required. Because most of the bores deposited will be more than 75 mm diameter, the type of honing machine used will normally be either the beam hone or large vertical hone.

The component should have been machined to within 0.1–0.2 mm of finished size. Experience suggests that CBN stones give best results where
Table 8.2 General recommendations for grinding wheels for welded coatings of cobalt base alloys of Group 4

<table>
<thead>
<tr>
<th>Purpose</th>
<th>Approximate peripheral speed/m s⁻¹</th>
<th>Recommended wheels</th>
</tr>
</thead>
<tbody>
<tr>
<td>Peripheral wheels</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Surface grinding</td>
<td>27.5</td>
<td>A60/V</td>
</tr>
<tr>
<td>Form grinding (use finer grits for sharper forms)</td>
<td>27.5</td>
<td>A100GV A120KV A150GV</td>
</tr>
<tr>
<td>Ring wheels</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mixed surfacing alloy and steel</td>
<td>19</td>
<td>A30DZB</td>
</tr>
<tr>
<td>Surfacing alloy only</td>
<td></td>
<td>A50BB</td>
</tr>
<tr>
<td>Segmental wheels</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mixed surfacing alloy and steel</td>
<td>19</td>
<td>A36FB</td>
</tr>
<tr>
<td>Surfacing alloy only</td>
<td></td>
<td>A46EB</td>
</tr>
<tr>
<td>Internal grinding</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bores over about 100 mm diameter</td>
<td>25</td>
<td>A54/V A54MV</td>
</tr>
<tr>
<td>External grinding</td>
<td></td>
<td></td>
</tr>
<tr>
<td>For wheels up to 305 mm</td>
<td>27.5</td>
<td>A60/V</td>
</tr>
<tr>
<td>For wheels over 305 mm</td>
<td></td>
<td>A46/V</td>
</tr>
</tbody>
</table>

Wheels carrying the same marking from different manufacturers do not necessarily have the same cutting action. Check that the maker’s maximum safe speed is not exceeded. The wheel descriptions above are based on the recommendations of ANSI B74.13.1970 and BS4491.

Surface finish requirements are no better than 0.1 µm Ra, but for general use stones in F1F3D (Delapena) give satisfactory performance.

**Lapping**

Lapping is an operation where the complete combination of machine, material and type of abrasive needs to be considered as a total system. Coarse diamond or silicon grit, together with pressure, gives best stock removal rates if surface finish requirements are not too fine. Fine diamond powder (3–5 µm) or aluminium oxide powder (520 grit) is necessary, however, if surface finish requirements are better than 0.2 µm Ra. Very fine surface finishes take time to achieve, therefore a roughing operation can be advantageous before fine finishing. However, two machines will be required, since to change a machine from coarse to fine grit requires a complete wash down.

**Spark erosion**

Little needs to be said about machining by spark erosion, but it should be stated that, as with most other processes, quality of finish versus production time is the main consideration. The operation itself is one that causes few problems whether using the wire erosion technique or the more conventional copper or carbon electrode method. Imperfections in the deposit,
however, when using wire eroding can have a dramatic effect on the quality and may cause wire wander.

**Thermally sprayed coatings**

For turning softer coatings, established tooling, such as high speed steels and sintered carbides, gives the best results. Often the biggest problem is to ensure that porosity in the deposit is not sealed by surface deformation.

Successful machining starts at the design stage where adequate support must be given to the ends of the deposit and allowance made for elimination of any ragged edge from which cracking may propagate. Correct substrate preparation must be employed with, if necessary, use of bond coats and/or threading to improve coating adhesion. Selection of material, too, is important. It may, for example, be advisable to spray a thin coating of a softer alloy on to a hard deposit to facilitate final machining. It is important to select the surfacing material with consideration of the machining equipment available.

While a very low carbon steel provides good wear properties under lubricated conditions, increasing the hardness, which may be essential, decreases machinability and requires better tool tips. As the thickness of the deposit increases it becomes more desirable to select materials with low shrink characteristics, to reduce internal stresses which can cause cracking when the additional machining stresses are superimposed. Various bronze alloys are in regular use. If a low cost commercial bronze is satisfactory this may be used but it will not machine to as good a finish as an aluminium bronze. The latter shows excellent cohesion to the substrate, a coherent dense structure and very good wear resistance. A good finish is more difficult with sprayed nickel than with sprayed Monel although the resistance needed to some corrosants may dictate the choice of nickel.

Selection of spraying process may affect machinability but whatever the process, it is important that it is carried out correctly. Lack of control may lead to abnormal particle size, oxidation or other chemical effects, excessive porosity or loss of cohesive strength, any of which can cause deterioration in the quality of the machined surface and possibly porosity or cracking.

Sufficient working data have been established for machining and grinding of most commercially developed materials used for metal spraying. Some provide better finishes than others but commercial finishes within commercial tolerances can be achieved with all. However, the machinist should not use a tool tip tailored for cutting cast or wrought metal nor should the grinder select wheel grades or grit sizes normal for these materials.

**Turning**

All but the hardest deposits can be machined with high speed tooling but faster cutting rates are obtainable using cemented carbide tool tips as indicated in Table 8.3. Such tooling is not normally used for softer materials. Carbide tips provide better finishes as higher speeds may be used and some of the harder alloys, which otherwise require grinding, may be machined easily. The low coefficient of friction and high resistance to welding enables
# Table 8.3 Typical basic turning parameters for sprayed coatings

<table>
<thead>
<tr>
<th>Metal</th>
<th>High speed steel tools</th>
<th>Sintered carbide tools</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Surface speed/m min⁻¹</td>
<td>Feed/μm rev⁻¹</td>
</tr>
<tr>
<td></td>
<td>Roughing</td>
<td>Finishing</td>
</tr>
<tr>
<td></td>
<td>Roughing</td>
<td>Finishing</td>
</tr>
<tr>
<td>Pb, Sn, Babbitts</td>
<td>45–75</td>
<td>125–175</td>
</tr>
<tr>
<td>Al, Cu, brasses, bronzes</td>
<td>30–40</td>
<td>75–125</td>
</tr>
<tr>
<td>Nickel alloys</td>
<td>30–40</td>
<td>75–125</td>
</tr>
<tr>
<td>Low carbon steels</td>
<td>23–30</td>
<td>75–125</td>
</tr>
<tr>
<td>Medium carbon steels</td>
<td>15–23</td>
<td>75–125</td>
</tr>
<tr>
<td>High carbon steels</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Martensitic stainless steel</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Austenitic stainless steel</td>
<td>30–40</td>
<td>75–125</td>
</tr>
</tbody>
</table>

Carbide to cut freely with much less rake and clearance than is needed for ordinary tool steel, again assisting in providing a better finish. The oxides and carbides frequently encountered in sprayed coatings subject the nose of the cutting tool to much abrasive wear and the hardness and wear resistance of carbides again provides an advantage over ordinary tool steels. With harder deposits, use of the latter will lead to problems in maintaining dimensions.

Sprayed metal coatings are porous. For ordinary build-up work on machine components used in lubricated conditions, development of self-lubrication is advantageous as the oil tends to seal the pores. Thick coatings are relatively impervious but with thin coatings or parts used in active corrosion environments or under high pressure, such as hydraulic pressure rams, sealing is necessary. There can be advantages in sealing shafts and other machine parts even if they are not used under high pressure or in corrosive environments. Self-sealing coatings of tin, copper, aluminium or 18/8 stainless steel may be sealed by using high pressures and blunt tools, by rolling, or by peening. Such methods may be used to control surface finish and, at times, dimensional tolerances.

## Grinding

Grinding is often the preferred finishing method for the harder surfacing materials and for pure metals such as Cr, Nb, Ta and W. It is quite practical provided that robust equipment is employed, with carefully dressed wheels. A grinder that is too light for the work, or a badly dressed wheel causing vibration, will damage the coating and produce a poor finish.

The grinding wheels for sprayed coatings should be chosen to suit the coating. A wheel of coarse grained open structure and low bond strength may be suitable for a soft coating, but a wheel of denser, finer grain size is more suitable for the denser, harder coatings. Silicon carbide is frequently used except for deposits containing high concentrations of tungsten carbide.
Table 8.4 Typical wet grinding parameters for sprayed coatings

<table>
<thead>
<tr>
<th>Process</th>
<th>For all processes</th>
<th>Centreless grinding</th>
<th>Internal grinding</th>
<th>Cylindrical grinding</th>
<th>Surfacing grinding</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wheel speed</td>
<td>1650–2000 smpm*</td>
<td>Regulating wheel-angle°, R 2–4; F 1–2; rpm R 15–30; F 20–40</td>
<td>Table speed m/min 1 1.25–1.8</td>
<td>Fraction of wheel width advance per rev. R (\frac{1}{2} )–(\frac{1}{2} ); F (\frac{1}{2} )–(\frac{1}{2} )</td>
<td>Table speed m/min 1 12–30; Cross feed, mm R 0.15–0.30; F 0.04–0.08</td>
</tr>
<tr>
<td>Work speed</td>
<td>smpm</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Regulating wheel-angle°</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Table speed m/min</td>
<td>1 1.25–1.8</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fraction of wheel width advance per rev.</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>R (roughing)</td>
<td>F (finishing)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>12–30</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cross feed, mm</td>
<td>R 0.15–0.30; F 0.04–0.08</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*smpm = surface metres min⁻¹.
R = roughing, F = finishing.

General-purpose wheels approximate to the following specification: grit size 60; hardness medium soft; structure open; bond vitrified; and a silicon carbide wheel would have a typical BSI marking C60-K10-V, to which the manufacturer's prefix and suffix may be added.

The wheel specification may be changed depending on the particular job. Thus a softer wheel should be used if there is a large area of contact, if faster stock removal is required or if a wide wheel is to be employed with light pressure. The hardness should be increased to provide a finer finish, or if the area of contact is small and a narrow wheel is used with high pressure. As sprayed metals tend to 'load' the wheel, a relatively coarse grain and a low bond strength are needed so that the surface breaks down with comparative ease. A dull or glazed wheel or use of an unsuitable abrasive can exert a high shearing force, generating much local overheating which is aggravated by the comparatively poor thermal conductivity of the coating, leading to burning and heat checking.

Wet grinding is preferable to dry grinding and should be used whenever suitable equipment is available. It presents few problems provided the right wheels are used and the following precautions observed: always use light pressures; avoid glazing of wheels; use finer grits for finer finishes; use copious quantities of coolant (see Table 8.4). In dry grinding use substantial equipment to prevent excessive wheel vibrations (see Table 8.5).

The coating should always be sealed to ensure a better and cleaner initial ground finish. The sealer should be applied before grinding to prevent debris from the grinding operation entering the pores of the coating, and the smoother surface makes it much easier to wash clean afterwards. Removal of all the debris may be a critical factor in the life of a bearing used with a metal sprayed shaft.

**Other machining methods**

**Planing and shaping**

Sprayed coatings on flat surfaces should be finished on a shaper or planer using tools of similar geometry to those employed for turning. Speed and
Finishing of surface coatings applied by welding and thermal spraying

Table 8.5 Typical dry grinding parameters for sprayed coatings

<table>
<thead>
<tr>
<th></th>
<th>Wheel speed/ smpm</th>
<th>Work speed/ smpm</th>
<th>Other</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cylindrical grinding</td>
<td>1650-2000</td>
<td>15-21</td>
<td>Wheel advance*</td>
</tr>
<tr>
<td></td>
<td></td>
<td>21-30</td>
<td>R $\frac{1}{2}$, F $\frac{1}{2}$</td>
</tr>
<tr>
<td>Surface grinding</td>
<td>1650-2000</td>
<td>12-30*</td>
<td>Cross-feed, mm</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>R 1.6-3.2, F 0.4</td>
</tr>
<tr>
<td>Tool post-grinding</td>
<td></td>
<td></td>
<td>Traverse/mm rev$^{-1}$</td>
</tr>
<tr>
<td>Cu, bronze, carbon steels</td>
<td>1500-1800</td>
<td>9-11</td>
<td>R 0.8, F 3.0</td>
</tr>
<tr>
<td>Ni alloys</td>
<td>1500-1800</td>
<td>9-11</td>
<td>R 0.15, F 0.4</td>
</tr>
<tr>
<td>Martensitic stainless steel</td>
<td>1500-1800</td>
<td>33-38</td>
<td>R 0.15, F 0.4</td>
</tr>
<tr>
<td>Austenitic stainless steel</td>
<td>1500-1800</td>
<td>9-11</td>
<td>R 0.8, F 1.6</td>
</tr>
</tbody>
</table>

* Table not work speed.

**Fraction of wheel width per revolution of work.

Feeds should be related to turning practice. It is necessary to remove raised sections or overlapping flash by grinding, using light cuts, before machining commences.

Milling

A milling cutter rotating in a direction opposite to the traverse or work feed is cutting in an upward direction and care is needed to avoid lifting the coating from the substrate. The problem is aggravated in milling keyways and splines. Milling can be performed on flat surfaces provided raised or uneven areas are removed first and that very light cuts are taken. Cutting speeds are generally in the range 20-30 m/min$^{-1}$ but can be greatly increased for some materials, such as copper alloys. Feed advance is generally 0.2-0.5 mm rev$^{-1}$ for roughing and should be reduced according to the finish required.

Linishing

Linishing is often applied where smooth surfaces are desired but tolerances are not critical.

Polishing

Normal equipment and procedures are used. It must be appreciated that the treatment is being carried out on a metallic veneer, only mechanically bonded to the substrate. The softer the metal the better the polish, so tin, zinc, Babbitt and aluminium are easy to polish, the harder ‘red and yellow’ alloys are a little more difficult, while the even harder steels, Monel and nickel alloys present great difficulty.
Ensuring quality in surfacing

The control of quality in the manufacture of engineering components has always been important but, in recent years, this importance has become more widely recognised. There are several reasons for this emphasis on quality: components must often withstand more exacting service conditions, they must be produced to more demanding economic standards and they are affected by environmental and legislative factors.

The control of quality of a surface coating is no different in principle from the control of other engineering products but, in practice, coatings require specialised techniques. It is the aim of this chapter to consider these and to outline the main factors that should be taken into account to ensure quality of surfaced components. Examples of applications of the principles are given, drawn from weld surfacing and thermal spraying, which are the most widely applied coating processes for the reduction of wear and corrosion.

The quality plan

The deposition of a surface coating involves a series of interrelated stages. To achieve quality repetitively it is essential for each and every stage of this processing cycle to be controlled, with standards clearly defined and rigorously followed. Surface coatings, by their nature, pose problems in testing, and consequently good quality must be built into the deposition process, rather than attempting to test out poor quality after fabrication has been completed. This emphasis on good housekeeping throughout the processing cycle is the keynote to quality assurance in surfaced parts.

The sequence of operations needed for any particular job should be clearly laid down at the outset to form the 'quality plan'. A specification should be placed upon each operation in the process so that it may be carried out effectively and reproducibly.

The elements of a typical quality plan are listed in Table 9.1, and the individual items are discussed below. The preparation of the plan should involve consideration between all interested parties. Ideally the designer, processor and user should be consulted; only if this is done can the optimum results be expected. The plan should be appropriate to the component and its service requirements. It need not – in fact it generally will not – aim at perfection, which is rarely obtained and seldom justified. Rather, it should take account of the types of defect that are likely to occur in process-
Table 9.1  The quality plan

| Design | Can the component be redesigned to reduce the severity of environmental effects and also to make it more suitable for the proposed surfacing technique? |
| Component | Is the composition/structure suitable for the proposed surfacing operation?  
Can the part withstand the necessary preparation?  
Is the component surface free from defects? |
| Surfacing choice | What is the most suitable coating and deposition process taking into account:  
(a) service requirements;  
(b) economic criteria;  
(c) equipment and expertise available;  
(d) location of job, etc? |
| Surfacing procedure | Process parameters needed  
Control needed (manual, automated, operator skills)  
Pretreatment required  
Post-treatment required  
Finishing required |
| Quality assurance | Quality required for the application  
Defects that may be encountered in the chosen surfacing process  
Agreed tests to be used, and acceptance standards  
Tolerance to defects and permitted rectification  
Operator training and qualification requirements |

The above list is typical of the factors that should be considered in setting up the quality plan for a surfacing job; the list is not comprehensive and may contain more (or fewer) items, depending upon the application.

