Dental Materials and Their Selection
- 3rd Ed. (2002)

by William J. O'Brien
DENTAL MATERIALS AND THEIR SELECTION - 3rd Ed. (2002)

Front Matter

Title Page

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Quintessence Publishing Co, Inc


Library of Congress Cataloging-in-Publication Data

Dental materials and their selection / edited by William J. O'Brien.—
3rd ed.
p.; cm.
Includes bibliographical references and index.
ISBN 0-86715-406-3 (hardback)
1. Dental materials.
RK652.5. D454 2002
617.6'95—dc21
2002003731

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4350 Chandler Drive
Hanover Park, IL 60133
www.quintpub.com
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Acknowledgments

I would like to thank the many contributors to the third edition who took the time and used their expertise to keep this book current in a field that has seen many changes in recent years. Several contributors to the second edition are also recognized for their valuable contributions: Dr Pui L. Fan, Dr Evan H. Greener, Carole L. Groh, Dr Valerie A. Lee, and Dr Mathijs M. A. Vrijhoef.

I appreciate the many people who contributed data to the biomaterials properties tables, especially Drs Hal O’Kray and Abe Jarjoura, who provided major additions to the data on restorative and impression materials, respectively. Chris Jung contributed many excellent illustrations, and Elizabeth Rodriguiz was invaluable in preparing the manuscript. Finally, I want to acknowledge the staff at Quintessence for their expert assistance in helping me prepare the book for publication.

Introduction

In revising this book for a third edition, the current situation confronting academic dental materials was considered. On one hand, dental materials is one of the most popular subjects among those who pursue continuing education seminars and read the dental literature. On the other hand, most dental students think of dental materials as a basic science course, filled with facts and concepts that have little application to clinical dentistry. A perusal of current dental textbooks on restorative dentistry and prosthodontics reveals that such texts cover much of the subject matter formerly taught only in dental materials courses. This sign of our success in integrating dental materials into dental education and research is also a sign that the dental materials curriculum must continue to evolve to maintain its vital position as an intellectual leader in dental education. More and more of the traditional approach simply will not do for this third edition. Instead, we must seize the opportunity to move the field of dental materials education forward to tackle two major challenges in dentistry: the proliferation of products and techniques and the information explosion in science and technology. The recent proliferation of dental products may lead to improved patient care, but keeping up with the new technology is a challenge to dental materials specialists and educators.

Dental materials textbooks have evolved significantly over the past century. An early textbook on dental materials provided recipes for a handful of materials (three cements, amalgam alloys, gold foil, vulcanized rubber, and gold casting materials) and emphasized formulation, techniques, and crude testing. Then came the research and development period, when dental materials properties were optimized by the dentist according to the results of laboratory testing and ADA standards were developed. Dental materials have been further refined to offer simpler techniques for clinicians and to meet the increasing esthetic demands of middle-class patients in developed countries.

Another dimension to proliferation is the large number of products and techniques available for each type of material, which only intensifies the need for dentists to stay current with the literature. To ease this burden, publications such as Clinical Research Associates Newsletter, Dental Advisor, and Reality compile new information and provide monthly updates for dental practitioners. Perhaps the greatest drawback of proliferation is that many new materials are not sufficiently tested prior to full-scale marketing, thereby increasing the risk of clinical failures.

As a result of this product explosion, dental materials education has an opportunity to
become a more integral part of the overall curriculum, but to do so it must revise its approach to teaching. A long-standing problem is that dental materials courses are grouped with basic sciences, which tends to encourage memorization of facts rather than understanding of clinical application. A new approach would be more pragmatic, integrating problem-based learning and evidence-based dentistry with the traditional overview of clinical materials and materials science concepts, which is still important. Table 1

<table>
<thead>
<tr>
<th>Material / Estimated longevity</th>
<th>Indications</th>
<th>Contraindications</th>
<th>Strengths</th>
<th>Weaknesses</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amalgam, silver /14 y</td>
<td>Incipient, moderate-sized, and some large lesions in adolescents and adults</td>
<td>Large intracoronal restorations (cusp replacement); endodontically treated teeth</td>
<td>Good marginal seal; strength; longevity; manipulability; cariostatic activity</td>
<td>Objectionable color; stains tooth; marginal breakdown; alleged health challenges</td>
</tr>
<tr>
<td>Cast gold (inlays, onlays, and crowns) / 20 y</td>
<td>Large lesions; teeth requiring additional strength; teeth used in rebuilding or changing</td>
<td>Adolescents; high caries activity; persons who object to gold display</td>
<td>Reproduces anatomy well; onlays and crowns may increase strength of tooth; longevity; wears occlusally</td>
<td>Time required for placement; high fee; poor esthetics; thermal sensitivity</td>
</tr>
<tr>
<td>Material</td>
<td>Use</td>
<td>Advantages</td>
<td>Disadvantages</td>
<td></td>
</tr>
<tr>
<td>----------------------------------</td>
<td>----------------------------------------------------------------------</td>
<td>-----------------------------------------------------------------------------</td>
<td>--------------------------------------------------------------------------------</td>
<td></td>
</tr>
<tr>
<td>Ceramic crowns /15 y</td>
<td>Restoration of teeth requiring good appearance and moderate strength</td>
<td>Heavy occlusal stress; bruxism; fixed prosthesis longer than three teeth</td>
<td>Esthetics; no metal content; Have only moderate strength; require resin bonding for strength</td>
<td></td>
</tr>
<tr>
<td>Ceramic inlays and onlays (fired or pressed) / 10 y</td>
<td>Class 2 and 5 locations where high esthetics is desired</td>
<td>Teeth that are grossly broken down and require crowns</td>
<td>Esthetic potential extremely high; properly etched tooth and restoration may increase strength of tooth; onlays stronger than inlays</td>
<td></td>
</tr>
<tr>
<td>Compacted golds (gold foil, powdered gold, mat gold) / 24 y</td>
<td>Initial Class 3 and 5 lesions for patients of all ages</td>
<td>Periodontally unstable teeth; high caries activity; persons who object to gold display</td>
<td>Marginal integrity; longevity; Time-consuming; poor esthetics</td>
<td></td>
</tr>
<tr>
<td>Compomer / 10 y</td>
<td>Moderate to high caries activity; repair of crowns; pediatric</td>
<td>Occlusal stress; locations where color stability is necessary</td>
<td>Moderate fluoride release; easy to use; Color degrades</td>
<td></td>
</tr>
<tr>
<td>Glass ionomer / 8 y</td>
<td>Class 1 and 2 High caries activity; crown repairs</td>
<td>Areas of high esthetic need; areas of difficult moisture control</td>
<td>High fluoride release; Only fair esthetics; difficult and time-consuming to place</td>
<td></td>
</tr>
<tr>
<td>Hybrid ionomer / 10 y</td>
<td>High caries activity; repair of crowns; pediatric</td>
<td>Occlusal stress; locations where color stability is necessary</td>
<td>High fluoride release; tri-cured; sets in dark; Somewhat difficult to use; color degrades</td>
<td></td>
</tr>
<tr>
<td>Porcelain-fused-to-metal crowns / 20 y</td>
<td>Teeth that require full coverage and are subject to heavy occlusal forces; fixed prosthesis Class 1 and 2</td>
<td>Heavy occlusal stress; bruxism</td>
<td>Strength; good marginal fit; acceptable to excellent esthetic result; Appearance not as good as some others; possible wear of opposing teeth</td>
<td></td>
</tr>
<tr>
<td>Resin</td>
<td>Class 1 and 2 Bruxers and</td>
<td>Esthetics; may</td>
<td>Wear of</td>
<td></td>
</tr>
<tr>
<td>Material Type</td>
<td>Characteristics</td>
<td>Indications</td>
<td></td>
<td></td>
</tr>
<tr>
<td>---------------</td>
<td>-----------------</td>
<td>-------------</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Composite (Class 1, 2) / 10 y</td>
<td>Areas of high esthetic need; patients sensitive to metal</td>
<td>Clenchers strengthen tooth with acid-etch concept; restoration during service; no cariostatic activity; may cause tooth sensitivity if bonding agents are not used adequately</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Resin composite (Class 3, 4, 5) / 15 y</td>
<td>Incipient to large Class 3, 4, and 5 lesions</td>
<td>Teeth where coronal portion is nearly gone; Esthetics; ease of use; strength</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

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summarizes the characteristics and indications of current restorative materials. An understanding of the properties and behavior of materials is essential for selection and clinical service. Problem-based learning and evidence-based dentistry would be the links between basic science and clinical practice.

**Problem-Based Learning**

Problem-based learning is an approach that focuses on developing the skills a student will need as a practicing dentist. In the dental materials curriculum, this includes selecting restorative materials as part of treatment planning, explaining their application to patients, handling materials for optimal results, and correcting problems in their clinical performance. A well-designed dental materials course will present not only a materials science framework but also the most current information on available materials. It should emphasize the selection of competing materials for a given clinical situation, taking into account not only material properties but also factors such as patient goals and financial situations. The clinical scenarios that were introduced in the second edition of this book proved to be helpful exercises in choosing the most appropriate materials, and therefore their number has nearly doubled for this edition. They present many facts about materials yet promote an understanding of the clinical application. While experts may disagree with some of the outcomes of these scenarios, their purpose is to reinforce the rational decision-making process necessary for treatment planning.

**Evidence-Based Approach**

The concept behind evidence-based dentistry originated in medicine about 20 years ago, the premise being to base clinical decisions on factual evidence from scientific studies. In the area of dental materials, evidence-based dentistry is used to evaluate and determine the clinical application of new materials. A hierarchy of the different types of evidence available for assessing the clinical performance of new biomaterials is shown in Fig 2. The most rigorous type of evidence is published data from large-scale, long-term clinical trials. Publication in a peer-reviewed journal gives assurance that the design and results of a study have been reported according to acceptable statistical approaches. Because the life cycles of dental materials are growing shorter, both critical thinking and knowledge of basic materials science are necessary to make competent, rational choices. This new
The level of evidence needed to evaluate a new material or technology depends on the level of innovation or the level of risk as compared with the conventional material or technology. The higher the level of innovation and the greater the potential for harm or financial loss to the patient or dentist, the higher the standard for evidence. Materials and technology that are entirely new for a given application have the highest level of innovation and risk. For example, dentin etching for the purpose of bonding composite materials to dentin was highly innovative and had many risks when it was first introduced. The next, lower level of innovation includes major product changes in a conventional material, such as with high-copper amalgam alloys in the 1970s or hybrid ionomer cements in the 1990s. Decreasing levels of innovation and risk include minor product improvements (eg, more shades for a resin restorative material). The majority of "new products" fall into this category.

Each category in the hierarchy of evidence is described below.

**Large-scale, long-term clinical trials**

A well-designed clinical trial will have a clearly stated hypothesis about the clinical performance of a new material when compared with a control material. It will have a large number of subjects to be sufficiently definitive. A good design will also reduce subjectivity by using methods such as calibration of observers, double-blind procedures, and randomization, and the institutional review board of the organization will protect the study participants. These studies are indicated for adoption of brand new innovations and major product changes.

**Other clinical studies**

Other types of clinical study generally are not as decisive as full clinical trials, but they nevertheless provide valuable information. A cohort study would follow a group of patients who receive biomaterials, for instance, and record successes and failures related to their characteristics. Follow-up studies evaluate product longevity and causes of failure in patients who are treated in a clinic. A significant finding might be one in which a researcher discovers a high failure rate of a new biomaterial as patients return for replacement within a short period of time. Short-term clinical studies performed for new dental restorative materials by manufacturers are common and useful, but they miss long-term effects and less frequent problems that are usually only evident in larger groups of patients. Studies in this category are indicated for product improvements and new techniques.

**Animal experiments**

Several of the biocompatibility tests for new materials involve animal testing. Animal tests are valuable, but they are often difficult to interpret. Cytotoxicity screening tests with cell cultures will detect gross toxicity of a material, but subtle effects require expert interpretation. Animal tests are required for new compositions or techniques with questionable biocompatibility.

**Physical properties data**

The publication of physical properties data on new biomaterials is essential for predicting
successful performance, as compared with a standard material. However, since conditions in the body are highly complex, data from laboratory tests cannot always be extrapolated to clinical performance. For example, a new material may be strong when tested in the laboratory, but it may deteriorate more rapidly in body fluids and thus may not be an improvement when compared with a standard material. It is important to evaluate all applicable physical properties of a new material alongside its clinical trials. Physical properties data are usually necessary for minor improvements in materials, but they are insufficient for products with a higher level of innovation.

In vitro experiments

The biomaterials literature has many examples of laboratory experiments designed to simulate the clinical situation. For instance, the wear resistance of biomaterials is often assessed with toothbrushing machines that use thousands of cycles to simulate years of daily brushing. Although useful, these experiments are tricky to interpret. In one such study using a toothbrushing test, a new porcelain glaze was reported to be more resistant to wear than the current glazes. It was later disclosed that no dentifrice had been used. Thermocycling, marginal leakage, adhesion testing, and corrosion testing are a few examples of in vitro tests. They are useful for all new products and techniques.

Deductions from clinical experiments and scientific theories

Deductive reasoning is frequently used to support the superiority of new materials, but it can be unreliable without supporting data. One example is the conclusion that the caries rate will be reduced when fluoride-containing materials are used. Original clinical research on fluoride-containing silicate cements reported that these materials were associated with a low caries rate. The deduction that other fluoride-containing restorative materials provide equal caries protection is often unsupported by clinical data. Another example is the claim that a new high-strength ceramic will have a low clinical failure rate for posterior crowns. It may be strong, but dental laboratory fabrication and the oral environment may contribute to clinical failure. This type of evidence is useful during product development, but very speculative for new products.

Product literature from the manufacturer

There are too many fallacies and extreme claims in dental advertising for this to be a reliable source of evidence. Advertisements that provide references to the published literature are more reliable than those that do not cite published studies.

Popular media, rumors, and myths

None of these is reliable.

Recommended Reading


Chapter 1. A Comparison of Metals, Ceramics, and Polymers

Introduction

When a dentist considers the type of restoration to place in a patient's mouth, the choice may be between different varieties of the same material, for example, different types of amalgam, or between two kinds of the same basic material, such as two kinds of metal—amalgam and cast gold. With the rapid developments in dental materials over the past several years, it is more common that the dentist's choice is between two basic materials, such as between a metal amalgam and a polymer-and-ceramic composite, or between a metal crown and an all-ceramic crown.

A wide spectrum of properties is present within each basic material type; nevertheless, there is a "family resemblance" among the varieties of each material type. For example, although metals exhibit a wide range of strengths, melting ranges, and so on, they resemble one another in their ductility, thermal and electrical conductivity, and metallic luster. Similarly, ceramics can be characterized as strong yet brittle, and polymers tend to be flexible (low elastic modulus) and weak. These "family traits" of the three basic materials are more easily understandable, and thus more easily remembered, if we know the reasons behind them. In fact, simply understanding one key concept for each of the three basic materials gives us significant insight into how each class of materials behaves as a restorative dental material, as well as an idea of the potential of these materials if some of their limitations can be overcome. The relationships among the three basic materials is shown in Fig 1-1.

![Fig 1-1 A tree diagram classifying the three basic materials.](image-url)
Selection Among Various Materials

<table>
<thead>
<tr>
<th>Properties</th>
<th>Metals</th>
<th>Ceramics</th>
<th>Polymers</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Alloy Intermetallic compounds</td>
<td>Inorganic Crystalline Glasses</td>
<td>Rigid Rubbers</td>
</tr>
<tr>
<td>Hardness</td>
<td>Medium to hard</td>
<td>Medium Hard</td>
<td>Soft Very soft</td>
</tr>
<tr>
<td>Strength</td>
<td>Medium High</td>
<td>High</td>
<td>Low Low</td>
</tr>
<tr>
<td>Toughness</td>
<td>High Low</td>
<td>Low Medium</td>
<td>Very Low High</td>
</tr>
<tr>
<td>Elastic modulus</td>
<td>High High</td>
<td>High High</td>
<td>Low Low Low</td>
</tr>
<tr>
<td>Electrical Conductivity</td>
<td>High High</td>
<td>Low Low Low</td>
<td>Low Low Low</td>
</tr>
<tr>
<td>Thermal Conductivity</td>
<td>High High</td>
<td>Low Low Low</td>
<td>Low Low Low</td>
</tr>
<tr>
<td>Thermal Expansion</td>
<td>Low Low</td>
<td>Low Low</td>
<td>High High</td>
</tr>
<tr>
<td>Density</td>
<td>High High</td>
<td>Medium Medium</td>
<td>Low Low</td>
</tr>
<tr>
<td>Translucency</td>
<td>None None</td>
<td>Medium Medium</td>
<td>Low Low Low</td>
</tr>
<tr>
<td>Examples</td>
<td>Gold copper</td>
<td>Gypsum, zinc phosphates</td>
<td>Dental porcelain</td>
</tr>
<tr>
<td></td>
<td>Amalgam phases</td>
<td>Al₂O₃ SiO₂ Dental porcelain</td>
<td>Poly (methyl methacrylate Material)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 1-1 summarizes the general behaviors of the three basic materials discussed in this chapter: metals, ceramics, and polymers. Certain inherent properties of materials will
influence their selection for use in dentistry. For example, metals are inherently strong, in
general, and have good stiffness (modulus of elasticity). These properties would tend to
recommend them as restorative materials. On the other hand, metals conduct heat rapidly
and are opaque (nonesthetic), limiting their usefulness in restorative dentistry. Ceramics
and polymers are thermally insulating and tend to be more translucent. Hence, these
materials insulate the pulp from extremes of heat and cold and offer the potential of more
lifelike esthetics. They tend to have lower toughness than metals, however, and polymers
have much lower strength.

Because no one class of materials possesses all the desired properties, it is not surprising
that materials tend to be used in combination. The porcelain-fused-to-metal restoration
combines the strength and ductility of metal with the esthetics of dental porcelain. A
ceramic or polymer base is used to insulate the pulp from a thermal-conductive metallic
restoration. A high thermal-expansion, low-strength, low-elastic-modulus polymer is
reinforced with a low thermally expanding, high-strength, high-elastic-modulus ceramic
filler to form a dental resin composite material. An understanding of the advantages and
limitations of the various types of materials enables us to make selections based on the
best compromise of desired properties versus inherent limitations.

**Predicted Versus Actual Strengths**

It is possible to predict the strength of a material from the strengths of the individual
bonds between the atoms in the material. The values of strength obtained by such a
prediction are typically 1 million to 3 million pounds per square inch (psi), or about 7 to
21 GPa. Actual strengths of most materials are ten to 100 times lower.

Why do materials fail to exhibit the strengths one would expect from the bonds between
atoms? Why do ceramics break suddenly without yielding, whereas metals often yield
and distort to 120% or more of their original length before fracturing? Why are polymers
so much weaker and more flexible than metals and ceramics? Why do metals conduct
heat and electricity, whereas polymers and ceramics do not? As will be seen in this
chapter, many of the answers to these questions can be understood by knowing only a few
things about the structures of these materials. There is one key concept, for example, that
will not only explain the tendency for ceramics to be brittle, but will also explain all of
the methods used to strengthen ceramics. Similarly, one key concept will explain why
polymers expand about ten times as much as metals or ceramics when heated the same
amount, why polymers are generally weak, why they are ten times more flexible than
metals or ceramics, and why they tend to absorb water and other fluids.

**Ceramics**

*Introduction*

Consider a block of material as depicted in Fig 1-2(a)
Fig 1-2 Stress raisers and the effect of their shape on stress concentration. (a) If no stress raiser is present, the stress is constant across cross section A. (b) If a rounded notch is present, the stress is constant over most of the cross section. (c) As the notch becomes sharper, the stress concentration becomes greater.

If this block is stretched by applying a force, F, the stress at any point on cross section A is the same as the average stress, $\sigma_{\text{ave}}$. For example, if the cross-sectional dimensions of the block are $1/2 \text{ in} \times 1/2 \text{ in} = 1/4 \text{ in}^2 (1.27 \text{ cm} \times 1.27 \text{ cm} = 1.61 \text{ cm}^2)$, and a force of 3,000 lb (13 kN) is applied, the average stress along cross section A is 12,000 psi (83 MPa). However, if a semicircular groove were machined across one side of the block of material, as depicted in Fig 1-2(b), the stress at each point across a plane passing through this groove would not be the same as the average stress. The stress would be constant over most of the cross section, but near the groove, the stress would suddenly rise and reach a maximum right at the edge of the groove. This phenomenon occurs around any irregularity in a block of material. The groove or other irregularity is called a stress raiser. The stress around a stress raiser can be many times higher than the average stress in the body. The amount of increased stress depends on the shape of the stress raiser. For example, if the stress raiser in our block of material were a sharp notch rather than a semicircular groove, the stress would increase greatly at the tip of the sharp notch (Fig 1-2(c)). As the tip of the notch becomes smaller (ie, the notch becomes sharper), the stress concentration at the tip of the notch becomes greater.

The minute scratches present on the surfaces of nearly all materials behave as sharp notches whose tips are as narrow as the spacing between atoms in the material. Thus, the
stress concentration at the tips of these minute scratches causes the stress to reach the theoretical strength of the material at relatively low average stress. When the theoretical strength of the material is exceeded at the tip of the notch, the bonds at the notch tip break

![Diagram](https://via.placeholder.com/150)

**Fig 1-3** Role of stress raisers in achieving localized stresses as great as the theoretical strength of the material. The stress at the tip of the notch reaches the theoretical strength of the material even though the average stress is many times lower. As the most highly stressed bond breaks (a), the stress is transferred to the next bond (b).

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(Fig 1-3(a)). The adjacent bonds now are at the tip of the notch and thus are at the point of greatest stress concentration (Fig 1-3(b)). As the crack propagates through the material, the stress concentration is maintained at the crack tip until the crack moves completely through the material. This stress concentration phenomenon enables us to understand how materials can fail at stresses far below their expected strength. This situation exists in the cutting of glass. When glass is cut, a line is scribed on one surface with a diamond point or a hardened steel glass-cutting wheel. This scribed line is a very shallow scratch or crack in comparison to the thickness of the glass, but it acts as a stress raiser to concentrate the stress at the tip of the crack, as already described.
Understanding the effect of stress concentration is the key to understanding the failure of brittle materials, such as ceramics, which influences their selection as dental materials and dictates the design of restorations fabricated from these materials.

The tendency for ceramics to fail in a brittle manner at stresses that are far below the theoretical strengths of these materials can be understood in light of the concept of stress concentration at surface scratches and other defects. Most of the techniques for strengthening ceramics can also be understood by virtue of this concept.

* Actually, the exceeding of the theoretical strength of the material at the crack tip is a necessary but insufficient condition for crack propagation. The remaining condition involves a balance between the surface energy required to form the two new surfaces of the crack, and the elastic strain energy arising from the applied stress. This is called the Griffith energy balance, discussion of which is beyond the scope of this book.

**Clinical applications of ceramics**

Ceramics are inherently brittle and must be used in such a way as to minimize the effect of this property. Ceramic restorations must not be subjected, for example, to large tensile stresses, to avoid catastrophic failure. A method for reducing the influence of the brittleness of ceramics is to fuse them to a material of greater toughness (eg, metal), as is done with porcelain-fused-to-metal (PFM) restorations. Ceramics also may be reinforced with dispersions of high-toughness materials, as is the case with the alumina ($\text{Al}_2\text{O}_3$)-reinforced porcelain used in porcelain jacket crowns.

![Brittle fracture of ceramic (dental porcelain) due to mismatch in the coefficient of thermal expansion between porcelain and metal.](Photo courtesy of R. P. O'Connor, DMD.)

**Fig 1-4** Brittle fracture (arrows) of ceramic (dental porcelain) due to mismatch in the coefficient of thermal expansion between porcelain and metal. (Photo courtesy of R. P. O'Connor, DMD.)

*Figure 1-4* shows brittle fractures that occurred in the porcelain of two PFM crowns due to the mismatch in thermal expansion between the porcelain and metal.
Metals

Effect of ductility on stress concentration

As discussed in the previous section, stress raisers at the surface of a material can cause the stress in a localized region around the tip of the stress raiser to reach the theoretical strength of the material. When this happens in a brittle material, a crack propagates through the material, resulting in fracture (see the footnote on page 3). In a ductile material, something happens before the theoretical strength of the material is reached at the tip of the stress raiser that accounts for the tremendous difference in behavior between, for example, a glass and a metal. As discussed previously, the magnitude of the stress concentration at the tip of a notch, surface scratch, or other stress raiser is determined by the sharpness of the stress raiser. If a sharp notch or scratch is present in the surface of a brittle material, the stress concentration around this notch would be something like that shown in Fig 1-5(a).

![Fig 1-5](image)

**Fig 1-5** Rounding or blunting of stress raisers that occurs in ductile materials. Stress concentration is self-limiting in ductile materials because the region under greatest stress, the tip of the sharp stress raiser (a), yields to round or blunt the stress raiser and lower the stress (b).

If a stress raiser is present in a ductile metal, however, the material at the tip of the stress raiser deforms under stress so the sharp notch becomes a rounded groove, as shown in Fig
1-5(b). Because the tip of the stress raiser is now rounded rather than sharp, the stress concentration at the tip of this stress raiser is much lower. There are two important facts to recognize in this process:

1. As with brittle materials, the actual strengths of ductile materials are many times less than those predicted from strengths of bonds between atoms.

2. Unlike the behavior around the notch in a brittle material, the stress concentration blunts the sharp tip of the stress raiser, thus lowering the stress concentration effect.

**Mechanism of ductile behavior**

What, then, is responsible for the ductile behavior of a metal? Consideration of what is happening on an atomic level provides insights into the difference between brittle materials and ductile ones. A schematic of the arrangement of atoms in a piece of metal is shown in Fig 1-6.

![Fig 1-6](image)

**Fig 1-6** Tensile stress on a piece of material can be considered stress normal (n) (perpendicular) to plane A-A’, together with stress parallel (s) to plane A-A’. The stress parallel to plane A-A’ tends to cause the atoms along the plane to slide (shear) past each other.

If this piece of metal is subjected to a tensile stress as shown, this stress can be resolved into two components when considered relative to the plane A-A’. One component tends to move the rows of atoms on either side of the plane A-A’ apart from each other, and the
other component tends to cause the planes to slide past one another along the plane A-A’.

The component of the stress that tends to cause the planes to slide past one another is the one that causes a material to deform plastically. Scientists are able to calculate, from the bond strengths between the atoms, the stresses that would be required to make one plane of atoms slide past another plane; these stresses are 100 or more times higher than those actually observed. If, however, the bonds were to break one at a time and re-form immediately with the adjacent atom, one plane could move past the other at very low stress levels.

The mechanism of this process is shown in Fig 1-7.

Fig 1-7 (a) through (f) show how the shearing stress can cause a dislocation to
pass through the network of atoms, breaking only one row of bonds at a time. For the atoms along plane A-A' to slide past one another all at once would require enormous stress. The fact that metals yield to stresses much lower than expected is explained by the breaking of only one row of bonds (perpendicular to the page) at a time.

**Figure 1-7** (a) through (f) show how, by breaking and re-forming bonds, an extra plane of atoms can move along plane A-A' until this "ripple" in the crystal lattice passes completely through the material. Multiple repetitions of this process along many planes similar to A-A' allow a metal to yield to an applied stress without fracturing. This ripple in the lattice structure is called a **dislocation**, and it is responsible for the ductile behavior of metals.

Metals can be hardened and strengthened by a variety of treatments that make it more difficult for dislocations to move through the metal lattice. Alloying, cold-working, and formation of second phases in a metal are all ways of impeding dislocation motion. Some crystal structures of metals, such as intermetallic compounds, make it difficult for dislocations to move. The passage of a dislocation through the ordered structure of an intermetallic compound would result in an unfavorable atomic arrangement, so dislocations move only with difficulty.

With metals, it is important to remember that their ability to yield without fracturing, as well as all of the methods for making metals harder and stronger, is understandable in light of the concept of dislocations in the metal structure.

Other properties of metals, such as their electrical and thermal conductivity, can be understood as resulting from the **metallic bond**. In the metallic bond some of the electrons are free to move rapidly through the lattice of metal ions. This unusual aspect of the metallic bond enables metals to conduct heat and electricity.

The electronic structure of the metallic bond also accounts for the opacity of metals. **Figure 1-8** illustrates the metallic bond with its lattice of positively charged metal ion cores and electrons that are free to move between the ion cores.
Fig 1-8 Representation of the metallic bond showing the metal ion cores surrounded by free electrons. (After Lewis and Secker, 1965.)

Dislocations in ceramic materials and in intermetallic compounds

Why do ceramic materials not yield in the same manner as metals? The answer to this question involves consideration of two types of ceramic materials:

1. **Amorphous** materials (glasses) — Glassy materials do not possess an ordered crystalline structure as do metals. Therefore, dislocations of a crystalline lattice cannot exist in glassy materials. Thus, glasses have no mechanism for yielding without fracture.

2. **Crystalline** ceramic materials — Dislocations exist in crystalline ceramic materials, but their mobility is severely limited, because their movement would require that atoms of like charge be brought adjacent to one another, as seen in Fig 1-9. The energy required to do this is so large that dislocations are essentially immobile in crystalline ceramic materials.

**Intermetallic compounds**, unlike ordinary metal **alloys**, have a specific formula (e.g., Ag₃Sn, the main component of dental amalgam alloy powder) and an ordered arrangement of atoms. The movement of a dislocation through this ordered structure would produce a disruption of the order similar to that shown in Fig 1-9 for crystalline ceramic materials. Hence, dislocations move only with difficulty in intermetallic compounds, and this property renders them more brittle than ordinary metal alloys.
The alternating charges of an ionic structure (crystalline ceramic) do not allow dislocations to move along plane A-A'. If a dislocation were to pass through such a structure, it would result in ions of like charge coming into direct contact, which would require too much energy.

Clinical applications of metals

Metals are generally ductile and tough when compared to ceramics, although a few types of metals, such as dental amalgams, are markedly more brittle than others. This ductility allows the margins of castings to be burnished, orthodontic wires to be bent, and partial denture clasps to be adjusted. Figure 1-10 shows how the ductility of metal allows the wire clasp for a partial denture framework to be bent permanently to provide the desired retention. The ductile behavior of the partial denture alloy can be contrasted with the brittle behavior of the intermetallic material, dental amalgam, as shown in Fig 1-11.
Fig 1-10 Ductility of metal as illustrated by the adaptation (bending) of a partial denture wrought wire clasp.

Fig 1-11 Brittle fracture of an amalgam post and core that had supported a PFM crown. Set dental amalgam is a mixture of several intermetallic compounds. Intermetallic compounds tend to be brittle rather than ductile. (Photo courtesy of R. P. O'Connor, DMD.)

Polymers
Introduction

The behavior of polymers is fundamentally different from both ceramics and metals. To understand this difference, it is useful to consider the modulus of elasticity and the strength of polymers at a molecular level.

Modulus of elasticity

When chains of polyethylene are aligned parallel to one another and are subjected to a tensile stress along their long axes (Fig 1-12(a)), the stress required to stretch the atoms in the chains farther apart is surprisingly high. In fact, the modulus of elasticity of polyethylene when measured in this way is 30 million psi, about the same as steel! However, if the applied stress is perpendicular to the long axes of the chains (Fig 1-12(b)), the modulus of elasticity is only about 0.5 million psi. The high modulus of elasticity (30 million psi) in the first case results from the strong bonds between the atoms within polymer chains. The low elastic modulus (0.5 million psi) in the second case results from the weak bonds between atoms in adjacent chains.

Bulk polymers have their polymer molecules in a more random, tangled arrangement, and hence there are somewhat fewer secondary bonds between chains than when the chains are perfectly aligned. The tangling and coiling of the polymer molecules make polymers even more flexible, because of the lower stress required to straighten out a coiled molecule compared to the stress necessary to stretch the atoms in a molecule farther apart.
Fig 1-12 (a) Strong (primary) bonds between atoms within a polymer chain result in a high stiffness (modulus of elasticity) when aligned chains are stretched along their lengths. (b) Weak (hydrogen) bonds between atoms in adjacent chains result in a low modulus of elasticity when aligned chains are stretched perpendicular to their lengths.

**Strength**

The low strength of polymers when compared to ceramics and metals can also be understood in terms of the strong bonds within polymer chains and the weak bonds between polymer chains. If the rule-of-thumb value for theoretical strength, $0.1E$, is applied to the oriented polyethylene fiber shown in Fig 1-12(a), a value of 3 million psi is obtained for the theoretical strength of polyethylene. Typical bulk polymers, however, seldom have tensile strengths of more than 10,000 psi. The weak secondary bonds between polymer chains allow these chains to slide past one another at much lower stresses than those required to break the bonds within the chains.

**Thermal expansion**

The increase of the temperature of a material as the result of increased atomic vibration within it is a familiar concept. This atomic vibration is limited by the bonds between atoms in a material such that when strong bonds are present between atoms, the atoms
vibrate over a small amplitude, and when weak bonds are present, the atoms vibrate over a large amplitude. Ceramics and metals are characterized by strong bonds between atoms, and the secondary bonds play an insignificant role in their properties. As a result, most ceramics and metals expand relatively little when heated—that is, their coefficients of thermal expansion are relatively low. Polymers, however, are characterized by strong bonds within polymer chains and weak bonds between polymer chains. Thus, the vibration of carbon atoms within the polymer chain is restricted in the directions parallel to the long axis of the chain, but the atoms are free to vibrate in the two directions perpendicular to the long axis of the polymer chain. As a result, when a polymer is heated, the chains must move farther apart to allow for the larger-amplitude vibration, which occurs perpendicular to the long axes of the polymer chains. This phenomenon accounts for the large coefficient of thermal expansion exhibited by polymers. Figure 1-13 illustrates the different thermal expansion behaviors of a polymeric material and a crystalline material.

Fig 1-13 The different thermal expansion behaviors of a polymeric material and a crystalline material.

Water absorption

Because of the weak secondary bonds in polymer materials, water molecules are able to penetrate between the polymer chains in a process called water absorption. Water absorption has both positive and negative aspects. On the positive side, it is the main factor in correcting the 0.5% processing shrinkage that occurs during the fabrication of heat-cured denture bases. When the acrylic polymer absorbs water, the polymer molecules are forced apart slightly, causing the denture base to expand. This expansion during water absorption compensates for the processing shrinkage.

On the negative side, water is responsible for the hydrolytic degradation of polymers. In addition, the ions carried by the water may cause the polymer to break down more quickly or to become stained or malodorous.

Effect of cross-linking
Because many of the undesirable properties of polymers are due to weak bonds between polymer chains, it would seem that a way to improve them would be to link chains together with primary chemical bonds. In fact, this method, called cross-linking, is widely used to improve strength, resistance to water absorption, abrasion resistance, and other properties of polymers. Because of the small number of primary bonds in a given volume of polymer material as compared to ceramics or metals, however, polymer properties remain generally inferior even with cross-linking.

The key to understanding polymers is that the low strengths and elastic moduli of polymers, as well as many other distinctive properties, can be seen in light of the concept of strong bonds within polymer chains and weak bonds between polymer chains.

Clinical applications of polymers

Polymers have had limited use as restorative materials by themselves because of their low strength and high thermal expansion. Some of the unfavorable properties of polymers have been mitigated by the incorporation of inorganic fillers to form composite materials. Modification of the polymer resins themselves to make them hydrophobic has improved their resistance to water absorption. The properties of polymers may be further improved by increasing their crystallinity, but the bonds between chains in crystalline polymers are still only secondary bonds. The intrinsic weaknesses of polymers will probably continue to limit their use as restorative materials. The incisal fracture of a Class 4 microfilled composite restoration is shown in Fig 1-14. Microfilled resin composites tend to have considerably lower levels of the stronger inorganic filler, and so the stresses are borne more heavily by the weaker resin matrix. In Class 4 restorations, microfills generally have insufficient strength to withstand the stresses that may be encountered.

Fig 1-14 Fracture of a Class 4 microfilled composite restoration. The low filler content of most microfills leaves the polymer matrix bearing the majority of the stress. Polymers are generally too weak for bearing large stresses. (Photo courtesy of C. D. Smith, DDS.)

Glossary

alloy A material that exhibits metallic properties and is composed of one or more elements—at least one of which is a metal. For example, steel is an alloy of iron and
carbon, brass is an alloy of copper and zinc, and bronze is an alloy of copper and tin.

**amorphous**  Literally, "without form." Atoms in crystalline solids exhibit an ordered arrangement, whereas atoms in amorphous solids lack this long-range periodicity.

**ceramic**  In the broadest sense, a compound of metallic and nonmetallic elements. By this definition, materials ranging from aluminum oxide (Al₂O₃) to table salt (NaCl) are classified as ceramics. In dentistry, gypsum (CaSO₄·2H₂O), many dental cements (eg, zinc phosphate), and porcelains are examples of ceramic materials.

**cross-linking**  A method for making a polymer stronger and more rigid by creating chemical bonds between the molecular chains in the polymer.

**crystalline**  Having atoms or molecules arranged in a regular, repeating three-dimensional pattern. Metals are nearly always crystalline; ceramics and polymers can be crystalline or noncrystalline (amorphous).

**dislocation**  A defect in a crystal that is caused by an extra plane of atoms in the structure (see Fig 1-7). The movement of dislocations is responsible for the ability of metals to bend without breaking.

**intermetallic compound**  A chemical compound whose components are metals. The gamma phase of amalgam, Ag₃Sn, is an example of an intermetallic compound.