...ing and should evaluate the extent to which such defects can be tolerated for the application. This is particularly so when repairing existing components not previously surface coated.

**Design**

There are certain design principles for parts to be coated and, if these are followed, many potential problems can be avoided. Conversely, incorrect design adds to the difficulties of coating a component and in the extreme may preclude it entirely. Thus, incorrect blank dimensions can affect the finished thickness of the deposit and its uniformity and concentricity; these in turn can affect wear resistance and thermal stability in use. Sharp corners and recesses can lead to unsound deposits and, in weld deposition, to excessive local dilution of deposit material by the substrate.

These comments assume that design changes can be made. Although this should be so with original equipment, it may not be feasible to implement modification when surfacing it to be used for repair and reclamation; a compromise on design features is then inevitable. However, even minor modifications, e.g. chamfering sharp corners, should be considered, as they can have a disproportionate benefit in subsequent processing. Coating design is dealt with in detail in the chapters covering the various processes.
Component/substrate requirements

The surface to be coated must be suitable for the proposed coating and deposition method, and it must be appropriately treated before surfacing begins.

The composition of the component, and its metallurgical condition, have a direct and important bearing on the success of the finished job. The composition, especially with weld deposition must be chemically compatible with the proposed deposit materials. It is desirable, therefore, to know the composition of the component, either by a suppliers’ certificate of analysis or an in-house analysis. This requirement is perhaps rather less important with coatings that do not involve substrate fusion, but it may be important to check the metallurgical condition of the substrate to ensure that it is suitable, e.g. if it is too hard it may not be possible to achieve effective grit blasting in preparation for thermal spraying. The structural condition can be checked by simple metallographic examination, supported by hardness tests.

High quality deposition can be obtained only if the component surface is free from defects: porosity, cracks or inclusions (in or near the surface layers) lead to poor quality coatings. Soundness must therefore be checked carefully, possibly by dye penetrant or magnetic particle testing, or by radiography or ultrasonic inspection.

The substrate must be able to withstand any preparation for coating, as well as the conditions during deposition. Thus, where a spray coating system requires a clean, rough surface it will involve the removal of paint and loose rust by pickling, scraping or wire brushing, vapour degreasing and then grit blasting the components. To minimise blasting time and ensure adequate adhesion, grit is carried at high gas pressures (∼0.6 MPa) and the part must not distort under this treatment. For delicate components lower pressures may have to be used, but this may adversely affect coating adhesion.

The ability of the component to resist reasonable heat input may also be necessary, particularly if surfacing by one of the ‘hot’ processes. A degree of heat resistance may, in any event, be desirable, as heating of the substrate is often practised in ‘cold’ deposition processes to improve adhesion. Finally large differences in the coefficient of thermal expansion between component and surfacing material should, if possible, be avoided where problems from contraction stresses on cooling may arise.

Consumables for welding and thermal spraying

The material to be applied may be in the form of rod, wire, cord or powder, depending upon the process selected. In general, appropriate standards should be placed on composition and physical characteristics, e.g. rod diameter and powder particle size. It is necessary to ensure that any description used is adequate and unique to the material required: makers’ brand names and trade names may cover a range of somewhat different alloys. Most suppliers will supply a guarantee of conformity with (or a certificate of) analysis and this should be obtained; failing this the feed materials
Ensuring quality in surfacing

**Table 9.2** Methods for evaluation of thermal spraying powders

<table>
<thead>
<tr>
<th>Property</th>
<th>Method</th>
</tr>
</thead>
<tbody>
<tr>
<td>Powder sampling</td>
<td>Dividing and quartering powder</td>
</tr>
<tr>
<td>Apparent powder density</td>
<td>Weight of powder to fill Hall flowmeter</td>
</tr>
<tr>
<td>Powder flow rate</td>
<td>Time for powder to flow through Hall flowmeter orifice</td>
</tr>
<tr>
<td>Particle size analysis:</td>
<td></td>
</tr>
<tr>
<td>Sieve analysis</td>
<td>Mechanical sieving (sieves to BS410)</td>
</tr>
<tr>
<td>Subsieve analysis</td>
<td>Coulter counter technique. Sedimentation, elutriation, etc.</td>
</tr>
<tr>
<td>Particle shape</td>
<td>Low powered microscopy</td>
</tr>
<tr>
<td>Powder surface area</td>
<td>Nitrogen adsorption</td>
</tr>
<tr>
<td>BET measurement</td>
<td>Air permeability. Calculation for size distribution</td>
</tr>
<tr>
<td>Others</td>
<td></td>
</tr>
<tr>
<td>Chemical analysis</td>
<td>Standard tests</td>
</tr>
<tr>
<td></td>
<td>Hydrogen loss</td>
</tr>
</tbody>
</table>

should be checked in-house. In any event, the material must be clearly marked before it is issued for use on the shop floor.

Where the coating material is presented as a powder it is necessary to make sure that it will feed properly; this entails detailed evaluation of the powder. Chemical analysis, powder shape, size and distribution, apparent density and flow rate are measured and structure and cleanliness may also be assessed. Powder tests such as those listed in Table 9.2 are suitable.

These feed materials must be kept in suitable conditions while awaiting use. In particular, care must be taken to keep them dry; this is commonly achieved by storing in a heated cupboard.

Strict specifications must be placed on the gases that sustain the heating processes and transport the coating material. This is necessary not only for good deposit quality, but is often essential to ensure low maintenance costs for the equipment.

Grit blasting the surface before thermal spraying requires use of clean, fresh and sharp grit, with a minimum of fines. It must have sufficient strength to prevent premature fracture, so impairing the effectiveness of the blasting operation. Equally, it should be low in impurities that can be transferred to the component surface and interfere with the spraying operation. The air used must be clean, dry and oil-free; this is best achieved by an oil-free air compressor. After grit blasting, the prepared surface should be ‘blown off’ with dry, oil-free air, and should be sprayed as soon as practicable. The maximum length of time that can be tolerated before spraying varies according to circumstances but, in good workshop practice, about four hours would be considered the maximum delay.

**Surfacing practice**

Successful surfacing depends on the availability of suitable equipment, specification of appropriate operating parameters, and the use of a trained
and qualified operator. It is necessary to use reliable equipment, fitted with an adequate range of calibrated controls and instrumentation, properly maintained and with all calibrations regularly checked and recorded; also conforming to proper safety standards. Without these controls there can be little control of product quality.

An aspect of the growing use of mechanised surfacing methods is greater uniformity of coating quality. This is accompanied by economy in surfacing consumables and a reduction in subsequent finishing operations.

The parameters for the actual operation must be carefully specified to ensure efficient deposition, acceptable throughput and quality of coating. However, the processing conditions which give the highest efficiency or greatest throughput may not give the desired coating characteristics (or may not be acceptable in the substrate). A degree of compromise may thus be necessary in the choice of operating parameters for a particular job.

Heating coils and cooling jets may be used to control the thermal flux to the component. The specification may call for preheating before deposition, control of the interpass temperature, or heating after deposition. With all these operations suitable pyrometric equipment should be made available to monitor and record the temperatures.

Acceptable and uniform results from manual deposition are dependent upon operator skill. Proper training in the process is therefore essential, and there should be periodic requalification tests to ensure that skills are maintained.

In summary, it may be said that good practice in the actual process of deposition is the key to production of satisfactory coatings. A reliable, trained and resourceful operator can eliminate many variables in the process but there are additional requirements if reproducible quality is to be obtained: suitable calibrated equipment, specification (and following) of correct process parameters and maintenance of accurate and detailed records.

**Finishing**

The finishing operation can play a vital role in the quality of the coating (and on the economics of production). Heat treatment and stress relieving operations may be necessary and these must be accurately specified, controlled and recorded. Surface machining, grinding or polishing of the deposit may be necessary to meet tolerances or produce smooth finishes. If the deposit is very hard, diamond grinding may be required. Care must be taken to lay down specifications on the finishing consumables which will ensure that the grades of material, e.g. diamond or lapping compounds, are suitable for the operation.

Should it be necessary to seal porosity in a thermally sprayed coating, this would normally be carried out with varnish, epoxy resin or metal infiltration. The actual method of conducting the sealing can be vital to its success which, in turn, calls for use of a clearly defined and properly controlled procedure by trained personnel.
Table 9.3 Typical checklist for processing parameters (weld deposition)

<table>
<thead>
<tr>
<th>Component</th>
<th>Identification/description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Work required</td>
<td>Brief description</td>
</tr>
<tr>
<td></td>
<td>Finishing requirements</td>
</tr>
<tr>
<td>Substrate</td>
<td>Material</td>
</tr>
<tr>
<td></td>
<td>Specification</td>
</tr>
<tr>
<td></td>
<td>Condition</td>
</tr>
<tr>
<td></td>
<td>Preparation required</td>
</tr>
<tr>
<td>Preheat treatment</td>
<td>Preheat temperature/°C, minimum and maximum</td>
</tr>
<tr>
<td></td>
<td>Interpass temperature/°C, minimum and maximum</td>
</tr>
<tr>
<td>Description</td>
<td>Process</td>
</tr>
<tr>
<td></td>
<td>Consumables: rod or wire size/mm</td>
</tr>
<tr>
<td></td>
<td>Thickness: mm</td>
</tr>
<tr>
<td></td>
<td>number of runs</td>
</tr>
<tr>
<td></td>
<td>Welding parameters:</td>
</tr>
<tr>
<td></td>
<td>Electrode voltage, DC</td>
</tr>
<tr>
<td></td>
<td>Electrode polarity</td>
</tr>
<tr>
<td></td>
<td>Current/A</td>
</tr>
<tr>
<td></td>
<td>Voltage/V</td>
</tr>
<tr>
<td></td>
<td>Speed/mm.s⁻¹</td>
</tr>
<tr>
<td></td>
<td>Electrode angle</td>
</tr>
<tr>
<td></td>
<td>Weaving parameters</td>
</tr>
<tr>
<td>Post-heat treatment</td>
<td>Heat treatment temperature/°C</td>
</tr>
<tr>
<td></td>
<td>Heating rate/°C⁻¹</td>
</tr>
<tr>
<td></td>
<td>Soaking time/h</td>
</tr>
<tr>
<td></td>
<td>Cooling rate/°C⁻¹</td>
</tr>
<tr>
<td>Inspection and testing</td>
<td>Purpose</td>
</tr>
<tr>
<td></td>
<td>Methods/tests</td>
</tr>
<tr>
<td></td>
<td>Acceptance level</td>
</tr>
<tr>
<td>Special instructions</td>
<td>Detail (e.g. operator qualifications, equipment calibration)</td>
</tr>
</tbody>
</table>

The above list is not intended to be comprehensive but simply to illustrate the type of items to be included. A similar checklist, suitably modified, would be used for thermal spraying operations.

Checklist of operations

To ensure that all necessary information and instructions are recorded it is desirable to prepare a checklist to summarise requirements for any surfacing job; see Table 9.3.

Testing and inspection of coated components

The aim of testing surface coatings should be to ensure that the operation has been correctly carried out, and not to ‘test out’ poor components. This distinction may seem trivial but it implies a fundamental difference in approach.

There are many criteria by which coatings can be judged (Table 9.4) and many techniques are available to assess them (Table 9.5). Some of these can be applied without damaging the component, but others cannot; furthermore, responses to many of the techniques will be affected by the substrate.
Table 9.4 Potential criteria for engineering coatings

<table>
<thead>
<tr>
<th>Adhesion to substrate</th>
<th>Electrical properties</th>
<th>Structure</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chemical composition</td>
<td>Evenness/uniformity</td>
<td>Surface finish</td>
</tr>
<tr>
<td>Chemical reactivity</td>
<td>Frictional properties</td>
<td>Thermal properties</td>
</tr>
<tr>
<td>Cohesive strength</td>
<td>Hardness</td>
<td>Thickness</td>
</tr>
<tr>
<td>Covering power</td>
<td>Machinability</td>
<td>Wear resistance</td>
</tr>
<tr>
<td>Edge definition</td>
<td>Porosity</td>
<td></td>
</tr>
</tbody>
</table>

Table 9.5 Typical tests applicable to coatings

<table>
<thead>
<tr>
<th>Generally non-destructive</th>
<th>Generally destructive</th>
</tr>
</thead>
<tbody>
<tr>
<td>Visual examination</td>
<td>Bend testing</td>
</tr>
<tr>
<td>Dimensional checking</td>
<td>Adhesion testing</td>
</tr>
<tr>
<td>Dye penetrant testing</td>
<td>Peel testing</td>
</tr>
<tr>
<td>Magnetic crack detection</td>
<td>Proof machining</td>
</tr>
<tr>
<td>Eddy current testing</td>
<td>Chisel testing</td>
</tr>
<tr>
<td>Acoustic methods</td>
<td>Chemical analysis</td>
</tr>
<tr>
<td>Radiography</td>
<td>Metallography</td>
</tr>
<tr>
<td>Hardness testing</td>
<td>Porosity measurement</td>
</tr>
<tr>
<td>Ultrasonic testing</td>
<td></td>
</tr>
</tbody>
</table>

For any particular application only some of the criteria will be relevant. What is essential is that those involved – essentially the coater and the customer – decide the criteria to be used to judge any particular coating and agreement should be reached before the work starts.

The result of testing procedure can vary according to the training and skill of the operator and the techniques used. It is therefore desirable not only to define the test conditions precisely, but also to audit the procedure (and the operator) regularly. Different techniques produce deposits with different characteristics; this calls for differences in the detailed test procedure. This is illustrated below by describing testing of weld-deposited and sprayed coatings. These represent two distinct types of deposit: the former are fully dense and integral with the substrate but the latter can contain some porosity and may not be metallurgically bonded to the substrate.

Weld-deposited coatings

If deposition is manually controlled the welder will be responsible for initial checks that the correct amount of deposit has been applied to the correct place: this requires simple measuring devices, such as a caliper, rule and micrometer. That these are simple instruments highlights one general feature of good inspection: the tool or the instrument selected should be appropriate to both test and coating.

The inspector normally examines a welded deposit in the following sequence:
Table 9.6 Possible defects in weld-surfaced and thermally sprayed coatings.

<table>
<thead>
<tr>
<th>Defect</th>
<th>Characteristic</th>
<th>Possible causes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weld-deposited coatings</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cracks</td>
<td>Contraction</td>
<td>Low ductility or low tensile strength in coating material</td>
</tr>
<tr>
<td></td>
<td>Fine surface cracks</td>
<td>Unsuitable grinding wheel or insufficient cooling during grinding</td>
</tr>
<tr>
<td></td>
<td>Lifting at the interface</td>
<td>Poor welding technique with failure to disrupt oxide film</td>
</tr>
<tr>
<td></td>
<td>Lifting within substrate</td>
<td>Embrittlement of substrate during welding</td>
</tr>
<tr>
<td>Porosity</td>
<td>Associated with final molten pool</td>
<td>Faulty crater filling</td>
</tr>
<tr>
<td></td>
<td>Randomly dispersed</td>
<td>Flux contamination, faulty flame adjustment (gas welding), damp</td>
</tr>
<tr>
<td></td>
<td></td>
<td>electrode coverings (arc welding), etc.</td>
</tr>
<tr>
<td>Inclusions</td>
<td></td>
<td>Incorrect welding conditions or use of dirty substrate material</td>
</tr>
<tr>
<td>Thermally sprayed coatings</td>
<td>Excess porosity</td>
<td>Poor spraying practice</td>
</tr>
<tr>
<td>Structural defects</td>
<td>Oxide inclusions</td>
<td>Pick-up during spraying or excessive surface oxide on feed material</td>
</tr>
<tr>
<td></td>
<td>Unmelted particles</td>
<td>Incorrect powder quality or spraying conditions</td>
</tr>
<tr>
<td></td>
<td>Sooty deposits</td>
<td>Insufficient interval between melting and boiling points in coating material</td>
</tr>
<tr>
<td>Poor adhesion</td>
<td></td>
<td>Poor practice and particularly poor substrate preparation</td>
</tr>
</tbody>
</table>

1 Correct location and thickness of deposit.
2 Visual examination for freedom from obvious defects, such as cracks and porosity.
3 Verification of heat treatment records, correct issue and use of consumables.
4 Dimensional checks for size, distortion, etc.
5 Dye penetrant inspection as an aid to identification of cracks; care is needed in interpretation of the result.
6 Hardness and non-destructive testing according to the application. Use of radiographic examination is not widespread and tends to be used only for certain specialised applications, e.g. surfacing of some nuclear components.