**metal**  A crystalline material that consists of positively charged ions in an ordered, closely packed arrangement and bonded with a cloud of free electrons. This type of bond, called a metallic bond, is responsible for many of the properties of metals—electrical and thermal conductivity, metallic luster, and (usually) high strength.

**metallic bond**  One of the three types of primary (strong) chemical bond. (Ionic and covalent are the other two.) Metallic bonds involve the sharing of valence electrons among all atoms in the metal. Metallic bonds are responsible for many of the distinctive properties of metals, including electrical conductivity.

**modulus of elasticity**  A measure of the stiffness or flexibility of a material. A stiff material has a high modulus of elasticity, and a flexible material has a low modulus of elasticity. Also called Young's modulus. (See chapter 2 for more detail.)

**polymer**  A material that is made up of repeating units, or *mers*. Most polymers are based on a carbon (-C-C-C-) backbone in the polymer chain, although a silicone (-O-Si-O-Si-O-) backbone is important in many polymers.

**stress raiser**  An irregularity on the surface or in the interior of an object that causes applied stress to concentrate in a localized area of the object. Other things being equal, the sharper the stress raiser, the greater the localized stress around it.

**water absorption**  The penetration of water into the structure of a material. Water absorption by polymers can help offset the effects of processing or polymerization shrinkage (as in heat-processed denture bases), but it can also have detrimental effects, such as discoloration and leaching of unreacted components. For water to be absorbed into the structure of a material, it must first be *adsorbed* onto the surface. Because both phenomena occur when water is absorbed, the process is often referred to simply as *water*
sorption.

**Discussion Questions**

1. Although metals have had a long history of application as restorative materials, which properties of metals are causing a decline in their use? Which properties will promote their use for dental applications for many more years?

2. Which physical properties are related to the large differences in wear resistance of polymers and ceramics?

3. Although the strength values of materials based on testing and reported in research publications are often high, dental devices may fail at low levels of stress. Explain and indicate how materials need to be used in dentistry to achieve the full strength potential of the materials used.

4. Why are the mechanical properties of materials so important in restorative dentistry?

5. How does the atomic bonding of materials determine many of the observed properties of materials?

**Study Questions**

*(See Appendix E for answers.)*

1. Characterize the three basic materials (metals, ceramics, and polymers) in regard to modulus of elasticity, strength, ductility, and coefficient of thermal expansion.

2. Describe the relationship between the shape of a stress raiser and the concentration of stress around it.

3. Discuss the key concept for each of the three basic materials and show how it explains the properties of each.

**Recommended Reading**


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Chapter 2. Physical Properties and Biocompatibility

Introduction

Physical properties determine how materials respond to changes in their environments, and biocompatibility relates to the effect a material has on the tissue it contacts. The relationships among the various properties are classified according to the scheme in Fig 2-1.

Fig 2-1 A tree diagram classifying physical properties.

Mechanical Properties

Introduction

The concepts of stress, strain, modulus of elasticity, plastic deformation, and other properties were introduced in chapter 1. Because of the unfamiliarity and abstruse nature of many of these concepts, they will be discussed here in somewhat greater depth, and some additional concepts will be introduced as necessary to understand the physical
behavior of materials.

**Stress**

Consider again the block of material described in the previous chapter that was 1/2 in × 1/2 in (1.27 cm × 1.27 cm) in cross section and was subjected to a 3,000-lb (about 13 kN) load. As was pointed out in that example, the stress experienced by that block—determined by dividing the force by the cross-sectional area of the block—is 12,000 psi, or 83 MPa. Stress (σ) is the force (F) divided by the cross-sectional area (A):

\[
\sigma = \frac{F}{A}
\]

Now consider a similar block, but with smaller dimensions—1/4 in × 1/4 in (about 0.6 cm × 0.6 cm) in cross section (area of 1/16 in², or about 0.4 cm²). If this new block is subjected to the same 3,000-lb (13-kN) tensile load, the stress in this case is 3,000 lb (13 kN) divided by 1/16 in² (about 0.4 cm²) = 48,000 psi (about 330 MPa).

Thus, the usefulness of the concept of stress is apparent. It is not sufficient merely to state the load or force that is being applied to a dental material, because the stress that is produced in the material depends just as much on the cross-sectional area on which the load is acting as it does on the load itself. For instance, if the block that measured 1/4 in × 1/4 in in cross section is subjected to a load of 12,000 lb (about 53 kN) instead of 3,000 lb (about 13 kN), the stress is 48,000 psi (about 330 MPa). If the cross-sectional area is made 4 times smaller (1/4 as large), or if the load is made 4 times larger, the stress is increased by a factor of 4. Thus, the stress is said to be inversely proportional to the cross-sectional area and directly proportional to the load.

The basic types of stresses produced in dental structures under a force are tensile, compressive, and shear. All three are present in a beam loaded in the center. If the value of these stresses exceeds the strength of the material, the structure will fail. It is therefore important to know the strength values of materials. It is rare that an object will be subject to the pure tensile, compressive, or shear stresses experienced by test specimens in a materials testing laboratory. As shown in Fig 2-2, however, wherever bending forces are present, tensile stresses are present in critical areas, which could result in failure.
Fig 2-2 (a) Tensile stress \( (S_T) \) at the isthmus of a two-surface amalgam restoration. (b) Tensile stress at the occlusal surface of a beveled amalgam restoration. (c) Tensile stress at the junctions of a partial denture clasp. (d) Tensile and compressive stresses \( (S_C) \) in a soldering bridge. \( F = \) force; \( A = \) area. (From Mahler and Terkla, 1958. Reprinted with permission.)

Strain

When a block of material is subjected to a tensile stress as described in the preceding section, it temporarily becomes longer by a certain amount. This temporary increase in length is called strain. A few examples will illustrate how strain is described. Consider the block with the cross section of \( \frac{1}{4} \text{ in} \times \frac{1}{4} \text{ in} = \frac{1}{16} \text{ in}^2 \) \( (0.6 \text{ cm} \times 0.6 \text{ cm} = 0.4 \text{ cm}^2) \) and assume it has a length of 10 in \( (25.40 \text{ cm}) \) when no load is applied. If the length is measured while the 3,000-lb \( (13\text{-kN}) \) load is being applied and the new length is 10.016 in \( (25.441 \text{ cm}) \), the strain is computed by dividing the increase in length \( (0.016 \text{ in}, 0.041 \text{ cm}) \) by the original length \( (10.000 \text{ in}, 25.40 \text{ cm}) \) to obtain 0.0016. Strain is a dimensionless quantity because we are dividing unit length by unit length, but sometimes it is written as "\( \text{in/in} \)" or "\( \text{cm/cm} \)." Strain can also be expressed in a percentage, in which case the dimensionless value is multiplied by 100%. In the given example, the strain would be 0.16%.

If a piece of the same material with the same cross section but 11 in \( (27.940 \text{ cm}) \) long
were subjected to the same load (3,000 lb, about 13 kN), and the length of the block of material were measured while the load was being applied, it would be 11.0176 in (27.9847 cm). To compute the strain produced, the change in length (0.0176 in, 0.0447 cm) is divided by the original length (11.0000 in, 27.9400 cm); the strain is determined to be 0.0016 (or 0.16%), the same value obtained above for the block of material measuring 10 in (25.4 cm). Thus, it can be seen that strain is independent of the length of the specimen.

Now consider a bar of high-strength steel, \( \frac{1}{4} \) in \( \times \) \( \frac{1}{4} \) in = \( \frac{1}{16} \) in\(^2\) (0.6 cm \( \times \) 0.6 cm = 0.4 cm\(^2\)) in cross section and 10 in (25.4 cm) long, that is capable of supporting a tensile load of 6,000 lb (about 27 kN). If the length of this bar were measured while under this 6,000-lb (27-kN) load, it would be 10.032 in (25.4813 cm). If the strain is computed as before, the change in length (0.032 in, 0.0813 cm) divided by the original length (10.000 in, 25.4 cm) is found to be 0.0032 or 0.32%. Hence, it can be seen that if the cross-sectional area of the block of material is kept the same and the applied load is doubled, the strain experienced by the material is doubled.

**Elasticity**

The preceding examples showed that both stress and strain are directly proportional to the load applied when the cross-sectional area is kept the same. Hence, if the load is doubled, both the stress and the strain likewise will be doubled. It can be seen that the ratio between stress (\( \sigma \)) and strain (\( \varepsilon \)) is the same, thus:

\[
\frac{\sigma}{\varepsilon} = \frac{2\sigma}{2\varepsilon} = \frac{3\sigma}{3\varepsilon}
\]

The property of having a constant ratio of stress to strain is called **elasticity**, and the constant that is the ratio of stress to strain is called the **modulus of elasticity**. To gain a clearer understanding of the concept of elasticity and to see what it means for two materials to have different moduli of elasticity, it is helpful to consider the analogy of two springs of different stiffnesses, as shown in Fig 2-3. Spring 2 is stiffer than spring 1; therefore, when they support equal weights, the stiffer spring is extended a smaller amount. Increasing the load from 1 lb (4.4 N) to 2 lb (8.9 N) causes each spring to extend twice as much as it did under the 1-lb (4.4-N) load. Subjecting both springs to a 3-lb (13-N) load causes each one to extend 3 times as much as it did under a 1-lb (4.4-N) load, and so on. It can be seen that, irrespective of the load applied for either spring 1 or spring 2, the ratio of the extension to the load is a constant for that spring. However, the constants for the two springs are different. In this analogy, the extension of the spring corresponds to strain, the load or weight on the spring corresponds to stress, and the spring constant corresponds to the modulus of elasticity.

It is obvious that more weight cannot be added to a spring indefinitely and still have the extension increase proportionately. At some point the spring will become "stretched out" and will not return to its original size when the weight is removed. In the same way, when a material is stressed above a certain point, stress is no longer proportional to strain. The highest stress at which stress is proportional to strain is called the elastic limit, or the **proportional limit**. Because this stress is difficult to determine precisely, as it would entail looking for an infinitely small deviation from proportionality, it is customary to designate a certain permanent deformation or offset (usually 0.002, or 0.2%) and to report the **yield strength** of the material at this strain. Thus the terms "elastic limit" and "proportional limit" are synonymous, whereas "yield strength" has a slightly different
meaning. The yield strength of a material is always slightly higher than the elastic limit.

If a material continues to have more and more weight applied to it, it will of course eventually break. If the material is being stretched (tensile loading), the stress at breakage is called the ultimate tensile strength (UTS). When many metals are stressed above their proportional limits, they undergo a process called work hardening, and actually become stronger and harder. But with the increased strength provided by work hardening comes increased brittleness. It is thus very important to avoid overdoing it when bending a wire or a partial denture clasp—too much bending of the metal back and forth will make it harder and more brittle.

The modulus of elasticity is an inherent property of the material and cannot be altered appreciably by heat treatment, work hardening, or any other kind of conditioning. This property is called structure insensitivity, because it is not sensitive to any alteration to the structure (meaning the microstructure) of the material. The modulus of elasticity is one of the few properties that is not sensitive to any alteration of the structure of the material. The yield strength of a material, for example, is sensitive to work hardening and will increase with increasing amounts of work hardening.

Fig 2-3 The concept of modulus of elasticity using the analogy of springs. Spring 1 is a flexible spring, representing a material with a low modulus of elasticity. Spring 2 is a stiff spring, representing a material with a high modulus of elasticity. When the same load is
applied to both springs, they stretch different amounts. The weight applied to the spring is analogous to the stress ($\sigma$), the amount the spring stretches is analogous to the strain ($\varepsilon$), and the stiffness of the spring is analogous to the modulus of elasticity ($E$).

**Plasticity**

When the elastic or proportional limit is exceeded in a material, it is said to exhibit plastic behavior. (The term "plastic" means "moldable," but the term has come to be associated with polymers.) Materials that experience a large amount of plastic behavior or permanent deformation are said to be **ductile**. Materials that undergo little or no plastic behavior are said to be **brittle**. If stress is plotted against strain, with stress as the ordinate (y axis) and strain as the abscissa (x axis), a diagram such as the one in Fig 2-4 is the result. It can be seen that stress is proportional to strength up to the elastic (or proportional) limit. Above this stress, stress and strain are no longer proportional. If a material is stressed into the plastic region or region of permanent deformation and then the stress is removed, the material will have a permanent set, or deformation. If stress is continuously increased, the material will undergo more and more plastic deformation and will ultimately fracture. The highest stress achieved during this process is called the ultimate tensile strength of the material, as mentioned previously. The total strain at fracture (elastic strain + plastic strain) is called the **elongation**. Elongation of metals is important in several situations in dentistry. When one is burnishing the margin of a crown, the property that most comes into play is the elongation, although the yield strength is also important. For example, gold alloys are generally easy to burnish because they have a lower yield strength. Nickel-chromium and cobalt-chromium alloys may have adequate elongation for burnishing, but their yield strengths are so high that they make burnishing difficult. Another example of the importance of elongation is the bending of a partial denture clasp to adjust the retention.

The entire area under the stress-strain curve is a measure of the energy required to fracture the material, or its **toughness**. The area under only the elastic region of the stress-strain curve is a measure of the ability of the material to store elastic energy (the way a compressed spring does). This is called the **resilience** of the material.

Tabulated data for (Young's) modulus of elasticity, yield strength, ultimate tensile strength, percent elongation, and other mechanical properties are listed in **Appendix A**.
Measurement of stress and strain on an object being stretched. In the region labeled "elastic deformation," stress is proportional to strain, whereas in the region labeled "plastic deformation," stress and strain are no longer proportional (strain increases faster than stress does). The highest stress at which stress and strain are still proportional is called the proportional limit, and the maximum stress just before the object breaks is called the ultimate tensile strength. The total amount that the object stretches (ie, the total strain), which is the sum of the elastic deformation and the plastic deformation, is called the percent elongation.

**Fatigue**

When materials are subjected to cycles of loading and unloading, as during mastication, they may fail due to fatigue at stresses below the ultimate tensile strength. Usually, small cracks at the surface or within the material gradually grow larger upon cycling, and eventually the material fails. Fatigue curves, such as the one shown in Fig 2-5, show the experimentally determined number of cycles to failure at different stress levels. The higher the stress placed on and off the material, the fewer the number of cycles until failure.
Viscous flow

Many dental materials are in a fluid state when they are formed, and **viscous flow** phenomena are important considerations. Our understanding of the behavior of impression materials and amalgam involve **viscoelastic** phenomena. When shear stress-strain rate (flow rate) plots are obtained, they enable viscous materials to be classified in several ways. A newtonian fluid shows a constant viscosity, \( \eta \), which is independent of strain rate:

\[
\eta = \frac{\sigma}{\varepsilon} = \text{constant}
\]

where \( \eta \) is the viscosity in poise, \( \sigma \) is the shear stress acting on the fluid, and \( \varepsilon \) is the strain, or flow rate.

Actual fluids differ from these in their flow responses to the level of stress applied. Plastic fluids (eg, putty) don't flow at all until a minimum stress is applied. Pseudoplastic fluids (eg, fluoride gels) show an instantaneous decrease in apparent viscosity or consistency (become more free-flowing) with increasing shear rate. Dilatant fluids (eg, fluid denture base resins) show an increase in rigidity as more pressure is applied. Thixotropic fluids (eg, fluoride gels, house paints) flow more freely when vibrated, shaken, or stirred than if allowed to sit undisturbed. Formulation of the various suspensions and gels used in dentistry to impart one or more of these properties can make them easier to handle. For example, impression materials that do not run off impression trays but are less viscous under pressure in a syringe have a definite advantage.

Viscoelastic behavior

Many materials, including elastic impression materials, waxes, and even hardened amalgam, show a combination of elasticity and viscous flow. **Figure 2-6** is a model of how a viscoelastic material acts under stress, the so-called spring-and-dashpot (or Voigt) model. (A dashpot is an oil-filled cylinder with a loosely fitted piston—most automobile shock absorbers are of a dashpot design.) As the model is stretched, the spring component
on the left stretches and the piston of the dashpot also moves through the viscous liquid. The strain under stress and after release of the stress is time-dependent. The strain gradually builds up until the release of stress, and then it gradually goes back to zero as the spring element returns to its original length but is dampened by the dashpot element. Elastic impression materials are viscoelastic; initially they are strained upon removal from the mouth and require a short period of time to recover before models or dies are poured.

\[ a \]

\[ b \]

Fig 2-6 (a) Voight model for viscoelastic behavior. (b) Response of (a) to stress. Strain increases with time and decreases after force (F) is released at \( t_1 \), but is slow due to the damping effect of the dashpot or the elastic action of the spring element. (From O'Brien and Ryge, 1978. Reprinted with permission.)

**Hardness**

The resistance of a material to indentation or penetration is called **hardness**. The technical definition of hardness varies little, if at all, from the familiar definition, with the possible exception that the technical definition carries with it the connotation of a method of measurement. Most of the methods for measuring hardness consist of making a dent in the surface of a material with a specified force in a controlled and reproducible manner and measuring the size of the dent. For the measurement methods discussed in this section, as the hardness increases, the respective hardness number increases also. For example, Fig 2-7 indicates the relative positions of a variety of dental and other materials on a Knoop hardness scale ranging from soft on the left to hard on the right. The positions of the different materials on the line indicate their relative hardnesses.

Brinell hardness number (BHN)

The indenter for the Brinell hardness test is a small, hardened steel ball, which is forced into the surface of a material under a specified load. This indentation process leaves a round dent in the material, and hardness is determined by measuring the diameter of the
dent. Brinell hardness values for selected metals are listed in Appendix A.

Vickers hardness number (VHN) or diamond pyramid hardness (DPH)

The Vickers hardness indenter is a square, pyramid-shaped diamond, which leaves a square, diamond-shaped indentation in the surface of the material being tested. Hardness is determined by measuring the diagonals of the square and taking the average of the two dimensions. The Vickers hardness test is also called the diamond pyramid hardness test.

The Brinell and Vickers hardness tests are both used to measure the hardness of dental alloys, and hardness values on alloy packages are expressed in either Vickers or Brinell hardness numbers. To compare Vickers and Brinell numbers, use the following relationship:

$$\text{VHN} = 1.05 \times \text{BHN},$$

or

$$\text{BHN} = \frac{\text{VHN}}{1.05}$$

For example, if a package of casting alloy lists a Vickers hardness of 242 and you want to compare it with a different brand that lists the Brinell hardness, you would divide the VHN by 1.05 to obtain a BHN of 230. Diamond pyramid (Vickers) hardness values for several types of materials are listed in Appendix A.

Knoop hardness (KHN)

The indenter for the Knoop hardness test is also made of a diamond, but its outline is somewhat different from the Vickers indenter; although it is diamond shaped, one diagonal is much longer than the other. Only the long diagonal is measured to determine the Knoop hardness number. Knoop hardness numbers for a variety of materials are listed in Appendix A.

Rockwell hardness (R_A, R_B, . . . R_G)

The Rockwell hardness test is used primarily for determining the hardnesses of steels and is the most widely used hardness test in the United States. The Vickers, Brinell, and Knoop hardness tests are more commonly used for dental materials, however. Rockwell uses different hardened steel balls or diamond cones and different loads. Each combination forms a specific Rockwell scale (A, B, and C scales are the most common). The different scales are used for materials of different hardness ranges.

Shore A durometer

The Shore A hardness test is used to measure the hardness of rubbers and soft plastics. The Shore A scale is between 0 and 100 units, with complete penetration of the material by the indenter yielding a value of 0, and no penetration yielding a value of 100. Shore A hardness values for several polymeric materials are listed in Appendix A.
Thermal Properties

Heat flow through a material

Metals tend to be good conductors of heat, and this property must be taken into consideration when placing metallic restorations. Dentin is a thermal insulator (poor conductor of heat); thus, when a sufficient thickness of dentin is present, the patient feels no sensitivity to heat and cold through a metallic restoration. However, when only a thin layer of dentin remains, some thermal protection must be provided for the pulp. A good rule of thumb in determining the thickness of cement base necessary in a given situation is to visualize how much dentin would have to be present in the excavation site so no base would be necessary and then to apply base up to this level. The rate at which heat flows through a material is expressed as thermal conductivity or thermal diffusivity. The difference between these two terms is described in the following subsections.

Thermal conductivity

Thermal conductivity \((k)\) is a measure of the speed at which heat travels (in calories per second) through a given thickness of material (1 cm), when one side of the material is maintained at a constant temperature that is \(1^\circ\text{C}\) higher than the other side. Thermal conductivity is expressed in units of cal/cm \(\cdot\) sec \(\cdot\) \(^\circ\text{C}\). Thermal conductivity values for a variety of materials are listed in Appendix A.

Thermal diffusivity

Whereas thermal conductivity gives an idea of the relative rates at which heat flows through various materials, it fails to take into account the fact that various materials require different amounts of heat (calories) to raise their temperatures an equal amount. For example, 1 gram of water requires 1.000 calorie to raise its temperature \(1^\circ\text{C}\), whereas 1 gram of dentin requires only 0.28 calorie, and 1 gram of gold requires only 0.031 calorie to produce a \(1^\circ\text{C}\) temperature increase. Thus, thermal conductivity alone will not tell us, for instance, how rapidly the interior surface under a crown will heat up when the exterior surface is heated. To know how quickly the interior of the crown will approach
the temperature of the exterior, we need to know the thermal diffusivity of the alloy. The thermal diffusivity \( (h) \) of a material (expressed in units of mm\(^2\)/sec) is dependent on its thermal conductivity, heat capacity \( (C_p) \), and density \( (\rho) \):

\[
h = \frac{k}{(C_p \times \rho)}
\]

The relative thermal diffusivities of several dental materials are shown in Fig 2-8, and tabular data on a variety of materials are included in Appendix A.

**Thermal expansion**

There are several situations in dentistry in which the thermal expansion of materials is important. Some restorative materials have coefficients of thermal expansion that are markedly different from tooth structure. In such cases, temperature fluctuations that occur in the mouth can cause **percolation** at the tooth-restoration interface as the restoration contracts and expands. The porcelain and metal in a porcelain-fused-to-metal (PFM) restoration must contract at the same rate upon cooling from the porcelain firing temperature if the buildup of large residual stresses is to be avoided. The cooling of a denture base from the processing temperature to room temperature is primarily responsible for the processing shrinkage that occurs. The thermal expansion behavior of dental wax, gold alloy, investment, and so on are all important in producing properly fitting castings. **Figure 2-9** illustrates the relative values of **linear coefficient of thermal expansion** for tooth, amalgam, composite, and acrylic resin. The diagram is only schematic—the expansion has been magnified to make it visible; the actual thermal expansion would be too small to see. The thermal expansion coefficients, or the fractional changes in length per degree celsius, are given in parentheses. Tabular thermal expansion data on a variety of materials are included in Appendix A.
Fig 2-9 Relative thermal expansions of several restorative materials and tooth structure. The relative amounts the materials expand when heated the same amount are represented by the dashed lines. The amount of expansion is magnified to make it visible for comparison of the materials.

Electrical and Electrochemical Properties

Electrode potentials

An electrochemical series is a listing of elements according to their tendency to gain or lose electrons in solution. The series is referenced versus the potential of a standard hydrogen electrode, which is arbitrarily assigned a value of 0.000 volts. If the elements are listed according to the tendency of their atoms to lose electrons, the potentials are termed oxidation potentials. If they are listed according to the tendency of their ions to gain electrons, the potentials are termed reduction potentials. Reduction potentials at 25°C and 1 atmosphere of pressure are termed standard electrode potentials (Table 2-1).

<table>
<thead>
<tr>
<th>Half-reaction</th>
<th>( \Delta E^0 ) (volts)</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \text{Li}^+ + e = \text{Li} )</td>
<td>-3.04</td>
</tr>
<tr>
<td>( \text{K}^+ + e = \text{K} )</td>
<td>-2.93</td>
</tr>
<tr>
<td>( \text{Ca}^{2+} + 2e = \text{Ca} )</td>
<td>-2.87</td>
</tr>
<tr>
<td>( \text{Na}^+ + e = \text{Na} )</td>
<td>-2.71</td>
</tr>
<tr>
<td>( \text{Mg}^{2+} + 2e = \text{Mg} )</td>
<td>-2.37</td>
</tr>
<tr>
<td>( \text{Al}^{3+} + 3e = \text{Al} )</td>
<td>-1.662</td>
</tr>
<tr>
<td>( \text{Zn}^{2+} + 2e = \text{Zn} )</td>
<td>-0.762</td>
</tr>
<tr>
<td>( \text{Cr}^{3+} + 3e = \text{Cr} )</td>
<td>-0.744</td>
</tr>
<tr>
<td>( \text{Fe}^{2+} + 2e = \text{Fe} )</td>
<td>-0.447</td>
</tr>
<tr>
<td>( \text{Ni}^{2+} + 2e = \text{Ni} )</td>
<td>-0.257</td>
</tr>
<tr>
<td>( \text{Sn}^{2+} + 2e = \text{Sn} )</td>
<td>-0.1375</td>
</tr>
<tr>
<td>( \text{Pb}^{2+} + 2e = \text{Pb} )</td>
<td>-0.1262</td>
</tr>
<tr>
<td>( \text{Fe}^{3+} + 3e = \text{Fe} )</td>
<td>-0.037</td>
</tr>
<tr>
<td>( \text{H}^+ + e = \text{H} )</td>
<td>0.000 (reference)</td>
</tr>
<tr>
<td>( \text{Cu}^{2+} + 2e = \text{Cu} )</td>
<td>+0.342</td>
</tr>
</tbody>
</table>
Metals with a large positive electrode potential, such as platinum and gold, are more resistant to oxidation and corrosion in the oral cavity. If there is a large difference between the electrode potentials of two metals in contact with the same solution, such as between gold and aluminum, an electrolytic cell may develop. If this occurs in the mouth, the patient may experience discomfort.

The exact nature of tarnish and corrosion of restorative materials in vivo is extremely complex and involves much more than electrode potentials of materials. A possible source of the corrosion of gold alloys is accidental contamination of the surface with copper during pickling, and subsequent tarnishing in saliva (Fig 2-10).

![Fig 2-10](image)

**Electrical resistivity**

**Electrical resistivity** measures the resistance of a material to the flow of an electrical current. The relationship between the resistance (R) in ohms, the resistivity (ρ), the length (l), and the cross-sectional area (A) is as follows:

\[ R = \rho \left( \frac{l}{A} \right) \]

Electrical resistivity values for several materials are given in Table 2-2.

**Table 2-2** Electrical resistivity of several materials
<table>
<thead>
<tr>
<th>Material</th>
<th>Resistivity (ohm-cm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Copper</td>
<td>$1.7 \times 10^6$</td>
</tr>
<tr>
<td>SiO₂ (glass)</td>
<td>$&gt; 10^4$</td>
</tr>
<tr>
<td>Human enamel</td>
<td>$\sim 4 \times 10^6$</td>
</tr>
<tr>
<td>Human dentin</td>
<td>$\sim 3 \times 10^4$</td>
</tr>
<tr>
<td>Zinc phosphate cement</td>
<td>$2 \times 10^5$</td>
</tr>
<tr>
<td>Zinc oxide-eugenol</td>
<td>$5 \times 10^9$</td>
</tr>
</tbody>
</table>

The low resistivity of metallic restorative materials is responsible for discomfort to the pulp if dissimilar metals generate a voltage. The insulating properties of cements help to reduce this problem.

**Biocompatibility**

A material that is designed to be implanted in the body for the repair or replacement of a diseased or damaged tissue is called a **biomaterial**. Biomaterials must be **biocompatible**. All materials used in medicine and dentistry interact with tissues, producing changes in both the materials and the surrounding tissues; there is no such thing as an "inert" material.

Several factors affect the biocompatibility of a material. The chemical composition and structure of the material affect the type and quantities of chemical species released. The relative abundance of chemical species in a material will not reliably predict the amounts or even the proportions of the chemical species released. For this reason, the biocompatibility of a material is usually determined empirically. The characteristics of the site of use also affect a material's biocompatibility. For example, the presence of many factors in the oral environment affect the release of components from dental materials. Chemical factors, such as bacterial metabolic products, water, enzymes, and polar and nonpolar solvents, can accelerate the release of components. Mechanical factors in the site of use can also affect the biocompatibility of a material. For example, under most conditions, high-molecular-weight polyethylene elicits little if any tissue reaction when implanted. When it is used for the acetabular (socket) side of a total hip replacement, however, it can generate wear debris over time that produces an inflammatory response that eventually causes loosening of the prosthesis and failure of the implant.

Evidence on a new product for use in the body must first be presented to government agencies that document possible adverse reactions the material might cause. This information includes the history of the material, its chemical composition, and results from a series of biocompatibility tests. A product that is a new version of an old composition used successfully in similar applications may be classified under the Federal Drug Administration (FDA) classification of 510(k), which requires less extensive testing for premarket approval. Another strategic qualification is whether the new product has temporary or long-term contact with the body. The FDA has different categories for materials with short-term contact, such as an impression material, than for materials used for long-term service, such as a cemented crown or an implant. The more potential for harm, the greater the burden of testing required. The testing required depends on many factors, including the results of initial minimal screening tests. A brief description of these tests is provided in **Appendix B**.

**Glossary**
biocompatible A material is considered biocompatible if it does not produce harmful or toxic reactions in the tissues it contacts or adverse systemic reactions as a result of elements, ions, and/or compounds it releases.

biomaterial A material that is designed to be implanted in the body for the repair or replacement of a diseased or damaged tissue.

brittle A material that tends to fracture without appreciable plastic deformation is said to be brittle.

compressive stress Two forces applied toward one another in the same straight line.

ductility Ability of a material to be plastically strained in tension.

elasticity Ability to sustain deformation without permanent change in size or shape.

electrical resistivity Ability of a material to resist conduction of an electric current.

elongation Overall deformation (elastic + plastic) as a result of tensile force application.

fatigue Tendency to fracture under cyclic stresses.

hardness Resistance to permanent indentation on the surface.

linear coefficient of thermal expansion Change in length per unit original length for a 1°C temperature change.

modulus of elasticity Stiffness of a material within the elastic range. Numerically, it is the ratio of stress and strain.

percolation The pumping of oral fluids in and out at the tooth-restoration interface as the restoration contracts and expands with temperature changes. Percolation occurs when the thermal expansion coefficient of the restoration is markedly different from that of tooth structure.

proportional limit (elastic limit) The maximum stress at which the straight-line relationship between stress and strain is valid.

resilience Energy needed to deform a material to the proportional limit.

shear stress Two forces applied toward one another but not in the same straight line.

strain (nominal) Change in length per unit original length.

stress (nominal) Force per unit area.

tensile stress Two forces applied away from one another in the same straight line.

thermal conductivity The quantity of heat passing through a material 1 cm thick with a cross section of 1 cm², having a temperature difference of 1°C.
**thermal diffusivity** Measure of the heat transfer of a material in the time-dependent state.

**toughness** Amount of energy needed for fracture.

**ultimate tensile strength** The maximum strength obtained based on the original dimensions of the sample.

**viscoelastic** Having both elastic and viscous properties.

**viscous** Resistant to flow (referring to a fluid).

**work hardening** The increase in strength and hardness, with accompanying decrease in ductility, that occurs in a ductile metal as it is plastically deformed. Also called *strain hardening*.

**yield strength** Strength measured at the stress at which a small amount of plastic strain occurs. Also called yield point.

**Discussion Questions**

1. What is galvanic action in the mouth and how can it be minimized?

2. Why is thermal diffusivity more relevant to insulation of the pulp than thermal conductivity?

3. Explain why fixed partial dentures may fail under tensile stresses when under biting forces that appear to be compressive?

4. How is the leakage of oral fluids around a composite restoration related to the coefficient of thermal expansion and temperature changes in the mouth?

5. Why is time so important in the behavior of viscoelastic materials such as impression materials and waxes?

6. Why is there a hierarchy of biocompatibility tests?

7. Why can biocompatibility tests be difficult to interpret?

**Study Questions**

*(See Appendix E for answers.)*

1. Discuss why a knowledge and understanding of the physical and mechanical properties of biomaterials is important in dentistry.

2. Explain why the thermal diffusivity of a material is more applicable to behavior in vivo than is its thermal conductivity.

3. Draw and label a stress-strain diagram including the proportional limit, yield point, and ultimate tensile strength.

4. Define hardness and its relation to other mechanical properties.
5. Compare materials \(a\) and \(b\) on the basis of the two stress-strain curves below, obtained by pulling the materials in tension.

![Stress-strain curves](image)

6. Why are laboratory biocompatibility tests run prior to clinical trials?

**Recommended Reading**


Chapter 3. Color and Appearance

Introduction

Among the important factors that influence esthetic appearances of restorations are color, translucency, gloss, and fluorescence. Each of these factors, as perceived by an observer such as a dentist, technician, or patient, is influenced by (1) the illuminant (light source), (2) the inherent optical parameters of the restorative materials that dictate the interaction of the light from the illuminant with the material, and (3) the interpretation of the observer (Fig 3-1). An understanding of these factors in restorative materials, and proper consideration and communication of this information, will greatly assist the choice and fabrication of esthetic restorations.
Fig 3-1 Illuminant, object, and observer interaction.

**Illuminant**

*Introduction*

Light waves emitted from the illuminant interact with the object and are perceived by the observer. The color content of the illuminant interacts with the object being perceived. The color content of the illuminant is the intensity of light emitted at each wavelength (spectral distribution). This is dependent on the type of illuminant. The wavelengths of light are associated with hues, commonly referred to as color (Fig 3-2).

Fig 3-2 Wavelength and hue. (Adapted from Billmeyer and Saltzman, 1960.)

White light contains a mixture of wavelengths (colors). It is dispersed into components when it passes through a prism, as shown in Fig 3-3.
Different illuminants have different intensity distributions, or spectra, with respect to wavelength, as illustrated in Figs 3-4a and 3-4b.
Illuminants are sometimes described by their color temperature (in degrees Kelvin). This is based on the equivalence of the illuminant as compared to a radiating body with a temperature equal to the color temperature.

The characteristics of an illuminant are important in color evaluation because the intensity distribution, with respect to wavelength, identifies the light spectrum available to interact with the object and then be perceived by the observer. In any color description the type of illuminant used needs to be defined. Furthermore, the appropriate illuminant needs to be used in shade selection and shade matching. Illuminants that closely approximate daylight are preferred because color considerations for restorations, when seen under these illuminants, are close to those seen under natural light. Several "daylight" sources intended for use in dental operatories are available. Representative spectral distributions of some of these illuminants are shown in Figs 3-5a and 3-5b.
Fig 3-5a Spectral distribution of Neylite (J.M. Ney), a device used for dental shade matching.

Fig 3-5b Spectral distribution of Vita-Lite (Duro Test), a fluorescent lamp for dental operatories.

Color-rendering index

Another parameter used in characterizing an illuminant is the color-rendering index. Color rendering of a light source is the effect that the source (illuminant) has on the color appearance of objects in comparison with their color appearance under a reference source. The color-rendering index is a measure of the degree to which the illuminant can impart the color of an object as compared to the reference source. A color-rendering index of 100 is considered ideal. For an adequate color-matching environment, the illuminant should have a color-rendering index of 90 or above (Preston et al, 1978). A summary of color-rendering indices of some illuminants used in dentistry is shown in Table 3-1.

<table>
<thead>
<tr>
<th>Illuminant (manufacturer)</th>
<th>Color-rendering index</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chroma 50 (General Electric)</td>
<td>92</td>
</tr>
<tr>
<td>Chroma 75 (General Electric)</td>
<td>94</td>
</tr>
</tbody>
</table>
Surroundings in a dental operatory may modify the actual light reaching the object. Colors of walls, clothing, and soft tissues such as lips contribute to the color of the light incident on teeth, shade guides, and restorative materials.

**Object**

**Introduction**

The inherent color property of an object is its characteristic interactions with the light from the illuminant. These interactions include reflection, transmission, and the absorption involved in both processes.

**Reflection**

A material gains its reflective color by reflecting that part of the spectrum of light incident upon it and absorbing the other parts of the light spectrum. A blue surface reflects only the blue part of the light spectrum and absorbs all other colors (Fig 3-6).

![Fig 3-6 Reflected color. A blue object reflects only blue light and absorbs all other colors.](image)

A **white object** reflects all incident wavelengths. A **black object** absorbs all wavelengths and reflects none. An object also appears black when no light is reflected from it; for example, a blue object appears black when viewed in red light. Materials of different **reflected color** have different **color reflectance** spectra (Fig 3-7). Reflection spectra of objects are usually obtained using a spectrophotometer.
Mixing of reflected colors

To obtain a desired esthetic appearance for a restoration, it is often necessary to use more than one colorant. Each colorant has its own characteristic reflected color because of its reflection spectrum and its absorption of other parts of the spectrum. The mixing of two colorants with different reflected colors, and therefore different absorptions, results in a reflection of the part or parts of the light spectrum common to both colorants. In other words, each colorant will absorb the part of the light spectrum as if it were by itself. The parts of the light spectrum that are not absorbed by either colorant are, therefore, the resultant reflected color of the mixture of the two.

Transmission

A translucent material gains its transmitted color by the resultant spectrum it transmits. As a beam of light passes through a translucent material (eg, glass), it can be changed by the color of the material or scattered by fine particles, giving a milky appearance. Wavelengths that are not transmitted are absorbed. For example, a green filter transmits light in the green wavelength region of the light spectrum and absorbs all other wavelengths.