As incorrect grinding can cause surface cracking of hard deposits, a final dimensional check and dye penetrant test are useful, after finish machining and grinding.

In seeking to set the appropriate quality standard it is necessary to evaluate the more commonly occurring defects in weld deposition (Table 9.6). Excluding dimensional errors these are discontinuities and are of three main types: cracks, porosity and inclusions.

Cracks may be of various kinds. Contraction cracks arise as a consequence of low ductility, and possibly low tensile strength of any hardfacing alloys,
which make them unable to withstand the stresses upon cooling of the surfaced component. Such cracking may be made worse by stress raising defects in the coating and by post-weld straightening to rectify any distortion. The presence of contraction cracks may be accepted as a necessary consequence of the use of the most wear-resistant deposit material and, for some applications, they may not impair service performance.

Their presence rarely involves the risk of the deposit breaking away from the substrate, provided there is no hardening of the HAZ and the welding operation has given a satisfactory bond to the substrate. However, such cracks would not be tolerated in the sealing surfaces of valves and sealing rings, in surfaces subject to erosion, and in surfaces that must give corrosion, as well as wear, protection to the substrate.

Fine surface cracking or checking can occur during grinding of hard deposits if grinding conditions are unsuitable or there is insufficient coolant. Such grinding cracks are usually shallow but may propagate with thermal or mechanical stress; they may therefore be subject to the same restrictions as contraction cracks.

The deposit lifting from the substrate constitutes another type of cracking. Cracks may occur at the interface between the deposit and the substrate and may stem from a poor welding technique resulting in a failure to develop a good bond, or to failure to disrupt surface oxide films before welding begins. Alternatively, cracks may occur in the substrate just below the interface with the coating; these are the result of embrittlement of the substrate during welding and can be avoided by ensuring that a correct thermal treatment is given. As the deposit may detach itself during service, such ‘lifting’ defects cannot be accepted for any application.

Porosity can have various causes. If associated with the final molten pool it may be the result of faulty crater filling. When randomly dispersed it can be a result of, for example, flux contamination, faulty flame adjustment (in gas welding) or damp electrode coverings (in arc welding). It can also be caused by chemical reaction in the molten pool. Porosity may range from a few specks to visible spongy metal, so that a form of graded specification should be used; the degree of acceptance can then be related to the application. It may also be possible to have acceptance standards that vary according to their location on the component. Any such variations should be shown in the acceptance standards.

For some applications visual examination for porosity may be sufficient; for more critical service dye penetrant examination is used. It may be unnecessary, and probably uneconomic, to specify a ‘zero bleed’ acceptance standard; moreover, grinding of certain hardfacing alloys may tend to pluck out hard, needle-shaped carbides which dye penetrant testing will pick up. For these reasons the standard for, and interpretation of, dye penetrant testing must be appropriately set.

Inclusions can range from a size which is obvious in general examination to that which may only be detected by dye penetrant testing. They may be caused by the use of a dirty substrate material, but more usually they arise from incorrect welding parameters or technique. Their identification, location and acceptance standard can conveniently be grouped with those criteria for porosity.
Where defects are found the designer will decide whether the deposit can be rectified. If weld deposition has originally been used it would be normal to use welding again. The following points should be taken into account when making such repairs:

1. Repairs should be made at the proof grinding stage where there is usually still sufficient machining allowance to deal with any further distortion or oxidation.
2. The same metallurgical conditions should generally be specified as for the original deposit.
3. The defective area must be removed completely down to sound metal, verified by dye penetrant testing. A shallow rounded depression should be produced and not one that is narrow and deep or has sharp corners. If the substrate metal has to be replaced, the profile should be restored by a compatible electrode (and not by a hardfacing alloy) before recoating.

**Thermally sprayed coatings**

The dimensions of the coated product must be correctly held to the customer's specification and these – as with welded deposits – must be checked. Again, a micrometer is usually sufficient for the accuracy demanded, although electronic displacement instruments (based upon eddy current or magnetic techniques) are coming into use for certain coating/substrate combinations.

The quality of the coating is vital. As some coatings are less dense and less homogeneous, and as adhesion tends toward mechanical rather than a metallurgical mechanism, some NDT techniques can be less reliable. Greater emphasis must consequently be given to destructive techniques such as adhesion tests and metallography. Destructive testing on a small percentage of the components is acceptable where large numbers of similar articles are being processed. However many thermal spraying jobs involve coating of small numbers of components, and often only one might be processed in reclamation work. In these circumstances it is necessary to coat a test standard or coupon. The difficulty is then to ensure that the coating on the components is comparable. This requires not only consistency in spraying conditions, but also an appreciation of the effects of, for example, differences in component section thickness and cooling rate on deposit structure and properties.

Metallographic examination is widely used to assess deposit quality, but care must be taken in interpretation of the section and in interpretation of the results to avoid misleading conclusions. At its best such examination can show up defects, give an indication of porosity, and highlight features of the interface. Equally, care must be exercised in the preparation of samples for examination, especially if sections must be cut to reveal the deposit/substrate interface.

Hardness, being easily determined, is commonly measured on coatings but its usefulness is limited. Tests of the Brinell or Vickers type give a general indication of coating quality, but the presence of porosity will lead
to an apparently low hardness compared with that of a similar solid material. Additionally, the presence of a hard substrate that supports a thin soft surface film will give a hardness that is affected by deposit thickness. These aspects cast doubt upon the absolute use of hardness measurements but may be useful for comparative purposes. Microhardness may also be used as a comparative check that the correct deposition conditions have been achieved.

The importance of coating adhesion and cohesion is such that a large number of methods of measurement have been tried. Many, such as chisel and scratch tests, are of a qualitative or semi-quantitative nature and are used with a skilled operator only for comparison of coatings. Others, however, set out to be quantitative and many testpiece designs and pulling arrangements have been tried. Figure 9.1 shows a typical example of testpiece design. To assess cohesion within the coating two accurately machined tubes, which have been sprayed on their outer surfaces, may be pulled apart. To determine adhesion, two cylinders are used, one being coated and resin bonded to the other; these are then pulled apart. There are, however, several limitations in tests of this type. If adhesives are used, the result may be affected by the type of adhesive, the degree of penetration into a porous coating and the conditions used during curing. Further, the adhe-
sion of some of the best coatings produced by high velocity spray systems is greater than the cohesive strength of the best adhesives. This makes it impossible to give a quantitative measurement of adhesion in the best coatings. The tests may in any event be affected by alignment of the specimen in the test machine and by strain rate variables between tests. Such tests must thus be interpreted with caution; at best they represent a degree of consistency so that gross variations in reading may indicate poor production conditions. For these reasons there has been continuing interest in the development of a reliable non-destructive method to measure coating adhesion. Despite much effort no fully satisfactory method has yet been proved, but recent work based upon use of an ultrasonic method gives promise of being developed into a useful technique.

The NDT methods to assess general coating quality have had only limited success with sprayed deposits. Surface finish measurements may be useful to indicate batch-to-batch differences in coatings: it has been suggested that surface roughness can be developed as a quick and reliable method of checking coating quality. Crack detection by magnetic or dye penetrant techniques often fails to detect isolated defects in the general presence of porosity. Eddy current and ultrasonic inspection are little used for sprayed coatings, which tend to be heterogeneous and give a high noise-to-signal ratio. Radiography can be used but with some uncertainty in interpretation.

Table 9.6 includes defects that may occur in sprayed coatings. Metallography will highlight many of the defects that can arise from bad spraying procedures: excessive porosity, high oxide content, the presence of unmelted particles and possibly poor bonding of coating to substrate. Most sprayed deposits have pores arising from shrinkage and degassing after deposition; the former gives acicular shaped pores, the latter more rounded ones. However, poor spraying practice can lead to a condition of gross porosity which must be avoided. Oxide inclusions can arise from pick-up during spraying – especially in the oxygas or arc methods – or from excessive surface oxide on a (powder) feed material. The latter may be controlled by attention to quality and particle size during powder manufacture. Where the oxide content from pick-up is unacceptably high it may be reduced by use of a shroud to protect the workpiece; this is quite common. Alternatively, deposition may be carried out in an inert gas chamber. The presence of unmelted particles indicates incorrect powder quality and/or spraying conditions. One of the advantages of using a solid, e.g. wire, feed is that the coating material must be melted before it is projected at the substrate. There is a need for a reasonable temperature interval between the melting and boiling points of the coating material, otherwise excessive evaporation can occur which will lead to a ‘sooty’ type of loosely bonded deposit. It should be emphasised that these defects will not occur with good spraying practice and, indeed, usually cause few problems.

Inadequate adhesion of coating to substrate is a defect that cannot be tolerated. There are certain deposits that inherently give problems with bonding to certain substrates and an intermediate (or bond) coat may be used to overcome problems. In practice, however, poor adhesion most commonly stems from poor substrate (component) preparation and even a slight departure from the recommended practice – especially in grit blasting – can
give poor results. If such problems exist the defective coating must be stripped and the part recoated; at the same time the operating practice should be critically reviewed. If a coating is subjected to a finishing operation its ability to withstand this is an empirical test of deposit adhesion.

A proven track record of a particular substrate-coating material and its application process can be a useful starting point when considering possible solutions to a user's problem on a different type of component.

Conclusions

The aim of this chapter has been to present the view that enduring quality of surface components depends not upon sophisticated and expensive test methods, but on simple, but strict, control of each stage of the processing cycle. Starting at design, each stage should be carefully considered, defined and carried out as specified.

Faulty materials or design, or incorrect surfacing practice, can produce a poor deposit that may be difficult to detect. The philosophy must therefore be that poor quality cannot be tested out; good quality must be built in so that inspection then becomes a verification that good practice has been observed.
Safe working in surfacing

The Health and Safety at Work etc. Act 1974 (HSAW) puts the duty on employers to provide premises, plant and systems of work that are safe and without risk to health. It also charges employees to take reasonable care for their own health and safety, and for that of others. They are required to co-operate with their employer as far as necessary to comply with his or her duties under the act, and they are required not to intentionally misuse or damage anything provided for their safety.

Since the Act, several Regulations and Approved Codes of Practice have been published. These are mandatory, and give detailed guidance on compliance with the HSAW in their specific areas. Guidance literature from the Health and Safety Executive (HSE) should be regarded as good practice; an accident that arises because working practice fell short of the standards laid down in HSE guidance could be seen as less than due diligence.

In addition, all equipment now marketed within the European Union (EU) must be 'CE' marked. The CE mark is the manufacturer's guarantee that the equipment complies with the safety requirements, both in terms of mechanical safety and electrical safety. In addition it will have had an assessment of its ability to operate safely within an environment of electromagnetic radiation, and its own emissions of electromagnetic radiation into the environment. The significance of these measures is to ensure that equipment with electronic control does not interfere with other such equipment, or itself be subject to interference resulting in dangerous malfunction.

This chapter briefly describes hazards encountered in surfacing and the steps that should be taken to overcome them and achieve safe operation. It is divided into three sections: surfacing by welding and thermal spraying, vacuum deposition and electrodeposition.

The bibliography at the end of this chapter lists the principal UK legislation and standards. Other countries have a similar range of documentation. A detailed treatment of working practice is given in The Welding Institute's book Health and Safety in Welding and Allied Processes. However, the current (fourth) edition was published in 1991, so references in that book to the legal framework in the UK have been largely superseded.
Surfacing by welding and thermal spraying

Compressed gases

The HSE has recently produced a booklet, 'The safe use of compressed gases in welding, flame cutting and allied processes', HS(G) 139, which contains a great deal of useful advice on this topic.

Storage

Cylinders containing compressed gases (Table 10.1) must always be handled with care and stored in well-ventilated areas, out of direct sunlight, and away from the working area whenever possible. The store should be secure, to prevent unauthorised removal. There should be a strict 'no-smoking' policy in the area. Cylinders should be segregated, according to the following rules:

1. Oxygen and other oxidants must be separated from any fuel gas by a distance of not less than 3 m, or by a wall that can resist fire for not less than 30 min.
2. Acetylene and LPG cylinders must not be stored in the same area.
3. Hydrogen must be segregated from other gases, and stored not less than 1 m away from them, or separated by a fireproof wall.

Fire brigades should be kept informed of the location of cylinder stores.

At no time should a cylinder be left free-standing. It should always be tethered securely. Serious leg injuries can result from being struck by a cylinder when it topples over and, if a regulator is fitted, it can shear off in
the fall. If the valve is open the cylinder will accelerate due to the release of the gas inside, and could kill someone in its path. Where portable welding equipment is in use, cylinders must be carried on a properly designed trolley.

**Acetylene cylinders**

Acetylene under pressure can undergo a chemical reaction that releases heat and tends to increase the pressure. This reaction can be triggered off by a flame entering the cylinder from a flashback or by general overheating of a cylinder, for example in a fire. For this reason it is strongly advised that flashback arrestors are fitted to all cylinders. A hot acetylene cylinder should be taken out of doors and sprayed with water to cool it and keep it cool, which should halt the internal chemical reaction. The suppliers should be advised of this immediately.

Safety devices (fusible plugs and/or bursting discs) may be provided to limit pressure increase, but when these operate there will be a massive release of highly flammable gas. If they fail, pressure may increase to the point at which the cylinder bursts.

**Equipment**

Regulators and other equipment must be suitable for the gas, with undamaged connectors, and a working pressure rating equal to or exceeding the maximum cylinder filling pressure. It is particularly important to review these, since the cylinder pressure has increased in several steps in recent years. Flame equipment should always be fitted with a flashback arrestor on both the oxygen and the fuel lines.

Regulators should be inspected annually by a competent person, and both regulators and flashback arrestors should be replaced every five years.

Gas hoses and their connectors must be inspected regularly to ensure that they remain in good condition. Particular attention must be paid to trailing hoses. If repairs are necessary they must be made with the correct connectors and hose clips. Check equipment for leaks after assembly using a leak detector fluid or Teepol and water.

All equipment, particularly that used for oxygen, must be kept clean and free from oil or grease. Oxygen at high pressures can react explosively with oil or grease.

**Key points**

- Check that the right gas is being used to suit the equipment and job.
- Do not smoke when handling or connecting fuel gas or oxygen cylinders.
- Secure cylinders so that they cannot fall over.
- Store and use acetylene and propane cylinders upright; use carbon dioxide cylinders upright.
- Protect cylinders from heating, impact or accidental contact with arcs.
- Keep connections clean and dry.
- Use correct regulators and fittings.
Do not use cylinders as work supports or as rollers.
Examine equipment for damage, check for leaks (with Teepol solution for flame equipment).

Fire
Almost all employers are required by law to have formal plans to deal effectively with emergencies, to ensure the safety of personnel. In practice this is normally the provision of fire alarms (which can be a simple hand bell for a very small workplace), suitable fire-fighting equipment, and safe means of rapidly leaving the building. Personnel need to be trained in the procedure to be followed, and the employer must establish that everyone is accounted for.

Flames and arcs
Use of flames and arcs in welding, preheating, etc., presents an obvious fire hazard and appropriate precautions should be observed. No combustible materials should be allowed to be used or stored near the working area and workbenches. Partitions and other fixtures should be made of fireproof materials. The possible hazards from weld spatter and from heated overspray particles should not be overlooked.

Metal powders
Finely divided metal powders may constitute a fire or explosion hazard if allowed to collect in sufficient quantity. There is also a health hazard arising from the inhalation of fine powders. The work space should be designed to reduce dust accumulations and should be cleaned regularly. Ducting should be designed so that it can be cleaned when necessary. Procedures for cleaning must give due consideration to the risk of inhalation of powders, and take preventive measures.

Key points
Avoid excessive release of fuel gas or oxygen into the atmosphere.
Remove flammable materials from the working area.
Where adjacent flammable material cannot be removed, station an observer to deal with any outbreak of fire.
Check for fire about an hour after work has finished.
Check emergency escape routes.
Use a tray to catch overspray where possible, preferably sand or water filled.
Ensure that appropriate firefighting equipment is at hand and that all personnel know how to use it.
Vessels with residual flammable material

A used or contaminated vessel, tank, drum, pipe or other closed installation that has at any time contained a flammable liquid or dust is liable to explode if a surfacing process is operated on it or in it. The risk can be removed only by thorough cleaning or filling with an inert gas.