Observer

The observer receives the light reflected or transmitted by the object and then interprets the results. In many cases, the observer uses the human eye as the detector. Eye responses vary among individuals. The human eye response varies with wavelength. It is most sensitive in the green color region (Fig 3-8). The human eye is best in detecting color differences by comparison.
The detection of color by the human eye results from stimuli received by cone-shaped cells in the retina. Color blindness—the inability to distinguish certain colors—is due to abnormalities in these cells. Constant stimulus of one color decreases the response of the eye to that color. This is sometimes known as color fatigue. After removal of the stimulus, a complementary color image may persist.

Other detectors may be used as an observer in place of the human eye. These are usually photodetectors, such as spectrophotometers or colorimeters. The response of photodetectors varies among types, and they differ from the response of the human eye. Color-measuring devices are designed to minimize the effect of photodetector responses.

In color measurements and parametric color determinations, the CIE (Commission Internationale de l'Eclairage) Standard Colorimetric Observer is often referenced.

**Color Systems**

*Introduction*

**Color systems** are used to describe the color parameters of objects. The following are examples of some color systems used in describing the color of dental materials.
**Munsell color system**

The Munsell color system uses a three-dimensional system with hue, value, and chroma as coordinates (Fig 3-9).

![Munsell color system diagram](image)

**Fig 3-9** Munsell color system with hue, value, and chroma as coordinates.

- **Hue** is commonly referred to as color; examples are blue, yellow, and red. Hue is also associated with the wavelengths of the light observed.

- **Value** is the lightness or darkness of a color. A tooth of low value appears gray and nonvital. Value is the most important color factor in tooth color matching.

- **Chroma** is a measurement of color intensity; that is, the amount of hue saturation in a color. An example is a beaker of water containing one drop of colorant; it is lower in chroma than a beaker of water containing ten drops of the same colorant.

**CIE color systems**

The CIE tristimulus values system uses three parameters, X, Y, and Z, which are based on the spectral response functions defined by the CIE observer. A CIE chromaticity diagram is also sometimes used to define color.

Another CIE color system (CIE L*a*b*) uses the three parameters L*, a*, and b* to define color. The L*, a*, and b* values can be calculated from the tristimulus X, Y, and Z values. The advantage of this color system is that its arrangement is an approximately uniform three-dimensional color space whose elements are equally spaced on the basis of visual color perception. A unit change in each of the three color parameters is...
approximately equally perceived. The quality $L^*$ correlates to lightness, similar to value in the Munsell system. The $a^*$ and $b^*$ coordinates describe the chromatic component (Fig 3-10).

![Fig 3-10 CIE and Munsell color arrangements. (From Seghi et al, 1986. Reprinted with permission.)](image)

The color difference, $\Delta E^*$, in the CIE $L^*a^*b^*$ system is defined as:

$$\Delta E^* = \sqrt{(\Delta L^*)^2 + (\Delta a^*)^2 + (\Delta b^*)^2}$$

where $\Delta L^*$, $\Delta a^*$, and $\Delta b^*$ are the differences between the CIE $L^*a^*b^*$ color parameter of two samples. An advantage of $\Delta E^*$ is that it can serve as a tolerance for color matching. Clinical color matching between teeth and restorations may be rated according to $\Delta E^*$ values, as given in Table 3-2, based on clinical studies. Shade guides are held to a tolerance of a $\Delta E^*$ of 1 according to an American Dental Association (ADA) standard. Although these $\Delta E^*$ values can serve as approximate tolerances, some individuals perceive color differences as low as 0.5, whereas others do not see differences of 4. This is often a source of disagreement among patients, dentists, and laboratory technicians.

**Color Measurements**

Apart from visual comparison using color standards such as Munsell color chips, color measurements can be made using either spectrophotometric or colorimetric methods.

**Spectrophotometers** measure the amount of light reflected at each wavelength. A double-beam spectrophotometer compares the responses from the object and a reference standard. From the spectral response, color parameters for the object can be calculated. Spectrophotometric measurements have been used to evaluate the color parameters for
restorative resins, denture teeth, porcelains, shade guides, and color changes in dental materials. Figure 3-11 shows one model of a spectrophotometer.

![Figure 3-11: Spectrophotometer for laboratory measurement of color parameters (Macbeth 7000, Gretag-Macbeth.)](image)

**Colorimeters** measure the amount of light reflected at selected colors (e.g., red, green, blue). The selections are based on the CIE tristimulus value standard observers. There are several instruments available, and applications of their measurement methods for dental materials have been reported (Powers et al, 1980; O’Brien et al, 1983; Stanford et al, 1985). The instruments give readings in tristimulus values (X, Y, Z) or CIE L*a*b* values.

The major technical problem in using such instruments to measure the color parameters of teeth is the **edge effect**. Dental enamel is a translucent material that scatters the incident light from the instrument at many angles. As a result, the light reflected back into the instrument for analysis is not reliable and presents major errors in shade selection.

A new approach to tooth shade selection is the use of photography in combination with a spectrophotometer. Although accurate instrumental measurements of color parameters of translucent materials are difficult to use, color measurements on photographs are routine. This procedure of photocolorimetry involves the following steps:
1. The clinician or assistant holds the 3 or 4 closest-matching shade guide teeth next to the patient's teeth and photographs them with a 35-mm camera under a balanced light source (Fig 3-12).

![Fig 3-12 Photograph of patient's teeth compared with the closest-matching shade guide teeth.](image)

2. The photograph is sent to the dental laboratory along with the impression. The laboratory technician measures the color of the patient's teeth and shade guide teeth in the photograph. A computer program then aids in selecting the closest match.

This approach has been tested in a clinical study (Groh et al, 1999) in comparison with visual matching of 30 patients. The difference in color parameters between shades selected by experienced observers and those selected by photocolorimetry resulted in a $\Delta E^*$ of 1.88, which is within the color tolerance of a $\Delta E^*$ of 2.00 for shade guide teeth. In another study (O'Brien et al, 1983), the accuracy of matching one Vita shade guide was significantly higher with photocolorimetry than with visual matching by dental faculty and technicians.

Several chairside colorimeters have been introduced in the past 3 years (Fig 3-13). These instruments determine tooth shade by direct application to a patient's tooth. Their accuracy is difficult to assess and depends on the angle at which the instrument is positioned and other operator variables. A dental professional who can match shades well (eg, two shade guides at 90%) is still the standard, but technology is improving. High prices of these instruments ($6,000 to $15,000) limit adoption.
Metamerism

The change in color matching of two objects under different light sources is called metamerism. Two objects that are matched under one light source but not under other light sources form a metameric pair (Fig 3-14). They have different color reflectance curves. An example of metamerism is when a shade-guide tooth matches the tooth under fluorescent light but not under incandescent light. Metamerism results from possible differences in illumination between the dental clinic and the dental laboratory, causing poor matching in a fabricated restoration, such as a porcelain crown.

Standardization of illuminations (usually similar to the patient's environment) in color matching diminishes the effect of metamerism in shade matching. The ideal situation, of course, is to have the objects possess the same color reflectance curve. The objects are then an isomeric pair; they are color matched under all light sources.
Fig 3-14 Metamerism. Objects 1 and 2 match under light source A but do not match under light source B.

Translucency

The translucency of an object is the amount of incident light transmitted and scattered by that object. A high translucency gives a lighter color appearance. A more translucent material will show more of the backing in the color and appearance. Translucency decreases with increasing scattering within the material.

The opposite of translucency is opacity. Light scattering in a material is the result of scattering centers that cause the incident light to be scattered in all directions. Examples of scattering centers are air bubbles and opacifiers such as titanium dioxide. Another example is the filler particles in a resin composite matrix. The effect of scattering is dependent on the size, shape, and number of scattering centers. Scattering is also dependent on the difference in refractive indices between the scattering centers and the matrix in which the centers are located.

Measurements of translucency may be performed using transmission spectrophotometers, reflection spectrophotometers, light meters, or colorimeters. Measurements of translucency of dental porcelains and human enamel are published in the dental literature (Brodbelt et al, 1980; Brodbelt et al, 1981; O'Brien et al, 1985; Spitzer and ten Bosch, 1975).

Gloss

Surface gloss is the optical property that produces a lustrous appearance. Contrast gloss, or luster, is the proportion of specular reflection to diffuse reflection (Fig 3-15). Another consideration for gloss is the amount of collimated incident light that is specularly reflected. In specular reflection, the angle of incidence is equal to the angle of reflectance. When the incident beam is scattered by the object, there is a decrease in gloss as a larger portion of the incident beam is diffuse-scattered.

Gloss is an important appearance property of dental restorative materials. A high surface gloss is usually associated with smooth surfaces. In resin composite, surface gloss decreases with increasing surface roughness. Differences in gloss between multiple
restorations or between restorations and teeth can be easily detected even in colors that are matched. In addition, high gloss reduces the effect of a color difference, because the color of the reflected light is more prominent. In a restorative material, high gloss also lightens the color appearance.

Fluorescence

Fluorescence is the emission of light by an object at wavelengths different from those of the incident light. The emission ceases immediately upon removal of the incident light. Natural teeth fluoresce in the blue region when illuminated by ultraviolet light. Dental porcelains are also fluorescent under ultraviolet light. The quality of the fluorescence depends on the brand of porcelain, some of which fluoresce in colors different from those of natural teeth.

Double Layer Effects on Esthetics

The color of a tooth is strongly influenced by the thickness of the enamel and the color of the underlying dentin. Similar considerations apply to restorations that are of layered structure. Examples are porcelain restorations that are composed of body porcelain over an inner opaque porcelain (Fig 3-16) and resin composites over more opaque resins. In these layered structures, esthetic appearances are no longer simple considerations of the factors previously described but also involve diffuse reflectance and the relation between the translucency and thickness of the outer layer and the color and reflectance of the inner layer. The outer translucent layer acts as a light-scattering filter over the inner layer (Fig 3-17). As the thickness of the outer layer increases, the effect of the inner layer is diminished. Similar situations also exist when the translucency of the outer layer decreases. Considerations of models of diffuse reflectance in dental porcelain systems using the Kubelka-Munk equation have shown excellent agreement between experimentally observed and theoretically calculated color parameters (O'Brien et al, 1985; Ragain and Johnston, 2001).
**Fig 3-16** Optical considerations for a porcelain-fused-to-metal restoration. (From O'Brien et al, 1985. Reprinted with permission.)

**Fig 3-17** Double layer effect. (From O'Brien, 1985. Modified with permission.)
Dental Shade Guides

Shade guides are used in determining the color of natural teeth so that artificial substitute restorations will possess similar color and esthetics. Preferred properties in a shade guide include logical arrangements and adequate distribution in color space, matching with natural teeth, inherent consistency among shade guides, and matching between shade guides and the dental materials such as porcelains and resin composites or denture teeth. Not all these properties are met by the shade guides currently available. Furthermore, not all shade guides are fabricated from the dental materials to which they will be matched.

Most shade guides use a designation to denote the shade and color. The same designation may not be comparable among brands. The distribution of shades within a shade guide is not necessarily evenly partitioned in color space. Some attempts have been made to use a more logical approach and an even distribution in shade guides, but only a limited number follow this approach. Recently a new shade guide (Vita 3D Guide, Vita) has been introduced that is based on hue, value, and chroma in a logical arrangement.

Shade Matching in the Dental Operatory

Shade matching is a complex situation. It is important to remember the triadic interactions of illuminant, object, and observer described previously. Considerations should also include metamerism, gloss, translucency, and fluorescence. Recognition of the factors influencing shade match improve the result of the match.

The most important factor in shade matching is the illuminant. This is the lighting in the dental operatory used in shade matching. A color-corrected light source with a color temperature of 5,500 K and a color-rendering index of 90 or above is recommended. If possible, the shade matching should also be checked under a different light condition, for example, a warm white fluorescent light. In cases where the patients may have specific requirements, such as extensive activity under certain lighting conditions, shade-matching checks under those conditions are also recommended. The color environment of the dental operatory is another important factor in shade matching. A neutral, light gray background color reduces modification of color perception.

Some recommendations in shade matching are as follows:

1. Shade match under lights of similar spectral distribution and intensity, both in the dental operatory and the laboratory. Lighting conditions should be similar to daylight. A dental ring light is one such option (Fig 3-18).
2. Consider the effects of translucency and position. A high translucency and a more distal position in the natural dentition cause a darker gray appearance.

3. Use the manufacturer's shade guide for fabricating the restoration.

4. Follow the manufacturer's recommendations for preparing the surface of the tooth for shade matching. A diffuse-reflection condition is usually used.

5. Remove the individual shade tab from the guide and hold it close to the tooth for shade matching.

6. The surface texture of the restoration should match that of the remaining dentition as closely as possible.

**Communication of Shade-Matching Information to the Laboratory**

The shade tab used for shade matching should be transmitted to the laboratory and should be specifically for the material to be used to fabricate the restoration. The type of lighting used for shade matching should be included in the communication to decrease the effect of metamerism. The prescription should accurately transmit the shade information as well as any characterization. An example of a map indicating the approximate shade, its depth, and information about its location is shown in Fig 3-19.
The sequential steps in the overall transmission of shade-matching information and the final clinical acceptance of the fabricated restoration have been described by O'Brien (1987). A schematic illustration is shown in Fig 3-20. The source spectrum is the color and translucency information of the tooth to which the fabricated restoration is to be matched. The shade taking is the encoding process. The information is decoded in the dental laboratory. The simulation is the fabricated restoration, which is then delivered to the dentist observer for evaluation and acceptance. If necessary, additional correction data to enhance the matching are communicated back for further simulation until the final simulation is reached. This is the fabricated restoration with necessary corrections to attain a clinically acceptable degree of shade matching to adjacent teeth.
**Fig 3-20** Schematic illustration of transmission of shade-matching information.

**Glossary**

**black object** An object that reflects no incident color lights.

**chroma** Color intensity.

**color content** Relative intensity at each wavelength.

**color fatigue** Decrease in response to one color due to constant stimulus.

**colorimeter** A device used to measure the amount of light reflected at selected colors.

**color reflectance** Relative amount of each color reflected.

**color system** A three-dimensional system for defining color.

**edge effect** Lateral scattering of a light beam upon entering a translucent material; causes an error in color measurement.

**fluorescence** Glow of an object when illuminated.

**gloss** Shininess; relative amount of light reflected.

**hue** Commonly called color (eg, blue, yellow). It is associated with wavelength.

**isomeric pair** Two objects that match color under different light sources.

**metameric pair** Two objects that exhibit metamerism.

**metamerism** Change in color matching of two objects under different light sources.

**opacity**

\[
\text{Opacity} = \frac{\text{Intensity of incident light}}{\text{Intensity of transmitted light}}
\]

High opacity is associated with low intensity of transmitted light.
**reflected color** Color reflected by an object.

**refractive index**

\[
\text{Refractive index } (n) = \frac{\text{Velocity of light in air}}{\text{Velocity of light in medium}}
\]

Refractive index of dental porcelain is 1.5. Refractive index of tin dioxide (an opacifier) is 2.0.

**spectrophotometer** A device used to measure the amount of light reflected at each wavelength.

**translucency** Amount of light transmitted; the rest of the light is scattered.

**value** Lightness or darkness of a color; a measurement of the amount of gray.

**white object** An object that reflects all incident color lights.

**Discussion Questions**

1. How would a dental clinic need to be changed in order to be a better environment for color matching?

2. Why is it not necessary that a dental restoration match adjacent tooth structure exactly? About how far off in color can it be?

3. Why is it necessary to send the selected shade guide sample to the dental laboratory rather than just the shade guide number?

4. Besides color, which other appearance properties are important in creating an esthetic restoration?

**Study Questions**

*(See Appendix E for answers.)*

1. Name the coordinates required to define a color in a color system.

2. A porcelain crown may appear gray and nonvital; which variable is involved?

3. Why does a green object appear black in blue light?

4. What factors affect color appearance?

5. How does translucency affect color appearance?

6. What does gloss do to color appearance?

7. What is metamerism?

8. How does metamerism affect the appearance of dental restorations?
9. Why do dental porcelains appear different under different lighting environments?

**Recommended Reading**


Chapter 4. Gypsum Products

Introduction

In general, the term gypsum products refers to various forms of calcium sulfate, hydrous and anhydrous, manufactured by the calcination of calcium sulfate dihydrate (CaSO\(_4\) · 2H\(_2\)O), which occurs as the mineral gypsum. Calcination can be controlled to produce partial or complete dehydration. Gypsum products can also be obtained by calcining "synthetic" or "chemical" gypsum, a by-product of the manufacture of phosphoric acid. Industrially, all these materials are known as gypsum plasters.

Although not directly employed in dental restorations, gypsum products are important accessory materials used in many clinical and laboratory procedures. Their correct use contributes to the success of these procedures. They are classified by the International Standards Organization (1998) into five types:

Type 1: Impression plaster
Type 2: Plaster
Type 3: Stone
Type 4: Stone, high-strength, low-expansion
Type 5: Stone, high-strength, high-expansion

This classification is illustrated in Fig 4-1. Both types of plaster are based on ordinary commercial gypsum plaster (plaster of Paris), while the three types of stone are based on high-strength gypsum plasters. Types 4 and 5 stones are commonly referred to as die stones. Table 4-1 lists typical materials.

<table>
<thead>
<tr>
<th>Type</th>
<th>Example</th>
</tr>
</thead>
<tbody>
<tr>
<td>Impression plaster (ISO Type 1)</td>
<td>Impression Plaster (Modern Materials Manufacturing)</td>
</tr>
<tr>
<td>Plaster (ISO Type 2)</td>
<td>Snow White Plaster No. 1 (Kerr/Sybron)</td>
</tr>
<tr>
<td>Stone (ISO Type 3)</td>
<td>Microstone (Whip Mix)</td>
</tr>
<tr>
<td>Stone, high-strength, low-expansion</td>
<td>Vel Mix Stone (Kerr/Sybron)</td>
</tr>
</tbody>
</table>
The term gypsum products can be extended to include gypsum-bonded investments, used as mold materials for casting dental gold alloys, because calcium sulfate hemihydrate is an essential component, forming the bond in the set material. These materials will be dealt with separately later in this chapter.

![Fig 4-1 Classification of gypsum products.](image)

### Plaster and Stone

Both plaster and stone are the result of partial dehydration of gypsum, which produces calcium sulfate hemihydrate (CaSO$_4$·$^{1/2}$H$_2$O). Differences in properties result from differences in the physical nature of the powders, which in turn are a result of differences in manufacturing methods.

### Chemistry

In the temperature range 20°C to 700°C, which is of importance in the dental manipulation of gypsum products, three phase transformations occur in the calcium sulfate-water system. The first two represent the two stages in the dehydration of gypsum, and these are followed by a further transformation into anhydrous calcium sulfate:
It is not possible to give unequivocal temperatures for these transformations; the ranges given summarize the results of several different determinations.

For the first transformation, from dihydrate to hemihydrate, the temperature limits were determined by weight loss measurements on specimens heated isothermally in dry air for 2.5 years (Andrews, 1951) and would represent an equilibrium value. Thermogravimetric measurements of the temperature for the second transformation, calcium sulfate hemihydrate to hexagonal calcium sulfate, have involved isothermal heating for periods only up to a maximum of 22 days (Khalil et al, 1971; Weiser et al, 1936), so the true equilibrium temperature for this transformation is probably a little lower than shown. Gay (1965a,b) has identified hexagonal calcium sulfate by x-ray diffraction in specimens of hemihydrate heated in the range of 75°C to 105°C. Hexagonal calcium sulfate is unstable below about 80°C, and, if cooled to lower temperatures and exposed to the environment, it rapidly rehydrates to form the hemihydrate. The temperature range given for the third transformation, hexagonal to orthorhombic calcium sulfate, is based on measurements made on specimens heated at a rate of 5 K/min (Earnshaw and Mori, 1985), so again the true equilibrium temperature for this transformation would be lower than shown here. Orthorhombic calcium sulfate is the stable anhydrous form in this system and exists as the mineral insoluble anhydrite.

Theoretically, hemihydrate is the stable hydrous form of calcium sulfate only in the approximate temperature range of 45°C to 90°C. It exists as a metastable phase under dry conditions at lower temperatures, including ambient, although it has been shown that hydration can occur if particles are exposed to the atmosphere under conditions where the water vapor pressure is high (Torrance and Darvell, 1990).

**Manufacture**

*Introduction*

From the transformation temperatures given in the previous section, it can be seen that calcium sulfate hemihydrate would be produced by heating gypsum to temperatures in the range of 45°C to 90°C. However, at these temperatures the reaction is slow; even at 90°C, substantially complete conversion takes about 12 hours (Khalil et al, 1971). Therefore, in commercial processes temperatures higher than this are used for shorter times. The stable phase at these higher temperatures is hexagonal calcium sulfate, so the initial product of calcination is partly or very largely this anhydrous form.
However, on cooling to temperatures below 80°C and exposure to the atmosphere, the hexagonal calcium sulfate rehydrates to form the hemihydrate.

*Plaster of Paris*

This is the traditional hemihydrate plaster produced by the dry calcination of ground gypsum in open containers (pans, kettles, or rotary kilns) at temperatures in the range 120°C to 180°C. In the absence of liquid water, there is no opportunity for reorganization of crystal morphology, so although the crystal structure of the final product after exposure to air is that of calcium sulfate hemihydrate, the powder particles retain the rough irregular shape of the original ground gypsum. Loss of water under dry conditions leaves parallel channels in these particles (Fig 4-2), so hemihydrate plasters produced by dry calcination are powders with a low apparent density, a high relative surface area, and a poor packing ability.

![Fig 4-2 Particles of plaster. (Original magnification × 400.)](image)

*Medium- and high-strength plasters (stones)*

Plasters giving a stronger set mass are manufactured by wet calcination. Here sufficient liquid water is present to allow through-solution conversion, so recrystallization produces dense prismatic crystals of hexagonal calcium sulfate. These rehydrate to the hemihydrate on cooling in air, but this secondary conversion cannot be accompanied by recrystallization. The final powder particles are pseudomorphic, having the monoclinic crystal structure of hemihydrate, but retaining the hexagonal crystal habit of the anhydrous calcium sulfate precursor (Fig 4-3).
Hemihydrate powders produced by wet calcination therefore have a higher apparent density and a smaller relative surface area than those resulting from dry calcination. A controlled amount of grinding rounds off the crystals and produces a proportion of fines, both factors improving the packing ability of the powder and further increasing the apparent density. Particles of a typical dental stone are shown in Fig 4-4.
Medium-strength plasters

These are typified by Hydrocal (United States Gypsum), which is the basis of many ordinary dental stones. It is produced by autoclaving lump gypsum in superheated steam, at a pressure of 117 kPa, giving a temperature of 123°C, for 5 to 7 hours. After drying at 100°C or higher and grinding, a hemihydrate powder is produced that gives a set mass considerably stronger than that produced by dry-calcined materials.

High-strength plasters

Modifications to the wet process yield hemihydrate powders giving even stronger set products. These are typified by Densite (Georgia-Pacific) and Crystacal (British Gypsum). Here wet calcination takes place in the presence of crystal habit modifiers, producing crystals that are shorter and thicker than those resulting from autoclaving in steam. After controlled grinding, these powders have even higher apparent densities and yield even stronger set masses than Hydrocal-type stones: they form the basis of most dental die stones. Densite is produced by boiling lump gypsum in a 30% calcium chloride solution, and Crystacal is produced by autoclaving finely ground gypsum in the presence of small amounts (< 1%) of sodium succinate.

Apparent density

The apparent density of a powder is the reciprocal of its bulkiness and so gives a measure of its packing ability. The low apparent density of hemihydrate powders produced by dry calcination is caused in part by the rough irregular shapes of the individual particles (see Fig 4-2). A more important factor, however, is a high adhesion of particles to their neighbors, caused by a high surface free energy resulting from crystal imperfections and adsorption of gases during calcination (Gregg, 1965). This adhesiveness of particles makes it likely that they will stick together at first contact, establishing bridges in the powder and creating a structure with many voids.
and a low apparent density. In contrast, powder particles produced by wet calcination are smooth and dense (see Fig 4-4). They show less crystallographic strain and therefore have a lower surface free energy. As a result, they have a better packing ability and a higher apparent density.

The Setting Process

The setting reaction

When the hemihydrate powder is mixed with water in the correct proportions, it forms a thick slurry. The hemihydrate is sparingly soluble in water (6.5 g/L at 20°C), so only a small amount can dissolve. Initially, therefore, the mix is a two-phase suspension of hemihydrate particles in a saturated aqueous solution. The stable hydrate at temperatures below 40°C is the dihydrate (gypsum), which is even less soluble (2.4 g/L at 20°C) than the hemihydrate. The aqueous phase is therefore supersaturated with respect to the dihydrate, which crystallizes out at suitable nucleation centers in the suspension. These gypsum crystals normally are acicular in habit and often radiate out from nucleation centers in the form of spherulitic aggregates.

These nucleation centers may be impurities (eg, residual gypsum particles), particles of gypsum added as seeds to accelerate setting, or strained areas on undissolved hemihydrate particles. The consequent depletion of calcium and sulfate ions in the aqueous phase allows more hemihydrate to go into solution and, in turn, to precipitate out as gypsum. The setting process is therefore one of heterogeneously nucleated recrystallization, characterized by a continuous solution of hemihydrate, diffusion of calcium and sulfate ions to nucleation centers, and the precipitation of microscopic gypsum crystals. The setting reaction is the reverse of the first stage of dehydration and so is exothermic. It is represented by the following equation:

\[ 2\text{CaSO}_4 \cdot \frac{1}{2}\text{H}_2\text{O} + 3\text{H}_2\text{O} \rightarrow 2\text{CaSO}_4 \cdot 2\text{H}_2\text{O} \]

Water requirement

The differing water requirements of plaster, stone, and high-strength stone are mainly the result of differences in the apparent density of the powder. The factors that promote adhesiveness of the particles in the dry powder persist when they are suspended in water (Ridge, 1961). For this reason dry-calcined plaster, with its low apparent density, produces a flocculated suspension and needs a relatively high proportion of mixing water to give a mix of workable viscosity. Water/powder (W/P) ratios of 0.5 to 0.6 are usual. Hemihydrate powders produced by wet calcination, because of their higher apparent densities, require less mixing water; typical W/P ratios are 0.30 to 0.33 for ordinary dental stones and 0.18 to 0.23 for high-strength stones.

In setting, 100 g of hemihydrate combines with 18.6 g of water. Therefore, at the completion of the reaction in normal mixes there is always some excess unreacted water (as a saturated solution of calcium sulfate) remaining in the set mass. This residual water weakens the cast. It can be removed by low-temperature drying but leaves microscopic porosity that weakens the dry cast. Both wet and dry strengths of
the set material depend on the relative amount of unreacted water remaining after setting, and so on the W/P ratio of the original mix. The relative amount of residual water is least in high-strength stone, which therefore gives the strongest set mass.

**Stages in setting**

The setting process is continuous from the beginning of mixing until the setting reaction is complete, by which time the material has reached its full wet strength. However, important physical changes can be recognized during this process. Initially there is a continuous aqueous phase present, and the mix is a viscous liquid, exhibiting **pseudoplasticity** so that it flows readily under vibration; in this stage the mix has a glossy surface giving **specular** reflections. As the setting reaction proceeds, gypsum crystals continue to grow at the expense of the aqueous phase, and the viscosity of the mix increases. When the clumps of growing gypsum crystals interact, the mix becomes plastic; it will not flow under vibration but can readily be molded. At this time the glossy surface disappears as the aqueous phase is drawn into pores formed when the growing gypsum crystals thrust apart. Continued crystal growth converts the plastic mass into a rigid solid, weak and friable at first but gaining strength as the relative amount of solid phase increases.

These four stages may be designated (1) fluid, (2) plastic, (3) friable, and (4) carvable.

**Volume changes during setting**

The setting reaction causes a decrease in the true volume of the reactants, and under suitable conditions this contraction can be observed early in the setting process, when the mix is still fluid. However, once the mix begins to attain rigidity (marked by the loss of surface gloss) an **isotropic** expansion is observed, resulting from growth pressure of the gypsum crystals that are forming. There is therefore a decrease in apparent density of the mix during the latter stages of setting, accompanied by the formation of microscopic voids separating individual crystals in the aggregate.

The initial contraction is unlikely to affect the important dimensions of a gypsum cast, because in the still-fluid mix it will occur mainly in a vertical direction. Gravity will keep the mix adapted to the anatomical portion of an impression. The expansion that is observed after the mix attains rigidity takes place in all directions and will affect the dimensions of the cast. The point at which the initial contraction ceases is used as zero in laboratory measurements of effective setting expansion.

**Rate of the setting reaction**

Within wide limits, the rate of hydration during setting is independent of the W/P ratio (Lautenschlager et al, 1969). However, the rate at which the associated physical changes described earlier occur is highly dependent on the W/P ratio of the mix, because these changes result from interaction of clumps of gypsum crystals growing from nucleation centers in the slurry. Thick mixes (low W/P ratios) harden more quickly because available nucleation centers are concentrated in a smaller volume; interaction of the growing solid phase occurs earlier and is more effective in promoting expansion.
The effect of additives

Many salts and colloids are known to alter the setting characteristics of hemihydrate plasters by their effect on the rate of the setting reaction. They have been used in the formulation of dental plasters and stones for many years, mainly on an empirical basis, because their modes of action are not always completely understood.

Finely powdered gypsum is an efficient accelerator, which acts by providing seeds for heterogeneous nucleation. In low concentrations soluble sulfates and chlorides are accelerators, apparently acting by increasing the rate of solution of the hemihydrate. However, salts of relatively low solubility, such as sodium chloride and sodium sulfate, act as retarders in higher concentrations, because as setting proceeds the amount of free water in the mix decreases and the concentration of the additive increases. When its limit of solubility is exceeded, the salt precipitates on nuclei of crystallization, thus poisoning them. Acetates, borates, citrates, and tartrates are retarders, which may act by nuclei poisoning, by reducing the rate of solution of hemihydrate, or by inhibiting the growth of dihydrate crystals. Reaction of some additives with hemihydrate can occur; soluble tartrates and citrates precipitate calcium tartrate and citrate, respectively. Colloids are effective retarders, presumably acting by nuclei poisoning.

Many accelerators and retarders reduce the setting expansion, sometimes by changing the crystal habit of growing gypsum crystals from acicular to a more compact form and inhibiting spherulite formation (Koslowski and Ludwig, 1984); both factors reduce the effect of the crystals' growth pressure. This is accompanied by a reduction in the strength of the set material.

The Microstructure of Cast Gypsum

Introduction

The set material consists of a tangled aggregate of monoclinic gypsum crystals, usually acicular in shape, with lengths in the range of 5 to 20 μm. The aggregate exhibits two distinct types of inherent porosity on a microscopic scale:

1. Microporosity caused by the presence of residual unreacted water. These voids are roughly spherical and occur between clumps of gypsum crystals.

2. Microporosity resulting from growth of gypsum crystals. These voids are associated with setting expansion and are smaller than the first type; they appear as angular spaces between individual crystals in the aggregate.

The effect of W/P ratio

The relative amounts of both types of porosity are affected by the W/P ratio of the mix, but in opposite ways:

1. A low W/P ratio (thick mix) leaves less residual water in the set mass and so decreases the amount of the first type of porosity.
2. A low W/P ratio increases the effect of crystal growth during setting, because available nucleation centers are concentrated in a smaller total volume of mix; interaction of growing gypsum crystals occurs earlier and is more effective so the amount of the second type of porosity is increased.

3. At any W/P ratio, the total proportion of inherent porosity in the set mass is the sum of these two types. The effect of the first type predominates, so for any given plaster or stone there is always a decrease in the total inherent porosity of the set mass (ie, an increase in apparent density) as the W/P ratio of the mix is reduced. Inherent porosity represents about 40% of the total cast volume at a W/P ratio of 0.50 and about 20% at a W/P ratio of 0.25 (Lautenschlager and Corbin, 1969).

Typical microstructures of casts made from plaster mixed in a W/P ratio of 0.50 and high-strength stone mixed in a W/P ratio of 0.25 are shown in Fig 4-5. The second type of microporosity described here (resulting from crystal growth) can be seen in both micrographs, while the first type (resulting from residual water) shows clearly in the micrograph of set plaster, where the voids occur between spherulitic clumps of gypsum crystals. The higher apparent density of the set high-strength stone is obvious.

Fig 4-5 Fracture surface of cast gypsum. (Scanning electron micrographs, original magnification × 3,000.) (A) Plaster, W/P = 0.50; (B) high-strength stone, W/P = 0.25. (From Bever, 1986. Reprinted with permission.)
**Properties**

*Rate of setting*

**Manipulation time**

Recognition of the physical changes occurring in the mix during setting is important in the manipulation of plaster and stone.

1. When casting (e.g., pouring casts or dies), manipulation must be completed before the mix loses fluidity. The change is marked by the disappearance of the glossy surface from the mix.

2. When molding (e.g., taking impressions or jaw registrations, articulating casts, flasking wax pattern dentures), manipulation must be completed before the mix loses plasticity and enters the friable stage. There is no recognized objective method of measuring this time.

**Setting time**

An arbitrary setting time (initial set) can be determined by using suitable penetrometers (e.g., Gillmore or Vicat initial needles; both give approximately the same initial setting time). Measured in this way, the initial set is a guide to the time when the rigid material is strong enough to handle and, in particular, when it can be carved or trimmed to the final shape.

If a dental manufacturer specifies a setting time, it will be a Gillmore or Vicat initial set. Both the setting reaction and the increase in strength continue for some time after this initial set. Gillmore and Vicat final needles may be used to establish a final setting time, but this is not usual in dental technology.

**Control of rate of setting**

1. *The use of additives.* In formulating dental products, manufacturers adjust the rate of setting of raw hemihydrates by adding accelerators and retarders, often as a balanced mixture. Typical accelerators are potassium sulfate and potassium sodium tartrate ("Rochelle salts"). Typical retarders are sodium citrate and sodium tetraborate decahydrate (borax). The action of accelerators and retarders has already been discussed.

2. *W/P ratio.* It has already been pointed out that changing the W/P ratio has a marked effect on the rate at which the physical changes associated with setting of the mix occur. These changes take place more rapidly as the W/P ratio is reduced. Manipulation and setting times are thus directly proportional to W/P ratio.

**Setting expansion**

Typical values for setting expansion for the five types of dental gypsum plaster are given in Table 4-2. These are the results of laboratory measurements on five widely used materials, setting in air.
Table 4-2 Linear setting expansions of typical dental gypsum products (setting in air)

<table>
<thead>
<tr>
<th>Type</th>
<th>W/P ratio</th>
<th>Setting expansion (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Impression plaster</td>
<td>0.60</td>
<td>0.13</td>
</tr>
<tr>
<td>Plaster</td>
<td>0.50</td>
<td>0.30</td>
</tr>
<tr>
<td>Stone</td>
<td>0.30</td>
<td>0.15</td>
</tr>
<tr>
<td>Stone, high-strength, low-expansion</td>
<td>0.20</td>
<td>0.10</td>
</tr>
<tr>
<td>Stone, high-strength, high-expansion</td>
<td>0.19</td>
<td>0.26</td>
</tr>
</tbody>
</table>

The observed expansion that occurs when dental plaster or stone sets is a volumetric one. In dental testing a value is determined for linear setting expansion, and the assumption is made that the expansion is isotropic. This assumption is not always justified; if restraint is imposed in some directions but not others (eg, by a rigid impression), setting expansion can be far from isotropic.

### Effect of immersion

Gypsum products exposed to additional water while setting (eg, by immersion) show a greater expansion than when setting in air, a phenomenon commonly (but inaccurately) called hygroscopic expansion. When expansion begins, externally available water is drawn into pores forming in the setting mass, and this maintains a continuous aqueous phase in which crystal growth takes place freely. Under dry conditions this additional water is not available, and as expansion occurs the aqueous phase in the mix is reduced to a film over the growing gypsum crystals. Surface tension forces in this liquid film restrain further crystal growth, thereby reducing the observed setting expansion of the mix (Mahler and Ady, 1960). Thus the so-called hygroscopic expansion is simply an enhanced setting expansion that occurs in the presence of additional water.

### Control of setting expansion

1. **Use of additives.** A low setting expansion is desirable in applications where dimensional accuracy is important (eg, in impression taking and pouring working casts and dies). Many accelerators and retarders of setting also reduce setting expansion. Manufacturers can reduce setting expansion and at the same time control setting time by adding a balanced blend of accelerator and retarder to the raw hemihydrate. Typical combinations are potassium sulfate-borax and potassium sodium tartrate- sodium citrate.

These additives also reduce the strength of the set material. This is not a disadvantage in impression plasters, but in stones, strength as well as dimensional accuracy is important; formulation of the latter materials therefore involves striking a compromise between a desirable reduction in setting expansion and an undesirable reduction in strength.

Normally die stones in particular are formulated to have a low setting expansion. As their name indicates, they are used to make cast gypsum dies on which restorations such as inlays, crowns, and fixed partial dentures are constructed by the indirect method, and they are required to reproduce the prepared teeth with high dimensional...
accuracy. Such materials constitute the ISO Type 4 high-strength, low-expansion stones, which have a setting expansion less than 0.15%. These would be used where the investment expansion is sufficient to compensate completely for the casting shrinkage of the alloy. There has now been introduced another category of die stone: ISO Type 5, high-strength, high-expansion, with a setting expansion in the range of 0.16% to 0.30%. The higher expansion results from a lower concentration of modifying additives, so these materials also generally achieve higher strengths when set. They are intended for use in casting techniques when investment expansion alone may not be high enough to compensate fully for alloy shrinkage. Here the setting expansion of the die stone is intended to contribute to the total mold expansion by producing an oversized pattern (Fusayama, 1964), but it is important that the impression be soft and flexible enough for isotropic setting expansion to occur. In an impression made from a stiff elastomeric material in a closely fitting rigid tray, setting expansion will be likely to occur preferentially in a vertical direction, producing a distorted die.

2. W/P ratio. For any given gypsum product, reducing the relative amount of aqueous phase in the mix allows more effective interaction of growing gypsum crystals during setting, thus increasing the setting expansion. Setting expansion is therefore inversely proportional to the W/P ratio. Because of their lower water requirement, the raw hemihydrates used to produce stones and die stones have a higher inherent setting expansion in normal mixes than does plaster. This effect is masked, however, by the additives used in their formulation (Fig 4-6).

![Fig 4-6 Linear setting expansion in air and in water of various types of gypsum products. (A) Plaster (W/P = 0.50); (B) unmodified stone (Hydrocal B, W/P = 0.32); (C) a commercial stone based on Hydrocal (W/P = 0.32); (D) unmodified high-strength stone (Densite K5, W/P = 0.24); (E) a commercial high-strength stone based on Densite (W/P = 0.24).]
**Strength**

Cast gypsum is a brittle material and so is weaker in tension than in compression. Materials with the highest compressive strengths are the most brittle, and their tensile strengths are proportionally lower. For set plaster, the tensile strength is about 20% of the compressive strength; for set high-strength stone, it is about 10%. Since fracture of cast gypsum typically occurs in tension, tensile strength is a better guide to fracture resistance. However, compressive strength gives a better indication of surface hardness. Values for tensile and compressive strengths of five typical dental gypsum plasters are shown in Table 4-3.

<table>
<thead>
<tr>
<th>Type</th>
<th>W/P ratio</th>
<th>Tensile strength (MPa)</th>
<th>Compressive strength (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Wet</td>
<td>Dry</td>
</tr>
<tr>
<td>Impression plaster</td>
<td>0.60</td>
<td>1.3</td>
<td>—</td>
</tr>
<tr>
<td>Plaster</td>
<td>0.50</td>
<td>2.3</td>
<td>4.1</td>
</tr>
<tr>
<td>Stone</td>
<td>0.30</td>
<td>3.5</td>
<td>7.6</td>
</tr>
<tr>
<td>Stone, high-strength,</td>
<td>0.20</td>
<td>6.1</td>
<td>10.6</td>
</tr>
<tr>
<td>low-expansion</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Stone, high-strength,</td>
<td>0.19</td>
<td>6.2</td>
<td>12.6</td>
</tr>
<tr>
<td>high-expansion</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*The materials tested were the same as those in Table 4-2.

1. **The effect of W/P ratio.** In general, strength properties are inversely related to W/P ratio and so to the total amount of inherent porosity (Fig 4-7). Therefore when maximum strength is required, a given material should be mixed in as low a W/P ratio as practicable. The limiting factor is the viscosity of the mix, because it increases with decreasing W/P ratio and can become so high that the ability to pour sound casts is prejudiced.

In high-strength stones mixed even in normal W/P ratios, there is little water present in excess of the theoretical W/P ratio (0.186) required for complete conversion of hemihydrate to dihydrate. Since the setting reaction depends on diffusion of ions, it may not proceed to completion in standard mixes; in fact, Donnison et al (1960) have shown that in normal mixes the hydration may be up to 3% short of completion and that the proportion of residual hemihydrate increases as the W/P ratio decreases. Residual hemihydrate particles are often seen in micrographs of set high-strength stone. The content of unreacted hemihydrate becomes much greater in very thick mixes, but as Fig 4-7 shows, this does not adversely affect the strength until the W/P ratio is well below that theoretically needed for complete hydration. Such thick mixes would be impossible to manipulate successfully in most dental applications.

With any plaster or stone, using a low W/P ratio to obtain maximum strength properties also gives an increased setting expansion, which must be accepted. But in
applications where dimensional accuracy is more important than strength (eg, impressions), higher W/P ratios can be used.

![Graph](image)

**Fig 4-7** The effect of W/P ratio on the wet tensile strength of a high-strength stone. (From Selby, 1979.)

2. The effect of drying. Removal of all uncombined water from cast gypsum by low-temperature drying approximately doubles strength properties (see Table 4-3), but there is no strength increase until the last 2% of free water is removed. This strength increase on drying is reversible; soaking a dry cast in water reduces its strength to the original level.

Gypsum is stable only below about 40°C. Drying at higher temperatures must be carefully controlled; loss of water of crystallization occurs rapidly at 100°C or higher and causes shrinkage and a reduction in strength.

**Solubility**

Because gypsum is sparingly soluble in water, long-term immersion is contraindicated. If dried gypsum casts have to be soaked, it is better to use a saturated solution of calcium sulfate.

**Disinfection of Gypsum Casts**

To prevent cross-infection, the practice of disinfecting impressions is becoming increasingly common. However, prolonged immersion in disinfectant solutions can cause unacceptably large dimensional changes in hydrocolloid (Bergman et al, 1985; Olsson et al, 1987) and polyether (Johnson et al, 1988) impressions. Moreover, during subsequent clinical procedures, casts or dies can become reinfected with pathogenic organisms, which can then be transferred to technical staff. As an alternative, the
addition of disinfectants to the mixing water when casts are poured has been investigated; 5% phenol (McGill et al, 1988) and 2% glutaraldehyde (Ivanovski et al, 1995) have proved to be effective and did not adversely affect the properties of the set material. However, both are known tissue irritants. Dental stones that contain a disinfectant are available (Donovan and Chee, 1989; Schutt, 1989). Alternatively, casts and dies may be treated by immersion in a disinfecting solution after each clinical stage.

Autoclave sterilization of casts has been suggested (Whyte and Brockhurst, 1996). Some loss of strength and surface hardness and an increase in dimensions occur, but it is claimed that under carefully controlled conditions the casts retain adequate properties for ordinary laboratory use.

**Gypsum-Bonded Investments**

**Introduction**

Gypsum-bonded investments are the mold materials most commonly employed in the casting of dental gold alloys with liquidus temperatures no higher than 1,080°C, which are normally used for gold inlays, crowns, and fixed and removable partial dentures. Because of their tendency to decompose at high temperatures, these investments are not suitable for casting high-melting gold alloys, palladium alloys (used for copings in alloy-ceramic restorations), or most base-metal alloys, such as nickel-chromium and cobalt-chromium. Gypsum-bonded investments are classified by the International Standards Organization (1990) as:

Type 1: Thermal expansion type, for casting inlays and crowns

Type 2: Hygroscopic expansion type, for casting inlays and crowns

Type 3: For casting complete and partial denture bases

This classification is illustrated in Fig 4-8, and typical materials are listed in Table 4-4. Some materials, classified by their manufacturers as "universal," are claimed to be suitable for casting all gold-alloy restorations.

**Fig 4-8** Classification of gypsum-bonded investments.
Table 4-4 Typical gypsum-bonded investments

<table>
<thead>
<tr>
<th>Type</th>
<th>Example</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inlay investment, thermal expansion</td>
<td>Cristobalite Inlay (Kerr/Sybron)</td>
</tr>
<tr>
<td>(ISO Type 1)</td>
<td></td>
</tr>
<tr>
<td>Inlay investment, thermal expansion,</td>
<td>Cristoquick (GC)</td>
</tr>
<tr>
<td>rapid heat (ISO Type 1)</td>
<td></td>
</tr>
<tr>
<td>Inlay investment, hygroscopic</td>
<td>Beauty-Cast (Whip Mix)</td>
</tr>
<tr>
<td>expansion (ISO Type 2)</td>
<td></td>
</tr>
<tr>
<td>Denture investment (ISO Type 3)</td>
<td>R&amp;R Gray (Dentsply/Ransom and</td>
</tr>
<tr>
<td></td>
<td>Randolph)</td>
</tr>
</tbody>
</table>

Composition

All gypsum-bonded investment powders consist basically of a refractory filler and a binder. There may also be small amounts (less than 5%) of important modifying agents present.

Refractory

The refractory component is either cristobalite or quartz (or occasionally a mixture of the two), usually present in the range of 55% to 75%. Both of these components are polymorphs of silica (SiO₂).

Binder

In the investment powder, the binder is calcium sulfate hemihydrate, either plaster or stone. When the investment sets, the silica is unaffected; the hemihydrate binder combines with water to form dihydrate (gypsum) as in the setting of other gypsum products. The setting reaction is therefore the same as that shown in the equation given previously. The set investment consists of fine particles of silica embedded in an interlocking aggregate of smaller acicular gypsum crystals (Fig 4-9). The continuous porosity characteristic of cast gypsum provides the necessary mold venting to assist in the production of sound castings.
Modifying agents

Small amounts of modifying agents are added to many commercial gypsum-bonded investments. These may be accelerators or retarders to control the rate of setting, reducing agents such as powdered graphite or copper to protect embedded gold alloy components in "casting-on" techniques, or additives to increase the thermal expansion of the investment. Typical additives of the last type are boric acid and soluble halide salts, particularly those of alkali or alkaline earth metals.

**Effect of composition on setting and thermal behavior**

Refractory

Although silica is referred to as the refractory component, cast gypsum itself is sufficiently heat-resistant to be used as a mold material, because it can be heated to temperatures as high as 1,000°C without decomposing (Gutt and Smith, 1967). In fact, adding silica reduces its heat resistance, because in such mixtures at temperatures above 900°C, the following reaction occurs:

\[ \text{CaSO}_4 + \text{SiO}_2 \rightarrow \text{CaSiO}_3 + \text{SO}_3 \]

At high temperatures, the sulfur trioxide liberated by this reaction causes rapid corrosion of the casting.

Crystalline silica is used in dental investments not to improve their heat resistance but to control dimensional changes on heating. When heated, cast gypsum shows a marked contraction, and because both quartz and cristobalite have high expansions when heated, they offset the shrinkage of the binder and can provide a positive
thermal expansion if this is needed.

Silica exists in 22 different condensed phases. Five of these are amorphous, and 17 are crystalline; the latter are the polymorphs of silica. Of this group, only one phase, low-temperature quartz, is thermodynamically stable at normal temperature and pressure. Two more, tridymite S-1 and low-temperature cristobalite, exist under normal atmospheric conditions as metastable (but actually long-lived) phases.

Only two polymorphs of silica are of importance in dental investments: quartz and cristobalite. Quartz is a common mineral; cristobalite occurs naturally as a rare mineral but is normally manufactured by prolonged heating of quartz at high temperatures to induce the appropriate slow inversion. Both quartz and cristobalite exist in low-temperature (α) and high-temperature (β) phases, and in both materials the change between low- and high-temperature phases is rapid and readily reversible on cooling. This change is known as the high-low inversion.

In both quartz and cristobalite the high-temperature phase is less dense than the low, so the α→β inversion is accompanied by an isothermal expansion. This is added to the normal thermal expansion to give a large overall expansion at high temperatures.

As in all crystalline materials, single crystals of quartz and cristobalite show anisotropy of physical properties, including their coefficients of thermal expansion. However, in dental investments they are used as a fine powder with random orientation of individual particles, so in the aggregate they behave isotropically, and properties such as the coefficient of thermal expansion have a uniform average value.

Typical linear thermal expansion curves for powdered quartz and cristobalite are shown in Fig 4-10. Specimens were made by densely compacting the powders into aggregates with a minimum amount of binder in the interstices, so the thermal expansion of the specimens was essentially that of the silica component in dental investments.

The temperature of the α→β inversion of cristobalite varies according to the source of the sample and its previous thermal history; in the manufactured cristobalite used in dental investments it is about 250°C (Fig 4-10, curve A). The inversion gives an isothermal expansion of about 1.3%, and the total expansion at 700°C is about 2.2%. Dental investments designed to have a high thermal expansion usually contain cristobalite as the refractory.
Fig 4-10 The linear thermal expansion of cristobalite (curve A) and quartz (curve B). The specimens were aggregates of fine powders, united with a minimum amount of binder; their thermal behavior was thus similar to that of the refractory component of dental casting investments. Cristobalite undergoes inversion at about 250°C, quartz at 573°C.

Quartz (Fig 4-10, curve B) undergoes its $\alpha \rightarrow \beta$ inversion at 573°C, producing an isothermal expansion of about 0.7%. The high-temperature form has a negative coefficient of thermal expansion, and the overall total expansion at 700°C is about 1.6%.

Binder

When the set investment is heated, the cast gypsum binder shows a marked contraction, which occurs in several stages. In the early stages of heating, a contraction accompanies loss of water of crystallization, as the dihydrate reverts to hemihydrate and then to hexagonal calcium sulfate. A large contraction then accompanies its $\gamma \rightarrow \beta$ transformation, which occurs in the approximate temperature range of 300°C to 400°C. A further large contraction begins at about 650°C, which is probably the result of densification by sintering. The total linear contraction of cast gypsum prepared from plaster (W/P = 0.50) and heated to 700°C can be as high as 3%; the corresponding shrinkage for cast gypsum prepared from a high-strength stone (W/P = 0.25) is about 1%. The latter is usually preferred as a binder in gypsum-bonded investments, not only because of its lower shrinkage on heating but because its superior strength when set allows a higher concentration of silica to be used in the investment, thereby reducing still further the effect of binder contraction.

Effects of varying composition

Within practical limits, increasing the proportion of silica in the investment powder
increases the manipulation time (given by the time of loss of fluidity) and the initial setting time (Fig 4-11),

**Fig 4-11** The effect of silica content on the manipulation time (curve A) and setting time (curve B) of experimental investments. Manipulation time was determined by the time of loss of gloss, which indicates loss of fluidity; setting time by the Vicat initial set. The investments were mixtures of cristobalite and a high-strength stone, and all specimens were mixed in a W/P ratio of 0.40.

the setting expansion, both in air and in water, and the thermal expansion (Fig 4-12)
Fig 4-12 The effect of silica content on the setting expansion in air (curve A), the setting expansion in water (curve B), and the thermal expansion (curve C) of experimental investments (compositions and W/P ratio as in Fig 4-11). Setting expansions were recorded 2 hours after mixing, and thermal expansions after heating to 700°C.

and reduces the compressive strength (Fig 4-13). The rate of the setting reaction itself is unchanged; the increases in observed manipulation and setting times occur because the particles of refractory filler interfere with the interlocking of growing gypsum crystals, making this less effective in developing a solid structure. The compressive strength of the set investment is reduced for the same reason. Setting expansion is increased when interlocking of growing gypsum crystals is inhibited by the refractory particles, because more of the crystal growth is directed outward. Thermal expansion is increased because, at any temperature, it is given by summing the binder contraction and the refractory expansion; increasing the proportion of the expanding component increases the observed expansion.
The effect of silica content on the compressive strength of experimental investments (compositions and W/P ratio as in Fig 4-11). Tests were made on wet specimens, 2 hours after mixing.

Effects of modifying agents

Important modifying agents are those added to some commercial investments to increase their thermal expansion. These include boric acid and soluble halide salts of alkali or alkaline earth metals. All of them act mainly by reducing the two large contractions of the gypsum binder that occur on heating to temperatures above 300°C.

Boric acid, when heated above 150°C, forms a viscous liquid with a composition intermediate between metaboric acid and boron oxide; this viscous liquid impedes the evaporation of the last traces of water, delaying the $\gamma \rightarrow \beta$ transformation of calcium sulfate (Mori, 1986). The presence of the viscous liquid phase also reduces the high-temperature contraction that results from sintering, because it stabilizes the original contacts formed between the gypsum crystals and silica during setting (Mori, 1982).

The presence of halide anions greatly reduces the first major shrinkage and completely eliminates the second. The reason for this is not yet understood. If alkali metal or alkaline earth cations are also present (eg, as halide salts), the effect of the halide anion is nullified at temperatures above 650°C and a rapid contraction occurs; this is probably the result of accelerated sintering. This large high-temperature shrinkage of the binder is not observed in gypsum-bonded investments containing alkali metal or alkaline earth halides, because at a concentration of 50% or more of silica the silica particles in the set investment form a continuous "skeleton" that resists overall shrinkage.

The presence of modifiers added to increase the thermal expansion also affects the strength changes of the investment that occur on heating, again because of their effects on the calcium sulfate binder.

On heating, gypsum-bonded investments without these additives show a rapid
increase in compressive strength of about 100% in the range of 100°C to 175°C; this is the result of drying and is analogous to that occurring in cast gypsum. Between 175°C and 225°C there is an equally rapid decrease in compressive strength, attributable to the dehydration reaction dihydrate $\rightarrow$ hemihydrate, bringing the strength back to about that of the original wet specimens. Relatively minor strength fluctuations occur during subsequent heating to higher temperatures, attributable to (1) further phase changes in the binder, (2) the $\alpha\rightarrow\beta$ inversion in the refractory, and (3) sintering of the binder (Ohno et al, 1982). Investments of this type, heated to temperatures in the range of 670°C to 700°C, show compressive strength changes ranging from +10% to -40%, compared with the wet strength at ambient temperatures.

Investments containing boric acid, when heated to the same temperatures, show increases in compressive strength ranging from +40% to +50%, probably the result of the effect of the viscous liquid phase stabilizing contacts between calcium sulfate and silica particles. Investments containing halides of alkali metals and alkaline earths (e.g., sodium, barium, and strontium chlorides) show a marked strength decrease on heating to 700°C, ranging from -50% to -85%. This is probably the result of the increased sintering contraction of the binder at temperatures over 650°C. The silica skeleton resists shrinkage of the investment as a whole, so the high sintering contraction occurs independently in the binder, reducing the strength of its bond with the silica particles.

Properties

Particle size of the powder

The particle size affects the smoothness of the mold cavity surface (and thus of the casting) and also the inherent porosity of the mold (and thus the venting of the mold cavity). Only the particle size of the refractory filler is of practical importance. It is the major constituent and remains unchanged in the set investment. The gypsum crystals formed during setting of the binder are much smaller than the silica particles (see Fig 4-9).

Excessive surface roughness of the casting can interfere with its fit; a refractory with a fine particle size ensures a smooth mold surface and a smooth casting. In gypsum-bonded investments, venting of the mold cavity is normally provided by the continuous porosity inherent in the set material. This is at a maximum when packing of the silica particles is least dense, which in turn is achieved by ensuring that the particle size is uniform.

Therefore, the refractory powder used in making the investment should have a uniform, fine particle size. A particle size of no more than 75 um is usual.

Rate of setting

The same physical changes that occur during the setting of gypsum plasters can also be recognized when gypsum-bonded investments set.

Manipulation time  Investing the wax pattern or pouring the investment cast must be completed while the mix is still fluid. As with gypsum plasters, loss of fluidity is
indicated by the disappearance of the glossy surface from the mix.

**Setting time**  This is usually given as a Gillmore or Vicat initial set. It indicates when the investment is strong enough for the sprue base and sprue former to be removed. Preferably mold heating should be delayed until setting expansion is complete (usually between 1 and 2 hours from the start of mixing).

**Expansion**

Inlay investments currently available have total expansions, measured under laboratory conditions, in the range of 1.5% to 2.5%. Denture investments gain increased strength by having a higher content of hemihydrate binder, at the expense of the refractory component, so their total expansions are less; the lower figure for the expansion range is about 1.3%. It is, however, by no means certain that such figures show the expansion of the mold cavity that can be expected under practical conditions.

Modern methods of casting small restorations can be classified into two groups:

1. In **hygroscopic expansion techniques**, all or most of the mold expansion is gained when the investment sets; the setting expansion is greatly increased by exposure to additional water. Thermal expansion of the investment is low.

2. In **thermal expansion techniques**, both setting and thermal expansion contribute importantly to mold expansion.

**Setting expansion**

1. **Setting expansion in air.** Linear setting expansions of most inlay investments, measured under dry conditions, are in the range of 0.1% to 0.6%.

2. **Setting expansion in water (hygroscopic setting expansion).** Inlay investments show a much greater setting expansion if exposed to additional water during setting (e.g., by immersion in a water bath). The phenomenon is the same as that described for gypsum plasters, but the effect on setting expansion is much greater because of the presence of the refractory particles (see Fig 4-12). If no constraint is imposed, the linear setting expansion of inlay investments can be as high as 4% (Earnshaw, 1969a). But even a small restraining force causes a large reduction, and when measured under usual laboratory conditions, the hygroscopic setting expansion of most inlay investments is in the range of 0.3% to 2.0%. Investments specifically designed for use in hygroscopic expansion techniques have setting expansions, when immersed in water, of at least 1.3%.

3. **The effect of the casting ring liner.** In most casting techniques the investment mold sets and is heated in a casting ring made of heat-resistant alloy, which has a much lower thermal expansion than the investment mold. The need for a soft ring liner, to eliminate or at least reduce restraint to investment expansion by the ring, was first recognized by Souder (Hollenback, 1962), who advocated the use of asbestos tape to provide the necessary cushioning. Since asbestos readily absorbs water, the liner was prewetted to prevent its absorbing water from the unset investment mix.
The technique of lining a casting ring with wet asbestos was first described in 1930 by Taylor and coworkers, and from then until recent times, the use of a wet asbestos ring liner has been a standard procedure. This makes additional water available to the setting investment and causes an increased setting expansion. Even when the investment mold sets in air, as in thermal expansion techniques, some hygroscopic setting expansion occurs. Investments used in thermal expansion techniques have a relatively high silica content, so the increase in setting expansion produced by exposure to water is high (see Fig 4-12). Because this high setting expansion is uncontrollable and likely to be anisotropic, techniques have been developed in which a dry, waterproofed asbestos ring liner is used (Fusayama, 1959, 1964) or the use of the rigid casting ring with its liner is avoided (Finger and Jorgensen, 1980).

Attention has been drawn to the danger that asbestos fibers in casting-ring liners could cause asbestosis or mesothelioma (Priest and Horner, 1980). It is claimed that the acceptable threshold limit for asbestos fibers in air can be considerably exceeded when castings are removed from asbestos-lined casting rings (Yli-Urpo et al, 1982). For this reason, and because of the increasing unavailability of asbestos products, asbestos as a casting-ring liner has been almost completely replaced by alternative materials. There are two types: (1) cellulose, which readily absorbs water and, like asbestos, must be prewetted, and (2) ceramic materials which at atmospheric pressure will not absorb water and are normally used dry. The ceramic materials are made from fibers of an aluminosilicate glass derived from kaolin; these are formed into sheets by means of standard paper-making techniques (Barnard, 1981). The major components in the glass are alumina (47% to 65%) and silica (38% to 50%), which makes the material highly heat-resistant. On the other hand, the cellulose materials burn if heated in air, and at a burnout temperature of 700°C they disappear completely from the casting ring. However, if liners are kept short of both ends of the ring, the investment mold is retained in place during casting.

Cellulose liners have a water uptake similar to asbestos and a similar effect on the setting expansion of the investment.

Although ceramic liners absorb negligible amounts of water when immersed at atmospheric pressure, they absorb it readily under vacuum. This occurs during vacuum investing; since the previously dry liners obtain the water from the investment mix, the W/P ratio of the unset investment is greatly reduced. The combination of a low W/P ratio and a wet liner considerably increases the investment's setting expansion; the total expansion is higher than when prewetted asbestos or cellulose liners are used with normal W/P ratios.

Thermal expansion  After it has set, the investment mold is heated to the recommended temperature for casting. This is necessary to dry the investment, to melt and burn out the wax pattern, to oxidize residual carbon from the mold, and to prevent premature freezing of thin sections when the alloy is cast. Heating also causes thermal expansion of the mold and therefore of the mold cavity.

1. Hygroscopic expansion techniques. With these techniques, the mold is heated to about 480°C, at which temperature its thermal expansion is relatively low (Fig 4-14).
Investments for use in these techniques are based on a quartz refractory that does not undergo inversion until heated to 573°C (see Fig 4-10).

Fig 4-14 Setting and thermal expansion of a typical investment of the hygroscopic expansion type (Whip Mix Beauty-Cast, W/P = 0.30). The specimen was immersed in room-temperature water during setting and then heated to 480°C. This investment is based on quartz and contains a halide, which gives it a small positive thermal expansion, uniform over a range of about 150°C. The dashed continuation of the curve shows the effect of heating the investment to 700°C.

A mold temperature as low as 480°C has the advantage of ensuring a fine grain structure in the solidified casting, though this is not an important consideration if the alloy contains grain-refining elements. It has the disadvantage that carbon remaining from burnout of the wax pattern is oxidized very slowly. Excessive carbon deposits interfere with venting of the mold cavity, so prolonged heating at the casting temperature (preferably 1 hour) is needed to ensure that casting defects caused by inadequate venting ("back-pressure porosity") do not occur.

2. Thermal expansion techniques. With thermal expansion techniques, the mold is usually heated to about 700°C in order to gain maximum mold expansion. A higher mold temperature should not be used or breakdown of the calcium sulfate binder can occur in the presence of carbon, thereby liberating sulfur dioxide. The carbon may be present as graphite added to the original investment powder as a reducing agent or may simply be carbon residue from burnout of the wax pattern. The reaction, which involves reduction of the calcium sulfate binder, takes place in two stages:
\[
\begin{align*}
\text{CaSO}_4 + 4\text{C} & \rightarrow \text{CaS} + 4\text{CO} \\
3\text{CaSO}_4 + \text{CaS} & \rightarrow 4\text{CaO} + 4\text{SO}_2
\end{align*}
\]

The sulfur dioxide formed by the second reaction causes sulfide formation on the gold-alloy casting, resulting in discoloration and embrittlement of the alloy (O'Brien and Nielsen, 1959).

Investments for thermal expansion techniques may contain either cristobalite or quartz as the refractory; occasionally a mixture of both is used. Cristobalite investments have the advantage that thermal expansion is fairly constant over the temperature range of 400°C to 700°C, but have the disadvantage that the large expansion at 250°C, resulting from the \(\alpha\rightarrow\beta\) inversion of cristobalite, can cause mold cracking if heating is not carefully controlled (Fig 4-15, curve A). It should be slow in the temperature range of 230°C to 270°C while the inversion of cristobalite is occurring throughout the mold.

Quartz has a lower thermal expansion than cristobalite (see Fig 4-10), so most quartz investments for use in thermal expansion techniques contain additives to increase their thermal expansion. The expansion on heating is more gradual, so control of heating rate is not as important, but maximum expansion is available only in the temperature range of 600°C to 700°C (Fig 4-15, curve B).

**Fig 4-15** Setting and thermal expansion of two typical investments of the thermal expansion type (curve A: Kerr Cristobalite Inlay, W/P = 0.38; curve B: R&R Gray, W/P = 0.25). With both materials the specimens were surrounded by a wet ring liner while setting and were then heated to 700°C. The former material is an inlay investment based on cristobalite. The latter material is described as a universal investment but is most commonly used for casting dentures and is based on quartz, with the addition of sodium chloride to increase its thermal expansion.
At a burnout temperature of 700°C, residual carbon is oxidized rapidly, so venting of the mold cavity is not a problem.

3. **The effect of a wet liner.** In thermal expansion techniques, if a wet ring liner is used, not only the setting expansion but also the subsequent thermal expansion can be affected. Once the investment has set, the extra water it absorbed from the liner has the same effect as if it had been added during mixing (i.e., an increased W/P ratio). Therefore, the thermal expansion may be reduced.

The same considerations apply to denture investments. The investment cast forms part of the final mold, and its expansion is the major factor determining the accuracy of the casting. It sets in contact with a duplicating impression, usually an aqueous gel, which, like a wet ring liner, makes free water available to the setting investment with the same effects as those already described.

**Control of expansion**

1. **Composition.** The manufacturer adjusts investment expansion by the choice of refractory and binder and sometimes by the use of suitable additives.

2. **W/P ratio.** Decreasing the W/P ratio increases both setting expansion (in air or water) and thermal expansion. The W/P ratio must be carefully controlled for reproducible mold expansion. Changes in expansion can most easily be effected by changing the W/P ratio (Fig 4-16).

![Fig 4-16](image)

**Fig 4-16** The effect of changing the W/P ratio on the setting, thermal, and total expansions of a typical investment of the thermal expansion type, set against a dry ceramic liner without vacuum investing, and then heated to 700°C.
3. Period of exposure to water. In hygroscopic expansion techniques, additional control can be obtained by varying the length of time the setting investment is exposed to an aqueous environment. This may be achieved by reducing the time for which the setting investment is immersed in a water bath, or by adding controlled amounts of water to the top of the investment mix in the ring instead of immersing it. In the latter case, if only a small volume of water is added, it will be completely absorbed by the investment before setting is complete, and subsequent expansion will be reduced. Varying the amount of added water varies the setting expansion proportionally.

Strength

The mold must be strong enough to withstand stresses at ambient temperatures (eg, during removal of the sprue former) and when heated to the recommended mold temperature for casting (eg, during rapid entry of molten alloy). Strength properties of investments are usually determined by testing in compression. Inlay investments have wet compressive strengths mostly in the range of 2 to 6 MPa. Because of the need to make a working cast, denture investments have higher compressive strengths when set, mostly in the range 9 to 14 MPa. In both investment types, changes in the compressive strength on heating to the recommended casting temperature vary according to the presence of additives to control investment thermal expansion. Unless it is known whether an additive is present in an investment, and if so what it is, it is impossible to predict strength at the casting temperature from measurements made at ordinary ambient temperatures. Little information is available on the compressive strength of gypsum-bonded investments at the casting temperature (hot strength). In one study on a limited number of materials (Earnshaw, 1969b), it was found that the hot strength of four investments without additives was in the range of 1.8 to 9.4 MPa; for three investments containing boric acid it was 6.3 to 13.3 MPa; and for three investments containing sodium chloride it was 3.1 to 5.2 MPa.

Rapid-heat investments

Although it was stated earlier that investments based on a cristobalite refractory require slow heating while the $\alpha \rightarrow \beta$ inversion is occurring, some "rapid-heat" investments have been introduced, which are placed immediately after setting into a furnace preheated to 700°C. At least some of these contain cristobalite as the refractory. The recommended technique is to place the mold, 30 minutes after the pattern is invested, into the preheated furnace for an additional 30 minutes; the casting is then made. A setting and thermal expansion curve for a typical example is shown in Fig 4-17.
Setting and thermal expansion of a rapid-heat investment of the thermal expansion type (GC Cristoquick, W/P = 0.33). As the manufacturer instructed, the investment was allowed to set against a dry ceramic liner for 30 minutes, then was immediately transferred to a furnace preheated to 700°C.

Little information is available on these materials. It can be seen in Fig 4-17 that setting expansion is still occurring rapidly at 30 minutes; measured under ordinary conditions, it is not complete until 2 hours after mixing and measures 1.1%. The rapid rate of expansion at 30 minutes means that precise timing of placement of the mold in the furnace is critical if reproducible mold expansion is to occur.

Such a drastic heating program could be expected to cause severe thermal cracking in an ordinary cristobalite investment. Measurements on a mold in a lined inlay ring showed that the periphery of the investment mass reached 250°C within 6 minutes of entering the hot furnace, while the center was at only 110°C. The center of the mold did not reach 250°C until 4 minutes later. Both periphery and center had reached a maximum of 690°C within the 30-minute heating period.

The indicated temperatures shown in Fig 4-17 were recorded by a thermocouple whose hot junction was embedded in the center of a specimen that had about the same thickness as an inlay mold. The expansion caused by the inversion of cristobalite, shown on the graph as beginning at 110°C and finishing at 170°C, obviously took place when enough of the outer parts of the specimen had reached 250°C to produce a volume change.

These rapid-heat investments save laboratory time and simplify the casting procedure, as the furnace is maintained at 700°C instead of being repeatedly heated and cooled. Despite the curtailed setting expansion, the investment's total expansion under these conditions was 1.95%, more than enough to compensate for the casting shrinkage of ordinary dental gold alloys.
Casting accuracy

In all casting procedures, after the molten metal or alloy has filled the mold, a volumetric contraction occurs in the liquid and then in the solid casting as it cools. In addition, with almost all metals and alloys there is a volumetric contraction during solidification.

In dental casting, thermal contraction of the liquid alloy, and its solidification contraction, do not affect the dimensions of the casting because, under the influence of the casting force, continued feeding of liquid alloy occurs from the excess in the sprue and button. Interruption to the flow of liquid during solidification, which may be caused by premature freezing of an incorrectly designed sprue, will not cause an overall contraction of the casting, but will lead to localized shrinkage porosity.

Therefore, thermal contraction of the solidified casting, as it cools from solidus to ambient temperature, remains as the sole cause of the observed casting shrinkage. Although this is a volumetric contraction, in dental technology dimensional changes are usually studied on a linear basis. It is assumed that these changes occur isotropically—an assumption that is frequently unjustified.

The fact that the observed casting shrinkage is thermal contraction of the solid alloy can be confirmed, at least for pure gold, by comparing measured values for its linear casting shrinkage with a value calculated from its coefficient of linear thermal expansion (\( \alpha \)) and the temperature difference between its freezing temperature and ambient.

Results of direct measurements of the linear casting shrinkage of gold are shown in Table 4-5. A theoretical value can be calculated as follows:

\[
\begin{align*}
\alpha_{(20^\circ\text{C}-900^\circ\text{C})} &= 16.7 \times 10^{-6}/\text{K} \\
\text{Melting temperature} &= 1,064^\circ\text{C}
\end{align*}
\]

If 20°C is taken as an average ambient temperature, the solidified gold cools through 1,044 K. Total linear contraction per unit length is given by 1,044 \times 16.7 \times 10^{-6} = 0.0174, which converts to 1.74%.

Alloying gold will alter the value for \( \alpha \) and for solidus temperature. Calculations similar to that given previously have been made on a group of 12 high-noble-metal casting alloys, and the calculated linear casting shrinkages varied from 1.65% to 1.80% (Finger and Jorgensen, 1980). Thus an average value of 1.7% would apply to gold and high-gold alloys.

<table>
<thead>
<tr>
<th>Reference</th>
<th>Specimens (mm)</th>
<th>Linear casting shrinkage (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hollenback and Skinner (1946)</td>
<td>Cylindrical rod (5.6 \times 25)</td>
<td>1.67 ± 0.02 (8)</td>
</tr>
<tr>
<td>Earnshaw (1960)</td>
<td>Cylindrical rod (6.3 \times 60)</td>
<td>1.74 ± 0.03 (6)</td>
</tr>
</tbody>
</table>


Nakai et al (1980) Rectangular prism (3 × 5 × 20)

*Values are shown as mean and standard deviation, with number of tests in parentheses.

It is conventionally assumed that compensation for this casting shrinkage is provided by mold expansion. Calculations of available compensation, however, must be based on measurements of investment expansion made under conditions that reproduce those obtained in the casting ring (Earnshaw, 1988).

Another factor that must be considered is the effect of the ring liner on investment expansion. Absorbent liners (asbestos and cellulose) vary in thickness, water uptake, and compressibility when wet. In thermal expansion techniques, some may not have sufficient compressibility to accommodate all of the increased setting expansion provided by the availability of extra water, so that restriction of expansion could occur in a diametral direction in the ring (Morey and Earnshaw, 1992). Thus they affect to a varying degree the available mold expansion. Ceramic liners also vary in thickness and compressibility, and if vacuum investing is used, they vary in the amount of water they absorb from the investment mix. Again they affect mold expansion in the ring to a varying degree (Earnshaw and Morey, 1992).

The choice of casting-ring liner can thus affect, often to a significant extent, the dimensional accuracy of the casting. Figure 4-18 shows the relative inaccuracy produced in full-crown castings made in a high-gold alloy with a gypsum-bonded cristobalite investment of the thermal expansion type, used with vacuum investing. The only variation in technique was the casting-ring liner used. All the liners were water absorbent, and all were prewetted under controlled conditions. Three were asbestos, and three cellulose.

Fig 4-18 The effect of different prewetted ring liners on the relative inaccuracy of gold-alloy full-crown castings. The castings were made with a cristobalite inlay investment of the thermal expansion type used with vacuum investing. C = control
castings made without a liner; GA, KA, and SH = castings made with asbestos liners; and AU, BE, and WM = castings made with cellulose liners. The height of each bar shows the mean of at least 10 castings, while the narrow superimposed bar shows the standard deviation.

Figure 4-19 shows the results of similar experiments made with five different dry ceramic liners used with and without vacuum investing. These two graphs show the great variation in casting inaccuracy produced when the same investment is used with different casting-ring liners. This variation in inaccuracy is often unpredictable; in these experiments, only when the ceramic liners were used with vacuum investing was there any correlation between measured liner properties and casting inaccuracy.

Fig 4-19 The effect of different dry ceramic ring liners on the relative inaccuracy of gold-alloy full-crown castings. Conditions were the same as in Fig 4-18, except that castings were made with and without vacuum investing. C = control castings made without a liner; DK, GC, and KF = castings made with ceramic liners supplied for dental use; and K1 and K2 = castings made with liners cut from commercial insulating material of the same type, supplied in sheets nominally 1 mm and 2 mm thick.