Key points

- Obtain a certificate from a competent person that any used or contaminated vessel is free from risk of explosion before preparing to work on it.
- Where a ‘permit-to-work’ system is in operation, ensure that all instructions are understood and obeyed.

Exposure to substances

With all welding and spraying operations fume, dust and toxic gases are liable to be produced. There will probably be both respirable fume (particle diameter <5\,\mu m), which can get deep into the lungs, and nuisance particles (diameter ≥5\,\mu m), which will affect the upper respiratory tract only. Several metals produce toxic fume – these include cadmium, lead, chromium, nickel and cobalt. Some fume, especially from stainless steel, can trigger asthma in susceptible persons.

Fumes and dust can arise from:

- airborne particles of spray material;
- chemical compounds of the surfacing material and the parent metal such as oxides.

Toxic gases can also be released as:

- gaseous products of decomposition of fluxes and electrode coverings;
- products of combustion of the gases used, e.g. carbon monoxide, carbon dioxide, oxides of nitrogen;
- breakdown products from degreasing fluids, or residues such as paint on the workpiece.

The Control of Substances Hazardous to Health (generally known as COSHH) 1994 laid down the guidelines for exposure to substances that can harm health. The employer is required to assess the hazards, and use measures to control exposure – with a preference for containing the problem to prevent exposure altogether. Personal protective equipment may well be necessary to supplement these measures, but is not to be used instead of proper control of fume and dust.

Many of the substances used in surface coatings are harmful to health. Two categories can be identified: those for which there is a Maximum Exposure Limit (MEL) and those for which there is an Occupational Exposure Standard (OES). In the case of the MEL, it is a legal requirement to reduce the exposure of the worker to the substance to the lowest level which is reasonably practicable, and in no case to expose him/her to a level in excess of that stated. For substances with an OES, exposure should be kept...
Table 10.2 Exposure limits for inhalable fume or dust from substances common in welded or sprayed surface coatings

<table>
<thead>
<tr>
<th>Substance</th>
<th>Maximum exposure limits (mg m⁻³) for an 8 hour period</th>
<th>Occupational exposure standards (mg m⁻³) of fume for an 8 hour period</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cadmium</td>
<td>0.025*</td>
<td>Copper fume</td>
</tr>
<tr>
<td>Chromium (and Cr(VI))</td>
<td>0.05*</td>
<td>Iron oxide</td>
</tr>
<tr>
<td>Cobalt (and soluble compounds)</td>
<td>0.1</td>
<td>Manganese</td>
</tr>
<tr>
<td>Nickel (insoluble compounds)</td>
<td>0.1*</td>
<td>Zinc oxide</td>
</tr>
<tr>
<td>Nickel (soluble compounds)</td>
<td>0.5*</td>
<td>Welding fume (in general)</td>
</tr>
</tbody>
</table>

* The oxides of these metals are also listed as 'may cause cancer by inhalation'.

Table 10.3 Exposure limits (OES) for some toxic/asphyxiant gases (mg m⁻³)

<table>
<thead>
<tr>
<th>Substance</th>
<th>Carbon dioxide</th>
<th>Carbon monoxide</th>
<th>Ozone</th>
<th>Nitrogen dioxide</th>
<th>Nitric oxide</th>
<th>Phosgene</th>
<th>Chlorine</th>
<th>Hydrogen chloride</th>
</tr>
</thead>
<tbody>
<tr>
<td>CO₂</td>
<td>9150</td>
<td>58</td>
<td>0.4*</td>
<td>5.7</td>
<td>31</td>
<td>0.08</td>
<td>1.5</td>
<td>7.6*</td>
</tr>
</tbody>
</table>

* Figure quoted is for a 15 minute exposure, remaining figures are for 8 hour exposures.
† Some toxic by-products which can arise when common degreasing solvents are present near welding arcs.

below the maximum stated, as it is believed that no harm will come to a person exposed to levels up to that maximum on a regular eight hour working day basis.

Note that zinc and copper can both give rise to ‘metal fume fever’, which is an acute illness brought about by exposure to fume. The symptoms are similar to influenza, and recovery is usually complete. It is also possible for a person to become sensitised, so that very low exposures can trigger the illness.

Tables of MELs and OESs are published annually in the Health and Safety Executive document EH40, ‘Occupational exposure limits’. It also lists those substances that are believed to be carcinogens. Table 10.2 gives the 1997 values for some of the dust and fume that may be encountered in coating by welding or spraying. Table 10.3 gives the corresponding data for gases and solvents.

Suppliers of consumables are required to provide information about their products, and generally make it freely available. Where any doubt exists as to the danger of a particular material or of the effectiveness of ventilation and extraction systems, the concentration should be measured. Guidance is given in the HSE document EH54.

Ventilation

Adequate ventilation of the working area is necessary to remove fumes. The position of ventilation inlets should be arranged so that the fumes are taken
away from the operator. The exhaust must be placed so as to avoid recirculation of fumes to work areas. Practical advice on local exhaust ventilation is available in HSE document EH55. Fume extraction systems must be properly maintained, and tested at intervals not exceeding 14 months (more frequently for processes concerning lead). Records of such examinations must be kept for not less than five years.

In addition to extraction it may be necessary to provide for dust collection, especially with materials having a low OES, or an MEL. In thermal spraying shops, use of water-washed spray booths is now common practice for catching the overspray and controlling the amount of dust generated, but other methods are also available and each case must be judged individually.

**Personal dust and fume protection**

This can be used to supplement ventilation to give adequate reduction of dust and fume. The options available are:

- dust respirator with replaceable filter;
- disposable dust respirator;
- air fed helmet;
- helmet with fan and filter (positive pressure-powered dust respirator).

**Key points**

- Keep degreasing fluids and vapour away from the work area.
- Check for possible toxic hazards from parent material or consumables.
- Where necessary use a local fume extractor repositioned as work proceeds.
- Supplement fume extraction with personal protection where necessary.
- Encourage good housekeeping and cleanliness.

**Work in enclosed spaces**

If surfacing is to be carried out in an enclosed space special care is needed because of the risk of a build up of fume or gas. There is also the possibility of oxygen-depleted atmospheres existing within confined spaces. Work in confined spaces carries a significant risk of a fatal incident, and this work is now subject to regulations. Please refer to the Confined Spaces Regulations, 1997, and the associated Approved Code of Practice, *Safe Work in Confined Spaces*. It is also advised that HSE Guidance Note GS5 is consulted. It will always be necessary to establish the safety of, and monitor, the atmosphere within the space. Adequate arrangements must be made for the safe rescue of a worker who is in difficulties, and this must be rehearsed. Adequate training must be undertaken.

**Noise**

High velocity oxyfuel, plasma spraying and, to a lesser but still significant extent, wire spraying operations give rise to intense high frequency noise which can cause damage to hearing. Sound attenuating screening
is advised for the former two processes for protection of adjacent personnel.

Wherever it is difficult to carry on a conversation with a person standing approximately 2 m away, it is likely that the noise level is above 85 dB(A). When the noise reaches 85 dB(A) employers are required to offer hearing protection to their employees. If it exceeds 90 dB(A) hearing protection becomes mandatory, and it is necessary to undertake a proper noise survey to analyse the amplitude/frequency spectrum of the noise. This survey is necessary in order to be able to select hearing protection which adequately attenuates the noise in the various frequency ranges.

Welders also require hearing protection since, for instance, hand grinding emits noise in the order of 108 dB(A), and this constitutes an unacceptable exposure to noise in the workplace.

Places where the noise levels exceed the two action levels (85 and 90 dB(A)) should have prominent warning signs reminding personnel of their duty to wear hearing protection.

**Key point**

- Wear suitable ear protection when exposed to loud noise, or when working in areas where the warning signs show that hearing protection must be worn.

**Radiation**

All surfacing processes that make use of a flame or an electric arc emit radiation in the form of heat, visible light and ultraviolet radiation.

**Radiation to the eye**

The ultraviolet radiation from an arc, and to a lesser extent from a flame, causes a painful irritation of the outer surface of the eyeball known as arc eye. Exposure for only a few seconds to the light from a high current welding arc can produce severe discomfort, starting a few hours after the exposure. Fortunately most cases recover completely in a day or two with no permanent ill effects.

The visible light is usually too intense to allow comfortable viewing. The potential hazard here is to the retina, the light-sensitive surface at the rear of the eye, which could be permanently damaged by viewing a bright source such as an arc. Blue light is the most harmful, but the infrared radiation may also be intense enough to be harmful in rare circumstances.

Protection of the operator is afforded by a suitable filter which cuts out ultraviolet and infrared and reduces visible light to a comfortable level. A suitable specification is BS EN 169, which identifies filters by a shade number: the higher the number the darker the filter. Table 10.4 gives recommended shades. Switchable filters, which have more than one transmittance value, are specified in BS EN 379. It is customary to protect the filter against spatter damage with clear plastic cover sheets on either side.
Table 10.4 Protective viewing filters for use when surfacing

<table>
<thead>
<tr>
<th>Process</th>
<th>Conditions</th>
<th>Filter shade number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gas welding/spraying/fusing</td>
<td>&lt;70 litres acetylene h⁻¹</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>70–200 litres h⁻¹</td>
<td>5</td>
</tr>
<tr>
<td></td>
<td>200–800 litres h⁻¹</td>
<td>6</td>
</tr>
<tr>
<td></td>
<td>over 800 litres h⁻¹</td>
<td>7</td>
</tr>
<tr>
<td>Manual metal arc welding</td>
<td>&lt;80 A</td>
<td>9 or 10</td>
</tr>
<tr>
<td></td>
<td>80–175 A</td>
<td>11</td>
</tr>
<tr>
<td></td>
<td>175–300 A</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td>300–500 A</td>
<td>13</td>
</tr>
<tr>
<td></td>
<td>&gt;500 A</td>
<td>14</td>
</tr>
<tr>
<td>MIG welding</td>
<td>&lt;100 A</td>
<td>10</td>
</tr>
<tr>
<td></td>
<td>100–175 A</td>
<td>11</td>
</tr>
<tr>
<td></td>
<td>175–300 A (steel)</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td>175–250 A (light alloys)</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td>300–500 A (steel)</td>
<td>13</td>
</tr>
<tr>
<td></td>
<td>250–350 A (light alloys)</td>
<td>13</td>
</tr>
<tr>
<td></td>
<td>&gt;500 A (steel)</td>
<td>14</td>
</tr>
<tr>
<td></td>
<td>350–500 A (light alloys)</td>
<td>14</td>
</tr>
<tr>
<td></td>
<td>&gt;500 A (light alloys)</td>
<td>15</td>
</tr>
<tr>
<td>TIG</td>
<td>&lt;20 A</td>
<td>9</td>
</tr>
<tr>
<td></td>
<td>20–40 A</td>
<td>10</td>
</tr>
<tr>
<td></td>
<td>40–100 A</td>
<td>11</td>
</tr>
<tr>
<td></td>
<td>100–175 A</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td>175–250 A</td>
<td>13</td>
</tr>
<tr>
<td></td>
<td>&gt;250 A</td>
<td>14</td>
</tr>
</tbody>
</table>

According to the conditions of use, the next greater or the next smaller scale number may be used.
Source: Based on BS EN 169: 1992.

In addition to radiation, a filter also prevents stray particles of coating materials, etc., from entering the eye and burning or otherwise injuring the outer surface. There is now a requirement for impact resistance in protective helmets, etc.

For light duty gas processes where face protection (see next section) is not required, the filters may be used in suitable goggles. Instrument panels, dials, etc., should be positioned so that they do not reflect light from the arc to the operator or anybody else.

Walls, ceilings and working surfaces should be painted so as to reduce amounts of ultraviolet radiation. This can be achieved by paint of almost every colour except metallic, but blue should be avoided to minimise visible light hazards. Pastel shades of matt emulsion paint are effective and easily applied. The polished surfaces of metal or glass, gloss paint finishes and water in wet booths reflect an image of the arc which is often distracting and may be harmful. This should be avoided by careful positioning of the work.

Protection of the eyes of others nearby may be given by fixed or portable screens or by personal goggles or face shields. Screens can take a number of forms:
1. Opaque solid (hardboard or sheet metal).
2. Opaque flexible (fireproof heavy canvas).
3. Transparent or translucent flexible plastic, transmission corresponding to viewing filters.
4. Transparent uncoloured flexible or rigid plastic or glass. Most such materials transmit only diffused light, greatly attenuating infrared and ultraviolet.

Likewise clear plastic or glass spectacles, goggles or face shield will protect against most very brief or minor exposures to ultraviolet light.

Radiation to the skin

The intense ultraviolet from the electric arc will rapidly cause ‘sunburn’ on any unprotected skin, and indeed can cause severe blistering after a very short exposure.

A helmet or hand shield is used, with the appropriate filter, to protect the face. Particular attention must be paid to the hands and neck and it should be noted that normal clothing does not necessarily give adequate protection. For instance with the high intensity emission from a plasma torch, and from any process where high currents are used it is possible to experience ultraviolet burns through a nylon or thin cotton shirt. Hence proper overalls which fasten up to the neck, gloves and face masks (see above) should be worn. Fabrics should be close woven cotton, which will not readily catch fire. White overalls should not be worn when arc processes are in use because they reflect too much light. The standard BS EN 470-1 describes the requirements for welder’s protective overalls.

With welding it may be necessary to provide additional operator protection against infrared radiation if work is to be carried out on large workpieces with high preheat temperatures.

Laser radiation

Surfacing can be carried out using a laser that supplies enough infrared energy (2–10 kW) in the form of a focused beam of radiation to melt powder to form a coating on the surface of a parent metal. The beam can burn eyes or skin even after reflection from a dull or diffuse surface. The usual safety procedure is to enclose the whole work zone in material that will not transmit infrared or be seriously damaged or set on fire by accidental impingement of the beam. Metal is often used, with clear plastic windows (even clear plastic strongly absorbs infrared), or the whole enclosure may be made of clear plastic for demonstration or experimental work. It is usual to install an interlocking system such that the beam is automatically shut off if the integrity of the enclosure is broken.

Some equipment uses an auxiliary low power visible light (red) laser beam to mark the beam path for system alignment. This visible laser could damage the retina of the eye if it is allowed to enter the pupil for any length of time, and care is needed not to look along the beam path during the alignment.
Key points

- Use helmet, hand shield or goggles fitted with a suitable filter and clear protective sheet.
- Check that helmet, hand shield or goggles do not admit light as a result of damage.
- Wear appropriate protective clothing to prevent burns.
- Warn bystanders before striking an arc or operating a laser whose beam is not enclosed.
- Screen the working area to protect others working in the vicinity.

Electric shock

All electrical equipment must be properly installed by competent personnel and must be regularly inspected and maintained. Equipment casings must be earthed in the normal manner with the usual mains earth. Particular attention must be given to trailing leads especially primary (mains) leads to ensure that their insulation and end connections remain in satisfactory condition.

It should be noted that there are several quite distinct generations of welding equipment. These are described in the HSE booklet HS(G) 118, in which advice is given about connection of the equipment and earthing.

High frequency supplies used to start the arc in TIG and plasma processes can cause deep burns if they are allowed to contact the skin. The equipment should never be operated with the covers removed, exposing the circuitry.

Key points

- Check what type of welding equipment is installed, and consult HSE guidance for advice on connection and earthing.
- Check for damage to insulation on electrode holders, torches, guns, cables and connectors.
- Keep work area dry.
- Provide emergency means close to the worker to cut off the current when out of reach of the controls.
- When working where a shock could cause a fall, provide guard rails or other protection.
- Check that equipment casing is correctly earthed on installation.
- Switch off equipment before cleaning or dismantling guns.

Surface preparation

Abrasive blasting

Abrasive blasting is the most commonly used method of surface preparation. A major hazard would result from inhalation of any silica in the abrasive particles, so commercial blasting media do not contain free silica (in the form of sand for example) and its use is not permitted under UK law. Instead materials such as aluminium oxide, chilled iron or steel grit or
shot, bauxilite or glass beads are used in various proprietary formulations.