Although with gypsum-bonded investments mold expansion is the major factor affecting relative casting inaccuracy, mold strength at the casting temperature must also be considered. The average value of 1.7% for alloy thermal contraction represents the inherent casting shrinkage of the alloy—the shrinkage that would occur in a casting that was free to contract without constraint. Under practical conditions the casting is enclosed in a mold, and in all but the simplest castings its shape allows some interlocking between investment and mold. In the early stages of cooling the mold has a lower rate of thermal contraction than the casting; the thermal contraction of a typical high-gold alloy, cooling from its solidus temperature to 500°C, would be
0.74%, while for a typical cristobalite inlay investment, cooling from 700°C to 500°C, it would be 0.20%. If the investment has a high hot strength, alloy thermal contraction will be opposed by the mold; the degree of constraint will depend on the casting shape and is usually directional. At temperatures near its solidus the alloy is weak and plastic, and mold constraint can cause anisotropic thermal contraction. When the alloy has cooled enough to attain rigidity, a strong mold can cause plastic deformation by hot-working the alloy. Both factors cause anisotropic contraction and distortion of the casting.

A constant pattern of distortion of gold-alloy mesio-occluso-distal inlays has been reported (Teteruck and Mumford, 1966) and can be attributed to directional restraint of alloy shrinkage by the mold (Earnshaw, 1969b). The gypsum-bonded investment used in Teteruck and Mumford's experiments had a compressive strength at 700°C in the range of 4 to 6 MPa.

Therefore, although with gypsum-bonded investments the total investment expansion that occurs in the casting ring is the major factor affecting the casting inaccuracy, the hot strength of the mold must be considered. For the casting to reproduce accurately the size and shape of the original pattern, the investment expansion must compensate fully for the thermal contraction of the solid alloy, the ring and ring liner must allow the potential expansion of the investment to be achieved isotropically, and the investment at the burnout temperature must not be so strong that it restricts the shrinkage of the alloy in any direction. On the other hand, it should not be so weak that it may fracture during casting. A safe minimum compressive strength at 700°C is 1.8 MPa, and this allows substantially isotropic alloy contraction (Morey and Earnshaw, 1995).

**Storage of Gypsum Products**

When hemihydrate powders are exposed to the atmosphere, water vapor is adsorbed; the extent to which this occurs depends on the prevailing water vapor pressure (Torrance and Darvell, 1990). If sufficient water is adsorbed to form a liquid film, and if this becomes thick enough to function as a solvent, the setting reaction begins on the surface of the powder particles, causing deterioration of the material. Gypsum crystals are formed that act as nucleation centers when the powder is mixed with water, and accelerate the setting rate. But if exposure of the powder continues, surface hydration will form a layer of gypsum on each particle, hindering the access of water to the hemihydrate, and the subsequent setting rate is retarded. If exposure continues over a long period, eventually the material will not set properly, because insufficient unreacted hemihydrate remains to form a coherent set mass. This is equally likely to occur in gypsum-bonded investments and will have the same effects.

A long-standing recommendation to store gypsum products at relative humidities not greater than 70% (Farmer and Skinner, 1942) appears satisfactory only at temperatures of 25°C or lower, but of uncertain validity at higher temperatures. To avoid deterioration, especially at high temperatures and high relative humidities, bulk gypsum products should be stored in closed bins, in a cool dry area of the laboratory. "Topping up" of bins should not be done over old stock. Packaging of small quantities of powder, in waterproof containers, is a preferable method of storage in the
laboratory. Many die stones and gypsum-bonded investments are now supplied by their manufacturers packaged in this way.

**Glossary**

**acicular (of a crystal)** Slender, needlelike.

**apparent density (of a powder)** The mass of a sample divided by the volume it occupies, measured at a specified degree of compaction. The measured volume includes all interstitial spaces and porosities, so powders with a poor packing ability show a low apparent density.

**calcination** Prolonged heating of a substance at some temperature below its melting temperature.

**crystal habit (crystalline form)** The external geometric shape of a crystal.

**crystal structure** The regular three-dimensional arrangement of atoms within a crystal. This internal atomic arrangement is characteristic of a particular crystalline solid, but may or may not be the same as the external shape of individual crystals.

**inversion** A temperature-dependent change from one polymorphic form to another. **Rapid (high-low) inversions** involve only an alteration in bond angles within the crystal structure (shear or displacive transformation). This change requires only a small activation energy (excess energy required for the change to occur), so it occurs rapidly on heating to the critical temperature and is rapidly reversible on cooling. An example is the α→β inversion of quartz or cristobalite. **Slow inversions** involve breaking of interatomic bonds and atomic diffusion to form a new crystal structure (reconstructive transformation). In a ceramic material in which a significant amount of the interatomic bonding is covalent, such as crystalline silica (Jastrzebski, 1976; Sosman, 1965), such a transformation involves a high activation energy, so prolonged high-temperature heating is needed for complete conversion. Because the transformation is so sluggish, the new high-temperature phase is retained during cooling to ambient temperature, where in most materials the rate of atomic diffusion is extremely slow. The high-temperature form therefore persists indefinitely as a metastable phase. An example is the quartz → cristobalite inversion.

**isotropic** Occurring equally in all directions. (Antonym: **anisotropic**.)

**nucleation center** In an aqueous solution, a region where spontaneous, orderly deposition of ions or molecules forms nuclei that continue to grow, beginning the process of crystallization. Normally this process involves heterogeneous nucleation, where nucleation centers are provided by the deposition of ions or molecules on existing solid surfaces, such as solid particles (seeds) already present in the solution.

**nuclei poisoning** The inactivation of nuclei of crystallization by the deposition of foreign material on their surfaces.

**polymorphism** The existence of an element or chemical compound in more than two different crystalline forms.
**pseudomorphic** A crystalline material whose crystal habit differs from the normal shape that would be dictated by its crystal structure.

**pseudoplasticity** The behavior of non-Newtonian fluids in which the rate of shear increases more than in proportion to the shearing stress (shear-thinning liquids). Thick mixes of plaster or stone are typical examples; they appear to be very viscous when at rest but flow readily when subjected to a shear stress (eg, stirring or vibration).

**refractory** Capable of resisting high temperatures.

**relative surface area (of a powder)** The total surface area of a given mass.

**specular (of reflections)** Mirrorlike.

**spherulite** A spherical aggregation of needle-shaped crystals radiating out from a common center.

**water/powder (W/P) ratio** The mixing proportions of plaster or stone, expressed as a decimal fraction. If 100 g of plaster is to be mixed with 50 g of water, the W/P ratio is 0.50.

**water requirement** The amount of water that must be added to a given mass of plaster or stone to produce a mix of suitable viscosity.

**Discussion Questions**

1. How many different applications do gypsum products have in dentistry, as well as other fields?

2. Why are storage conditions and shelf life important considerations in the use of gypsum products? Why do these present special problems for the dental services of large organizations such as the armed forces?

3. Why is it essential to follow directions to achieve the higher strengths possible with improved die stones?

4. What are the main methods of reducing bubbles in gypsum casts and molds?

**Study Questions**

*(See Appendix E for answers.)*

1. What differences in chemical composition are there in the powders of plaster, stone, and die stone?

2. What physical differences are there in the powders of plaster, stone, and die stone? How are these differences related to the methods of manufacture of the powders?

3. Give typical W/P ratios for plaster, stone, and die stone. Why are these different?
For each type, assuming that the setting reaction goes to completion, what percentage of the mass of the set material would consist of free water?

4. What chemical and physical changes accompany the setting of gypsum products?

5. What is the difference between the microstructures of gypsum casts made from plaster and stone? What is the cause of this difference, and what is its effect on strength properties?

6. What is meant by the setting time of gypsum products? What is the practical significance of (a) the loss of surface gloss, (b) the Gillmore initial set, and (c) the Vicat initial set?

7. Explain how the manufacturer of a dental gypsum product adjusts its rate of setting.

8. For a given gypsum product, what is the effect of increasing the W/P ratio on the rate of setting? Explain.

9. Theoretically, the setting of gypsum products should be accompanied by a volumetric contraction. Why is a setting expansion observed in practice?

10. Why do gypsum products show a greater setting expansion in water than in air?

11. What is the practical significance of the setting expansion of dental gypsum products?

12. Give typical values for the linear setting expansion in air of impression plaster, plaster, stone, and low-expansion and high-expansion die stones.

13. Explain how the manufacturer of a gypsum product controls its setting expansion.

14. What practical limits are placed on the extent of this control?

15. For a given gypsum product, what is the effect of increasing the W/P ratio on setting expansion? Explain.

16. What is the practical significance of the tensile strength and the compressive strength of dental gypsum products?

17. For a given gypsum product, what is the effect of increasing the W/P ratio on tensile strength and compressive strength? Explain.

18. What is the effect of drying on the tensile strength and the compressive strength of cast gypsum? Explain.

19. What constituents are likely to be present in a gypsum-bonded investment powder? Give the function of each.

20. Within normal limits, what effect does increasing the proportion of refractory
filler in an investment have on (a) rate of setting, (b) setting expansion in air, (c) setting expansion in water, (d) thermal expansion, and (e) compressive strength?

21. How does the investment set?

22. What is the practical significance of microscopic porosity formed in the investment mass during setting?

23. What are the requirements for the particle size of an investment powder? Give reasons.

24. Why is the particle size of the refractory filler more important than that of the binder?

25. What is the most important property of an investment? Why?

26. What is a hygroscopic expansion technique?

27. What is a thermal expansion technique?

28. In hygroscopic expansion techniques, why is prolonged heating of the mold at the burnout temperature necessary?

29. How can the user of a dental investment discover whether it contains cristobalite or quartz as a refractory? What is the practical importance of this information?

30. Why is control of the heating rate more important with a cristobalite investment than with a quartz investment?

31. Why is an accurate furnace pyrometer more important with a quartz investment than with a cristobalite investment?

32. Why should gypsum-bonded investments not be heated above 700°C?

33. In thermal expansion techniques, what is the effect of a wet liner on the investment's setting expansion?

34. What is the effect of increasing the W/P ratio on (a) the investment's setting expansion in air, (b) its setting expansion in water, and (c) its thermal expansion.

35. In hygroscopic expansion techniques, how can mold expansion be controlled by the user?

36. In thermal expansion techniques, how can mold expansion be controlled by the user?

37. What is the practical significance of the compressive strength of the investment at the casting temperature?

38. How should gypsum products be stored in a dental laboratory? How can incorrect
storage conditions affect these materials?

**Recommended Reading**


Morey EF, Earnshaw R (1992). The fit of gold-alloy full-crown castings made with


Chapter 5. Surface Phenomena and Adhesion to Tooth Structure

Introduction

Surface phenomena include surface tension, wetting, adsorption, capillary action, and adhesion. Applications include capillary penetration around restorations, dentures, and teeth, and adhesion to tooth structure by sealants and restorative materials.

Surface Energy

Introduction

Atoms and molecules at the surfaces of liquids and solids possess more energy (surface energy) than do those in the interior. In the case of liquids, this energy is called surface tension. As illustrated in Fig 5-1.

![Fig 5-1 Distribution of vacancies (open circles) within isolated capillary liquid. More vacancies are present at the surface (a) as compared to the bulk (b). Fewer vacancies are found at the liquid-solid interface (c). (From O'Brien, 1970. Modified with permission.)](image)

The molecules at the surface are farther apart due to loss of molecules by evaporation. From Fig 5-2 it can be seen that this greater average separation leads to a net attraction between molecules and a higher energy of attraction. This results in a surface contractile force or surface tension, which causes the liquid to form drops and to exhibit a surface skin that resists extension or penetration.
The surface energies of several substances are given in Table 5-1. It can be seen that the surface energies of oxides and metals are greater than those of liquids. In general, the higher the bond strength of a substance, the greater the surface energy. Since metallic bonds are much stronger than the van der Waals bonds of liquids, metals have higher surface energies. The units of surface energy are ergs/cm², but the surface tension of liquids is often expressed in the equivalent units of dynes/cm. The total surface energy of a system is the product of the surface energy of the material and the total area. Therefore, a high total surface energy exists if a material is finely divided (powder or colloid) to provide a large surface area, especially if the material has a high surface energy per unit area (eg, metals or ionic crystals).

**Table 5-1** Surface energies of various substances

<table>
<thead>
<tr>
<th>Substance</th>
<th>Surface energy (erg/cm²)</th>
<th>Temperature (°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water</td>
<td>72</td>
<td>20</td>
</tr>
<tr>
<td>Benzene</td>
<td>29</td>
<td>20</td>
</tr>
<tr>
<td>Olive oil</td>
<td>36</td>
<td>20</td>
</tr>
<tr>
<td>Saliva</td>
<td>56</td>
<td>23</td>
</tr>
<tr>
<td>NaCl crystal</td>
<td>300</td>
<td>25</td>
</tr>
<tr>
<td>Dental porcelain</td>
<td>365</td>
<td>1,000</td>
</tr>
<tr>
<td>Copper, solid</td>
<td>1,430</td>
<td>1,080</td>
</tr>
<tr>
<td>Silver, solid</td>
<td>1,140</td>
<td>750</td>
</tr>
</tbody>
</table>
**Sintering**

*Sintering* is a densification process in which finely divided particles are heated in contact. The firing of porcelain is a sintering process used for forming denture teeth and porcelain-fused-to-metal crowns. The driving force for this process is the reduction of total surface area and, thus, total surface energy.

**Wetting**

*Wetting* is defined in terms of the degree of spreading of a liquid drop on a solid surface. The **contact angle** (θ), formed by the liquid surface and the interface separating the liquid and solid, is used as a measure of the degree of wetting (Fig 5-3).

![Wetting Diagram](image)

**Fig 5-3** Wetting. Low contact angle indicates good wetting (*left*); high contact angle indicates poor wetting (*right*). (From O'Brien and Ryge, 1965b. Reprinted with permission.)

A 0-degree contact angle indicates complete wetting, and low values correspond to good wetting. Values above 90 degrees indicate poor wetting. **Table 5-2** gives contact angles for several systems.

**Table 5-2** Contact angles of liquids on solids

<table>
<thead>
<tr>
<th>Solid</th>
<th>Liquid</th>
<th>Contact angle (degrees)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amalgam alloy</td>
<td>Water</td>
<td>80</td>
</tr>
<tr>
<td>Silicate cement</td>
<td>Water</td>
<td>10</td>
</tr>
<tr>
<td>Acrylic</td>
<td>Water</td>
<td>75</td>
</tr>
<tr>
<td>Teflon</td>
<td>Water</td>
<td>110</td>
</tr>
<tr>
<td>Ag₃Sn</td>
<td>Mercury</td>
<td>140</td>
</tr>
<tr>
<td>Gold alloys</td>
<td>Porcelain enamel</td>
<td>40-50</td>
</tr>
<tr>
<td>Nickel alloys</td>
<td>Porcelain enamel</td>
<td>80-100</td>
</tr>
<tr>
<td>Hydron</td>
<td>Water</td>
<td>0</td>
</tr>
<tr>
<td>Etched enamel</td>
<td>Pit and fissure sealants</td>
<td>0</td>
</tr>
</tbody>
</table>
Good wetting promotes **capillary penetration** and **adhesion** and indicates strong attraction between the liquid and solid surface molecules. Good wetting is important in soldering and is a factor in better denture retention. A more natural appearance is achieved if restorative materials are wetted by a thin film of saliva. **Hydrophobic** substances are those that exhibit high contact angles with water (eg, Teflon, silicone coatings) ([Figs 5-4 and 5-5]).

**Fig 5-4** Hydrophobic film of fluorinated polymer on enamel giving high contact angle. (From O'Brien, 1973a.)
Fig 5-5 Poor wetting shown by porcelain enamel drops on nongold alloy at 1,040°C. The furnace thermocouple is seen above the drops. (From O'Brien and Ryge, 1965a. Reprinted with permission.)

Adsorption

To reduce surface energy, atoms and molecules that are mobile will concentrate at high energy surfaces. For example, finely divided charcoal will adsorb quantities of several gases; detergent (or soap) molecules will concentrate at the surface of water, leading to a large reduction in surface energy. Adsorption is strongest when there is large energy saving (high surface energy material and large surface area) and slows down as the surface is covered.

Adsorption occurs only at the surface, whereas absorption involves penetration and uptake by the interior of the material (as in swelling of hydrocolloid impression materials in water). Gold foil adsorbs gases readily and must be degassed before use.

Colloids

Introduction

Colloids contain material that is present in particles larger than ordinary atoms or molecules but still invisible to the unaided eye (ie, 10 to 10,000 Å). There are three types of colloidal systems.

Insoluble dispersed particles

These are called lyophobic, since the materials are insoluble in a liquid medium (eg, sulfur in water). The fine particles acquire electrical charges that keep them in suspension. Table 5-3 gives a classification of the types of lyophobic systems. As one example of such a system, colloidal gold is used to form a gold coating on alloys to be bonded to porcelain. Although lyophobic systems may last for many years, they are unstable and can be precipitated by electrical methods (smokes) or gravity (emulsions).

Table 5-3 Classification of lyophobic colloids
<table>
<thead>
<tr>
<th>Dispersed phase</th>
<th>Continuous phase</th>
<th>Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>Solid</td>
<td>Liquid</td>
<td>Sol</td>
</tr>
<tr>
<td>Solid</td>
<td>Gas</td>
<td>Aerosol (smoke)</td>
</tr>
<tr>
<td>Liquid</td>
<td>Liquid</td>
<td>Emulsion</td>
</tr>
<tr>
<td>Liquid</td>
<td>Gas</td>
<td>Aerosol (fog)</td>
</tr>
<tr>
<td>Gas</td>
<td>Liquid</td>
<td>Foam</td>
</tr>
<tr>
<td>Gas</td>
<td>Solid</td>
<td>Foam</td>
</tr>
</tbody>
</table>

**Large molecules**

These systems are true solutions in which the dispersed molecules are of colloidal dimension. They are stable, but the large size of the macromolecules present gives the solutions properties similar to those of lyophobic systems.

Dental materials consisting of such solutions include agar and alginate hydrocolloid impression materials. These materials aggregate to form gels, due to van der Waals bonding between the long chain molecules. Gels undergo imbibition and syneresis (absorption and exudation of solvent) with resulting swelling and shrinkage, respectively. This behavior is responsible for the dimensional instability of these impression materials. As with other colloidal systems, the liquid state is called the sol state.

**Association colloids**

These are aggregates of smaller molecules that achieve colloidal size. Surface active agents, such as soaps and detergents, are examples. Each molecule consists of a long-chain hydrocarbon with a small charged polar group at one end (eg, sodium palmitate). The aggregates formed by these molecules, called micelles, often are spherical; in aqueous systems, the hydrocarbon ends gather in the center of the micelle and the polar groups are exposed on the outside. This system is useful in cleaning because grease and other organic films are dissolved in the interior of the micelles and held in suspension.

**Capillary Penetration**

**Introduction**

The surface energy of a liquid creates pressure that drives the liquid into crevices, narrow spaces, and thin tubes. Saliva penetration around restorations (leakage) and around teeth are important examples. Capillary penetration of saliva is also partially responsible for denture retention.

**Capillary rise**

If a glass tube is immersed in a liquid (Fig 5-6(a)), the capillary rise, $h$, is given by the formula:
where \( \gamma \) is the surface tension, \( \theta \) is the contact angle, \( r \) is the tube radius, \( d \) is the liquid's density, and \( g \) is the gravitational constant, 980 dynes/g. If the contact angle is less than 90 degrees, as in water-glass and saliva-enamel systems, elevation of the liquid occurs. However, if the contact angle exceeds 90 degrees (eg, mercury on glass or water on Teflon), depression takes place and pressure must be applied to force the liquid into the space (Fig 5-6(b)). If the capillary liquid is not connected with a reservoir of liquid, isolated capillaries (isocaps) are formed (Fig 5-7). The liquids in isocaps exert adhesive force on the walls of the capillary, which can be strong for thin layers. This capillary adhesion is a factor in denture retention.

\[
h = \frac{2\gamma \cos \theta}{rdg}
\]

**Fig 5-6** Effect of contact angle on capillary penetration: (a) capillary elevation with concave meniscus (low contact angle); (b) capillary depression with convex meniscus (high contact angle). \( \Delta P \) is the capillary pressure. (From O'Brien et al, 1968. Modified with permission.)

**Fig 5-7** (a) Capillary liquid in contact with a reservoir. (b, c) Isolated capillaries. (From O'Brien and Fan, 1975. Modified with permission.)
To increase the penetration of saliva around acrylic dentures, surface coatings of silica are applied to reduce the contact angle. The main factor here, however, is maintenance of a small gap distance (ie, close adaptation between denture and mucosa), as shown in Fig 5-8.

**Fig 5-8** Capillary space between denture and mucosa.

---

**Penetration coefficient**

The rate of movement of a liquid into a capillary space is related to the surface tension (\(\gamma\)), contact angle (\(\theta\)), and viscosity (\(\eta\)), as given by the penetration coefficient (PC):

\[
PC = \frac{\gamma}{2\eta} \cos \theta
\]

Therefore, a liquid with low viscosity, high surface tension, and low contact angle (ie, good wetting) will penetrate faster than one with the opposite combination of properties. This is important in adhesives such as pit and fissure sealants, which must penetrate into surface roughness and crevices quickly for good bonding.

**Adhesion**

**Introduction**

Adhesion is the attachment of materials in contact that resists the forces of separation. Adhesive phenomena are critical in many areas of dental materials, including the bonding of porcelains to metals and the adhesion of resins to tooth structure. Several types of adhesive bonds may be identified according to the classifications in Fig 5-9. Mechanical adhesion depends on mechanical interlocking of the two phases and may include microscopic attachments, as in the case of resin bonding to etched enamel or hoop stresses of a porcelain around a metal core. Chemical adhesion relies on chemical bonding between two phases. Diffusion bonding results when one phase
penetrates by diffusion into the surface of a second phase and forms a hybrid layer, which is a composite of the two materials.

**Fig 5-9** Classification of bonding mechanisms.

Several factors affect the strength of an adhesive bond:

1. **Cleanliness.** The surfaces to be attached should be free of debris and contamination.

2. **Penetration of surface.** Liquid adhesives (eg, sealants and bonding agents) must penetrate into crevices created by acid etching of enamel and dentin. The penetration coefficient of the liquid was discussed in the preceding section.

3. **Chemical reactions.** The formation of strong chemical bonds across an interface will increase the number of attachment sites. This is believed to occur between porcelain enamels and the oxides of tin, indium, and iron formed on the surfaces of alloys containing high proportions of precious metals. On the other hand, a weak compound may be formed by a chemical reaction, resulting in a weak boundary layer (eg, certain oxides) rather than attachment sites.

4. **Shrinkage of adhesive.** Liquid adhesives solidify by processes such as solvent evaporation and polymerization, and shrinkage results. The adhesive may then pull away from the substrate, or stresses may be created that weaken the bond. Shrinkage is toward the center of the adhesive mass (**Fig 5-10**). However, shrinkage in light-cured systems occurs toward the light source.
5. **Thermal stresses.** If the adhesive and substrate have different thermal expansion coefficients, changes in temperature will produce stresses in the bond. For example, porcelain enamels are bonded to alloys at high temperatures and then cooled to room temperature. Close matching of the thermal expansion coefficients of porcelain enamel and alloy is required to minimize stresses.

6. **Corrosive environment.** The presence of water or corrosive liquid or vapor will often lead to deterioration of an adhesive bond. For example, acrylic resins will initially adhere to clean unetched tooth enamel, but the bond deteriorates upon storage in water.

**Adhesion to enamel**

Enamel is highly mineralized tissue composed of hydroxyapatite (about 96%), water (about 4%), and collagen (about 1%). Treatment with 35% to 50% phosphoric acid results in selective demineralization of exposed enamel rod ends, leaving a surface with increased area and high energy (Fig 5-11).
High surface energy permits efficient wetting by the hydrophobic resin, which penetrates to form tags (Fig 5-12) and provides bond strength through mechanical interlocking. Contamination of the dry, etched surface by saliva or water, or incomplete removal of the etching agent or dissolved minerals, adversely affects the long-term stability of the bond. The **acid-etch technique** produces shear bond strengths during laboratory testing of 20 to 22 MPa, on average above those produced at the composite-enamel interface by shrinkage due to polymerization (~18 MPa).
Acid etching of enamel is a widely accepted clinical procedure and has increased the life of resin composite restorations by decreasing the possibility of marginal staining, secondary caries, and postoperative sensitivity due to sealing of the enamel margins. The acid-etch technique allows the use of a more conservative cavity preparation and, to a certain extent, restores fracture resistance to the bonded structure.

*Adhesion to dentin*

Human dentin is composed of hydroxyapatite (45%), water (25%), and organic matrix (30%). Bonding to dentin is routinely achieved by etching and the diffusion of hydrophilic primers, followed by application of bonding resins. Composite restorative materials are then bonded to the resin layer; the resin-dentin interface has laboratory bond strengths of 22 to 35 MPa.

The stages of conditioning and resin impregnation are illustrated in Fig 5-13. In the first stage, the conditioning dissolves the smear layer of debris on the dentinal surface and in tubules, partially decalcifies the dentin to an optimum depth of around 5 um, and opens dentinal tubules. In the second stage, hydrophilic primer coupling agents such as 2-hydroxyethyl methacrylate (HEMA), 4-methoxyethyl trimellitate anhydride (4-META), and glutaraldehyde are applied and penetrate into both the tubules and the decalcified intertubular dentin. The primer stabilizes collagen and facilitates the penetration of the bonding resins, which are bisphenol glycidyl methacrylate (bis-GMA) or urethane dimethacrylate (UDMA) monomers that are applied and polymerized.
There are two main types of conditioner components in current use. First there are the strong acids (phosphoric and citric acids). These acids nonselectively attack the dentin and the smear layer and expose the tubules (Fig 5-14). Other acids (10% maleic acid) are used to dissolve both inorganic and organic tissues, but they are less aggressive. The second type of conditioner is weak acids (eg, polyacrylic acid), which are used to dissolve only the smear layer without demineralizing dentin.
The layer of resin penetration is known as the **hybrid zone** and may extend from 1 to 5 µm deep (Fig 5-15). If there is a layer of decalcified dentin below the hybrid zone, it acts to weaken the bond due to the resin being absent. This can be avoided by limiting the depth of etching with shorter etching times or less concentrated acids. Excessive drying of the dentin layer before primer application is not desirable, because it leads to collapse of the surface collagen layer and reduces primer diffusion. Lateral penetration into peritubular dentin may also take place from resin that has penetrated first into the tubules. **Figure 5-16** shows a scanning electron micrograph (SEM) of a fractured hybrid layer with tags that have been plucked from the tubules during bond testing. Resin tags in tubules are not strongly anchored due to polymerization shrinkage and dentinal fluid.

**Fig 5-14** Dentin after deep etching with 20% phosphoric acid for 2 minutes. (Original magnification × 5,000.)

**Fig 5-15** Cross-sectional view of resin-etched bovine dentin interface showing resin (R), the hybrid zone (H), and demineralized dentin (DD). (From Wang and Nakabayashi, 1991. Reprinted with permission.)
Fig 5-16 Scanning electron micrograph of a fractured resin-dentin interface showing resin tags plucked from tubules. DD = demineralized dentin. (From Wang and Nakabayashi, 1991. Reprinted with permission.)

Bond-strength values of materials to enamel and dentin are given in Table 5-4. The laboratory bond-strength values to dentin vary significantly more than those reported for enamel. Therefore, bonding to dentin is less reliable.

<table>
<thead>
<tr>
<th>Bonding couple</th>
<th>Bond strength (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acid-etched enamel + resin (eg, bis-GMA)</td>
<td>16-20</td>
</tr>
<tr>
<td>Enamel + glass ionomer</td>
<td>5</td>
</tr>
<tr>
<td>Dentin + glass ionomer</td>
<td>1-3</td>
</tr>
<tr>
<td>Dentin + glass ionomer (light cured)</td>
<td>3-11</td>
</tr>
<tr>
<td>Enamel + NPG-GMA</td>
<td>2-5</td>
</tr>
<tr>
<td>Dentin + 4-META</td>
<td>2</td>
</tr>
<tr>
<td>Dentin + mordant ion + 4-META</td>
<td>18</td>
</tr>
<tr>
<td>Dentin + phosphate esters of bis-GMA</td>
<td>3-5</td>
</tr>
<tr>
<td>Acid-etched enamel + phosphate esters of bis-GMA</td>
<td>20</td>
</tr>
<tr>
<td>Dentin + maleic acid/HEMA + HEMA/bis-GMA</td>
<td>18</td>
</tr>
<tr>
<td>Acid-etched enamel + HEMA/bis-GMA</td>
<td>23</td>
</tr>
<tr>
<td>Dentin + isocyanate</td>
<td>4</td>
</tr>
<tr>
<td>Dentin + glutaraldehyde/HEMA</td>
<td>18</td>
</tr>
</tbody>
</table>

*Derived from McCabe (1990).
A major difference between commercial products is the number of components that need to be applied in sequence. Several multicomponent products contain an etchant (conditioning agent), primer, and bonding resin. According to CRA (2000), there are four types of dentin adhesive products:

Type 1: Etchant applied and rinsed off to remove smear layer; primer and adhesive applied separately

Type 2: Etchant applied and rinsed off to remove smear layer; primer and adhesive applied in a single solution

Type 3: Self-etching primer applied to dissolve smear layer and not rinsed off; adhesive applied separately

Type 4: Self-etching primer and adhesive applied as a single solution to dissolve the smear layer

Single-component products (Fig 5-17) include a gel etchant and a one-bottle component that contains the primer and resin (bis-GMA). This appears to be the direction for future developments since it is simpler than the multicomponent "chemistry sets."

Fig 5-17 A one-step dentin adhesive. A combination of the two components contains etchant, primer, and initiators. (One-Up Bond F; courtesy of J. Morita.)
**Acid-Etch Technique** The method of etching enamel with an acid to produce a roughened surface for resin bonding.

**Adhesion** Surface attachment of two materials (as opposed to cohesion, which is the bonding within a single material).

**Capillary Penetration** Movement of a liquid into a crevice or tube because of capillary pressure.

**Colloid** Material in which a constituent in a finely divided state that is invisible to the eye but capable of scattering light is suspended. Although colloids are not true solutions, many have long-term stability.

**Contact Angle** Angle formed between the surface of a liquid drop and the solid surface on which it rests. Usually designated by the Greek letter \( \theta \) and used as a measure of wetting.

**Hybrid Zone** A layer of dentin that contains resin. Produced by etching and resin diffusion.

**Hydrophobic** Water-repellent; showing a high contact angle.

**Isocaps** Isolated capillary bridges formed between solid surfaces.

**Lyophobic Colloid** An unstable colloid made from insoluble dispersed material in fine suspension.

**Micelles** Aggregated molecules formed in association colloids.

**Penetration Coefficient** Combination of the properties of viscosity, surface tension, and contact angle that promotes rapid capillary penetration.

**Sintering** Densification process in which solid particles are fused together, usually at high temperatures.

**Surface Energy** Extra energy that atoms or molecules on the surface of a substance have over those in the interior. The units are erg/cm\(^2\).

**Surface Tension** The surface energy of a liquid.

**Wetting** The spreading of a liquid drop on the surface of a solid.

**Discussion Questions**

1. Why is wetting by an adhesive so important for bonding to tooth structure or any other material?

2. Why is the strength of enamel and dentin a limiting factor in the adhesive bond?
strength?

3. What is the essential role that hydrophilic primer coupling agents play in the formation of the hybrid layer?

4. How could the geometry of a restoration affect the final bond strength to tooth structure?

**Study Questions**

(See *Appendix E* for answers.)

1. Why does surface tension exist in liquids?

2. How does wetting affect capillary penetration of a liquid?

3. How do absorption and adsorption differ?

4. Name three colloidal systems of importance to dental materials.

5. Which properties affect the penetration rate of liquids into capillaries?

6. In terms of innovation, would the bonding of restorative materials to dentin by etching and forming a hybrid layer be classified as a major, medium, or minor product improvement?

**Recommended Reading**


Dent 15:1094-1100.


Chapter 6. Polymers and Polymerization

Introduction

Polymers have a major role in most areas of dentistry. Their distinctive properties allow a range of clinical applications not possible with other types of materials. The most widely used impression materials (alginate, polyethers, polysulfides, and silicones) are polymers. A polymeric matrix with a particulate ceramic filler is the most commonly used anterior esthetic restorative material. Additional applications include denture teeth, cements, dies, provisional crowns, endodontic fillings, tissue conditioners, and pit and fissure sealants. However, the primary use of polymers in terms of quantity is in the construction of complete dentures and the tissue-bearing portions of partial dentures. This chapter introduces concepts related to polymer composition and properties and concludes with a discussion of those polymers currently used in the construction of complete and partial dentures, liners, and tissue conditioners.

Polymers

Composition

A polymer is a molecule that is made up of many units (poly = many; mer = unit). The oligomer is a short polymer composed of two, three, or four mer units. A mer is the simplest repeating chemical unit of which the polymer is composed, and is often the basis for naming the material. Thus, polystyrene is a polymer composed of styrene units.

Monomers (mono = single) are the molecules that unite to form a polymer, and the process by which this occurs is termed polymerization. If monomers of two or more different types are joined, copolymers are formed. Copolymers may be either random (mers do not appear in specific order) or block (large numbers of one type of mer appear arranged in sequence). Atoms along the length of any polymer are joined through strong, primary covalent bonds.

Molecular weight

The degree of polymerization is defined as the total number of mers in a polymer molecule. The molecular weight of a polymer molecule is the sum of the molecular weights of the mers of which it is made. Typical polymer molecules may be composed of thousands to millions of mers. Often, a distribution of molecular sizes is present in a material, and the reported molecular weight is the average molecular weight. The particular distribution is the result of conditions present during
polymerization. Variations in conditions have a pronounced effect on the properties of the final material.

**Spatial structure**

There are three basic spatial structures of polymers: linear, branched, and cross-linked (Fig 6-1). Linear and branched molecules are discrete but are bonded to one another through weak, physical bonds. Upon heating, the weak bonds break and the ability of the chains to then slide past one another results in a softened material. Upon cooling, the bonds reform and hardening occurs. Materials that are able to undergo this process are termed **thermoplastic**. Examples include polystyrene, polyvinyl acrylics, and poly(methyl methacrylate) (PMMA).

![Fig 6-1](image.png)

**Fig 6-1** Polymer chains. (a) A linear chain; (b) a branched chain; (c) copolymer chains, linear (upper) and branched (lower); (d) cross-linking chains.

Cross-linking results in the formation of a network structure of covalently bonded atoms; primary linkages occur between chains, and the polymer actually becomes a single giant macromolecule. The spatial structure that allows chain sliding upon heating is not present in cross-linked materials. Cross-linked polymers therefore do not undergo softening upon heating and are termed **thermosets**. Typical examples are silicones, cis-polyisoprene, bisphenol A-diacrylate, and cross-linked poly(methyl methacrylate).

**Properties**
Many factors affect the properties of polymers, including the chemical composition of the chain, its degree of polymerization, and the number of branches and/or cross-links between polymer chains. In general, longer chains and a higher molecular weight result in the polymer's increased strength, hardness, stiffness, and resistance to creep along with increased brittleness (Fig 6-2). Resin composites, for example, have a highly cross-linked matrix, in which a large number of strong covalent linkages between chains transforms the molecules into a rigid, very high-molecular-weight material. The resulting increased strength and stiffness contribute to the ability of this material to withstand occlusal stresses during function.

![Fig 6-2](relation_strength_polymer_molecular_weight.png)

**Fig 6-2** Relation between strength and polymer molecular weight. (From Mark, 1962.)

In contrast, elastomeric impression materials are composed primarily of individual coiled chains with just a few cross-links. This type of molecular structure permits the large-scale uncoiling and recoiling of chains that give these materials high flexibility.

The amount of crystallinity present in a polymer affects properties. Materials that are highly crystalline have atoms with a very regular arrangement in space and are stronger, stiffer, and absorb less water than do noncrystalline materials. Few dental polymers are crystalline. Most are amorphous, meaning that the atoms of which they are composed have irregular arrangements in space. Amorphous polymers are often called glassy polymers.

Small plasticizer molecules, when added to a stiff uncross-linked polymer, reduce its rigidity. When small molecules surround large ones, the large molecules are able to move more easily. A plasticizer therefore lowers the glass-transition temperature ($T_g$) of the polymer, so a material that is normally rigid at a particular temperature may become more flexible. The glass-transition temperature is the temperature at which a polymer ceases to be glassy and brittle and becomes rubberlike. The temperature of a polymer, as shown in Fig 6-3, has a strong effect on its strength properties.
Fig 6-3 Effect of temperature on the tensile properties of a polymer. The temperature below which brittle failure occurs is called the glass-transition temperature. (From Berry and Bueche, 1962.)

Finally, during polymerization a volumetric decrease occurs, resulting in shrinkage (up to 21% for unfilled acrylic resins, 6% for denture resins, 1% to 3% for resin composites) and the production of internal stresses. A change in shape, often called warpage, may occur when the polymer is reheated. Additionally, polymers have varying abilities to absorb water. A small amount of expansion may occur during this process.

**Polymerization**

*Introduction*

Most polymerization reactions are of two types: **addition polymerization**, in which no by-product is formed, and **condensation polymerization**, in which a low-molecular-weight by-product such as water or alcohol is formed. Materials that set by addition polymerization include poly(methyl methacrylate), used in the construction of dentures, and bis-GMA, a common component of the matrix of resin composites. Materials that set by the condensation mechanism include polysulfide rubber and some silicone rubber impression materials.