Although the abrasive is therefore non-hazardous it will have considerable nuisance value as well as being detrimental to other machinery in the same workshop. Any blast cabinet installation should be fitted with an adequate dust collector and effective door seals which should be maintained in good working order. Safety door interlock switches should be fitted on automatic installations and are desirable on manual equipment. A recommended maintenance schedule is shown in Table 10.5.

Electrostatic charges may build up on gun or workpiece if either is not earthed but the gun will normally have its own earth lead with the supply hoses, and the work is readily earthed except where it is held in the operator’s gloved hand, in which case a lead can be taken from a convenient point, usually at the armhole to a ring fitting outside the glove.

With work on a larger scale where the operator enters the cabinet or for open site work the operator must use gauntlets, overalls and a suitable helmet providing eye protection and a supply of fresh air (0.17 m³/min⁻¹).

**Key points**

- Use only approved abrasive media.
- Avoid excessive contamination of media.
- Check seals regularly.
- Check dust collectors regularly.
- Ensure adequate fresh air supply and protective clothing for operators working within cabinets or on open site.

**Solvent degreasing**

Solvents available for industrial degreasing may be hazardous because of flammability and/or toxic vapours. UK regulations specify the hazard control measures to be adopted. Non-flammable solvents such as carbon tetrachloride, trichloroethylene and perchloroethylene evolve vapours which are themselves toxic, and will also be decomposed by heat or ultraviolet light from an arc, with the evolution of even more toxic gases such as phosgene.

111 Trichloroethane is being phased out, since it has a deleterious effect on the ozone layer. This has meant that many users have reverted to

<table>
<thead>
<tr>
<th>Table 10.5 Typical maintenance checks for a grit blasting unit</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Daily</strong></td>
</tr>
<tr>
<td>Seals and toggle catches</td>
</tr>
<tr>
<td>Media, for fill and contamination</td>
</tr>
<tr>
<td>Nozzle and air jet for wear</td>
</tr>
<tr>
<td>Gun hose connectors</td>
</tr>
</tbody>
</table>
trichloroethylene, even though it is significantly more harmful to health. It is recommended that this substance is only used within a closed degreasing system, sited away from any welding or other heat source. Several alternative, much less hazardous, degreasing preparations are now available on the market, and it is recommended that these should be evaluated.

Ventilation should effectively remove fumes: where degreasing must be carried out near surfacing operations, the general ventilation flow should lead away from the surfacing area towards the degreasing bay. The exhaust must be placed to avoid recirculation of fumes to work areas.

**Key points**
- Keep degreasing fluids away from the work area.
- Ensure that degreasing vapour does not reach surfacing operations.
- Ensure safe disposal of used solvents.

**Grinding**

Preparation by grinding is subject, in particular, to the Abrasive Wheels Regulations. Machines must be marked with their operating speeds, and where practicable guards and rests to be fitted. They must be maintained in a safe condition. Wheels must be chosen to be adequate for the machine and the application, and may only be fitted by competent persons, specifically trained to a recognised level of competence, and appointed for the task by their employer.

Eye protection such as to BS EN 166 is required for all grinding operations.

**Key points**
- Use suitable eye protection when deslagging, chipping or grinding.
- Ensure abrasive wheels are correctly fitted.

**Vacuum deposition**

**PVD techniques**

All these processes involve vacuum equipment, to which very little hazard is attached. Rotary pumps should not exhaust directly into the work room, because there is often a discharge of fine oil droplets from such pumps during the initial pumping down stage. Diffusion pumps pose few dangers other than the risk of thermal burn from the heater. Some vacuum gauges operate at potentials of a few kilovolts, but they are generally well protected by the manufacturer.

Liquid nitrogen is sometimes used, and the operator should be adequately protected against splashes by wearing gloves and goggles, or preferably a face mask. When pouring, and in general operation, there should be ample ventilation to prevent build-up of asphyxiant gas. Care is needed to avoid
risk arising from the evaporation of nitrogen and condensation of oxygen in open containers leading to an oxygen-rich liquid.

There will be some special risks arising from reactive gases when reactive techniques are used, but these risks are often specific to the gases involved and the supplier's advice should be sought and closely followed.

There is a risk of electrical burn and shock when RF or ionising processes (e.g. ion sputtering, plating) are used; in both considerable power is available and fatal injury can result. Prevention of accidents is a matter of design. In particular, there should be provision of interlocking to prohibit access to 'live' equipment, and a suitable operating procedure should be adopted.

**CVD techniques**

Nearly all CVD techniques involve high temperatures and toxic or carcinogenic gases such as carbonyls or organo-metallic compounds. The manufacturer's safety data sheets must be consulted, and a full risk assessment of the hazards arising from their use must be addressed before they are used. The composition of the exhaust gas must also be taken into account.

**Electrodeposition**

*Workflow*

The objective of an electroplating installation is to apply a specific finish to a variety of surfaces. These surfaces can be reactive to various chemicals and a thorough knowledge of the initial, intermediate and final surface conditions is required if work is to be accomplished safely and competently. For example, a plating shop might have to cope with materials such as cadmium, already present on the surfaces of items sent to them for plating. These materials might then enter the effluent cycle and have to be handled by the worker, and special protective measures will have to be considered for these cases.

Cyanide-containing chemical residues on the surfaces of parts do not produce atmospheric hazards. If, however, such surfaces are immersed in acid solutions the cyanide radical is reduced to volatile hydrocyanic acid, which presents acute inhalation dangers. Most deaths from cyanide poisoning result from such accidental mixing of substances.

It is thus extremely important that the chemical state of the surface and its processing cycle are known. Equally important, however, is the layout of the processing cycle in the plating shop. If not designed carefully, this can create problems of dangerous cross-contamination, gas evolution or work spoilage.

*Engineering*

Although often neglected, the condition of engineering appliances in the plating shop can contribute significantly to the hazards. Critical areas are
Safe working in surfacing

construction, transmission and extraction. Problems are often caused or aggravated by the wet or corrosive conditions that prevail in the plating shop and for these reasons its construction and the equipment in it are important. It must also be remembered that emission to the atmosphere or environment through an opening in the roof or leaking draining system constitutes environmental pollution.

An electroplating shop contains probably the most aggressive components of the process industry in as much as they are fluid, corrosive and dynamic. As combating the effects of these is often quite difficult, great stress is laid on containment through proper selection of materials of construction, sources of energy and process materials. Again the emphasis is on prevention rather than cure.

The most dangerous combination is corrosion caused by the emission of process liquids and gases. These are often variable in composition, concentration, temperature and locality. The materials of containment capable of withstanding these conditions are often so expensive that they cannot be considered realistically for a low cost environment such as a plating shop. A compromise must often be made, relying on various forms of cladding, painting, coating or impregnation. Again, a thorough knowledge of characteristics of construction materials greatly assists in selection, economical use and effective maintenance of process equipment. Time and expertise used in selecting proper materials, prevention of ingress of aggressive materials (either liquid or gas) into sensitive working areas of control installations and purging or extracting spent solutions or gases will help to maintain equipment efficiently.

Process materials

Electroplating and other finishing processes involve the required conversion of base surfaces into functional surfaces where the required function is physical or chemical. This conversion occurs mainly by immersion in various alkaline, acid and salt solutions with or without the passage of current. Chemical compounds that are used to form the electrolytes are the primary source of danger in the electroplating department and methods must be devised to combat potential hazards from handling these.

Typical dangers emanating from handling chemicals can be chronic or acute, irrespective of the mode of entry and the target organ, and the degree of their insidious effect can usually be described in the form of danger signs, risk phrases and safety phrases. These signs and phrases are incorporated in the Classification, Packaging and Labelling (CPL) Regulations 1994 which categorise the effects of most of the commonly used chemicals according to their behaviour in animal or material tests. Thus, according to the animal trials the effect on skin or internal organs is characterised as very toxic, toxic, corrosive, harmful or irritant, whereas in material trials the effect is determined as explosive, oxidising, extremely flammable, highly flammable or flammable. As a rule, one symbol designating the most acute environmental effect and one symbol designating the most acute effect on tissue or organ are displayed on the label attached to the receptacle containing the material. In addition the appropriate risk and safety phrases describing the environ-
mental or toxicological aspects of the materials are also displayed. The phrases are standardised in the EU countries and can be decoded even if only the code, e.g. R25, S25, is quoted. In addition the substance identification number must be quoted describing the worst ecological or toxicological hazards. All this information, and a great deal more, is supplied by the manufacturer in the safety data sheet for the substance.

Storage and preparation of working solutions

Because of the hazardous nature of solutions involved in electrodeposition, consideration must be given to the elimination of risk. Personal hygiene and good housekeeping measures are vital preventive measures. The chemical store should be laid out with chemical classes carefully segregated, and effective means used to prevent spills from becoming mixed on the floor. There should be an adequate number of staff trained in first aid measures appropriate to the substances used.

Key points

- Do not store acids or alkalis together as excessive quantities of heat or toxic gases might be emitted on mixing.
- Do not add water to acid as the localised heat evolution might cause 'spitting' or ejection of acid. Always add acid to water.
- Allow good drainage in the storage area and separate acid drainage from alkaline. The storage area should have mains water and rubber or plastic hose connected to the mains through a tap.
- Ensure adequate mechanical assistance in materials' handling and dispensing facilities in the storage area as containers can be heavy.
- The first aid box should be available in the storage area or in the immediate vicinity. A first aider should be on call during mixing or draining and should be trained in procedures required for dealing with possible emergencies, e.g. the first aider should be familiar with the cyanide emergency procedure and with treating hydrofluoric acid burns.
- Allow adequate ventilation in the storage and dispensing area to prevent concentration of toxic or harmful substances.

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An introduction to local exhaust ventilation HS(G)37.
Industrial robot safety HS(G)43.
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Part 7: 1984 Safety requirements for installation and
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BS679 Replaced by EN169.
BS697: 1986 Spec. for rubber gloves for electrical purposes. See also BS EN 60903: 1993.
BS1542: 1982 Equipment for eye, face, and neck protection against non-ionising radiation arising during welding and similar operations.
BS1651 Replaced by BS EN 374, 388, 407, 420.
BS2092 Replaced by BS EN 166, 167, 168.
BS2653 Replaced by BS EN 470-1.
BS3510: 1968 A basic symbol to denote the actual or potential presence of ionising radiation.
BS4031: 1966 Specification for X-ray protective lead glasses.
BS4094: Recommendations for data on shielding from ionising radiation.
BS4275: 1974 Recommendations for the selection, use and maintenance of respiratory protective equipment.
BS4803: 1972 See BS EN 60825.
BS5330: 1976 Method of test for estimating the risk of hearing handicap due to noise exposure.
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BS5531: 1978 Safety in erecting structural frames.
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BS EN 349: 1993 Safety of machinery: minimum gaps to avoid crushing parts of the human body.
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BS EN 388: 1994 Protective gloves against mechanical risks.
BS EN 407: 1994 Protective gloves against thermal risks.
BS EN 420: 1994 General requirements for gloves.
BS EN 585: 1995 Gas welding equipment: pressure regulators for gas cylinders used in welding, cutting and allied processes, up to 200 bar.
BS EN 730: 1995 Equipment used in gas welding, cutting and allied processes. Safety devices for fuel gases and oxygen or compressed air.

British Standards may be obtained from the British Standards Institution, Customer Services, telephone +44 (0)181 996 7000, fax +44 (0)181 996 7001.
Chapter 11

Industrial applications of engineering coatings

Use of engineering coatings to improve the wear life of components in industry is worldwide and the circumstances prevailing at each site, which can differ widely, play a significant role in choice of coating process, and material. Factors such as working conditions, availability of specialised subcontract facilities nearby or of on-site skills, equipment and materials are all important, as is a policy of preventative maintenance or reaction to breakdown.

Previous experience of site conditions is an important factor in deciding the course of action to take and the information in earlier chapters is intended to assist in the decision-making process. There is thus no single answer to most problems and the examples given in this chapter must be considered with this in mind. It is fair to say, however, that the majority of applications described have developed in circumstances where there is a wide choice of possible solutions. Some readers will have found alternative or better answers to specific problems.

To illustrate the widespread use of engineering coatings, applications have been grouped under a series of industrial headings, with a selection of typical examples under each heading. The lists are not intended to be exhaustive and new applications are being developed continuously. Many industries share similar problems and a study of how other industries have solved their difficulties can often provide the practising engineer with ideas to apply in his/her own sphere of activity.

The aircraft industry

Air travel is a competitive business in which running costs must be tightly controlled while safety and reliability must not be compromised. Military aircraft demand high reliability and exceptional performance and the widespread use of engineering coatings in both types of aircraft, including their engines, is a good indication of their importance and usefulness (Fig. 11.1 and 11.2).

The engine is clearly a major part of the aircraft and application of coatings to engine parts has been practised for over 50 years. In the early days of piston engines, exhaust valves were (and still are) weld surfaced on their seal faces and often all over the crown to maximise life and reliability.
when working at high temperatures and in the presence of corrosion by the products of burnt fuel.

The special demands of gas turbine engines have provided a great opportunity for surface coating as the combination of special properties required, high temperature strength/corrosion resistance/bearing properties, etc., are often impossible to provide from a single material. Many engine parts are fabricated from thin sections of material that would present problems of distortion if welded. This has led to the extensive use of thermal spraying, a process that as noted earlier can also apply non-metallic coatings, and has allowed application of thermal barrier materials which effectively reduce the operating temperature of the metallic alloy core of the part.

The fact that thermal spraying can be carried out at low temperatures (compared with welding) means that optimum properties are retained in the substrate material and that reprocessing of the coating can often be carried out after a period in service.

Electroplating is used to protect many parts against atmospheric corrosion and thermally sprayed polymers also find application for the same purpose.

Typical applications are given in Table 11.1.

**Chemical and petroleum industries**

The chemical and petroleum industries cover a wide range of activities, from extraction of raw materials to production of highly refined end products. Thus all forms of wear are found, with special emphasis on corrosion, erosion and also high temperatures in some applications.
Table 11.1 Aircraft industry: typical applications

<table>
<thead>
<tr>
<th>Component</th>
<th>Function</th>
<th>Surfacing material</th>
</tr>
</thead>
<tbody>
<tr>
<td>Engines</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intake ducts, engine seals</td>
<td>Sliding, fretting wear</td>
<td>Tungsten carbide/cobalt</td>
</tr>
<tr>
<td>Compressor blade contact faces</td>
<td>Impact, fretting wear</td>
<td>Tungsten carbide/cobalt</td>
</tr>
<tr>
<td>Combustion chamber</td>
<td>Thermal barrier</td>
<td>Magnesium zirconate</td>
</tr>
<tr>
<td>Turbine blades</td>
<td>Hot corrosion</td>
<td>M–Cr–Al–Y</td>
</tr>
<tr>
<td>Turbine blade shrouds</td>
<td>Fretting wear</td>
<td>Chromium carbide/NiCr</td>
</tr>
<tr>
<td>Nozzle guide vanes</td>
<td>Thermal barrier</td>
<td>Multilayer</td>
</tr>
<tr>
<td>Knife edge seals</td>
<td>Abrasive wear</td>
<td>Tungsten carbide/cobalt</td>
</tr>
<tr>
<td>Airframes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>De-icing heaters</td>
<td>Electrical resistance</td>
<td>High speed steels</td>
</tr>
<tr>
<td>Flap and slat tracks</td>
<td>Abrasive wear</td>
<td>Tungsten carbide/cobalt</td>
</tr>
<tr>
<td>Rivets</td>
<td>Atmospheric corrosion</td>
<td>Nylon</td>
</tr>
<tr>
<td>Ground installations</td>
<td>Atmospheric corrosion</td>
<td>Austenitic stainless steels</td>
</tr>
</tbody>
</table>

To meet these conditions of service special materials are required, which if used for one piece construction of components would involve high costs. Surfacing, using appropriate materials applied to the exposed areas of cheaper substances, enables costs to be minimised, life to be increased and often allows resurfacing to be carried out after a period of use (Fig. 11.3–11.6).

Selection of surfacing materials may need careful study, for example small
changes in operating temperature or composition of the chemical environment can have a significant effect on component life. Certain points need to be kept in mind, for example:

1 Corrosion – assuming the selected coating material is considered to have satisfactory corrosion resistance in the medium and at the temperature involved, the question may arise as to its electrode potential compatibility with the substrate and any other constructional materials. Porosity in thermally sprayed coatings may need to be sealed.