The three stages in the free-radical addition polymerization reaction are described in the following subsections. They may be accelerated by heat, light, or small amounts of peroxides.
Initiation

The initiation step involves the production of free radicals, which will encourage a polymer chain to begin growing (Fig 6-4). Free-radical molecules have chemical groups with unshared electrons. In chemically activated systems, free radicals are generally produced by the reaction of an organic peroxide initiator and an amine accelerator. In light-activated systems, the scission of camphorquinone results in the production of two molecules with one unshared electron each. Whatever the means of production, the free radicals attack the double bonds of available monomer molecules, resulting in the shift of the unshared electron to the end of the monomer and the formation of activated monomer molecules.

1. Initiation

\[
\text{Benzoyl peroxide} \rightarrow \text{Free radicals (R)} + \text{carbon dioxide}
\]

2. Propagation

3. Termination

![Fig 6-4 Three stages of addition polymerization of methyl methacrylate.](Image)
Propagation

Activated monomers attack the double bonds of additional available monomers, resulting in the rapid addition of monomer molecules to the free radical. This second stage, propagation, continues as the chain grows in length.

Termination

Termination of the growing free radical may occur by several mechanisms and can result in the formation of branches and cross-links.

Small amounts of inhibitors, such as hydroquinone, may be added to the monomer to increase storage life. Hydroquinones react with free radicals, thereby decreasing the rate of initiation.

Denture Base Polymers

Introduction

The polymeric denture base can consist of either a simple stiff base on which the teeth are arranged, or a sandwich of stiff base and a resilient liner to provide greater retention and comfort. When the tissue underlying a loose denture is traumatized due to the constant motion of the hard plastic over the mucosa, a viscoelastic gel known as a tissue conditioner can be molded onto the fitting surface of the denture in situ so the tissue can heal and an accurate impression of the untraumatized fitting surface can be taken prior to making a new, better-fitting denture. A classification is shown in Fig 6-5.

![Denture Base Polymers Classification](image)

**Fig 6-5** A classification of denture bases, liners, and tissue conditioners.
Advances in denture design have been mediated by the materials available at the time. In the 1800s the art of hand carving ivory and wooden denture bases resulted in dentures retained by mechanical devices such as springs. Goodyear's invention of vulcanized rubber, vulcanite, in 1839 provided not only a thermoplastic material that could be molded accurately, but also one that did not biodegrade and was strong enough to withstand masticatory forces for many years. Vulcanite's intrinsic dark color and opacity meant that the translucency and reflectivity of living mucosa was impossible to mimic.

With the commercial availability of man-made polymers in the early 1930s came an opportunity to apply some of the optically brilliant polymers to dentistry. It was the need for custom fit that delayed their use, however, since polymerization of the free monomer, with its inherent massive volume shrinkage, did not lend itself easily to the vulcanizing techniques used at the time. It was the adoption of the dough technique first described in the mid-thirties that made the use of acrylics in dentistry possible. In the dough technique, a liquid component (monomer) is mixed with a powder component (polymer). The monomer wets the polymer to a dough-like consistency, which is packed into the mold prior to polymerization. Adoption of the new denture bases was rapid in America. However, in Europe, change was forced upon the profession by rubber shortages during World War II. By the end of the war the use of vulcanite for dentures had almost ceased. After the war, resins developed for aircraft production and the burgeoning plastics industry were offered for use as denture base materials, but the simplicity of the dough technique and the lifelike results have sustained acrylics as the market leader to the present day. Table 6-1 gives a comparison of current denture base materials.

<table>
<thead>
<tr>
<th>Material</th>
<th>Advantages</th>
<th>Disadvantages</th>
<th>Ideal properties</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heat cured</td>
<td>Good appearance, high glass-transition temperature, ease of fabrication, low capital costs, good surface finish</td>
<td>Free-monomer content or formaldehyde can cause sensitization, low impact strength, low stiffness</td>
<td>Good appearance, high flexural strength, high impact resistance, high stiffness</td>
</tr>
<tr>
<td>Heat cured, rubber reinforced</td>
<td>Improved impact strength</td>
<td>Reduced stiffness</td>
<td>Low free-monomer content, good adhesion with teeth and liners</td>
</tr>
<tr>
<td>Heat cured, fiber reinforced</td>
<td>High stiffness, very high impact strength, good fatigue life, polypropylene fibers</td>
<td>Carbon and Kevlar fibers, poor color, poor surface</td>
<td>Low solubility, low water uptake, dimensional stability, dimensional accuracy</td>
</tr>
</tbody>
</table>
Composition and manufacture

Heat-cured acrylic

The bulk of denture base acrylic is supplied these days in the form of a free-running powder and a liquid. Originally the powder was produced by grinding blocks of PMMA. However, it was soon found that smoother, more consistent doughs resulted from the use of a spherical bead polymer. By suspending monomer liquid in water with the aid of either a surfactant or water-soluble polymer, the rate of polymerization can be well controlled by the cooling action of the surrounding water. The additives required to make the beads dough easily are fortunately very soluble in monomer globules and relatively insoluble in water. The globules can be thermally polymerized without a catalyst; however, benzoyl peroxide is usually added, partly to act as a catalyst in the polymerization of the beads, and partly so the beads become the source of the peroxy free radical during the polymerization of the dough in the dental flask (Fig 6-6).
To assist dough formation, a plasticizer is also incorporated into the bead polymer. For many years this was an external plasticizer, that is, one that resided between the polymer chains rather than being chemically attached to them. Any inert, nontoxic organic molecule will suffice; dibutylphthalate was used for many years. Today, internal plasticizers are used instead, containing various methacrylate or acrylate monomers. They locally soften the bead and allow the monomer to diffuse more rapidly into the bead during the dough stage. Pigment can be added to the bead either during polymerization or after, by ball milling. By this method, the manufacturer produces a PMMA bead mixture having a fairly wide molecular weight distribution with an average molecular weight on the order of 1 million. The all-important doughing characteristics of the bead are governed by particle size distribution, molecular weight distribution, and plasticizer content. The highest molecular weight distributions and lowest plasticizer content are favored, because they result in better physical and mechanical properties in the cured denture base.

The monomer used to form the dough is largely the same as that used to make the methyl methacrylate beads. Methyl methacrylate quickly diffuses into the polymer beads on contact, causing them to swell, and extracting some low-molecular-weight polymer into the monomer trapped in the interstices between the beads. As the beads swell, entanglements occur between the juxtaposed beads, and the bead-monomer mixture becomes a cohesive gel. The beads never dissolve completely (Fig 6-7), although the monomer infiltrates well into the core of each bead. In the swollen state, benzoyl peroxide can diffuse from the bead into the interstices, where later it will initiate the curing of the dough. In addition, cross-linking agent molecules, capable of diffusing into the beads, are present in the monomer phase.
The cross-linking agent confers two useful properties on the cured gel. It reduces the denture base's solubility to organic solvents, and it reduces the tendency of the denture base to craze (form precracks) under stress. However, cross-linking agents also are very important from a practical point of view because they help to keep the denture-making apparatus simple. For a monomer to be converted to a solid polymer, the polymer chain produced must achieve a certain minimal length. Polymers having molecular weights below 5,000 are liquid and viscous; resilient polymers need to achieve a minimum molecular weight of about 150,000. Without a cross-linking agent the flask would have to be airtight and the monomer flushed with nitrogen to achieve the molecular weight required without being inhibited by air. Free-monomer levels would then be unacceptably high, and the denture would have a lower stiffness level and a greater tendency to creep. The cross-linking agent accelerates the increase in the curing system's molecular weight and combats the effects of oxygen inhibition. However, excessive levels of cross-linking agent in the monomer result in denture bases that are brittle. The most common cross-linking agents are dimethacrylates, either ethylene glycol dimethacrylate or 1,4-butylene glycol dimethacrylate.

High-impact acrylic

High-impact acrylic denture base is also made by the heat-cured dough method; impact resistance arises from the incorporation of a rubber phase into the beads during their suspension polymerization. Certain rubbers will dissolve in methyl methacrylate monomer, notably copolymers of butadiene with styrene and/or methyl methacrylate. The rubber remains soluble in the monomer globule until the polymer content of the globule becomes too high and the rubber begins to precipitate out. The nature of this precipitation is complicated by the fact that some of the growing chains of PMMA may have become grafted to the butadiene rubber. This results in what is known as a phase inversion, resulting in dispersion throughout the bead of tiny islands of rubber...
containing small inclusions of rubber/PMMA graft copolymer (Fig 6-8). Why these inclusions improve the impact strength of the cured denture base will be discussed later. There are, however, many patents that describe the formation of the beads, and dentistry uses examples that are either beads with uniformly distributed rubber inclusions, or beads that have a core of rubber-included polymer covered by an outer shell of conventional polymer to give a more conventional dough formation. Beads that have no shell often gel very quickly and may entrap air as a result. The monomer used to get high-impact beads differs from conventional monomers in that it contains either very little or no cross-linking agent. Fortunately, the inclusion of rubber does have a craze-inhibiting effect, as will be explained later.

![Fig 6-8](image.png)

**Fig 6-8** Electron micrograph of high-impact denture base showing the size and shape of the polystyrene-butadiene rubber inversion phase.

**Autopolymerizing denture base**

The autopolymerizing, or pour-type, denture base is chemically similar to the heat-cured denture base except that a reducing agent is added to the monomer. The reducing agent is usually a tertiary aromatic amine, although barbituric acid derivatives also have been used. The reducing agent reacts with the benzoyl peroxide at room temperature to produce peroxy free radicals, which initiate the polymerization of the monomer in the denture base. There is wide variation among manufacturers in the molecular weights of the polymers in the beads. Some have average molecular
weights as low as 190,000. Cross-linker concentrations vary greatly in the monomer liquid, from 0% to 9%; interestingly, excess cross-linker is associated with high creep. The size, molecular weight, and plasticizer content are balanced to give a high penetration of monomer into the bead without too early an increase in viscosity of the mix to allow pouring of the acrylic into the mold and good wetting of the plastic teeth. This compromise is difficult to achieve and often results in high residual free-monomer contents and low cross-link densities. Manufacturers' attempts to achieve the best compromise result in the wide ranges of molecular weights and cross-linker concentrations found in materials, which inevitably result in large differences in physical and mechanical properties between the various products, especially where creep is concerned.

Injection-molded plastic

Injection-molded plastics have the advantage of consistent molecular weight, but the disadvantage of capital equipment costs, low craze resistance, and difficulties associated with attachment of teeth to the denture base. The plastics still offered for use as injection-molded denture base acrylic are polycarbonate and nylon. They represent a very small fraction of the market, although they offer a real alternative to metal dentures for patients sensitized to conventional methacrylate or to nickel or cobalt.

**Acrylic.** Acrylic is supplied as granules of low-molecular-weight (MW = 150,000) linear PMMA, with narrow molecular weight range and only a small amount of residual free monomer. Note there is no cross-linking, as this would increase the melt viscosity during molding. Plasticization is low and often results in stiffnesses slightly in excess of conventional heat-cured denture base, despite the low molecular weight.

**Polycarbonate.** This tough plastic is supplied as granules but is not suited to injection into damp molds. It has a high melt viscosity and may depolymerize explosively if overheated in the presence of water. Again, absence of cross-linking results in poor solvent resistance and craze resistance. The high melt viscosity exacerbates problems of tooth attachment.

**Nylon or polyamides.** This is a family of condensation polymers that result from the reaction of a diacid with a diamine to give a variety of polyamides whose physical and mechanical properties depend on the linking groups between the acid or amine groups. The first dental use of nylon was not a success because of the excessive water absorption of the type chosen, which resulted in excessive creep and some biodegradation. More recent work on glass-reinforced nyons with much lower water absorptions (eg, nylon 66) has produced more encouraging results. These nyons are either filled with specially coated glass beads or chopped glass fibers. The glass fibers increase the stiffness of the nylon to about that of a conventional heat-cured denture base from a stiffness of half that when only glass-bead reinforcement is used. Glass-fiber reinforcement should be used with care, and patients should be warned not to abrade the fitting surface so as to avoid exposing irritation-causing fibers.

Light-activated materials

In recent years, a light-activated system has come on the market and is now used for
many prosthetic applications. This material consists of a urethane dimethacrylate matrix with an acrylic copolymer and has a microfine silica filler. It is supplied in premixed sheet or rope form. A baseplate is made by adapting the material to a cast and polymerizing in a light chamber at 400 to 500 nm. Teeth are added to the base with additional material followed by a second light exposure. The system eliminates the need for flasks, wax, boil-out tanks, packing presses, and heat-processing units required for the construction of conventional dentures. The manufacturer claims a significant time savings in both the dental office and the laboratory.

**Fabrication and its effect on physical and mechanical properties**

When fabricating a denture base from polymers—by whatever means—certain physical and mechanical properties of the final polymer are important. The cured polymer should be stiff enough to hold the teeth in occlusion during mastication and to minimize uneven loading of the mucosa underlying the denture. The polymer should not creep under masticatory loads if good occlusion is to be maintained. The polymer must have sufficient strength and resilience to withstand not only normal masticatory forces, but sudden high stress caused by impact forces. The material should not deteriorate in the aqueous oral environment, and crazes should not form due to attack by solvents present in food, drinks, or medicaments. The cured polymer should be biologically inert and slow to foul when in contact with oral flora.

During fabrication of a denture the physical and mechanical properties mentioned here can be influenced by cure conditions and choice of materials. Each cure cycle or fabrication technique is a compromise that attempts to optimize the properties thought important for a given application. For an allergic patient, low free-monomer content may be thought more important than stiffness. For a patient requiring a soft lining, stiffness is very important if the reduced cross-sectional area of the denture is not to cause stability or loading problems.

**Heat-cured denture base**

In the previous discussion of heat-cured denture base acrylic, the mechanism of doughing was described. When manipulating the dough prior to packing, the operator is aware of various stages of gelation: the initial melting of the beads with the monomer; the sandy stage; the formation of first entanglements as the outer layers of the bead swell; and, when the dough is pulled, the formation of stringy threads that can be drawn out as low-molecular-weight components dissolve into and thicken the interstitial monomer. In the last stage, the dough becomes elastic as monomer penetration reaches the core of the beads, plasticizing them and lowering their glass-transition temperature from an initial 125°C to well below room temperature; the beads have become rubber. Curing the dough before the monomer has diffused to the core of the beads may result in reduced flexural strength and a tendency for cracks to propagate along lines of weakness between the cross-linked interstitial phase and the linear, more pliable beads. Allowing the monomer and cross-linker to diffuse to the center of the beads results in a more homogeneous distribution of stress, and the formation of a tough, three-dimensional network within the existing amorphous bead polymer to form what is known as an **interpenetrating polymer network** (IPN) (see Fig 6-7).
The curing cycle is designed to raise the temperature to a point at which (1) sufficient benzoyl peroxy radicals are produced to overcome the scavenging effect of oxygen, and (2) polymer chains form by free-radical addition polymerization. Too rapid a rise in temperature produces large numbers of radicals—a radical avalanche—and, as a result, many growing polymer chains. These chains collide either with other radicals or with polymer chains, producing an increase in branching and cross-linking of the interstitial polymer. This in turn reduces toughness. The polymerization reaction itself is exothermic; if the rate of reaction is too high, heat builds up in the dough until the boiling point of the monomer is exceeded. Porosity of the final denture base results, with a subsequent loss of strength and esthetics and an increase in the possibility of fouling.

Slow cures result in much tougher denture bases, producing fewer cross-links and branches, and having a higher overall molecular weight between cross-links because fewer polymer chains grew at any one time. Free-monomer content is often lower also, because the steadier rise in internal viscosity of the curing polymer allows the monomer easier access to the growing free radicals. The cross-linker is more completely polymerized in heat-cured systems; this results in significantly lower creep values due to removal of the plasticizing effect of unreacted pendant cross-linker groups.

The shrinkage that occurs when a liquid monomer is converted to a solid polymer can be largely compensated for by keeping the curing dough under compression. There are, however, hybrid systems that deliberately start the polymerization of the dough on one side of the mold, while forcing uncured dough into the mold on the opposite side. As curing progresses across the mold, the mold space is kept filled, with a resulting improvement in dimensional accuracy. The advantage of this system is that there is less tendency to raise the bite by overpacking the mold with dough, which can occur with conventional flasks.

Heat-cured systems have one great advantage over autopolymerized and injection-molding methods: an increased rate of monomer diffusion at the higher temperature. This is most beneficial when acrylic teeth are used, for it leads to better wetting of the teeth by the dough and the formation of chemical welds between the teeth and denture base. Conversely, an increased temperature of cure can also result in the annealing of stresses that build up in the structure due to polymerization shrinkage. If this stress is not released, it can act as the foci for crazes or distortions caused by overzealous cleaning regimes.

Autopolymerizing acrylic

The pour technique for dentures, originally developed during the 1960s, has lost much of its popularity because of problems that stem from incorrect processing. In this technique, the acrylic is mixed to a liquid consistency, then poured into a sprued mold that consists largely of reversible hydrocolloid. The fitting surface of the mold consists of the plaster model itself; the acrylic teeth occupy their positions in the agar mold in the same way they do in a conventional plaster mold.

The pour-type mold itself has design weaknesses. The gelatinous agar cannot grip the
teeth as easily as the rock-hard plaster, and hence there is a greater tendency for the teeth to be displaced during the pouring of the acrylic. Prior to being placed in the mold, the teeth themselves have been part of the wax-up; any wax remaining on the teeth will prevent the monomer from wetting their surfaces. This problem is far less common when solution and diffusion of the wax can occur at the elevated temperatures of the heat-curing process.

The use of a hydroflask to produce an increased atmospheric pressure around the mold has two main advantages: (1) Most important, porosity caused by monomer boiling is prevented by the simple expedient of raising its boiling point; and (2) air included during mixing is compressed, raising the density of the cured resin and improving its transverse strength. However, the technician can do little to reduce free-monomer content to achieve high toughness, because this is built into the formulation when the manufacturer chooses the powder/liquid ratio, the cross-linker content, and the accelerator/catalyst ratio. In general, the creep of these products is greater than that of heat-cured acrylics.

Injection molding

The technician has little leeway when using injection-molded plastics. The mold should be dry to prevent the generation of steam during molding. Patience is required to ensure the melt has reached the right temperature and cools sufficiently after molding. Inadequate spruing will lead to underfilled molds, as can underheating the melt; overheating the melt can cause explosions, especially when polycarbonate is injected into moist molds.

Injection moldings rely almost totally on mechanical forces to retain the teeth. Low melt temperatures will cause strong forces to be put on the teeth during the injection phase and may dislodge some molars, even from plaster molds.

Depolymerization or oxidation from overheating the melt can result in porosity, loss of strength, color changes, and increased fouling.

Light-activated materials

Light-activated materials compare well to conventional heat-cured materials in terms of impact strength and hardness, but have a considerably lower elastic modulus. A denture constructed of light-cured material would therefore be expected to deform elastically to a greater extent than a heat-cured denture under the forces of mastication. However, transverse strength—a measure of the load required to fracture a thin strip of material in the transverse direction—is just slightly lower than that of the conventional material.

As a consequence of the higher-molecular-weight oligomers used in light-cured systems, polymerization shrinkage is smaller, about 3%, rather than the 6% shrinkage found in conventional systems. One study showed that denture bases processed by visible light fit better than conventional or quick heat-cured resins. Ideally, a total lack of polymerization shrinkage would allow the best fit.

Since light-activated materials contain no methyl methacrylate monomer, they may be
considered for use in those patients who have demonstrated a sensitivity. The formulation of light-activated denture bases contains a copolymer of urethane dimethacrylate and an acrylic resin along with silica fillers. Blue light is used to polymerize thin sheets of the plastic raw material in a light chamber.

**Future improvements in polymeric denture bases**

The properties of polymeric denture bases that are closest to being offered as improved features of commercial products are radiopacity, impact strength, and stiffness.

Radiopacity in denture bases would be a desirable attribute. Denture wearers, be they motorists, members of the security forces, or athletes, can endure serious complications from relatively minor traumatic incidents if their denture fractures and a portion of it either is ingested into the lungs or intestines or, in more serious incidents, is driven through the skull into the brain pan. Fragments of radiolucent denture base are difficult to find even when sophisticated ultrasound techniques are used, and their presence is often suspected only after a secondary infection sets in.

The use of radiopaque salts and fillers often reduces esthetic properties and strength, and organometallic components have often proved too toxic for use. Bromine-containing organics can give good esthetics but often lack the heat stability necessary for heat processing or have to be added in such high quantities that the bulky bromine groups overplasticize the resulting acrylic denture base, causing creep, water adsorption, and stiffness problems. However, by phase-separating a bromo-polymer additive within the bead phase, it has been shown that sufficiently high glass-transition temperatures of 110°C and stiffnesses of 2.0 GPa (290,000 psi) can be achieved, while at the same time preserving esthetic properties and achieving high levels of radiopacity.

High-impact denture base materials can be prepared using inversion-phase separated polymer beads. However, a 50% improvement in impact strength (2.1 J/m) could be achieved, in combination with good esthetics and radiopacity, by the formation of a three-phase bead consisting of (1) PMMA, (2) styrene butadiene rubber, and (3) the poly(2,3-dibromopropyl methacrylate). Processing conditions must be well controlled.

The availability and quality of high-modulus fibers is improving quickly, as is the variety of materials from which the fibers are made. Early experiments with glass fibers resulted in failure because of the irritating nature of the fibers that protruded from the finished surfaces. Carbon fibers had no such irritant effects and greatly increased impact strength and flexural stiffness of the denture base for little material expense. However, the carbon fibers are black, so their use must be restricted to the lingual aspects of a denture. Kevlar fibers (poly-p-phenylene terephthalamide) have stiffnesses of 90 GPa (13 million psi), are straw-colored, and are not easy to pack. They can, however, greatly enhance the mechanical properties of the denture. As with carbon fibers, they are disappointing esthetically and need to be restricted to lingual aspects of the denture, although they can be extended to the midline of the teeth without being noticeable. This is an advantage, for if this were done with carbon fibers it would create a black shadow beneath the teeth in the plastic gum work.
Incorporation of the fibers into dentures produces conventional dentures that can withstand uncommonly rigorous treatment. When the fibers are combined with bis-GMA, the flexural strength is such that they do not break in the conventional three-point flexural test apparatus, yet they have Young's moduli of 30 GPa (4 million psi), a figure comparable with ceramics. Lingual bars for partial dentures could be made using such materials, and, perhaps, with some changes in partial denture design, much less obtrusive polymeric partial dentures could be fabricated with only the clasps being made of metal.

**Permanent Soft Lining Materials**

*Introduction*

Permanent soft lining materials are resilient polymers used to replace the fitting surface of a hard plastic denture, either because the patient cannot tolerate a hard fitting surface or to improve retention of the denture. Because the lining is soft, its dimensional stability is important, as are its durability and resistance to fouling. However, because by definition soft lining materials are above their glass-transition temperature when in the mouth, such physical phenomena as water absorption, osmotic presence of soluble components, and biodegradability play a greater role in the clinical success of a liner than they do in the glassy polymers used as denture bases.

Acrylics and silicones are the two main families of polymers used commercially as soft liners, though other rubbers have been used in limited clinical experiments. Table 6-2 gives a comparison of soft lining materials.

**Table 6-2** A comparison of soft liners and tissue conditioners

<table>
<thead>
<tr>
<th>Material</th>
<th>Advantages</th>
<th>Disadvantages</th>
<th>Ideal properties</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acrylic</td>
<td>High peel strength to acrylic denture base</td>
<td>Poor resilience, loses plasticizer in time, some buckle in water, some can be polished if cooled, reasonable resistance to damage by denture cleansers</td>
<td>High resilience, unaffected by aqueous environment and cleansers, good bond to denture base, good abrasion resistance</td>
</tr>
<tr>
<td>Silicone (RTV)</td>
<td>Resilience</td>
<td>Low tear strength, low bond strength to dentures, attacked by cleansers, buckle in water, poor abrasion resistance</td>
<td>Biocompatible, antifouling properties, good dimensional stability</td>
</tr>
<tr>
<td>Silicone heat cured</td>
<td>Resilience</td>
<td>Low tear strength, poor abrasion resistance</td>
<td></td>
</tr>
</tbody>
</table>

Materials:

- **Acrylic**
- **Silicone (RTV)**
- **Silicone heat cured**
aqueous environment and cleansers than RTV tissue conditioners.

<table>
<thead>
<tr>
<th>Rheological and viscoelastic properties</th>
<th>Low cohesive strength</th>
<th>Low under constant force</th>
<th>Rheological and viscoelastic properties almost ideal</th>
</tr>
</thead>
<tbody>
<tr>
<td>Can be applied chairside</td>
<td>Affect by cleansers</td>
<td>Resilient at high rates of deformation</td>
<td></td>
</tr>
<tr>
<td>Dentures fit well</td>
<td>Alcohol can sting inflamed mucosa</td>
<td>Remain viscous for several days</td>
<td></td>
</tr>
<tr>
<td>Can record freeway space</td>
<td></td>
<td>Have a high tack to aid retention to denture base</td>
<td></td>
</tr>
</tbody>
</table>

**Acrylic soft liners**

The acrylics consist of either highly plasticized intrinsically glassy polymers or soft acrylics that have a natural glass-transition temperature at least 25°C less than that of the mouth. The plasticizer used to soften the acrylic can either be unbound to the acrylic and hence free to diffuse out during use, resulting in a loss of resilience, or it can be reacted into the cured matrix of the acrylic. The latter method is preferred because it should increase the clinical life of the soft liner; unfortunately, in practice, such acrylics are hard to formulate.

The reactive plasticizer often has a much lower rate of polymerization than the acrylic monomer. The result is a form of phase separation that leads to an uneven uptake of water by the soft liner. Water accumulates in the plasticizer-rich phase, and soluble impurities in the polymer create an osmotic pressure that causes it to swell and distort. Therefore, although such internally plasticized acrylics have been produced commercially, they have often been withdrawn after distortion problems have been noted in practice, often due to insufficient curing of very finely balanced formulations.

The plasticized acrylics are based on copolymer beads consisting mainly of ethyl methacrylate; both n- and isobutyl methacrylate can be used, as can 2-ethoxyethyl methacrylate. However, the latter monomer is used mainly in the liquid component. The beads are copolymerized with acrylates, which, in general, have much lower glass-transition temperatures than their methacrylate homologues, but unfortunately have very unpleasant odors. So that the beads are free-flowing, the bead polymers have glass-transition temperatures slightly above room temperature. The monomer usually contains the plasticizer, which is a large phthalate ester. The monomer must swell sufficiently for the plasticizer to enter the beads. The plasticizer is then trapped inside the beads as the monomer polymerizes and the mean free path between the polymer chains decreases. The monomer can be methyl methacrylate, although its $T_g$, even when plasticized, is rather high; n- or isobutyl methacrylate is preferred because it has a much lower $T_g$ when polymerized. Some manufacturers use isobutyl methacrylate to produce a balance of properties such that in ice water the liner can be polished like denture base acrylic, while the liner remains resilient in the mouth.
Water can, of course, be used as the plasticizer in hydrophilic liners. This was the principle behind the hydroxyethyl methacrylate soft liners. However, the water is not present when the denture is fabricated, and its uptake leads to a swelling of the liner, which must be compensated for. Water-swollen liners also allow ions into their matrices, which can subsequently crystallize and thus harden the lining as well as cause osmotic pressure effects, as in the polymerizable plasticizers.

Silicone soft liners

The silicones used as soft linings can be divided into two types: room-temperature vulcanizing (RTV) and heat curing. The resilience of silicones makes them at first seem to be the ideal soft lining materials. However, silicones have poor tear strength, no intrinsic adhesion to acrylic denture base, and, if not properly cured, a tendency to osmotic pressure effects. The RTV silicones' greatest drawback is their lack of adhesion, which is especially a problem around the edges of the attachment between acrylic and silicone. Heat-cured silicones have in their formulation a siloxane methacrylate that can polymerize into the curing denture base and into the heat-cured addition silicone. The RTV silicones use a condensation cross-linking system based on organo-tin derivatives such as those used in impression rubbers. Their degree of cross-linking is lower and their serviceability is low as a result, with frequent reports in the literature of swelling and buckling during use and excessive sensitivity to denture cleansers. The rupture strength of some RTV silicones is known to deteriorate considerably when exposed to water for long periods. The heat-cured silicones achieve a greater degree of cross-linking and have much longer clinical lifetimes.

Temporary Soft Liners and Functional Impression Materials

Temporary soft liners, or tissue conditioners, need only survive in the mouth for a few weeks—although some are so well formulated as to remain resilient and in place for many months. However, it is their viscoelastic properties that are important, specifically their ability to flow under masticatory and linguistic forces, spreading the load on the mucosa evenly. When first mixed they flow easily, recording such voids as mean freespace. They soon become highly viscous, however, and thereafter only respond to persistent forces, such as changes in the shape of the mucosa beneath the denture. In this way swollen mucosa traumatized by ill-fitting dentures can recover while the denture, with its tissue conditioner lining, adapts to any lack of fit driven by masticatory forces.

The materials used for the temporary soft liners were many when the technique was first developed, and included many nontoxic, puttylike materials, such as plasticine and chewing gum. Modern materials are exclusively acrylic gels. An acrylic gel can be made by mixing swellable acrylic beads with alcohol. Poly(methyl methacrylate) is unsuitable for this purpose, although poly(ethyl methacrylate) or its copolymers with acrylates have proved most popular. To maintain the softness of the gel in the mouth, plasticizer is added to the alcohol, which diffuses into the polymer beads and lowers their glass-transition temperature to well below that of the mouth. The beads are made of low-molecular-weight polymer and have a high tack when swollen. This has the advantage of increasing the cohesive strength of the gel and causing it to be well retained by the denture base acrylic.
There is no polymerization or curing reaction involved in the setting of the gel, just the entanglement of outer polymer chains of juxtaposed beads. The rate of gelling is increased by lowering the molecular weight of the bead, reducing its size, or increasing the amount of acrylate in its copolymers. The alcohol content of the liquid also can be used to control the rate of gelling, as can the size of the plasticizer molecule; the more alcohol used, the faster the rate of gelling. However, because the alcohol diffuses out of the gel and is only partially replaced by water, high-alcohol-content gels tend to harden much faster than others. Practitioners should be aware of the alcohol content of these products, first because they sting when initially inserted, and second because they give false positive results if the patient is given a breathalyzer test.

**Glossary**

**addition polymerization** A polymerization process involving free radicals in which no by-product is formed as the chain grows.

**condensation polymerization** A polymerization process in which a by-product, such as water or alcohol, is formed as the chain grows.

**copolymer** A polymer consisting of two or more types of mers, or units, joined together.

**crazing** Minute surface cracks on polymers; precursors to crack growth and subsequent failure of the material.

**cross-linking agent** A monomer having two or more groups per molecule capable of polymerization. When polymerized, each active group is capable of incorporation in a growing polymer chain, causing either a loop in the chain or a cross-link between two chains.

**degree of polymerization** The total number of mers in one polymer molecule.

**denture base** Materials used to contact the oral tissues and support artificial teeth.

**glass-transition temperature (T_g) (softening temperature)** The temperature at which the polymer ceases to be a glass (ie, fractures in a brittle manner) and becomes a rubber or leather (ie, tends to permanently deform under a load too small to cause fracture).

**glassy polymer** An amorphous polymer that behaves as a brittle solid.

**hydrophilic liner** A soft liner that is readily wet by water.

**interpenetrating polymer network (IPN)** A combination of two polymers in network form, at least one of which is synthesized and/or polymerized in the immediate presence of the other. An IPN can be distinguished from simple polymer blends, blocks, and grafts in two ways: (1) An IPN swells but does not dissolve in solvents; and (2) creep and flow are suppressed.
**monomer** The basic repeating chemical molecule of a polymer (eg, polyethylene is a long chain of ethylene monomer units).

**molecular weight** The sum of the molecular weights of the mers of which the polymer is made.

**oligomer** A polymer made up of two, three, or four monomer units.

**phase inversion** The inversion of the internal and external phases in an emulsion, eg, the change of an oil-in-water emulsion to a water-in-oil emulsion.

**plasticizer** A small molecule that, when added to a polymer, lowers its glass-transition temperature and increases the rate at which solvents penetrate the polymer.

**polymer** A molecule made up of thousands or millions of repeating units. Polymers may be linear, branched, or cross-linked.

**polymerization** The process by which monomers unite to form a polymer.

**soft lining material (soft liner)** A soft polymer used as a thin layer on the tissue-bearing surface of a denture.

**thermoplastic** A polymer that softens upon heating and rehardens upon cooling.

**thermoset** A polymer that is not able to undergo softening upon heating.

**tissue conditioner** A soft liner used to treat traumatized mucosa.

**Discussion Questions**

1. Why is higher impact strength an important advantage for denture base materials?

2. How does the heat-curing rate affect the porosity and strength of acrylic denture bases?

3. Although cross-linking can improve mechanical properties, how can it lead to problems in the bonding between acrylic teeth and the denture base?

4. Why do acrylic polymers still dominate the denture base market?

**Study Questions**

*(See Appendix E for answers.)*

1. What are the main types of denture base materials used?

2. What are the ideal properties of a denture base material?

3. What are the advantages and disadvantages of heat-cured acrylic denture base materials?
4. What are the advantages and disadvantages of auto-cured denture base materials?

5. What are the advantages and disadvantages of rubber-reinforced denture base materials?

6. What are the advantages and disadvantages of fiber-reinforced denture base materials?

7. What are the advantages and disadvantages of injection-molded denture bases?

8. What are the advantages and disadvantages of light-activated denture base materials?

9. What are the materials currently being used as soft liners?

10. What are the advantages and disadvantages of RTV silicones?

11. What are the advantages and disadvantages of heat-cured silicones?

12. What are the advantages and disadvantages of acrylic soft liners?

13. What are the ideal properties of tissue conditioners?

**Recommended Reading**


Chapter 7. Impression Materials

Introduction

Impression materials are used to make replicas of oral structures. All impression materials must be in a plastic or fluid state while the replica is being made. Physical change, chemical reaction, or polymerization convert these fluid materials into either elastic or nonelastic (i.e., plastic or brittle) negative replicas of the soft and/or hard tissues of the mouth. A model or cast material (e.g., high-strength stone) is poured into the impression and, upon setting, produces a positive impression of the tissues of interest. Nonelastic materials include impression plaster, impression compound, and zinc oxide-eugenol. Elastic materials in current use are agar hydrocolloid, alginate, polysulfide, condensation silicone, addition silicone, and polyether. These materials will be discussed separately. A classification key is shown in Fig 7-1.
The American Dental Association's (ADA's) Council on Dental Materials, Instruments, and Equipment is responsible for developing and disseminating specifications for dental materials, instruments, and equipment. The American National Standards Institute (ANSI) has made the Council the Administrative Secretariat of the American National Standards Committee MD156 for Dental Materials, Instruments, and Equipment. Specifications relating to dental impression materials can be found in published documents listed in Table 7-1.

**Table 7-1** ADA/ANSI specifications for dental impression materials

<table>
<thead>
<tr>
<th>Material</th>
<th>ADA/ANSI Specification No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dental impression compound</td>
<td>3</td>
</tr>
<tr>
<td>Zinc oxide-eugenol impression paste</td>
<td>16</td>
</tr>
<tr>
<td>Dental agar impression material</td>
<td>11</td>
</tr>
<tr>
<td>Alginate impression material</td>
<td>18</td>
</tr>
<tr>
<td>Nonaqueous, elastomeric impression material</td>
<td>19</td>
</tr>
<tr>
<td>Dental duplicating material</td>
<td>20</td>
</tr>
<tr>
<td>Dental gypsum products</td>
<td>25</td>
</tr>
</tbody>
</table>
Nonelastic Materials

Impression plaster

Plaster of Paris is seldom used as impression material now that elastomeric materials are available, but it can be used as a "wash" (a thin lining material placed over a stiffer base material or tray) material for edentulous impressions. The main component of impression plaster is calcium sulfate hemihydrate, which reacts with water to form calcium sulfate dihydrate. Manufacturers incorporate additives to adjust the setting time and setting expansion. The water/powder (W/P) ratio recommended by the manufacturer should be measured out carefully. The powder should be sprinkled into the water, allowed to sit for 30 seconds to wet the powder, and then mixed for the minimum time necessary to obtain a homogeneous mix. Impression plaster is rigid and will break rather than bend. The plaster must be stored in an airtight container because it will absorb water from the air and its setting time will be adversely affected.

Dental impression compound (Types I and II)

There are two types of dental compound as defined by the ADA. Type I is used for impression taking, and Type II is used for tray preparation. Although dental compound has fallen into disuse, it can be used for full-crown impressions (Type I), impressions of partially or completely edentulous jaws (Type I), and impression trays in which a final impression is taken with another material (Type II). Compound cannot be used to record undercuts since it is not elastic. Impression compound is available in either cakes or sticks in various colors from a number of manufacturers (Jelenko, Kerr, Mizzy, and Moyco).

Composition

Dental compound contains several ingredients. Natural resins, which comprise about 40% of the formulation, make the compound thermoplastic. Shellac is often used. Waxes (about 7%) also produce thermoplastic properties. Stearic acid (about 3%) acts as a lubricant and plasticizer. Fillers and inorganic pigments account for the remaining 50% of the formulation. Diatomaceous earth, soapstone, and talc are examples of commonly used fillers.

Thermal and mechanical properties

Dental compound is thermoplastic; it is used warm (45°C) and then cooled to oral temperature (37°C), at which it is fairly rigid. The setting mechanism is therefore a reversible physical process rather than a chemical reaction. Dental compound is limited by its thermal properties. Type I materials have a flow of at least 85% at 45°C and less than 6% at 37°C. Type II materials flow about 70% at 45°C but less than 2% at 37°C. Both types become quite plastic with only an 8°C rise in temperature. The thermal conductivity of dental impression compounds is very low. These materials do not conduct heat very well and therefore require heat soaking to attain a uniform temperature throughout the mass. When heated or cooled, they soften or harden quickly on the outside, but time is needed for the temperature to become uniform throughout the entire mass. If the impression is removed from the mouth before it has
cooled completely, severe distortion may occur. Since these materials contain resins and waxes, they have high thermal expansion and contraction coefficients. Contraction from oral temperature to room temperature may be as high as 0.3%. Therefore, the dimensions of the resulting impression could be significantly different than those of the mouth. Since compound has such a high viscosity, it is difficult to record details.

Manipulation

The correct use of compound as an impression material requires considerable judgment in handling. Compound is softened by heating over a flame or in a water bath. Care must be taken to prevent volatilizing ingredients over a direct flame. Kneading in water also may cause changes in composition and flow properties. The aim, therefore, is a thorough heating without excessive temperatures or long periods of storage in water. A room-temperature water spray is used to cool the impression in the mouth. Cooling must be continued until the entire mass is rigid to reduce plastic flow. Care must also be taken to prevent overheating and burning of tissues. Also, cooling water should not be too cold, in order to prevent thermal shock.

To ease separation of the die stone, the impression should first be softened by immersion in warm water.

Advantages

Dental impression compound is compatible with die and model materials and is easily electroplated to form accurate and abrasion-resistant dies.

Disadvantages

The handling of dental impression compound is very technique sensitive. If it is not prepared properly, volatiles can be lost on heating, or low-molecular-weight ingredients can be lost during immersion in a water bath. Also, excessive wet kneading can incorporate water into the mix and change the flow properties of the compound. Due to a high coefficient of thermal expansion, the dimensions of the impression are not likely to be the same as the dimensions in the mouth. These materials are nonelastic and may distort on removal from the mouth. The casts should be poured within 1 hour.

Troubleshooting

1. Distortion. If the material is not completely cooled, the inner portions of the impression will still be soft when the impression is removed, resulting in distortion. Also, if water has been incorporated as the result of wet kneading, the material could have excessive flow at mouth temperature, producing distortion during removal from the mouth. If the tray used to carry the compound to the mouth is too flexible, distortion can result. It is important to select a tray that is strong and rigid. A delay in preparing the stone cast also may cause distortion. The cast should be poured as soon as possible after the impression has been removed from the mouth.

2. Compound is too brittle or grainy. Prolonged immersion in the water bath will
cause low-molecular-weight components to leach out.

Disinfection

Dental impression compound can be disinfected by immersion in sodium hypochlorite, iodophors, or phenolic glutaraldehydes. The manufacturer's recommendations for proper disinfection should be followed.

**Zinc oxide-eugenol**

Zinc oxide-eugenol's main use as an impression material is for dentures on edentulous ridges with minor or no undercuts. It can also be used as a wash impression over compound in a tray or in a custom acrylic tray. Zinc oxide-eugenol is also used as a bite registration material.

Composition

This material is commercially available in a powder and liquid form and as two pastes. One paste, called the base or catalyst paste, contains zinc oxide (ZnO), oil, and hydrogenated rosin. The second paste, the accelerator, contains about 12% to 15% eugenol, oils, rosin, and a filler such as talc or Kaolin. These two pastes have contrasting colors so it can be determined when the pastes are thoroughly mixed.

These materials are supplied as a soft- or hard-set type. Equal lengths of the two pastes, or properly proportioned amounts of the powder and liquid, are mixed with a stiff spatula on a special oil-resistant paper pad or a glass slab. The mixed material is placed in a preliminary impression made from tray compound or tray acrylic. The setting time is shortened by increases in temperature and/or humidity. The set material does not adhere to set dental plaster or stone.

\[
\begin{align*}
\text{Eugenol} & \quad \text{OCH}_3 \\
\text{CH}_2 - \text{CH} & = \text{CH}_2 \\
\text{OH} & \\
\end{align*}
\]

Zinc oxide, in the presence of moisture, reacts with eugenol to form zinc eugenolate, which acts as a matrix holding together the unreacted zinc oxide:

\[
\text{ZnO (excess)} + \text{eugenol (liquid)} + \text{H}_2\text{O} \rightarrow \text{Zn eugenolate (solid)} + \text{ZnO (unreacted, solid)}
\]
The setting reaction is accelerated by the presence of water, high humidity, or heat. A dimensional change of only about 0.1% shrinkage accompanies the setting.

Impression materials are classified as hard- and soft-set according to the ANSI/ADA specification. The hard-set material sets faster (in about 10 minutes, compared to 15 minutes for the soft-set material), although the hard- and soft-set materials both begin to set in about 5 minutes. The hard-set material is more fluid before setting than the soft-set material; after setting, it is harder and more brittle.

Noneugenol pastes containing carboxylic acids (e.g., lauric or ortho-ethoxybenzoic acid) in place of eugenol are available to avoid the stinging and burning sensation experienced by some patients.

Mechanical properties

The hardness of zinc oxide-eugenol impression materials is determined using a Krebs penetrometer with a load of 100 g for 10 seconds. The hardness for Type I (hard-set) materials should be no greater than 0.5 mm and the hardness for Type II (soft-set) materials should be between 0.8 and 1.5 mm.

The shrinkage of these materials during the hardening process is approximately 0.1%. Subsequently, no additional dimensional change should occur.

Manipulation

These materials are usually mixed on a mixing pad with a spatula. Equal lengths of base and catalyst are extruded on the mixing pad. The components are mixed thoroughly with a stiff stainless steel spatula. Adequate mixing time is 45 to 60 seconds, after which the mix should appear streak-free. The pastes have an initial set time of 3 to 5 minutes, with the setting time decreasing as the temperature and/or humidity increases. The model or cast should only be made from gypsum-type plaster or stone. After the stone has set, the impression is immersed in warm water (60°C) to ease its removal from the cast.

The spatula may be cleaned by warming or by wiping with available solvents.

Advantages

The advantages of zinc oxide-eugenol include high accuracy of soft tissue impressions due to its low viscosity. The material is stable after setting, has good surface detail reproduction, and is inexpensive. It also adheres well to dental impression compound.

Disadvantages

The Disadvantages of this material are messiness and a variable setting time due to temperature and humidity. Eugenol is irritating to soft tissues. This material is nonelastic and may fracture if undercuts are present.

Troubleshooting
1. **Inadequate working or setting time.** This could result from excessive humidity and/or temperature. An increase in either of these variables results in decreased working and setting time. It is important to select a material that provides the required setting time.

2. **Distortion.** If the tray warps on standing, the impression will also become distorted. It is important to select a stable tray material.

3. **Loss of detail.** If there is loss of detail, the impression material may not be compatible with the stone used to prepare the cast, and/or there may be adhesion between the impression and the stone.

**Disinfection**

Zinc oxide-eugenol impressions can be disinfected by immersion in 2% glutaraldehyde or 1:213 iodophor solutions at room temperature. The manufacturer's recommendations for proper disinfection should be followed.

**Elastic Materials**

**Agar (reversible) hydrocolloid**

**Agar hydrocolloids** have been largely replaced by rubber impression materials but are still used for full-mouth impressions without deep undercuts, quadrant impressions without deep undercuts, and single impressions (less frequently). They can be used for fixed partial denture impressions because of their high accuracy.

**Composition**

Agar hydrocolloids are available in both tray and syringe consistencies. The material is supplied as a gel in plastic tubes and contains agar (12% to 15%) as a gelling agent, borax (0.2%) to improve strength, potassium sulfate (1% to 2%) to provide good surfaces on gypsum models or dies, alkylbenzoates (0.1%) as preservatives, and coloring and flavoring agents (traces) for ease of "reading" the impression and esthetics. The balance of the formulation (~85%) is water. The syringe consistency is prepared by increasing the water content and decreasing the agar content.

The material, supplied as a solid gel, can be converted to a sol (liquid) by heating; cooling a sol causes it to become a gel:

\[
\begin{align*}
\text{cool to} & \quad 43^\circ\text{C} \\
\text{agar hydrocolloid (hot)} & \xrightarrow{\text{heat to}} \text{agar hydrocolloid (cold)} \\
\text{(sol)} & \xrightarrow{\text{gel}} 100^\circ\text{C}
\end{align*}
\]

The gel-to-sol and sol-to-gel transformations are dependent on time and temperature. The liquefaction and gelation temperatures are different (the latter being lower), and the effect is called **hysteresis**. A typical value of the gelation temperature is 43°C (109°F).
Mechanical properties

The mechanical properties of agar hydrocolloids are given in Table 7-2. They are highly elastic (98.8%) and sufficiently flexible (11%) to give accurate impressions of teeth with undercuts. They are stronger when stressed quickly; therefore, rapid removal is recommended.

![Table 7-2 Properties of elastomeric impression materials*](image)

<table>
<thead>
<tr>
<th></th>
<th>Agar</th>
<th>Alginate</th>
<th>Polysulfide</th>
<th>Condensation silicone</th>
<th>Addition silicone</th>
<th>Polyether</th>
</tr>
</thead>
<tbody>
<tr>
<td>Elastic recovery (%)</td>
<td>98.8</td>
<td>97.3</td>
<td>96.9-94.5</td>
<td>99.6-98.2</td>
<td>99.9-99.0</td>
<td>99.0-98.3</td>
</tr>
<tr>
<td>Flexibility (%)</td>
<td>11</td>
<td>12</td>
<td>0.4-1.9</td>
<td>&lt; 0.10</td>
<td>&lt; 0.05</td>
<td>&lt; 0.05</td>
</tr>
<tr>
<td>Flow (%)</td>
<td>—</td>
<td>—</td>
<td>0.2-1.0</td>
<td>0.01-0.2</td>
<td>0.2-0.3</td>
<td></td>
</tr>
<tr>
<td>Reproduction limit (um)</td>
<td>0.4-0.5</td>
<td>0.2-1.0</td>
<td>25</td>
<td>25</td>
<td>25</td>
<td></td>
</tr>
<tr>
<td>Shrinkage, 24 hours (%)</td>
<td>—</td>
<td>—</td>
<td>0.4-0.5</td>
<td>0.2-1.0</td>
<td>0.01-0.2</td>
<td>0.2-0.3</td>
</tr>
<tr>
<td>Tear strength (g/cm)</td>
<td>700</td>
<td>380-700</td>
<td>2,240-7,410</td>
<td>2,280-4,370</td>
<td>1,640-5,260</td>
<td>1,700-4,800</td>
</tr>
</tbody>
</table>

*See end-of-chapter glossary for definitions of terms.

Manipulation

Agar requires a special water bath with three chambers for heating and water-cooled trays. The following sequence is used:

1. Heat in water at 100°C (212°F) for 8 to 12 minutes.
2. Store in water at 65°C (149°F).
3. Place in a tray (containing cooling coils) at 65°C (149°F).
4. Temper in 46°C (115°F) water for 2 minutes before taking the impression.
5. After seating the tray, cool it with water at no less than 13°C (55°F) until gelation occurs.
6. After the impression is removed from the mouth, wash it to remove saliva, which will interfere with the setting of the gypsum.
7. Shake off excess water and lightly blow off with air.
8. Disinfect the impression.
9. Pour mixed dental stone into the impression. If the impression is stored for a short time in 100% relative humidity, it should be washed as described in steps 6 and 7 to remove any exudate on the surface caused by *syneresis* (the exudation of water, accompanied by contraction) before pouring the model.
10. After the initial setting of the stone, store the gypsum model and impression in a humidor.

Agar impressions become less accurate during storage, and prompt pouring of gypsum casts is necessary. Table 7-3 lists the dimensional changes that occur upon storage under different conditions. If agar impressions must be stored, the minimum changes in dimensions are found upon storage in 100% relative humidity for no longer than 1 hour. However, the gel structure can absorb water, a process called imbibition, which is usually accompanied by expansion.

<table>
<thead>
<tr>
<th>Storage conditions</th>
<th>Dimensional change</th>
<th>Causes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Air</td>
<td>Shrinkage</td>
<td>Evaporation of water from gel</td>
</tr>
<tr>
<td>H₂O</td>
<td>Expansion</td>
<td>Imbibition and absorption of water</td>
</tr>
<tr>
<td>100% relative humidity</td>
<td>Shrinkage</td>
<td>Syneresis</td>
</tr>
<tr>
<td>Inorganic salt solutions</td>
<td>Expansion or shrinkage</td>
<td>Depends on relationship of electrolyte in gel and in solution</td>
</tr>
</tbody>
</table>

As the values in Table 7-4 indicate, agar materials have a long working time. Handling, however, offsets this convenience because of the need for storage tanks. Gelation, produced by circulating cool water through the special trays, also requires special equipment. Thermal shock produced by suddenly cooling the warm colloid may be painful to patients who have metallic restorations.

<table>
<thead>
<tr>
<th>Agar</th>
<th>Alginate</th>
<th>Polysulfide</th>
<th>Condensation silicone</th>
<th>Addition silicone</th>
</tr>
</thead>
<tbody>
<tr>
<td>Preparation</td>
<td>Boil, temper, store</td>
<td>Powder, water</td>
<td>2 pastes</td>
<td>2 pastes or paste-liquid</td>
</tr>
<tr>
<td>Handling</td>
<td>Complicated</td>
<td>Simple</td>
<td>Simple</td>
<td>Simple</td>
</tr>
<tr>
<td>Ease of use</td>
<td>Technique sensitive</td>
<td>Good</td>
<td>Fair</td>
<td>Fair</td>
</tr>
<tr>
<td>Patient reaction</td>
<td>Tedium, thermal shock</td>
<td>Pleasant, clean</td>
<td>Unpleasant, stains</td>
<td>Pleasant, clean</td>
</tr>
<tr>
<td>Ease of removal</td>
<td>Very easy</td>
<td>Very easy</td>
<td>Easy</td>
<td>Moderate</td>
</tr>
<tr>
<td>Working time (min)</td>
<td>7-15</td>
<td>2.5</td>
<td>5-7</td>
<td>3</td>
</tr>
<tr>
<td>Setting time (min)</td>
<td>5</td>
<td>3.5</td>
<td>8-12</td>
<td>6-8</td>
</tr>
<tr>
<td>Stability</td>
<td>1 h at</td>
<td>Immediat</td>
<td>1 h</td>
<td>Immediate</td>
</tr>
</tbody>
</table>
Contact with agar retards the setting of gypsum, resulting in dies and casts with poor surface finish. With older products, soaking the impression in a 2% potassium sulfate solution was necessary to achieve a smooth surface finish. Most agar products now contain potassium sulfate, which acts as an accelerator for the gypsum setting reaction, and soaking is no longer necessary.

Advantages

Agar impression materials are inexpensive, have no unpleasant odors, and are nontoxic and nonstaining. They do not require a custom tray or adhesives, and the components do not require mixing. These materials are hydrophilic and can be used in the presence of moisture, are able to displace blood and body fluids, and are easily poured in stone. The stone casts are easily removed from the hydrocolloid impressions.

Disadvantages

These materials require the use of expensive equipment and must be prepared in advance. They tear easily, must be poured immediately, are dimensionally unstable, can only be used for single casts, and cannot be electroplated. The surface of stone casts will be weakened by compositions containing borax.

Troubleshooting

Sometimes problems of distorted impressions or loss of detail may be encountered when using agar hydrocolloids. The following are factors that could lead to distortion:

1. **Slow removal from the mouth.** To avoid permanent deformation, the impression should be removed with a quick jerk.

2. **Removal from the mouth before the gel reaches a temperature of 37°C (98.6°F) or less.** Above this temperature the impression material will still be plastic. The cooling rate of hydrocolloid is dependent on the temperature of water circulating through the

---

<table>
<thead>
<tr>
<th>Wetting and ease of pouring</th>
<th>100% RH pour</th>
<th>e pour</th>
<th>Excellent</th>
<th>Pour</th>
<th>Fair to good</th>
<th>dry Good</th>
</tr>
</thead>
<tbody>
<tr>
<td>Die material</td>
<td>Stone</td>
<td>No</td>
<td>Stone</td>
<td>Stone</td>
<td>Stone</td>
<td>Stone</td>
</tr>
<tr>
<td>Electroplating</td>
<td>No</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Disinfection</td>
<td>Poor</td>
<td>Poor</td>
<td>Fair</td>
<td>Excellent</td>
<td>Excellent</td>
<td>Fair</td>
</tr>
<tr>
<td>Comparative cost</td>
<td>Low</td>
<td>Very low</td>
<td>Low</td>
<td>Moderate</td>
<td>High to very high</td>
<td>Very high</td>
</tr>
</tbody>
</table>

*See end-of-chapter glossary for definition of terms.*
3. **Cooling water that is too cold (<13 °C).** Rapid cooling of the impression may cause a concentration of internal stresses that may be subsequently released.

4. **Application of force on the tray during gelation.** After the load is removed, relaxation of stresses will occur.

5. **Delay in pouring the cast.** Waiting any length of time to pour the cast will result in shrinkage of the impression due to the loss of water.

6. **Instability of tray.** If there is loss of detail, it may be caused by movement of the tray before gelation is complete. Failure to keep the impression stabilized will result in a multiple impression of the oral structures.

**Disinfection**

Agar hydrocolloids can be disinfected by immersion in sodium hypochlorite, iodophors, or phenolic glutaraldehydes. The manufacturer's recommendations for proper disinfection should be followed.

**Alginate (irreversible) hydrocolloid**

**Alginate**s are the most widely used impression materials in dentistry. They are used for making impressions for removable partial dentures with clasps, preliminary impressions for complete dentures, and orthodontic and study models. They are not accurate enough for fixed partial denture impressions.

**Composition**

Alginate is supplied as a powder containing sodium or potassium alginate (12% to 15%) and calcium sulfate dihydrate (8% to 12%) as reactants; sodium phosphate (2%) as a retarder; a reinforcing filler (70%), such as diatomaceous earth, to control the stiffness of the set gel; potassium sulfate or alkali zinc fluorides (~10%) to provide good surfaces on gypsum dies; and coloring and flavoring agents (traces) for esthetics. The sodium phosphate content is adjusted by the manufacturer to produce either regular- or fast-set alginates.

The powder is mixed with water to obtain a paste. Two main reactions occur when the powder reacts with water during setting. First, the sodium phosphate reacts with the calcium sulfate to provide adequate working time:

$$2Na_3PO_4 + 3CaSO_4 \rightarrow Ca_5(PO_4)_2 + 3Na_2SO_4$$

Second, after the sodium phosphate has reacted, the remaining calcium sulfate reacts with the sodium alginate to form an insoluble calcium alginate, which forms a gel with the water:

$$\text{Na alginate} + \text{CaSO}_4 \xrightarrow{\text{H}_2\text{O}} \text{Ca alginate} + \text{Na}_2\text{SO}_4$$

(powder) (gel)
To avoid the inhalation problems of alginate dust, some materials have been introduced in a dustless version in which the powder is coated with a glycol (Identidust Free, Cadco; Jeltrate Plus, Caulk).

Some products contain a chemical disinfectant in the alginate powder due to the concern for infection control (Coe Hydrophilic Gel, GC America; Identidust Free; Jeltrate Plus). Two examples of these disinfectants are didecyl-dimethyl ammonium chloride and chlorhexidine acetate. When the quaternary ammonium compound is used, the detail reproduction and gypsum compatibility of the alginate improve. However, the impressions made from these materials should still be disinfected upon removal from the mouth.

Mechanical properties

Table 7-2 gives an elastic recovery value of 97.3% for alginates, which indicates less elasticity and therefore less accuracy than agar hydrocolloids and silicone and polyether impression materials. The compressive and tear strengths increase with increasing rates of deformation. The limit of reproduction is also lower, indicating that less fine detail will be obtained. Figure 7-2 compares the elasticity of alginates with the more accurate agar materials. Alginates have a higher permanent deformation upon stretching to pass over undercuts.

Fig 7-2 Illustration of agar hydrocolloids' greater accuracy due to their greater degree of recovery after deformation around undercuts. (From Roydhouse, 1962. Reprinted with permission.)
Manipulation

Although easy to use, care is required in handling alginate hydrocolloids. The powder, supplied in a can, should be shaken up for aeration and one scoop of powder is used for one measure of water. A powder scoop and a graduated cylinder for water are usually supplied with the product. With predispensed powder products, one packet of powder is used with the amount of water specified by the manufacturer. A lower W/P ratio increases strength, tear resistance, and consistency, and decreases working and setting times and flexibility. Cooling the water increases the working and setting times. Insufficient mixing results in a grainy mix and poor recording of detail. Adequate spatulation gives a smooth, creamy mix with a minimum of voids. One minute of thorough mixing for the regular-set material and 45 seconds for the fast-set material are generally recommended. Alginates have a relatively short working time of about 2.5 minutes (see Table 7-4) and set about 3.5 minutes after mixing. Alginates are as unstable as agar hydrocolloids, because both are gels, and they undergo shrinkage or expansion upon loss or gain of water. Storage in either air or water results in significant dimensional change; however, storage at 100% humidity results in the least dimensional change. Therefore, the cast should be poured soon after removal of the impression and cleaning (see Table 7-4). Alginates, like agar, retard the setting of the gypsum model and die materials when in contact. Potassium sulfate is added by the manufacturer to accelerate the setting of the gypsum and to obtain smooth model and die surfaces.

An alginate tray material can be combined with an agar syringe material to prepare impressions. These impressions take advantage of the agar hydrocolloid's detail reproduction and compatibility with gypsum qualities and at the same time minimize equipment needs. A simple heater can be used to prepare the syringe material, and the water-cooled trays are no longer necessary. The alginate is placed in a tray, the agar is syringed around the preparation, and then the alginate is seated on top of the agar. Care must be taken to select an agar-alginate impression pair with suitable bond strengths. It is best to select combinations recommended by the manufacturers. Best results are obtained when single-unit impressions are made by this technique.

Advantages

Alginate impression materials are inexpensive, easy to manipulate, pleasant tasting, able to displace blood and body fluids, hydrophilic, and easily poured in stone. They can be used with stock trays.

Disadvantages

Alginate impression materials are inexpensive, easy to manipulate, pleasant tasting, able to displace blood and body fluids, hydrophilic, and easily poured in stone. They can be used with stock trays.
Problems may sometimes be encountered when using alginate hydrocolloids. The following should serve as a guide for troubleshooting problems with these materials:

1. **Inadequate working or setting time.** The temperature of the mixing water may be too high. Generally, the temperature of the water should range between 18°C and 24°C (65°F and 75°F). If the mixture is incompletely spatulated, it may be inhomogeneous and may set prematurely. Under normal conditions, adequate spatulation requires 45 to 60 seconds. If the W/P ratio is too low as the result of incorrect dispensing, the setting time could be too fast. Improper storage of the alginate powder can result in deterioration of the material and shorter setting times.

2. **Distortion.** If the tray moves during gelation or if the impression is removed prematurely, the result will be a distorted impression. The amount and duration of compression should be considered. It is important, therefore, to remove the impression from the mouth rapidly. Since the weight of the tray can compress or distort the alginate, the impression should not be placed face down on the bench surface. If the impression is not poured immediately, distortion could occur.

3. **Tearing.** If the impression tears, it is possible that the impression was removed from the mouth before it was adequately set. Wait 2 to 3 minutes after loss of tackiness to remove the impression for development of adequate tear strength. Also, the rate of removal from the mouth may be a factor. Since the tear strength of alginate increases with the rate at which a stress is applied, it is desirable to rapidly remove the impression from the mouth. In addition, thin mixes are more prone to tearing than those with lower W/P ratios. The presence of undercuts also can produce tearing. Blocking out these areas will place less stress on the impression material during removal. It is also possible that there is not enough impression material; there should always be at least 3 mm of material between the tray and the oral tissues.

4. **Loss of detail.** If there is loss of detail, the impression may have been removed from the mouth prematurely. Multiple impressions of the oral structure will result if the material is still in the plastic state when removed.

5. **Consistency.** If the preset mix does not have the proper consistency (is either too thick or too thin), the W/P ratio is incorrect. Care must be taken to fluff the powder before measuring and not to overfill the powder dispenser. Vigorous spatulation and mixing for the full recommended time is required to avoid consistency problems caused by inadequate mixing. If hot water is used, the mix may become grainy and prematurely thick.

6. **Dimensional change.** If dimensional change is a problem, a delay in pouring the impression might be the cause. Such delays will result in a cast that is distorted as well as undersized, because alginate impressions lose water when stored in air.

7. **Porosity.** If the impression is porous, this could result from whipping air into the mix during spatulation. After the powder has been wetted by the water, the alginate should be mixed so as to squeeze the material between the spatula blade and the side of the rubber bowl.

8. **Poor stone surface.** If the set gypsum remains in contact with the alginate for too
long a period of time, the quality of the stone surface will suffer.

Disinfection

Alginate hydrocolloids can be disinfected by immersion in sodium hypochlorite or iodophors. The manufacturer's recommendations for proper disinfection should be followed.

*Polysulfide rubber (mercaptan)*

**Polysulfide** rubbers are widely used for fixed partial denture application, due to their high accuracy and relatively low cost. These materials are useful for multiple impressions when extra time is needed. The polysulfides are supplied in tubes of base paste and catalyst paste, which are mixed together. Polysulfides are available in low, medium, and high viscosities.

Composition

The base paste contains the polysulfide polymer, fillers, and plasticizers. Low-molecular-weight (~4,000 MW) polysulfide polymer, having both terminal and pendant (near the center of the polymer) mercaptan groups (-SH), is used:

\[
\begin{align*}
\text{C}_3\text{H}_5 \\
\text{HS} - \text{R}_n - \text{S} - \text{S} - \text{C} - \text{S} - \text{S} - \text{R}_n - \text{SH} \\
\text{SH} \\
\text{Mercaptan}
\end{align*}
\]

The content of the reinforcing fillers (eg, zinc oxide, titanium dioxide, zinc sulfide, and silica) varies from 12% to 50% depending on the consistency (light, regular, or heavy). The fillers and plasticizers control the stiffness of the paste. The accelerator or catalyst paste contains lead dioxide (30%), hydrated copper oxide or organic peroxide, as a catalyst; sulfur (1% to 4%) as a promoter; and dibutyl phthalate or other nonreactive oils (17%) to form a paste. The balance of the catalyst paste is inorganic fillers used to adjust the consistency and reactivity. Those materials containing an organic peroxide may have decreased dimensional stability due to evaporation of the peroxide.

The lead dioxide catalyzes the condensation of the terminal and pendant -SH with -SH groups on other molecules, resulting in chain lengthening and cross-linking. In the process the material changes from a paste to a rubber. The reaction is accelerated by increases in temperature and by the presence of moisture.
Note that this is a condensation polymerization with water as a by-product.

Mechanical properties

The values for the mechanical properties of these materials are summarized in Table 7-2. For elastic recovery, the polysulfides have values of about 96%, slightly lower than those for the other rubber impression materials (e.g., silicone and polyether). Values for flow range from 0.4% to 1.9%, indicating a tendency to distort upon storage. The flow tends to be highest for the light-bodied and least for the heavy-bodied materials. Light-bodied polysulfides have flexibilities of about 16%; regular-bodied polysulfides have values of about 14%; and heavy-bodied polysulfides have values of about 10%. This is an advantage over the stiffer addition silicones and polyethers because removal from undercut areas is easier. Polysulfides have the highest tear strength of the rubber materials, which allows their use in deep subgingival areas where removal is difficult.

Manipulation

These materials are mixed on a mixing pad with a spatula. Equal lengths of base and catalyst are extruded on a disposable mixing pad. The components are mixed thoroughly with a stiff tapered spatula. The catalyst is dark and the base is white so thorough mixing is readily observed by lack of streaks in the mix. Adequate mixing time is 45 to 60 seconds. The working time is about 5 to 7 minutes (see Table 7-4), which is adequate considering the mixing times. Both working and setting times are shortened by higher temperatures and humidity. A value of 0.45% is given for shrinkage after 24 hours. Although this is less than that of the condensation silicones, the cast or die should be poured within 1 hour of taking the impression. Since polysulfides take longer to set than silicones, they require more chair time. They stain clothing permanently. They can be electroplated; some products can be silverplated, but copperplating is not recommended.

Advantages

Advantages of polysulfides include a long working time, good tear strength, good flow before setting, good reproduction of surface detail, high flexibility for easier removal around undercuts, and lower cost compared to silicones and polyethers.

Disadvantages
Disadvantages of these materials include the need to use custom-made rather than stock trays due to a greater chance of distortion, a bad odor, a tendency to run down the patient's throat due to lower viscosity, and the lead dioxide materials that stain clothing. Polysulfides must be poured within 1 hour and cannot be repoured.

Troubleshooting

Sometimes problems may be encountered when using polysulfide rubber impression materials:

1. *Inadequate working time.* This could result from excessive humidity and/or temperature. An increase in either of these variables results in decreased working and setting times. An improper base-to-catalyst ratio could also produce inadequate working time (too much catalyst will reduce working time).

2. *Distortion.* A number of factors cause distortion, one of which is too much load. Because recovery after deformation is dependent on the amount and duration of loading, the impression should not be placed face down on the laboratory bench. Improper removal from the mouth could also cause distortion of the impression; removal should be rapid because permanent deformation is a function of duration of stress. If a bubble is located just below the surface adjacent to a preparation (internal porosity), the impression may distort.

3. *Loss of detail.* Premature removal from the mouth may be a cause of this. Removal prior to sufficient polymerization or before the material is sufficiently elastic will result in inaccurate registration of detail. This may also be the result of incomplete mixing; failure to incorporate all the catalyst into the base will result in incomplete polymerization of portions of the surface. Movement of the tray prior to the time of removal could also cause loss of detail. The impression tray should be held firmly until the elastic stage is attained.

4. *Surface bubbles or voids.* These could be caused by the incorporation of air into the mix. The impression material should always be mixed carefully with only the flat surface of the blade. If the material is partially polymerized prior to insertion in the mouth, voids may occur in the impression.

Disinfection

Polysulfide impressions can be disinfected by immersion in sodium hypochlorite, iodophors, complex phenolics, glutaraldehydes, or phenolic glutaraldehydes. The manufacturer's recommendations for proper disinfection should be followed.

*Condensation silicone rubber*

**Condensation silicone** rubber impression materials are used mainly for fixed partial denture impressions. They are ideal for single-unit inlays. These materials are supplied either as two-paste or paste-liquid catalyst systems. Condensation silicones are available in low, medium, high, and very high (putty) viscosities.

Composition
The base paste usually contains a moderately high-molecular-weight poly(dimethylsiloxane) with terminal hydroxy groups (-OH), an orthoalkylsilicate for cross-linking, and inorganic filler. A paste will contain 30% to 40% filler, whereas a putty will contain as much as 75%. The catalyst paste or liquid usually contains a metal organic ester, such as tin octoate or dibutyl tin dilaurate, and an oily diluent. A thickening agent is used when making catalyst pastes. Sometimes a catalyst will contain both the orthoalkylsilicate and the metal organic ester:

\[
\begin{align*}
\text{HO} & - \text{Si} - \text{O} - \text{H} & \text{RO} & - \text{Si} & - \text{OR} & \text{Sn} & \left[\text{O} - \text{C} - (\text{CH}_2)_{6} - \text{CH}_3\right]_2 \\
\text{CH}_3 & & & & & & \\
\end{align*}
\]

The metal organic ester catalyzes the reaction. One part of the polymerization involves chain extension by condensation of the terminal -OH group in a siloxane. The other part consists of cross-linking between chains by the orthoalkylsilicate molecules:

\[
\begin{align*}
\text{HO} & - \text{Si} - \text{O} - \text{H} & \text{RO} & - \text{Si} & - \text{OR} & \text{tin} & \text{octoate} & \rightarrow & - \text{Si} - \text{O} - \text{Si} & - \text{O} & - \text{Si} - \text{OH} \\
\text{CH}_3 & & & & & & \\
\end{align*}
\]

A volatile alcohol is formed as a by-product.

**Mechanical properties**

Accepted values for the mechanical properties of these materials are given in Table 7-2. An average value of 99% for elastic recovery is excellent. The flow of silicones is low; most values are less than 0.1%, indicating that less distortion is likely to be caused by light pressure on standing. The silicones are stiffer than polysulfides, as indicated by lower flexibility values in Table 7-2. The shrinkage in 24 hours ranges from 0.2% to 1.0%. About half the shrinkage takes place in the first hour, and it is greater than for polysulfides or polyethers. Polymerization and evaporation of the alcohol formed in the reaction are responsible for this high shrinkage. Accuracy is greatly improved by first taking an impression with a highly filled silicone putty and,
after setting, taking a second impression with a light-bodied silicone. Thus, the final total shrinkage is lower.

Manipulation

The manipulation of condensation silicones is the same as for polysulfides, except that the silicone material may be supplied as a base paste plus a liquid catalyst. When it is supplied in this form, one drop per inch of extruded base paste is usually recommended. The setting time (6 to 8 minutes) is less than that of the polysulfides, which offers some advantage in chair time savings. Electroplating is possible. Because of the high polymerization shrinkage, the cast or die must be poured as soon as possible. Higher temperatures and humidity shorten the setting time.

Advantages

Condensation silicones are clean, pleasant materials for the patient. They are highly elastic, and the setting time can be controlled with the amount of accelerator. The use of a putty-wash system improves accuracy and eliminates the need for a custom tray.

Disadvantages

These materials tend to be inaccurate due to shrinkage on standing and should be poured within 1 hour. They are very hydrophobic, require a very dry field, and are difficult to pour in stone.

Troubleshooting

1. *Inadequate working time.* This could result from excessive humidity and/or temperature. Although not as critical as polysulfide rubber, the setting times of silicone impression materials are influenced by temperature and humidity. Increases in these conditions tend to shorten both working and setting times. An improper base-to-catalyst ratio could also produce inadequate working time. Insufficient catalyst will result in prolonged setting times. Failure to polymerize in the predicted time may result from deterioration during storage.

2. *Distortion.* This may result from incorrect separation from impressed structures. Failure to remove the impression in a rapid, jerking motion may cause the impression to become permanently deformed. An impression can also become distorted if there is inadequate support of the impression after removal from the mouth. It may undergo permanent deformation if allowed to rest face down on the bench. Excessive delay in pouring the cast (30 minutes or more) also may result in dimensional changes. Possible shrinkage may result from continued polymerization and vaporization of volatiles in the silicone rubber. This shrinkage can be compensated for by use of the double impression technique. A preliminary impression is made with a very high-viscosity material (putty), providing space for the final impression, which is made with a low-viscosity material using the preliminary impression as the tray.

3. *Loss of detail.* Premature removal of the impression from the mouth could cause loss of detail. Failure to allow the impression material to set adequately will result in plastic deformation. Loss of detail also can be caused by incomplete mixing. Failure
to adequately incorporate all the catalyst into the base will result in incomplete polymerization. Loss of detail can also result from movement of the tray after the impression has been seated. Failure to maintain the tray in a stable position will result in a blurred impression.

Disinfection

Condensation silicone impressions can be disinfected by immersion in sodium hypochlorite, iodophors, complex phenolics, glutaraldehydes, or phenolic glutaraldehydes. The manufacturer's recommendations for proper disinfection should be followed.

Addition (vinyl) silicones

Addition silicones represent an advance in accuracy over condensation silicones. This has been achieved by a change in polymerization reactions to an addition type and the elimination of an alcohol by-product that evaporates, causing shrinkage. These materials are available as two-paste systems in four viscosities—light, medium, heavy, and putty—and a range of colors, allowing monitoring of the degree of mixing. Due to their high accuracy, these materials are suitable for fixed and removable partial denture impressions. They are rigid after setting and expensive, and therefore are not used for routine study models. Hydrophilic materials have been introduced that reportedly contain surfactants to improve the wetting characteristics compared to unmodified silicones. Monophase materials have been formulated with sufficient shear thinning to be used as both low-viscosity and high-viscosity materials.

Composition

These materials are based on silicone prepolymerst with vinyl and hydrogen side groups, which can polymerize by addition polymerization. They are therefore called vinyl or addition silicones. The setting reaction is produced by mixing one paste containing the vinyl-poly(dimethylsiloxane) prepolymer with a second paste that contains a siloxane prepolymer with hydrogen side groups. A platinum catalyst, chloroplatinic acid, is present in one of the pastes and starts the addition polymerization reaction as follows:

\[
\begin{align*}
\text{CH}_3 \\
\text{Si} \text{CH}_2 \text{CH}_2 \text{CH}_2 \text{Si} \text{CH}_3 & \quad \quad \text{H}_2 \text{PtCl}_6 \\
\text{Si} \text{CH}_3 & \quad \quad \quad \text{CH}_3 \text{SiCH}_2 \text{CH}_2 \text{SiCH}_3
\end{align*}
\]

Other reactions also occur that may release hydrogen gas, which on rare occasion can produce porosity. Some manufacturers include hydrogen absorbers in their formulations to eliminate this problem. Since no volatile by-products are produced in the reaction, addition silicones have a much greater dimensional stability than condensation silicones. Some manufacturers include a retarder for extending the working and setting times.
Mechanical properties

The working and setting times of addition silicones are faster than polysulfides; a retarder is often supplied to extend the working time. Addition silicones have excellent elasticity and show very low dimensional shrinkage upon storage (see Table 7-2). Therefore, addition silicones can be safely poured up later or sent to a dental laboratory. Addition silicones do have greater rigidity, however, and therefore it is difficult to remove the impression around undercuts, as indicated by the lower flexibility value. The tear strength of addition silicones is similar to that of condensation silicones, but less than that of polysulfides.