2 Abrasion – materials that depend on the formation of surface films to minimise the rate of attack perform badly if this film is continuously removed by abrasion or erosion.

3 Adhesive wear – some materials of high corrosion resistance such as austenitic steels are prone to adhesive wear and ‘pick-up’ or seizure can easily occur when rubbing against themselves and similar materials.

4 Temperature – the expansion coefficients of the material used may need to be watched, because of the big differences between some substrate materials and some coatings. Possible problems to arise include cracking of the surface coating, distortion in use (especially if used at high temperatures) and loss of running clearances.
11.5 Centrifugal separator screw surfaced with cobalt alloys to resist abrasion and corrosion on the diameter and faces of the flights (courtesy Deloro Stellite Group).

11.6 HVOF spraying, using a JP5000 torch on a ball valve for the petrochemical industry, with 0.4mm tungsten carbide/cobalt to resist wear. The surface will be ground and super finished before use (courtesy Eutectic Co Ltd).
Table 11.2  Chemical and petroleum industries: typical applications

<table>
<thead>
<tr>
<th>Component</th>
<th>Surfacing material</th>
</tr>
</thead>
<tbody>
<tr>
<td>Abrasion</td>
<td>Carbon steel and carbides</td>
</tr>
<tr>
<td>Oil well rock drill bits</td>
<td>Co-Cr-W high alloy</td>
</tr>
<tr>
<td>Oil well rock drill bit bearings</td>
<td>Carbon steel and carbides</td>
</tr>
<tr>
<td>Drill collars</td>
<td>Ni-Cr-Si-B</td>
</tr>
<tr>
<td>Mud pump sleeves, rods, plungers</td>
<td>Carbon steel and carbides</td>
</tr>
<tr>
<td>Tool joints</td>
<td>Carbon steel and carbides, or WC pads in Ni-Cr-Si-B matrix</td>
</tr>
<tr>
<td>Stabilisers</td>
<td>Ni-Cr-Si-B</td>
</tr>
<tr>
<td>Centrifugal separator screw flights</td>
<td>Ni-Cr-Si-B and carbides</td>
</tr>
<tr>
<td>Conveyor screws</td>
<td>Ni-Cr-Si-B and carbides, or carbon steel and carbides</td>
</tr>
<tr>
<td>Divertor spools</td>
<td>Ni-Cr-Si-B</td>
</tr>
<tr>
<td>Ball valves</td>
<td>Tungsten carbide/cobalt</td>
</tr>
<tr>
<td>Adhesive wear</td>
<td>Co-Cr-W medium alloy</td>
</tr>
<tr>
<td>Mixer seals</td>
<td>Co-Cr-W medium alloy or tungsten carbide/cobalt</td>
</tr>
<tr>
<td>Extrusion screw flights</td>
<td></td>
</tr>
<tr>
<td>Corrosion</td>
<td>Aluminium, zinc or polymer</td>
</tr>
<tr>
<td>Oil storage tanks, structural steelwork</td>
<td>Austenitic stainless steels, Ni-Cr-Si-B, or Co-Cr-W-Ni-Si-B alloys</td>
</tr>
<tr>
<td>Crude oil heat exchangers</td>
<td>Nylon</td>
</tr>
<tr>
<td>Tank heating coils</td>
<td>AUSTENTIC STAINLESS STEELS Ni-Cr-Mo-Al-Ti, Co-Cr-W low alloy, Co-Cr-W medium alloy, WC/cobalt</td>
</tr>
<tr>
<td>Thermometer pockets</td>
<td></td>
</tr>
<tr>
<td>Pipeline and flow control valves, valve trim</td>
<td>Ni-Cr-Si-B, Ni-Cr-Si-B, Ni-Cr-Mo-Al-Ti, Co-Cr-W low alloy, Co-Cr-W medium alloy, WC/cobalt</td>
</tr>
<tr>
<td>Erosion</td>
<td>As for pipeline valves</td>
</tr>
<tr>
<td>Valves</td>
<td>Austenitic or martensitic irons</td>
</tr>
<tr>
<td>Pump impellers</td>
<td>Co-Cr-W-Ni-Si-B alloys, WC/cobalt, chromium oxide</td>
</tr>
<tr>
<td>Pump shafts and sleeves</td>
<td></td>
</tr>
<tr>
<td>Pipe elbows and bends</td>
<td>Austenitic irons, Ni-Cr-Si-B.</td>
</tr>
</tbody>
</table>

Typical applications are given in Table 11.2. These examples are grouped according to the principal wear factor involved.

Earthmoving, agricultural, quarrying and mining

Earthmoving, agricultural, quarrying and mining industries use plant that suffers primarily from abrasive wear combined with varying degrees of impact (Fig. 11.7 and 11.8). The abrasive wear may involve high or low stress contact with the abrasive material.

Alloys used for hardfacing to resist this type of wear rely on the hardness of the metallurgical phases present in their structures and, in general, increasing the hardness confers increasing resistance to abrasive wear. However, resistance to impact tends to decrease with increasing hardness and in the extreme can lead to cracking or chipping of the surface coating.

Hardness acquired as a result of work-hardening through impact – as with
Excavator bucket for limestone, hardsurfaced by MMA using alloy steel deposit (courtesy ESAB Group (UK) Ltd).

Extensive use of engineering coatings extends the life of the toothed buckets on this excavation unit (courtesy ESAB Group (UK) Ltd).

Austenitic manganese steels – does not lead to the same loss of toughness as the raised hardness may imply, hence its extensive use on parts exposed to abrasion and impact loading.

Deposits of alloys of the harder austenitic matrix type may be subject to contraction cracking (relief cracking) if applied in thick layers or on large areas. Where extensive building up is required to restore worn areas, it is normal to restore the profile with a deposit of softer and tougher low alloy steel, which limits the amount of hardfacing to be subsequently deposited but provides adequate support.

Table 11.3 shows the general relationship between abrasion resistance and toughness/resistance to impact of the most important groups of material used in these industries. There is, of course, some overlapping between the various alloys in each group and between groups.

The presence of impact calls for a coating that has the strongest possible bond with its substrate and that can provide protection in depth. In consequence, such applications usually call for welded deposits. Thermally
Table 11.3 Relationship between abrasion resistance and toughness/resistance to impact (numbers refer to the classification system in Chapter 3)

<table>
<thead>
<tr>
<th>Group</th>
<th>Abrasion resistance increasing</th>
<th>Impact resistance increasing</th>
<th>Typical uses</th>
</tr>
</thead>
<tbody>
<tr>
<td>6 Tungsten carbide materials</td>
<td></td>
<td></td>
<td>Sinter plant, rock drills, brick and clay tools</td>
</tr>
<tr>
<td>2/5 Hi-Cr complex irons</td>
<td></td>
<td></td>
<td>Hot wear – sinter plant scrapers and screens</td>
</tr>
<tr>
<td>2/4 Hi-Cr martensitic irons</td>
<td></td>
<td></td>
<td>Ball mill liners</td>
</tr>
<tr>
<td>2/3 Hi-Cr austenitic irons</td>
<td></td>
<td></td>
<td>Shovel teeth, screen plates, bucket lips</td>
</tr>
<tr>
<td>2/2 Martensitic irons</td>
<td></td>
<td></td>
<td>Abrasion and adhesive wear</td>
</tr>
<tr>
<td>2/1 Austenitic irons</td>
<td></td>
<td></td>
<td>Crushers, pump casings and impellers</td>
</tr>
<tr>
<td>1/7 Austenitic Cr–Mn steel</td>
<td></td>
<td></td>
<td>Heavy impact, e.g. excavator buckets</td>
</tr>
<tr>
<td>1/6 Austenitic Mn steels</td>
<td></td>
<td></td>
<td>Heavy impact, e.g. crushers, hammers</td>
</tr>
<tr>
<td>1/2 Low alloy steels</td>
<td></td>
<td></td>
<td>Build-up materials, tractor parts</td>
</tr>
<tr>
<td>1/1 carbon steels</td>
<td></td>
<td></td>
<td>Build-up materials</td>
</tr>
</tbody>
</table>

sprayed deposits find application on some bearing surfaces not normally exposed directly to impact by the minerals being handled.

The alloy selected and also the pattern of weld beads applied to tools depend on working conditions. Cost-effective deposits do not necessarily need to provide all-over coverage; for example a ‘waffle’ or criss-cross pattern is used when handling wet sand or clay, which packs in between the beads and protects the substrate. A ‘dot’ pattern is used on difficult to weld substrates when overheating must be avoided. When handling rock a series of beads running parallel to the flow of material causes the spoil to rub on them and not on the substrate.

Typical applications are given in Table 11.4.

**Internal combustion (piston) engines**

The ubiquitous piston engine fitted with poppet valves depends on an effective gas-tight seal when the valves are shut to develop its rated power output.

The exhaust valve, surrounded by the hot products of combustion during the exhaust stroke, rises to high temperatures until it shuts and the heat is dissipated through the seat in the engine and the stem. Precise working conditions depend on the engine design, fuel used and operating conditions, but the valve seat surface is exposed to high temperatures, corrosion, thermal and mechanical shock. Temperatures are highest in highly rated
petrol engines and corrosion is greatest in diesel engines running on residual fuels.

The diameter of the valve head may be 25mm or less on small petrol engines, and up to 500mm or more on large diesel engines. A variety of steels and some nickel base alloys are used to provide adequate mechanical properties in the valve at the operating temperatures. It is frequently necessary to provide special coatings on the valve seat areas to minimise valve and seat wear, indentation of seat surface by hard particles produced in the combustion chamber, and to give the best possible protection against erosive wear from the high velocity exhaust gases.

The end of the valve stem is in contact with the valve actuation device, e.g. the tappet or rocker, and valve materials chosen for good high temperature strength and corrosion resistance may need local hardening on the stem tip to resist deformation and wear from the forces necessary to open the valve.

For valve seating surfaces in diesel engines it is normal to use welded deposits of Ni–Cr–Si–B or cobalt alloys, whereas for petrol engines Co–Cr–W–Ni alloy is widely used. Valve stem tips are often welded deposited with Co–Cr–W medium or high alloys.

The valve stem slides in a lubricated guide bush and electrodeposited hard chromium is normally used on this. Manufacture of new valves usually incorporates a mechanised welding operation to deposit the surfacing alloy using purpose-built machines based on the PTA process – as illustrated in Chapter 3 – or TIG/MIG, or less frequently by oxyacetylene torch.

As valve size increases, so does the cost of substrate materials, even when a low alloy stem is friction welded to the head. Thus valves with a crown diameter in excess of about 75mm are regularly restored by resurfacing, an operation often carried out manually in areas remote from more highly developed servicing facilities.

Other important applications of surfacing in the internal combustion
engine field include coated piston rings (Fig. 11.9) and spraying of piston crowns (Fig. 11.10).

Typical applications are given in Table 11.5.

Food

Food is essentially from two sources – animal and vegetable farming. However, much of agriculture, from the field to the table, is mechanised, and the industry therefore faces problems similar to those of the earthmoving and chemical industries, coupled with the overriding need for hygiene, which can, for example, involve the use of corrosive cleaning media.

Farm cultivation of vegetable material including grain, root and stem
crop faces problems of abrasive wear with some impact if soils are stony, and the surfacing materials appropriate to protect cultivation machinery are those discussed in the section of this chapter that deals with earthmoving and associated activities.

Many vegetable and dairy products such as beet and cane sugar, milk and cream are processed and the equipment is usually made of stainless steel for reasons of hygiene. The low hardness and low resistance to adhesive wear of this type of steel indicates the use of more wear-resistant engineering coatings in parts such as pumps and valves, using material of good wear resistance such as nickel and cobalt alloys; see Fig. 11.11.

Processing of meat, involving cutting or mincing operations, calls for durable, sharp cutting edges of corrosion-resistant material and, again, weld surfacing of corrosion-resistant substrates with surfacing by ferrous, nickel or copper alloys provides an effective answer.

Margarine substitutes for butter are produced from vegetable oils. These are extracted from a variety of plants such as seeds and nuts in expeller machines, which comprise a long steel screw running inside a close fitting barrel, rather like a mincing machine, and developing pressure that squeezes the oil from the seed. The residue of vegetable matter is used as a basis for cattle cake. To reduce wear in the screw and barrel, the screw flights are coated with wear- and corrosion-resistant ferrous or cobalt alloys and the barrels are lined with similar alloys by a specialised coating process designed for the purpose.

At the end of the food chain, unused food, together with miscellaneous waste such as woven materials, is frequently comminuted by a waste disposer and flushed into the sewage disposal system. To maintain the sharp cutting edges necessary for this type of duty, welded deposits of hard alloys resistant to corrosion, typically nickel or cobalt alloys are used, depending on the machine part and its working conditions.

Typical applications are given in Table 11.6.

**Forging**

Shaping of components in forgeable alloys is carried out at elevated temperatures by press, hammer or drop forging. This work calls for tools that
**Table 11.6** Food: typical applications

<table>
<thead>
<tr>
<th>Component</th>
<th>Surfacing material</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sugar cane cutter knives</td>
<td>Co-Cr–W low alloy, C steel and carbides</td>
</tr>
<tr>
<td>Shredder hammers</td>
<td>Hi-Cr austenitic irons, C steel and carbides</td>
</tr>
<tr>
<td>Grain crusher rolls</td>
<td>Cr steel</td>
</tr>
<tr>
<td>Pipeline valves and seats</td>
<td>Co-Cr–W low and medium alloy</td>
</tr>
<tr>
<td>Homogeniser valves and seats</td>
<td>Co-Cr–W high alloy</td>
</tr>
<tr>
<td>Pump shafts, sleeves, impellers</td>
<td>Various</td>
</tr>
<tr>
<td>Expeller screws</td>
<td>Co-Cr–W medium alloy, Hi-Cr austenitic irons</td>
</tr>
<tr>
<td>Expeller cage bars</td>
<td>Co-Cr–W low alloy, Co-Cr–W–Ni–Si–B alloys</td>
</tr>
<tr>
<td>Doctor’s knives</td>
<td>Co-Cr–W medium alloy</td>
</tr>
<tr>
<td>Can seaming rolls</td>
<td>Co-Cr–W high alloy, plus TiN by PVD</td>
</tr>
<tr>
<td>Packaging knives</td>
<td>Co-Cr–W medium alloy</td>
</tr>
<tr>
<td>Sewage shredder knives and pumps</td>
<td>Co-Cr–W medium and high alloy</td>
</tr>
<tr>
<td>Brewing containers</td>
<td>Polymer</td>
</tr>
<tr>
<td>Food mixer bowls</td>
<td>Nylon</td>
</tr>
</tbody>
</table>

11.11 Can seaming roll of Co–Cr–W low alloy before coating with TiN by PVD (courtesy Deloro Stellite Group).

resist deformation, abrasion by mill scale and cracking caused by thermal and mechanical stress.

As in other examples already referred to, this requires properties in the surface of tools that are not easy to provide from the materials used for the bulk of the tool. The tools are often quite massive and may be rendered unusable by a small amount of wear on a working face. Weld surfacing with appropriate alloys can prolong the initial life and provide the opportunity to carry out rebuilding of the surfaces when worn.

Selection of coating alloy depends on the working conditions of the tool. Nickel and cobalt alloys are used to provide good high temperature properties (Fig. 11.12 and 11.13). For example, a Ni–Mo–Cr–W alloy is resistant to thermal and mechanical shock and is used on drop forging dies. It workhardens in use which helps it to retain the tool profile without cracking. Tools used for press forging lack the impact required to develop workhardening so a harder alloy which is a modification of a Co–Cr–W low alloy is used.
11.12 Milling die cavity deposited with Ni-Mo-Cr-W alloy to finished size (courtesy Deloro Stellite Group).

11.13 Flash trimming die deposited with Co-Cr-W low alloy (courtesy Deloro Stellite Group).
Table 11.7 Forging: typical applications

<table>
<thead>
<tr>
<th>Component</th>
<th>Surfacing material</th>
</tr>
</thead>
<tbody>
<tr>
<td>Drop forging die impressions</td>
<td>Ni–Mo–Cr–W</td>
</tr>
<tr>
<td>Drop forging die flash lands</td>
<td>Ni–Mo–Cr–W</td>
</tr>
<tr>
<td>Press forging die impressions</td>
<td>Special cobalt alloys</td>
</tr>
<tr>
<td>Clipping beds</td>
<td>Co–Cr–W low alloy</td>
</tr>
<tr>
<td>Hot shears</td>
<td>Ni–Cr–Mo–Al–Ti, Co–Cr–W low alloy</td>
</tr>
<tr>
<td>Rotary forging hammers</td>
<td>Co–Cr–W medium alloy</td>
</tr>
<tr>
<td>Furnace skid rails</td>
<td>Special cobalt alloys</td>
</tr>
</tbody>
</table>

Cutting devices such as shears and clipping beds require a still harder alloy and a Co–Cr–W low alloy is used. Typical applications are given in Table 11.7.

Glass

Production of hollow glasswear such as drink bottles, jars and other containers uses a series of operations to mould or shape a 'gob' of glass that is sufficiently hot to be plastic. At the final shaping stage the tool in contact must extract sufficient heat from the glass to ensure that it retains its shape when ejected. Tools used include split moulds to form the outside of the container and plungers to develop the internal form. Other tools simultaneously form the shape of the neck and profiles to engage with the lid or other closure.

Tools are typically made of cast iron or bronze which gives higher thermal conductivity. When assembled to form a complete mould there are a number of joints between the individual parts. When new there is no visual evidence of these on the formed glass, but after a period of use, corners become worn or even chipped and this shows up on the finished glass container.

To obtain the longest life of the tools and to enable parts that ultimately do wear to be refurbished, a technique was developed using powder welding as described in Chapter 3. Worn corners are first chamfered to provide a small flat platform, perhaps 3mm wide, which is then built up with a self-fluxing alloy of Ni–Cr–Si–B series. One of the softer alloys of this family is used, enabling the original sharp corner to be restored by a hand operation such as filing.

Glass plungers used for piercing the glass gob are surfaced with a similar alloy, but the spray fused process is more suited to the coverage of larger areas than powder welding and has the advantage of giving smoother deposits on such parts that require less work to finish to size (Fig. 11.14).

Typical applications are given in Table 11.8.

Pulp and paper

Paper of many different qualities is manufactured using variations of a basic production process. It may be made from raw materials such as wood


<table>
<thead>
<tr>
<th>Component</th>
<th>Surfacing material</th>
</tr>
</thead>
<tbody>
<tr>
<td>Press/blow plungers</td>
<td>Ni–Cr–Si–B</td>
</tr>
<tr>
<td>Blow–blow plungers</td>
<td>Ni–Cr–Si–B</td>
</tr>
<tr>
<td>Blanks and blow moulds</td>
<td>Ni–Cr–Si–B</td>
</tr>
<tr>
<td>Guide plates and rings</td>
<td>Ni–Cr–Si–B</td>
</tr>
<tr>
<td>Neck rings</td>
<td>Ni–Cr–Si–B</td>
</tr>
<tr>
<td>Take-out jaws</td>
<td>Ni–Cr–Si–B, Ni–Cr–Si–B and carbides</td>
</tr>
</tbody>
</table>

11.14 Glass plungers coated with Ni–Cr–Si–B alloy 40 RC by spray fusing (courtesy Deloro Stellite Group).

fibres, or contain or consist of recycled waste material. The process involves removal of bark from tree trunks, breaking down the wood into chips, forming pulp mechanically/chemically and cleaning/modifying the fibrous structure of the pulp. It is then converted into paper or board and cut to size.

Viewed as a whole the processes involve wear of all types described in Chapter 2 and these occur in various combinations. Corrosion of plant occurs at several stages of manufacture, especially in chemical pulping and bleaching. It would not be unusual for normal materials of plant construction to have a short life in these circumstances and long-term protection is achieved by use of appropriate engineering coatings (Fig. 11.15–11.17).

A coating can also be used to improve traction on a face reel drum. This component has a series of concave recessions and narrow slots to stretch paper and remove excess water from the process. Tungsten
Rotary feed valve for digester manually deposited with Co-Cr-W medium alloy before machining to size (courtesy Deloro Stellite Group).

Paper mill cylinder, 11 m long and 1.5 m diameter metal sprayed with a mixed bond coat of carbon steel and tin-bronze materials and a final layer of mixed martensitic chromium steels and austenitic stainless steels (courtesy Metallisation Ltd).

carbide/cobalt, sprayed with the Axial III system, faithfully followed this profile, and produced a coating of the required roughness for the task (Fig. 11.18).

Use of such protection is not limited to paper manufacture; there are applications in cutting, folding, gluing and printing of paper and board which increase life and reduce downtime.

Typical applications are given in Table 11.9.
Industrial applications of engineering coatings

11.17 Paper drills plasma arc sprayed with tungsten carbide–cobalt material to a thickness of around 0.05mm (courtesy CSIRO, Australia).

11.18 A face reel drum coated with tungsten carbide/cobalt using the Axial III system to improve traction of the paper (courtesy Mettech Corporation).

Plastics and rubber

In the plastics and rubber industries a number of different operations are carried out, each of which has its own group of wear problems. They have been grouped together as there are similarities in certain operations which are employed in the two industries.

Both use extrusion machines that are similar to the extruders mentioned in the section on food industry, although there are significant differences in the design of the screw flights. Protection of the crest of the flights is carried out by weld surfacing using nickel or cobalt alloys, applied by manual and mechanised welding processes (Fig. 11.19). Thermally sprayed deposits tungsten carbide in cobalt are sometimes used. Barrel liners are coated by a spin casting process.
Table 11.9 Pulp and paper: typical applications

<table>
<thead>
<tr>
<th>Component</th>
<th>Surfacing material</th>
</tr>
</thead>
<tbody>
<tr>
<td>Debarking flails</td>
<td>Hi-Cr austenitic irons, C steel and carbides</td>
</tr>
<tr>
<td>Chipper knives</td>
<td>Co-Cr-W medium alloy</td>
</tr>
<tr>
<td>Chipper wear plates</td>
<td>Co-Cr-W low and medium alloys</td>
</tr>
<tr>
<td>Conveyor and feed screws</td>
<td>Hi-Cr austenitic irons, Co-Cr-W medium alloy</td>
</tr>
<tr>
<td>Pulper segments</td>
<td>Hi-Cr austenitic irons, Co-Cr-W high alloy</td>
</tr>
<tr>
<td>Pulper drive shafts</td>
<td>Co-Cr-W-Ni-Si-B alloys</td>
</tr>
<tr>
<td>Dirt trap cones</td>
<td>Ni-Cr-Si-B</td>
</tr>
<tr>
<td>Rotary feed valves</td>
<td>Co-Cr-W medium alloy</td>
</tr>
<tr>
<td>Shaft sleeves for finers, defibrators</td>
<td>Co-Cr-W-Ni-Si-B alloys</td>
</tr>
<tr>
<td>Disintegrator hammers</td>
<td>Co-Cr-W high alloy, C steel and carbides</td>
</tr>
<tr>
<td>Packaging knives</td>
<td>Co-Cr-W medium alloy</td>
</tr>
</tbody>
</table>

Extrusion screw for plastics weld deposited on the flight diameters before grinding to size (courtesy Deloro Stellite Group).

Plastic materials are produced in a semi-finished form as pellets from pelletiser plates and cutters whose cutting edges are coated with a welded or thermally sprayed deposit of tungsten carbide/cobalt using D-gun or Jet Kote processes (Fig. 11.20). Mixers of the Banbury type suffer wear on the internal surfaces of the body and on the rotors; these and the rotor seals are typically protected with welded deposits of cobalt alloys applied by MMA, MIG/MAG or SAW processes (Fig. 11.21).
The protection of biffer screens used in polymer production is difficult. The screens are perforated metal rings approximately 760 mm in diameter, inside which rotates a beater cross. Glass-filled polymer granules pass through the screen, causing wear. A solution to the wear shown in Fig. 11.22 on an unprotected type 321 stainless steel screen has been found by applying a coat of 55% tungsten carbide in a Ni-Cr-Si-B matrix by the spray fuse process. Figure 11.22(b) shows the wear after 21 days of use following application of the coating. The main problem in surfacing is control of distortion.

Typical applications are given in Table 11.10.

**Power generation**

Generation of electricity by thermal means accounts for a number of interesting uses for engineering coatings. Mention has already been made of such uses in gas turbine and diesel engines and this section is concerned with steam turbines coupled to fossil or nuclear fuel heat sources.

Coal fuel is widely used and for large boilers it needs to be pulverised, creating abrasive wear of the pulverisers. When burned, products of combustion cause hot corrosion and the ash content causes abrasion of boiler tubes and induced-draught fans.

Nuclear fuel creates its own problems for mechanisms that control the movement of fuel rods, etc., in the reactor coolant, which are subject to adhesive wear. Steam generated in the boiler/heat exchanger is admitted to the turbines through various control valves which operate at high temperatures and can suffer erosion of seating surfaces. Themally sprayed ratings, deposited using the high energy processes, such as Jet Kote, can prevent wear (Fig. 11.23). The turbine itself can suffer from wear and erosion. A
Plasma transferred arc surfacing of a plastics extrusion screw (courtesy Deloro Stellite Group Ltd).

Biffer screen handling glass filled polymer: (a) after 2 days’ service – unprotected stainless steel; (b) after 21 days’ service following coating with tungsten carbide in a Ni–Cr–Si–B matrix (courtesy ICI).

typical applications are given in Table 11.11.

Steelmaking/general metal forming

The operations required to convert iron ore into rolled steel products cause rapid wear of production plant, much of it due to the high processing temperatures used and the abrasive effects of mill scale.
Table 11.10  Plastics and rubber: typical applications

<table>
<thead>
<tr>
<th>Component</th>
<th>Surfacing material</th>
</tr>
</thead>
<tbody>
<tr>
<td>Extrusion and injection moulding</td>
<td>Austenitic irons, Ni–Cr–Si–B, Co–Cr– W</td>
</tr>
<tr>
<td>screw flights</td>
<td>medium alloy, WC/cobalt</td>
</tr>
<tr>
<td>Rotary chopper blades</td>
<td>Co–Cr–W medium alloy</td>
</tr>
<tr>
<td>Pelletiser plates and cutters</td>
<td>Co–Cr–W medium alloy WC/cobalt</td>
</tr>
<tr>
<td>Banbury mixer rotors, bodies,</td>
<td>Co–Cr–W medium and high alloy</td>
</tr>
<tr>
<td>doors and end plates</td>
<td></td>
</tr>
</tbody>
</table>

11.23  Steam turbine valve spindles coated by Jet Kote using chromium carbide and nickel chrome and with seat surfaces deposited with cobalt base alloys.

11.24  A stationary turbine being sprayed using a JP5000 torch to deposit chromium carbide/nickel chrome to a thickness of 0.25 mm, to reduce wear and erosion (courtesy Eutectic Co. Ltd).
Table 11.11 Power generation: typical applications

<table>
<thead>
<tr>
<th>Component</th>
<th>Surfacing material</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coal treatment plant</td>
<td>Hi-Cr austenitic irons, C steel and carbides</td>
</tr>
<tr>
<td>Boiler tubes</td>
<td>50:50 Ni-Cr</td>
</tr>
<tr>
<td>Induced draught fan blades</td>
<td>C steel and carbides</td>
</tr>
<tr>
<td>Hydroelectric turbine blades</td>
<td>Martensitic chromium steels</td>
</tr>
<tr>
<td>Turbine stop, throttle, intercept valve seats</td>
<td>Co-Cr-W low alloy, or Co-Mo type</td>
</tr>
<tr>
<td>Valve grinding surfaces</td>
<td>Co-Cr-W low alloy, Co-Cr-W-Ni-Si-B</td>
</tr>
<tr>
<td>Steam pipe expansion joints</td>
<td>Co-Cr-W low alloy</td>
</tr>
<tr>
<td>Stationary turbine</td>
<td>Chromium carbide/nickel-chrome (75/25)</td>
</tr>
</tbody>
</table>

11.25 Hot mill roll neck reclaimed with a thermally sprayed deposit of low alloy steel up to 25 mm thickness (courtesy Metallisation Ltd).

Use of engineering coatings has been practised for many years as a means of prolonging life between breakdowns and allowing repeated repairs to large and expensive items of plant (Fig. 11.25 and 11.26). Many of the applications call for thick coatings resistant to thermal and/or mechanical stress and weld-deposited coatings are usually best suited to this type of work. Arc welding is widely used because of the heavy sections of components to be built up.

In wire manufacture, drawing dies and drums can suffer from severe wear
Twin high velocity thermal spray Jet Kote guns applying a deposit of tungsten carbide-cobalt to the surface of a steelworks pickling line roll to a thickness of 0.08–1.5 mm and to a desired roughness between Ra3 and Ra12. Life improvements over an uncoated roll is about 4:1.

problems. Figure 11.27 shows a wire draw drum being surfaced by HVOF with tungsten carbide/cobalt, to protect it from wear.

Typical applications are given in Table 11.12.

**Textiles**

Both natural and synthetic fibres are used in textile production, which is carried out on high speed machines. These speeds lead to rapid wear caused by abrasion from the thread or adhesive wear between machine parts. Use of conventional lubricants is often not possible, because of the risk of staining the product, so surface coatings having good bearing properties can provide an ideal solution (Fig. 11.28).

Corrosion is not normally a problem except in the manufacture of artificial fibres that require coagulation in a corrosive solution. The continuous fibre is cut to form tow of a staple length in the coagulating solution and, for this purpose, knives must retain a sharp cutting edge by use of a corrosion resistant and hard alloy. For this Co–Cr–W medium alloys are used.

Some applications lend themselves to use of Co–Cr–W high alloys in the form of Co–Cr–W high alloy precast inserts, brazed into position, rather than surfacing.

Typical applications are given in Table 11.13.

**Timber**

Growing timber, particularly fast-growing tropical hardwoods, is often contaminated with abrasive mineral matter such as sand or tramp metal such as
### Table 11.12 Steelmaking/general metal forming: typical applications

<table>
<thead>
<tr>
<th>Component</th>
<th>Surfacing material</th>
</tr>
</thead>
<tbody>
<tr>
<td>Coal crushers</td>
<td>Hi-Cr austenitic irons, C steel and carbides</td>
</tr>
<tr>
<td>Ore crushers</td>
<td>Austenitic Mn steels, austenitic irons</td>
</tr>
<tr>
<td>Shovel teeth, bucket lips</td>
<td>Hi-Cr austenitic irons, austenitic irons</td>
</tr>
<tr>
<td>Sinter plant high wear areas</td>
<td>Hi-Cr complex irons, C steel and carbides</td>
</tr>
<tr>
<td>Furnace conveyor skidder bars</td>
<td>Co-Cr-W low alloy</td>
</tr>
<tr>
<td>Bell and hopper seals</td>
<td>Ni-Mo-Cr-W</td>
</tr>
<tr>
<td>Explosion valves</td>
<td>Co-Cr-W low alloy</td>
</tr>
<tr>
<td>Ingot lifting dogs</td>
<td>High speed steels, Co-Cr-W low alloy</td>
</tr>
<tr>
<td>Hot slab shears</td>
<td>Ni-Mo-Cr-W, Co-Cr-W low alloy</td>
</tr>
<tr>
<td>Roll neck abutment rings</td>
<td>Co-Cr-W medium alloy</td>
</tr>
<tr>
<td>Table rolls</td>
<td>Martensitic irons</td>
</tr>
<tr>
<td>Roll journals</td>
<td>Martensitic Cr steels</td>
</tr>
<tr>
<td>Down coiler rolls</td>
<td>Martensitic Cr steels, Ni-Mo-Cr-W</td>
</tr>
<tr>
<td>Bar mill entry guide rolls</td>
<td>Co-Cr-W medium alloy</td>
</tr>
<tr>
<td>Acid pump parts</td>
<td>Co-Cr-W medium alloy</td>
</tr>
<tr>
<td>Galvanising bath rolls, brushes</td>
<td>Co-Cr-W low alloy</td>
</tr>
<tr>
<td>Pickle line rolls</td>
<td>WC/Co</td>
</tr>
<tr>
<td>Tube piercer points</td>
<td>Ni-Mo-Cr-W, Co-Cr-W low alloy</td>
</tr>
<tr>
<td>Rail points and crossings</td>
<td>Low alloy steels, austenitic Cr and Cr-Mn steels</td>
</tr>
<tr>
<td>Crane brake drums</td>
<td>WC/Co</td>
</tr>
<tr>
<td>Roll bearing seals</td>
<td>Chromium oxide</td>
</tr>
<tr>
<td>Plating line seals</td>
<td>Tungsten</td>
</tr>
<tr>
<td>Wire draw drum</td>
<td>Chromium carbide/nickel-chrome</td>
</tr>
</tbody>
</table>

**11.27** Wire draw drum being sprayed by a JP5000 torch with tungsten carbide-cobalt to a depth of 0.5mm to protect against wear (Courtesy Eutectic Co Ltd).
Industrial applications of engineering coatings

**Table 11.13** Textiles: typical applications

<table>
<thead>
<tr>
<th>Component</th>
<th>Surfacing material</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thread guides, overrun rollers</td>
<td>Alumina, chromium oxide</td>
</tr>
<tr>
<td>Machine cams</td>
<td>Co–Cr–W high alloy</td>
</tr>
<tr>
<td>Looper and latch needles</td>
<td>Co–Cr–W high alloy</td>
</tr>
<tr>
<td>Winding spindles</td>
<td>WC/Co</td>
</tr>
<tr>
<td>Flattening rolls</td>
<td>WC/Co</td>
</tr>
<tr>
<td>Crimper rolls</td>
<td>Co–Cr–W medium alloy</td>
</tr>
<tr>
<td>Carpet trimming knives</td>
<td>Co–Cr–W medium alloy</td>
</tr>
</tbody>
</table>

11.28 Thread overrun roller and thread guides coated with ceramic by plasma spraying to protect against frictional wear (courtesy Sulzer Metco Ltd).

wire, nails and bullets. Tree trunks are usually cut with band saws running at high speed, the teeth of which need to have high resistance to abrasive wear and frictional heat.

Until the technique of tipping band saw teeth by weld surfacing was developed some years ago, saw mills tended to accumulate tree trunks which could not be cut economically with normal steel saws (high carbon steel). Hardfacing was originally carried out using an oxyacetylene torch and this is still practised. Special machines have also been developed to do the job automatically using arc welding methods.

The sequence of operations is shown diagrammatically in Fig. 11.29. The swaging operation is the same as that used in normal resharpening to give side clearance to the tooth. As the saw band is a high carbon steel and therefore tends to harden under the weld, a light tempering operation is carried out before each tooth is ground to give the usual cutting edge and angles. Other applications of engineering coatings to equipment handling wood are covered in the section of this chapter dealing with pulp and paper.

Typical applications are given in Table 11.14.

**Transport**

Use of engineering coatings in the transport industry falls into three categories – power units, other vehicle components such as suspension and brakes, and fixed permanent structures such as bridges.


**Table 11.14** Timber: typical applications

<table>
<thead>
<tr>
<th>Component</th>
<th>Surfacing material</th>
</tr>
</thead>
<tbody>
<tr>
<td>Veneer pressure bars</td>
<td>Co–Cr–W medium alloy</td>
</tr>
<tr>
<td>Band saw teeth</td>
<td>Co–Cr–W medium alloy</td>
</tr>
</tbody>
</table>

11.29 Hardfacing of saw teeth: (a) each tooth is swaged; (b) the dimple is filled by hardfacing and the tooth tempered; (c) tooth is ground until alloy front is correctly faced and sharp.

The working environment may be sea, land or air and the last is dealt with in the section on aircraft. Piston engines are covered in their own section and here we are concerned with road, rail and marine transport.

**Road transport: vehicle components**

Those areas of road vehicles exposed to the abrasive effects of grit and corrosion caused by salt can show severe deterioration in a short period of use. Manufacturers are increasingly improving the life of these areas by the use of zinc coatings and wax injection, but these have a limited resistance to abrasion. Thermally sprayed coatings of epoxy-based polymers are now applied to exposed areas such as wheel arches, bumpers and spoilers, and similar coatings are used as body coats on some vehicles. Chassis of heavy transport vehicles are typically sprayed with zinc while worn brake drums are reclaimed with arc sprayed deposits of Cr–Mo steel (Fig. 11.30 and 11.31).
11.30 Semi-trailer flame sprayed with zinc to a depth of about 0.1 mm, to be followed by the normal decorative paint finish, giving a life of about ten years without maintenance (courtesy Metallisation Ltd).

11.31 Reclamation of a passenger bus brake drum by arc spraying an initial coating of molybdenum, followed by a working layer of low alloy steel (courtesy Metallisation Ltd).

**Railways: vehicle components**

Coach coupler knuckles are protected from new with thermally sprayed and fused deposits of tungsten carbide in a matrix of self-fluxing Ni–Cr–Si–B alloy to resist dry rubbing and some impact (Fig. 11.32). Other parts successfully reclaimed include brake torque reaction tubes plasma sprayed with
Manufacture of high speed train couplings incorporates coatings on the rubbing surfaces exposed to unlubricated wear and impact. The coating is a thermally sprayed layer of Ni–Cr–Si–B alloy containing tungsten carbide, which is then furnace fused. The lifetime is raised typically from three months to six years or more (courtesy British Rail).

Rail traction motor shaft bearing seating reclaimed with a gas or arc sprayed deposit of martensitic chromium (13%) steel, before machining to size (courtesy British Rail).

tungsten carbide cobalt, and brake pad actuator pivots sprayed and fused with Ni–Cr–Si–B alloy.

Location surfaces for ball and roller journal bearings that suffer from fretting wear are reclaimed by thermal spraying; Fig. 11.33 shows a traction
motor armature spindle set up for arc spraying with a martensitic 13% chromium steel.

Arc spraying of copper on the surface of coach axles is used to create a brush path. The deposit is subsequently rolled to compact the deposit.

**Railways: track**

Points and crossings are subject to greater wear than track and are repaired with arc welded deposits of steels from the low alloy, or austenitic Mn or Cr–Mn group. Tools used to remove the hot flash from butt welded rails are surfaced with Co–Cr–W medium alloy.

**Marine transport**

The environment of saltwater, often with abrasive sand, shells, etc., in suspension, poses combined wear and corrosion problems for exposed equipment. Typical examples of the use of engineering coatings on such parts are the bearing surfaces of rudders, stabilisers and submarine hydroplanes. These bearing surfaces, which can be up to about 600mm diameter, are weld surfaced with an alloy similar to Co–Cr–W low alloy, giving a substantial improvement in life and avoiding regular and costly dry docking for inspection and possible replacement (Fig. 11.34).

The operating gear of fishing vessels suffers rapid corrosion and wear; sprayed and fused coatings of Ni–Cr–Si–B and Co–Cr–W–Ni–Si–B alloys find application on bearing surfaces while protection against seawater corrosion of other parts can be achieved with sprayed deposits of plastics.
Two interesting onboard ship reclamation jobs are the coating of worn diesel engine piston skirts (approximately 300 mm diameter) with aluminium bronze and of the worn skirt of a propeller shaft bearing, etc., arc sprayed with Monel metal.

Typical applications are given in Table 11.15.
Index

abrasive wear, 10, 309
control, 15
diagnosis, 14
and hardness, 11
test, 35
accessibility, 114
adhesive wear, 17, 309
control, 20
diagnosis, 21
agriculture, 284
air plasma spraying, 137, 309
aircraft, 279
aluminium, 5, 128
anodising, 173, 185, 309
arc wire spraying, 135, 309
austenitic chromium-manganese steel, 3, 48, 55
austenitic irons, 3, 48, 56
austenitic manganese steel, 3, 47, 55
austenitic stainless steel, 3, 33, 47, 54, 77
autocatalytic deposition, 171
barrel plating, 169
blank preparation, 94
bond coats, 123, 162, 309
brush plating, 170
buffer layers, 92
bulk welding, 67
carbon steels, 3, 33, 45
cavitation erosion, 24, 309
coating rate, 222
coating thickness, 222
cost, 222
design for, 223
plasma activated, 219
safety, 272
chromium, 6, 174
boride, 61
carbide, 129
ceramics, 5, 6, 129, 131
cermets and graded coatings, 129, 309
cemented carbides, 129
chemical and petroleum industry, 280
chemical vapour deposition, 194, 195, 215, 309
cobalt, 4, 49, 59, 180
alloys, 4, 6, 59, 90, 123, 127
-chrome carbide, 6, 184
complex irons, 56
compressed gas, 260
concentration cells, 32
consumables (tables), 3, 46
copper, 4, 5, 50, 60, 128, 179
tin, 4, 6, 50, 179
corrosion
fatigue, 34
gaseous, 25
liquid, 28
resistance, 28, 41
testing, 40
CoSHH, 263
cracks, 21, 33, 87, 99, 116, 117, 253, 254
defects, 87, 115
design
chemical vapour deposition, 223
electrodeposition, 186
physical vapour deposition, 223
polymer coatings, 233
quality assurance, 267
Index

thermally sprayed coatings, 157
welded coatings, 92, 94
detonation coating, 143, 310
Diamond jet, 145
dilution, 88, 310
distortion control, 97
drilling, 239
earth moving, 284
electric shock, 269
electrodepositon, 6, 167
brush plating, 170
design, 186
finishing, 192
materials, 174, 186
safety, 272
substrate materials, 192
surface preparation, 192
vat plating, 169
electrode potentials, 30
electroless deposition, 171, 310
electroslag, 72
electrostatic spray, 231
enclosed spaces, 265
erosion, 14, 24, 310
cavitation, 24
cavitation, control of, 25
spark, 241
test, 35
evaporation materials, 201
sources, 200
vacuum, 199, 310
fatigue, 21
contact, 21
corrosion, 34
surface, 21
ferrous alloys, 3, 123
finishing, 235, 310
fire, 262
flame spraying, 227, 310
fluidised bed coating, 230
flux cored arc welding, 66
food processing plant, 208
forging, 289
fretting, 19, 311
friction surfacing, 44, 75
adhesion, 76
applications, 81
galvanic cells, 30
series, 31
gas scattering deposition, 203
Gibbs free energy, 26
glass forming tools, 292
gold, 6, 183
grinding, 240, 271
cylindrical, 240
dry and wet, 244
parameters, 240
safety, 271
surface, 240
wheel selection, 241, 243
hard anodising, 173, 185
hardness and wear, 11
heat affected zones, 85, 118, 311
high chromium irons, 3, 56
high speed steels, 3, 47, 54
high stress abrasion, 11
high velocity oxyfuel spraying, 144, 311
hardening, 145
humidity cabinets, 40
HV2000, 145
hydrogen induced cracking, 33
inclusions, 117, 118, 254
inspection, 87, 116
internal combustion engines, 286
ion implantation, 195, 214, 311
materials, 215
ion plating, 195, 203, 311
materials and applications, 204, 208
reactive, 207
sputter, 208
irons, 3, 6, 48, 55, 56, 87, 180
Jet Kote, 145
JP5000, 145
lapping, 241
laser cladding, 44, 71
radiation, 268
lead, 6, 180
finishing, 245
lubrication, 19, 21, 33, 34
milling, 238, 245
allowances, 108
manual metal arc welding, 62, 312
marine transport, 307
martensitic chromium steels, 3, 46, 54
martensitic irons, 3, 56
masking, 163
materials for coatings (tables), 3, 46
electrodeposition, 174, 186, 192
evaporation, 201
ion implantation, 204, 208
polymer coating, 227
sputter coatings, 213
thermal spraying, 122 et seq
wear resistance, 38
weld surfacing, 45 et seq
mechanised surfacing, 101, 111
metal active gas welding, 65, 312
metal inert gas welding, 65, 312
dry and wet, 122
mild and severe wear, 18
milling, 238, 245
mining, 284
nickel, 3, 6, 57, 176
  alloys, 3, 49, 58, 91, 123, 126
    – boron, 178
    – phosphorus, 178
    – silicon carbide, 4, 6, 185
noise, 265
occupational exposure limits, 263
oxidation, 27
oxyacetylene, 69
oxyfuel gas spraying, 132
palladium, 184
paper, 292
personal protection, 265, 266, 267, 268,
  271
phosphating, 173
physical vapour deposition, 194, 195, 312
  design for, 223
  safety, 271
pin-on-disc test, 37
pitting, 23
plasma activated CVD, 219
plasma transferred arc, 64
plastics and rubber, 295
platinum group, 6, 183, 184
polishing, 245
polymer coatings, 5, 226
  design for, 233
  materials, 227
porosity, 117, 254
porous coatings, 257
post weld heat treatment, 85
Pourbaix diagram, 29
powder welding, 70, 312
power generation, 297
preheat, 84, 99, 312
pulp and paper, 292
quality, 246
  component/substrate requirements, 248
consumables, 248
finishing, 250
  in design, 247
  testing and inspection, 251
thermal spray, 255
welding, 252
quarrying, 284
radiation, 266
railways, 305
reactive ion plating, 207
relief cracking, 92, 99
residual stress, 84
resistance welding, 44, 73
resurfacing, 44
rhodium, 6, 184
road transport, 304
robots, 138
rolling fatigue, 21
running-in wear, 19
ruthenium, 184
safety, 259
  abrasive blasting, 269
acetylene, 261
compressed gas, 260
CoSHH, 263
chemical vapour deposition, 272
electric shock, 269
electrodeposition, 272
enclosed spaces, 265
fire, 262
grinding, 271
lasers, 268
noise, 265
physical vapour deposition, 271
radiation, 266
solvents, 270
salt spray cabinets, 40
scuffing, 19, 313
sealing, 165, 313
silver, 6, 182
solvents, 270
spark erosion, 241
spray fused alloys, 152, 156, 313
sputter coating, 195, 209, 313
  materials, 213
sputter ion plating, 208
steels,
  austenitic Cr-Mn, 3, 48, 55
  austenitic manganese, 3, 47, 55
  austenitic stainless, 3, 33, 47, 54
  carbon, 3, 33, 45
  high speed, 3, 47, 54
  low-alloy, 3, 33, 45, 46
  martensitic chromium, 3, 46, 54
  steelmaking, 298
stress/corrosion, 33
stress relief, 313
strip consumables, 67, 73
structure and wear, 11
submerged arc welding, 66, 313
Super D-Gun, 144
surface preparation, 313
surfacing practice, 246
testing and inspection, 251
textiles, 301
thermal spray coatings, 2, 119, 313
  adhesion to substrate, 119
  arc wire, 135
  air plasma, 137
  bond coats, 123, 162
  design for, 157
  detonation coating, 143
  ferrous alloys, 123
  grinding, 243
  high velocity oxyfuel, 144
  finishing, 245
  oxyfuel gas, 132
  planing and shaping, 244
polishing, 245
powder morphology, 122
quality, 255
sealing, 165
surface preparation, 161
thickness, 158
turning, 242
vacuum plasma, 140
wires, 122
three body abrasion, 11
timber, 301
tin, 6, 128, 181
   – nickel, 6, 182
Top Gun, 145
transport, 303
tungsten carbide, 4, 50, 60, 89, 129, 130
tungsten inert gas welding, 63, 313
turning, 237
vacuum evaporation, 199
vacuum plasma spraying, 140, 314
vapour deposition, 6
ventilation, 264
wear, abrasive, 10
   control, 15
   diagnosis, 14
   and hardness, 11
adhesive, 17
   control, 20
   diagnosis, 21
resistance
   material choice, 38
welding, 2
   adhesion, 42
   bulk welding, 67
   coating area, 109
   coating thickness, 42
cracks, 116
defects, 115
design, 92, 94
dilution, 43, 54, 88
distortion control, 97
electroslag, 44, 72
explosive, 74
flux cored arc, 43, 66
heat affected zone, 83, 85, 118
machining allowances, 108
manual vs mechanised, 111
metallurgical effects, 82
metal inert/active gas welding, 43, 65
manual, metal arc welding, 43, 62
oxyacetylene, 44, 69
patterns, 104
preparation, 92
powder, 44, 70
plasma transferred arc, 43, 64
quality, 252
resistance, 73
submerged arc welding, 43, 66
substrate defects, 87
surfacing, 42
temperature cycle, 84
thickness, 43, 93
tungsten inert gas welding, 43, 63