Manipulation

Addition silicones are as pleasant to handle as condensation silicones. Because there is the possibility of hydrogen release upon setting, finely divided palladium is added to some products to absorb the hydrogen and prevent bubbles from forming on stone die surfaces. If a product does not contain a hydrogen absorber, an hour should pass before pouring dies, and the impression should stand overnight before epoxy dies are poured. Addition silicones can be electroplated with both copper and silver.

Automatic mixers that provide quick, bubble-free mixes are available with several products (Fig 7-3).

![Fig 7-3](image)

Advantages

Addition silicones are highly accurate and have high dimensional stability after
setting. Recovery from deformation on removal is excellent. The material stays in the tray of reclined patients, does not stain clothing, has pleasant colors and scents, may be used with stock or custom trays, and can be copper- or silverplated. The materials may be poured 1 week after taking the impression, and multiple pours are possible.

Disadvantages

The Disadvantages are that the material is expensive—twice the cost of polysulfides; is more rigid than condensation silicones and difficult to remove around undercuts; has a moderate tear strength, making removal from gingival retraction areas somewhat risky; and may release hydrogen gas on setting, producing bubbles on die surfaces if an absorber is not in the product. Hydrophobic materials are difficult to electroplate and to pour in stone. Also, sulfur in latex gloves and rubber dams can inhibit polymerization.

Troubleshooting

1. *Inadequate working time.* This can result from excessive temperature. As the temperature increases, the working and setting times decrease. If the impression material does not set, the catalyst may have been contaminated. The platinum-containing catalyst becomes inactive after contacting certain substances, such as tin or sulfur compounds. If addition silicones are combined with condensation silicones, the material will not set. Components from these two systems are not compatible and cannot be mixed together.

2. *Loss of detail.* This could occur if unmodified addition silicones are used. These hydrophobic materials cannot displace any moisture or hemorrhage that is not removed prior to placement of the impression material.

3. *Porosity.* The stone surface may appear porous if hydrogen gas is evolved. If this is the case, it is recommended that the pouring of dies be delayed for at least 1 hour.

4. *Distortion.* This may occur if the polysiloxane adhesive does not provide adequate retention. Mechanical retention may be required in combination with the adhesive.

Disinfection

Addition silicone impressions can be disinfected by immersion in sodium hypochlorite, iodophors, complex phenolics, glutaraldehydes, or phenolic glutaraldehydes. The manufacturer's recommendations for proper disinfection should be followed.

*Polyether rubber*

Polyether rubbers are used for accurate impressions of a few prepared teeth without severe undercuts. Their high stiffness and short working time restricts their use to impressions of a few teeth. Polyethers are available in low-, medium-, and high-viscosity materials.
Composition

Polyethers are supplied as two-paste systems. The base paste contains low-molecular-weight polyether with ethylene-imine terminal groups:

\[
\begin{align*}
\text{CH}_2 & \\
\text{-N} & \\
\text{CH}_2 & \\
\end{align*}
\]

along with fillers, such as colloidal silica, and plasticizers:

\[
\begin{align*}
\text{H} & \text{O} & \text{O} & \text{H} \\
\text{CH}_3 & \text{-C-CH}_2 & \text{-C-} & \text{O} & \text{-R-O-C-CH}_2 & \text{-C-CH}_3 \\
\text{N} & \\
\text{CH}_2 & \text{-CH}_2 & \\
\end{align*}
\]

Polyether

The catalyst paste contains an aromatic sulfonic acid ester plus a thickening agent to form a paste along with fillers:

\[
\begin{align*}
\text{SO}_3 & \text{CH}_2 \text{CH}_3 \\
\end{align*}
\]

Sulfonic ester

When the base paste is mixed with the catalyst paste, ionic polymerization occurs by ring opening of the ethylene-imine group and chain extension. The reaction converts the paste to a rubber as follows:

\[
\text{polyether} + \text{sulfonic ester} \rightarrow \text{polyether rubber} \\
(\text{base paste}) (\text{catalyst}) (\text{impression})
\]

Mechanical properties

Polyethers are similar to addition silicones in properties. The early polyethers had short working and setting times and low flexibilities. Thinners were available to increase the working time and flexibility without any significant loss of other physical or mechanical properties. However, more recent formulations have a working time of 2.5 minutes and a setting time of 4.5 minutes (see Table 7-4). Shrinkage values of 0.3% in 24 hours place the polyethers at the upper end of the range for accuracy, but inferior to some addition silicones. Because this rubber absorbs water and changes dimensions, storage in water is not recommended. Elastic recovery values average 98.5%, between those for polysulfides and addition silicones (see Table 7-2). The
flow of polyethers is very low and contributes to accuracy. The flexibility also is low (ie, the stiffness is high). This quality causes some problems on removal of the impression from the mouth or the die from the impression. More rubber between the tray and the impression area is recommended to relieve this problem. Polyethers have low tear strength values.

Manipulation

The manipulation of polyethers is similar to that of polysulfides and silicones. Equal lengths of base and catalyst paste are mixed vigorously and rapidly (30 to 45 seconds), because the working time is short. They are easy to mix. The impressions can be readily silverplated to produce accurate dies. Precautions should be taken to mix the material thoroughly and to avoid contact of the catalyst with the skin or mucosa because tissue reactions have been observed.

A handheld gun-type mixer that provides quick, bubble-free mixes is available for one product (Permadyne Garant, ESPE). Also, an automatic mixing device (Pentamix, ESPE) (Fig 7-4) has been introduced for use with a polyether packaged in polybags (Impregum Penta).

Advantages

Advantages of polyethers include pleasant handling and ease of mixing. These materials are more accurate than polysulfide or condensation silicone impression materials. They have good surface detail reproduction and are easily poured in stone. If kept dry, they will be dimensionally stable for up to 1 week.

Disadvantages

Disadvantages include high cost, short working and setting times, and high stiffness after setting, which limit their use. Their bitter taste is objectionable to some patients.
Storage of polyether impressions is critical, as they will distort if stored in water or high humidity. They cannot be left in disinfectant solutions for long periods.

Troubleshooting

1. *Inadequate working time*. This can result from excessive temperature. An increase in temperature will decrease the working and setting times. An improper base-to-catalyst ratio (too much catalyst) will decrease the working time.

2. *Tearing*. The rigidity of this material may result in tearing upon removal of the impression from the mouth or of the die from the impression. Tearing may occur if the rubber thickness is inadequate (at least 4 mm).

3. *Distortion*. The impression may distort due to moisture absorption and/or plasticizer extraction. These phenomena may result in dimensional change when impressions are stored in high-humidity environments or are exposed to water. The use of the thinner may increase the water absorption. Delay in impression placement may cause distortion. The onset of setting will result in the formation of elastic properties, which will cause deformation upon removal from the mouth.

4. *Loss of detail*. This can be caused by incomplete mixing. Failure to obtain a homogeneous mix will result in incomplete polymerization. If problems are encountered with multiple dies, it may be the result of gingival tearing and/or swelling. The rigidity and the relatively low tear strength of the polyethers may result in progressive deterioration at the gingival margin of the impression. The impression may absorb enough moisture from the gypsum that multiple pours could produce swelling.

Disinfection

Polyether impressions can be disinfected by immersion in sodium hypochlorite. The manufacturer's recommendations for proper disinfection should be followed.

**Dental Duplicating Materials**

Dental duplicating materials are used to prepare duplicate casts for prosthetic appliances and orthodontic models. These materials are used to make an impression of the original dental cast. Agar hydrocolloid duplicating materials are used most frequently. Their composition has a higher water content than agar hydrocolloid impression materials, and subsequently, a lower agar content. Therefore, the compressive strengths for duplicating materials is generally lower than those of impression materials. Advantages of the reversible materials are that they can be reused a number of times and may be stored in the liquid state for use as needed. Disadvantages include the fact that immediate pours are necessary and degradation of the material will be accelerated by contamination. A classification key is shown in Fig 7-5.
Clinical Decision Scenarios for Dental Impression Materials

This section presents one approach for choosing impression materials for specific situations. This approach is based on consideration of the advantages and disadvantages of each material and takes into account the different practitioner requirements, such as skill level, cost factors, and so on, as applicable.

Each scenario includes (1) a description of the situation; (2) a list of the critical factors; (3) a list of the advantages and disadvantages of each material prioritized using the following codes: * = of minor importance, ** = important, *** = very important; and (4) an analysis of the situation and the decision reached.

---

Materials Polysulfides vs addition silicones in a dental school clinic

Description A dental school instructor has been asked to determine which impression material will be used in the undergraduate crown-and-bridge clinics. There is a fixed budget that, typical of universities, dictates the use of inexpensive materials. The students are slow and unskilled at their stage of training, so a material allowing long working time is desirable. In addition, the students have not yet developed their gingival retraction techniques to the point that they have a wide sulcus into which to inject their syringe material. Because of this, a material with a high tear strength is desirable so the gingival area will not tear off from the main part of the impression and remain in the sulcus upon removal of the tray. After review of the materials available, all are rejected except for the polysulfides (mercaptans) and the addition silicones (polyvinyl siloxanes).

Critical Factors Low cost, long working time, high tear strength
Polysulfides

Advantages

*** Longer working time  
*** Lower cost  
*** Good tear resistance

* High flexibility (easy to remove from undercuts)

*** Good accuracy

Disadvantages

* Disagreeable odor and taste  
* Can stain clothing  
* Low dimensional stability (should be poured within 1 hour)

* Longer setting time

Addition silicones

Advantages

* Good dimensional stability (can be mailed unpoured)

* Short setting time  
*** Very accurate

* Easy to mix and clean up  
* No disagreeable odor or taste

* Rigid material

Disadvantages

*** High cost  

* Low flexibility (harder to remove from undercuts)

*** Shorter working time

*** Lower tear resistance (may tear off in deep subgingival areas)
* Some emit hydrogen gas (pouring must be delayed)

**Analysis/Decision** Since the factors identified as important were low cost, high tear strength, and long working time, all advantages or Disadvantages pertaining to these properties are marked ***,** as is the property of good accuracy. This is essential in any impression material used for crown-and-bridge procedures. Because accuracy is an advantage of both materials, it cancels out and is not a deciding factor.

In this example, the factors the dentist is interested in are advantages for the polysulfides, but not for the addition silicones. This makes the choice easy—the polysulfides are selected.

The next step is to conduct a trial of the chosen material to see if this decision is well founded. For example, one half of the class takes impressions with polysulfide, and the other half uses the material previously used. If the polysulfide works well in the hands of students and is less expensive than the addition silicone, the trial confirms the decision.

**Materials** Polysulfides vs addition silicones in a busy practice

**Description** An experienced clinician in a busy practice has been using agar hydrocolloid in water-cooled trays but has heard that the new elastomeric impression materials have become more dimensionally stable than they were originally. She is interested in trying either a polysulfide or an addition silicone. She has always obtained very good impressions with the agar hydrocolloid material but has never liked having to interrupt her busy schedule to pour the impressions right away. She feels that her time could be better used if a technician could set the dowel pins and pour the impression, but she has not felt that her practice is large enough to justify an in-office laboratory technician. She has likewise been unable to send the agar hydrocolloid impressions to a laboratory for pouring because of the material's properties.

The clinician realizes that to use a polysulfide material, she should use custom trays due to the lack of dimensional stability of the material. However, she does not like to take the dental assistant away from chairside for the time required to make a tray because she lowers productivity by doing so. She would prefer a material with sufficient dimensional stability that it could be used in a disposable stock plastic tray. She is also looking for a material that will set much faster than the agar so she can decrease chair time for a crown impression.

**Critical Factors** Low cost, short setting time, good dimensional stability

**Polysulfides**

*Longer working time*
*** Lower cost

* Good tear resistance
* High flexibility (easy to remove from undercuts)

*** Good accuracy

Disadvantages

* Disagreeable odor and taste
* Can stain clothing

*** Low dimensional stability (should be poured within 1 hour)
*** Longer setting time

Addition silicones

Advantages

*** Good dimensional stability (can be mailed unpoured)
*** Short setting time
*** Very accurate

* Easy to mix and clean up
* No disagreeable odor or taste
* Rigid material

Disadvantages

*** High cost

* Low flexibility (harder to remove from undercuts)

** Shorter working time

* Lower tear resistance (may tear off in deep subgingival areas)
* Some emit hydrogen gas (pouring must be delayed)

Analysis/Decision  Here the decision is less obvious than in the previous example. The addition silicone seems to offer much of what the dentist desires, but it costs twice as much as the polysulfide. In this case, the decision might be in favor of the polysulfide due to the significant cost difference. After the trial period, however, the
dentist confirmed her suspicion that the polysulfide was not stable enough to use stock trays and it took 15 to 30 minutes for either her or her assistant to fabricate the custom trays. She also found that that use of "putty" material to line the stock tray greatly reduced the cost of the addition silicone. In addition, she found that placing dowel pins and pouring up the polysulfide took another 15 to 30 minutes of someone's time. The amount of time lost from chairside production more than offset the higher cost of the addition silicones, whose properties allowed her to use stock trays and to mail the impressions to her laboratory. Seeing this in her trial, the dentist reconsidered and changed to the addition silicone, which satisfied her needs.

Materials  Polysulfides vs addition silicones for "triple tray"

Description  The dentist has been using the polysulfides and is quite content but wishes to try a new impression technique that is gaining great favor among his peers. This method uses what is known as a "triple tray" and has the advantage for the single crown of obtaining the working impression, the opposing impression, and the bite relation all at the same time. This could significantly cut down the amount of time needed for each patient. The dentist finds, however, that the sides of the flimsy plastic trays sometimes spread apart when the dental arches impinge on them as the patient bites into position. The tray then attempts to spring back from this distortion when it is removed from the mouth. The recovery from the spreading requires that he use a material with sufficient rigidity that it can maintain its shape against this elastic recovery of the tray. If not, the tray recovery may distort the impression and result in a nonfitting crown.

Critical Factors  Low flexibility, good dimensional stability

Polysulfides

Advantages

* Longer working time

* Lower cost

* Good tear resistance

*** Good accuracy

Disadvantages

*** High flexibility (easy to remove from undercuts)

* Disagreeable odor and taste

* Can stain clothing

*** Low dimensional stability (should be poured within 1 hour)
* Longer setting time

Addition silicones

Advantages

*** Good dimensional stability (can be mailed unpoured)

* Short setting time

*** Very accurate

* Easy to mix and clean up

* No disagreeable odor or taste

*** Rigid material

Disadvantages

* High cost

*** Low flexibility (hard to remove from undercuts)

* Shorter working time

* Lower tear resistance (may tear off in deep subgingival areas)

* Some emit hydrogen gas (pouring must be delayed)

Analysis/Decision  In this case, the trial period showed that the addition silicone was the only feasible choice. The need for rigidity in the impression material for the triple tray is important enough that it becomes the deciding factor. Although other factors, such as accuracy, are important, unless the material is rigid enough to hold its shape, these other properties are meaningless.

Materials  Polysulfides vs addition silicones for full-arch impression

Description  In this situation, the dentist is a prosthodontist in an established specialty practice. He has a dental laboratory in his office and employs two dental laboratory technicians. He therefore has the capability of having custom trays made in his office and the impressions poured right away. Most of his patients have been referred from dentists who do the easier cases themselves. Consequently, his patients usually already have multiple crowns or fixed partial dentures in different areas of the mouth. He feels that full-arch impressions are better because in the more extensive cases he needs the accuracy in bite relations that can only come from full-arch registration. He has heard that addition silicones are very accurate and that they have better
dimensional stability than the material he has been using. This appeals to him for "pickup" impressions where a rigid material and better dimensional stability greatly enhance the probability of success. On the other hand, he is concerned that the stiffness of the addition silicone will make the removal of full-arch impressions from the mouth much more difficult. He is also concerned that the stiffness may pose a greater risk of inadvertently removing crowns or fixed partial dentures that his patients already have in other areas of the mouth.

**Critical Factors**  Good accuracy, high flexibility, good dimensional stability

**Polysulfides**

* Advantages
  * Longer working time
  * Lower cost
  * Good tear resistance

  *** High flexibility (easy to remove from undercuts)
  *** Good accuracy

* Disadvantages
  * Disagreeable odor and taste
  * Can stain clothing

  *** Low dimensional stability (should be poured within 1 hour)
  * Longer setting time

**Addition silicones**

* Advantages

  *** Good dimensional stability
  * Short setting time

  *** Very accurate

  * Easy to mix and clean up
  * No disagreeable odor or taste

  *** Rigid material
Disadvantages

* High cost

*** Low flexibility (harder to remove from undercuts)

* Shorter working time

* Lower tear resistance (may tear off in deep subgingival areas)

* Some emit hydrogen gas (pouring must be delayed)

Analysis/Decision  The high flexibility listed under the advantages of polysulfides is offset by the rigidity listed as an advantage under addition silicones. He found that the rigidity was desirable for pickup impressions, but the flexibility was desirable for regular impressions. He decided to try the addition silicone and found that it did, indeed, give very accurate and dimensionally stable impressions. His pickup impressions were better than they had ever been. However, in normal situations the rigidity of the material made removal of the full-arch impressions much more difficult. Some crowns and fixed partial dentures on which he was not working came off in the impression material due to its rigidity. He found, however, that he could compensate for this rigidity by filling the impression tray only partially full in the areas where he only needed the occlusal registration and filling it full in the area of the preparations.

Glossary

addition silicone A silicone polymer resulting from the free-radical polymerization of vinyl groups by a platinum catalyst.

agar An ingredient in agar impression material. It is usually extracted from seaweed as a polysaccharide.

alginate Alginate impression material containing salts of alginic acid.

cast material Material used to form casts (models or dies) from impressions.

condensation (in polymerization) In a condensation polymerization (eg, to form silicone impression materials), two molecules unite and one small molecule (eg, H₂O, ROH) is released as a by-product.

~Si-OH + R-O-Si~ → ROH + ~Si-O-Si~

condensation silicone A silicone polymer resulting from the condensation of terminal -OH groups by orthoalkylsilicates and releasing alcohol as a by-product.

elastic recovery The amount of rebound after a cylinder of material is strained 10% for 30 seconds.
**electroplate** The process of depositing metal from solution onto the surface of an impression using an electric current.

**eugenol** Oil of cloves. As a derivative of phenol, eugenol reacts as an acid with zinc oxide.

**flexibility** The amount of strain produced when a sample is stressed between 100 and 1,000 g/cm². A flexible material shows a higher value of flexibility than a stiff material.

**flow** The amount of shortening of a cylinder when placed under a light load for 15 minutes.

**gel** A colloid system in which the solid (eg, agar) and liquid (eg, water) are continuous phases. A gel is usually flexible.

**hydrocolloid** A colloid system in which the liquid phase is water. Agar impression material is a hydrocolloid (agar + water).

**hydrophilic** Having a strong affinity for water; can be readily wetted by water.

**hydrophobic** Resistant to wetting by water.

**hysteresis** The phenomenon of a gel's having a liquefaction temperature different from the solidification temperature of the sol.

**imbibition** The taking up of fluid by a colloidal system, resulting in swelling.

**plasticizer** A material that is added to increase flow.

**polyether** The polymer resulting from the ionic polymerization with ring opening of the ethylene-imine group and chain extension.

**polysulfide** The polymer resulting from the condensation of terminal mercaptan groups catalyzed by lead dioxide or other catalysts.

**silicone rubber** A polymer resulting from the formation of silicon-oxygen-silicon bonds (-Si-O-Si-). Silicone impression material is a silicone rubber.

**sol** A colloid system in which the solid phase is dispersed in the liquid phase. A sol usually has fluid properties.

**syneresis** The exudation of a liquid film on the surface of a gel.

**terminal group** A chemical group at the end of a molecule (eg, -SH).

**thermal conductivity** The quantity of heat passing through a body 1 cm thick with a cross section of 1 cm² when the temperature difference between the hot and cold sides of the body is 1°C.
thermoplastic The property of softening on heating and hardening on cooling.

working time Duration from the start of mixing to the time when a test rod leaves a permanent indentation in the material upon withdrawal.

Discussion Questions

1. Why are there so many impression materials in dentistry at the present time?

2. How could the movement of soft tissues during impression taking affect the accuracy of the final impression?

3. Since elastic impression materials are viscoelastic, wouldn't immediate pouring of a gypsum cast result in inaccuracy?

4. Why is it essential to sterilize an impression before sending it to a laboratory, and how is this done?

5. Why is the impression tray an important factor in obtaining a good impression?

Study Questions

(See Appendix E for answers.)

1. What components are present in dental compound, and what is the purpose of each?

2. What term describes the quality of dental compound that allows it to be repeatedly softened on heating and hardened on cooling?

3. What is the principal difference between the properties of impression and tray-type compounds?

4. What precautions should be taken when heating dental compound?

5. What importance does low thermal conductivity have on the clinical handling of dental impression compound?

6. What factors affect the flow of impression compound?

7. What are the reactive ingredients in a zinc oxide-eugenol material?

8. What is responsible for the setting of zinc oxide-eugenol impression paste?

9. What factors affect the setting time of zinc oxide-eugenol?

10. What are the functions of the various components in agar hydrocolloid?

11. What is meant by hysteresis in agar hydrocolloid?
12. What kinds of dimensional changes occur when an agar impression is stored in air, water, 100% relative humidity, or potassium sulfate solution?

13. What is the function of each of the components in alginate powder?

14. What factors affect the setting time of alginates?

15. What reaction is responsible for providing the working time of alginate, and what other reaction is responsible for the setting of alginate?

16. Why are alginate impression materials called irreversible hydrocolloids?

17. What effect does the water/powder ratio of alginates have on their properties?

18. What effect does spatulation of alginates have on their properties?

19. What effect does water temperature have on the working and setting times of alginates?

20. How do the properties of alginate impression materials compare with those of agar impression materials?

21. Describe the setting reactions for polysulfide, silicone, and polyether rubber impression materials.

22. What effect do proportioning and temperature have on the working and setting times of the three rubber impression materials?

23. What impression material may be used if you wish to use either a gypsum die, a silverplated die, or a copperplated die?

24. Compare the elastic recoveries of polysulfide, silicone, and polyether impression materials. How does elastic recovery affect clinical usage?

25. Compare the flexibilities of polysulfide, silicone, and polyether impression materials.

26. How would you disinfect your impression prior to sending it to the laboratory?

27. List the advantages of each of the following recent advances in impression materials: (a) addition silicones with hydrophilic properties; (b) single-viscosity, or monophase, addition silicones; (c) automatic mixers for addition silicones and polyethers.

**Recommended Reading**


Chapter 8. Polymeric Restorative Materials

Introduction

The first material developed for use as a direct esthetic restorative was silicate cement. Introduced in the late 1800s, the cement was prepared from an alumina-silica glass and a phosphoric acid liquid. Highly soluble in oral fluids, silicate deteriorated rapidly, yet remained the favored material until the early 1950s. Dissolution, discoloration, loss of translucency, and lack of adequate mechanical properties contributed to its eventual replacement. Its main advantage was the slow release of fluoride from the glass phase.

Self-curing unfilled acrylic resins were introduced around 1945 as a substitute for silicate cement and were in moderate use in the 1950s. These materials were related to denture base resins and were much less soluble and more color stable than silicates. They were easy to use, polishable, and had good initial esthetics. Their main problems were high shrinkage upon polymerization, large thermal dimensional change, eventual discoloration, and a high wear rate.

Resin composites, a combination of hard, inorganic filler particles bonded to soft dimethacrylate polymer, were introduced in the 1960s. As a consequence of the bonded filler phase, these materials had much better mechanical properties than did unfilled resins, approaching the properties of dentin and enamel. Originally intended for use in anterior Class 3, 4, and 5 restorations where esthetics are important, improvements have included light curing, bonding to tooth structure, and reduced wear. Continued development in wear resistance, dentin bonding, and reduced polymerization shrinkage has led to their increased use in posterior restorations.

Classification

A composite is a material composed of two or more distinct phases. Resin composites for dental use were formulated to combine the esthetics and ease of use of a polymerizable resin base with the improved properties to be gained from the addition of a ceramic filler. Resin composites are often classified according to the size of the ceramic filler particle (Fig 8-1 and Tables 8-1 and 8-2).

![Fig 8-1 The microstructure of resin composite materials.](image-url)
Table 8-1 Average properties of resin composites

<table>
<thead>
<tr>
<th>Property</th>
<th>Microfilled</th>
<th>Microhybrid</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inorganic filler content (vol %)</td>
<td>20-55</td>
<td>60-70</td>
</tr>
<tr>
<td>Thermal conductivity</td>
<td>Insulator</td>
<td>Insulator</td>
</tr>
<tr>
<td>Coefficient of thermal expansion (°C × 10^-6)</td>
<td>50-68</td>
<td>20-40</td>
</tr>
<tr>
<td>Hardness (Knoop)</td>
<td>22-36</td>
<td>50-60</td>
</tr>
<tr>
<td>Water sorption (mg/cm^2)</td>
<td>1.2-2.2</td>
<td>0.5-0.7</td>
</tr>
<tr>
<td>Compressive strength (MPa)</td>
<td>225-300</td>
<td>300-350</td>
</tr>
<tr>
<td>Tensile strength (MPa)</td>
<td>25-35</td>
<td>35-60</td>
</tr>
<tr>
<td>Young's modulus (GPa)</td>
<td>3-5</td>
<td>7-14</td>
</tr>
<tr>
<td>Polymerization shrinkage (%)</td>
<td>2-4</td>
<td>1.5-1.7</td>
</tr>
</tbody>
</table>

Table 8-2 Clinical characteristics and selection of resin composites

<table>
<thead>
<tr>
<th>Filler size</th>
<th>Microfilled</th>
<th>Microhybrid</th>
<th>Fine particle</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.01-0.12 um</td>
<td>0.01-0.2 um</td>
<td>0.5-3.0 um</td>
<td></td>
</tr>
<tr>
<td>Appearance</td>
<td>Good gloss, luster, and smoothness</td>
<td>Opaque appearance</td>
<td></td>
</tr>
<tr>
<td>Polishability</td>
<td>Highly polishable</td>
<td>Polishable</td>
<td>Semi polishable</td>
</tr>
<tr>
<td>Usage</td>
<td>Non-stress-bearing esthetic restorations*</td>
<td>Anterior and posterior restorations</td>
<td>Core buildups, conservative posterior restorations</td>
</tr>
</tbody>
</table>

*Only heavily filled microfilled materials may be used for posterior restorations.

Fine-particle resin composites contain ground glass or quartz particles 0.5 to 3.0 um in diameter, which occupy 60% to 77% of the composite by volume. Since the filler has a density greater than that of the polymer matrix, the fraction of the filler by weight is higher, about 70% to 90%. Particles present may be of uniform diameter or have a distribution of diameters, in which case smaller particles fit in the spaces between larger particles, and packing is more efficient.

Microfilled resin composites contain spherical colloidal silica particles 0.01 to 0.12 um in diameter. Colloidal silica is produced by vapor-phase hydrolysis of silicon compounds resulting in an average surface area of 200 m^2/g, which greatly increases the viscosity of the polymer matrix upon incorporation. Filler loading in these composites is therefore limited to about 20% to 55% by volume or 35% to 60% by weight, and low-molecular-weight organic diluents of low viscosity are often added to give the composite a workable clinical consistency. Filler content may be increased and properties improved by grinding a polymerized microfilled composite into particles 10 to 20 um in diameter and subsequently using these reinforced particles as filler along with colloidal silica. Heavily filled microfilled composites have a filler content of 32% to 66% by volume or about 40% to 80% by weight (Fig 8-2).
Hybrid (or blend) resin composites have a combination of colloidal and fine particles as filler. The colloidal particles fill the matrix between fine particles, resulting in a filler content of around 60% to 65% by volume. Hybrids (Figs 8-3a and 8-3b) currently dominate the market.

Microhybrid resin composites, the generic term for advanced composites, are a combination of a microfilled and fine-particle composite and are so called because of their small-diameter (0.4 to 1.0 μm) filler particles. They were recently introduced as all-purpose "universal" composites offering both esthetics and superior wear resistance for use in anterior and posterior applications (Fig 8-3b).
Composition and Reaction

Filler composition

Filler particles are of inorganic composition. In addition to quartz, fine-sized particles may be composed of barium or lithium aluminum silicate glasses; borosilicate glass; or barium, strontium, or zinc glasses. Particles in microfilled composites are colloidal silica.

Composites may be made radiopaque by incorporating elements of high atomic weight, such as barium, strontium, zirconium, or ytterbium, into the filler particle.

Organic matrix

The organic polymer matrix in currently available composites is most commonly an aromatic or urethane diacrylate oligomer such as bisphenol A-glycidyl methacrylate (bis-GMA) or urethane-dimethacrylate (UDMA), represented by the simplified formula:

\[
\begin{align*}
\text{CH}_2 & \quad \text{C} - \text{R} - \text{C} = \text{CH}_2 \\
\text{CH}_3 & \quad \text{CH}_3
\end{align*}
\]

where R may be any of a number of organic groups, such as methyl-, hydroxyl-, phenyl-, carboxyl-, and amide-.

The oligomers have in common reactive double bonds at each end of the molecule, which are able to undergo addition polymerization in the presence of free radicals. The oligomer molecules are highly viscous and require the addition of low-molecular-weight diluent monomers, usually triethylene glycol dimethacrylate (TEGMA), so a clinically workable consistency may be maintained upon the incorporation of the
filler.

**Coupling agents**

A bond between filler particle and matrix in the set composite is achieved by use of an organic silicon compound, or silane **coupling agent**. The silane molecule has reactive groups at both its ends and is coated on the filler particle surface by the manufacturer before mixing with the oligomer. During polymerization, double bonds on the silane molecule also react with the polymer matrix. A bond between filler and matrix allows the distribution of stresses generated under function. The net result is a material with strength properties greater than those of the particulate filler or the matrix separately. Bonding also enhances the retention of the filler particle during abrasive action at the composite surface. As a result, hard filler particles, in addition to soft matrix, are present to engage in abrasive wear with opposing enamel, for example, greatly improving the wear resistance of the material.

**Initiators and accelerators**

**Polymerization** of composites may be achieved by chemical means (self-cure) or by visible-light activation. Dual cure is a combination of light and chemical curing. In chemically activated systems, an organic peroxide initiator (or catalyst), upon reacting with a tertiary amine accelerator, produces free radicals that attack the double bonds of oligomer molecules and begin the process of addition polymerization.

Initiation of polymerization in light-activated systems depends on the scission of the initiator molecule, often camphorquinone, by visible light of appropriate wavelength. In the presence of an aliphatic amine accelerator, free radicals are produced and polymerization begins. For both systems, the following general reaction occurs:

\[
\text{Dimethacrylate} + \text{Initiator} + \text{Accelerator} + \text{Silane-treated particles} \rightarrow \text{Dental composite}
\]

Since **dimethacrylate** oligomers as well as dimethacrylate diluent monomers have reactive double bonds at each end of the molecules, polymerization results in a highly **cross-linked polymer**.

**Other ingredients**

Inorganic oxide pigments are added to composites in small amounts to provide a range of standard shades. Most often, four shades, ranging from yellow to gray, are supplied. In response to consumer interest, manufacturers now offer an extended range of 16 or 25 shades, as well as shade selections from the Bioform and Vita ceramic shade guides. Most manufacturers offer modifiers such as highly pigmented tints for mixing with standard shades, as well as opaquers and glazes.

Polymerization inhibitors and stabilizers are added to the composite to lengthen shelf
life.

**Composite Product Systems**

Composites are packaged as a two-paste system supplied in two jars, or as a single-paste system supplied in a syringe or in compules. An enamel acid etchant, along with a dentin bonding agent, may also be supplied by the manufacturer.

Chemically cured composites supplied as two pastes are typically used as resin cements or for core applications. Each jar contains dimethacrylate and filler; one jar also contains the peroxide initiator, and the other contains the amine accelerator. The initiator and the accelerator are kept separate until mixing. A few composites are offered as two-paste, dual-curing systems. Setting begins after the catalyst and base are mixed and can be accelerated by light curing.

Today's composites designed for restorative applications are supplied as single pastes in opaque, disposable syringes or in color-coded compules for use with a syringe. Light-activated composites are currently the most widely used systems available.

Commercially available curing units transmit light from a halogen lamp to the tooth surface by way of a curved quartz rod, a liquid-filled transmission tube, or a bundle of flexible quartz fibers attached to a fiber-optic handpiece. Ultraviolet light is generally filtered out at the light source.

The initiator present in most photocuring monomers is camphorquinone, which is sensitive to light with the spectrum shown in Fig 8-4. To initiate polymerization, curing lights must emit light within this spectrum, which is in the blue range. Filtered halogen lights produce a broad range of wavelengths within the camphorquinone spectrum and are the standard.

Other lights that have higher intensities for faster polymerization have been introduced. These include plasma arc lamps (PAC) and argon laser lights (Fig 8-4). Although more intense, not all PAC and laser lights have the broad spectrum of the halogen lamps. It is important to match the spectrum of a PAC light to the product being used. The PA 430 has been found to be more effective for certain resin pr

![Fig 8-4 Spectra of curing lights including halogen (HAL), plasma arc (PA), and argon laser, along with absorption spectrum needed for initiation with camphorquinone.](image-url)
Laser curing systems are available with continuous operation and programmed cycles. One program is called a stepping function, which cycles the light on and off to reduce possible overheating of the tooth.

As with ultraviolet light used in early curing units, blue light has the potential to cause retinal damage. Figure 8-5 shows a curing light for composite materials. Protective eyewear during operation of curing units is also available and recommended.

![Halogen blue curing light with variable cycles.](image)

**Properties**

**Setting time**

Composite systems that are chemically activated have setting times ranging from 3 to 5 minutes from the start of mixing. The setting time is determined at the time of manufacture by control of the concentrations of initiator and accelerator. However, studies show that even after a curing time of 24 hours, polymerization is incomplete and 25% to 45% of double bonds remain unreacted.

The setting time and the depth of cure of light-initiated materials depend on the intensity and penetration of the light beam. Polymerization is approximately 75% complete at 10 minutes after exposure to blue light, and curing continues for a period of at least 24 hours. At 24 hours, up to 30% of double bonds still remain unreacted.

**Polymerization shrinkage**
The occurrence of shrinkage during polymerization creates stresses (~18 MPa) at the tooth-composite interface that may exceed the strength of any bond between composite and enamel or dentin. Bond failure at the interface allows an influx of oral fluids and greatly contributes to the possibility of marginal staining, secondary caries, and postoperative sensitivity. In addition, stresses at the tooth-composite interface may exceed the tensile strength of enamel perpendicular to the enamel rods, resulting in fractures through the enamel along the interface.

Shrinkage is a direct function of the volume fraction of polymer matrix in the composite, and thus occurs to a larger degree in microfilled composites than in fine-particle composites or hybrids. Microfilled composites typically show setting contractions of 2% to 4% as compared to 1.0% to 1.7% for fine-particle composites (see Table 8-1).

The shrinkage problem can be partially overcome in two ways. First, incremental addition and polymerization of thin layers of a light-initiated material will result in decreased total setting contraction as opposed to bulk curing a single thick layer. However, although this method does result in lower stresses at the tooth-composite interface, studies show that marginal gaps may occur nonetheless.

A second approach involves the preparation of a composite inlay either directly in the mouth or indirectly as a laboratory procedure. In the latter procedure the inlay is heat processed (see Fig 8-6), allowing the degree of polymerization to approach 100%, and then cemented in the mouth with a thin layer of resin cement. The bulk of the resin composite cement layer needed is small, producing a very small amount of shrinkage and low interfacial stresses. Long-term studies are still underway, and composite inlays produced in this way are expected to show improved durability and increased wear resistance due to enhanced physical properties.

Fig 8-6 Composite laboratory curing unit (Targis Power; courtesy of Ivoclar North America).
**Thermal properties**

The organic polymer matrix has low thermal conductivity, and composites therefore provide good thermal insulation for the dental pulp. Thermal conductivities of all composites closely match those of enamel and dentin and are much lower than that of dental amalgam.

As a consequence of the weak physical bonds by which individual polymer molecules are held together, polymers have a marked tendency to expand and contract in response to temperature changes. In contrast, the highly inorganic content of tooth structure is affected to a much smaller degree. Dimensional changes in the composite resulting from thermal cycling in the mouth produce further strain on the bond at the tooth-composite interface, increasing the possibility of marginal percolation. This effect occurs to a larger extent with resin-rich microfilled composites than with fine-particle materials or hybrids.

**Water sorption and solubility**

The polymer matrix is able to absorb water, which is accompanied by some swelling of the composite, but this is not sufficient to counteract polymerization shrinkage. The uptake of water by composites has been correlated with decreases in surface hardness and wear resistance. As a result of their larger volume fraction of matrix, microfilled composites have higher water sorption values and therefore a greater potential for discoloration by water-soluble stains.

Solubility of resin composites ranges from 1.5% to 2.0% of the original material weight. The major detected leachable components in water include residual oligomer or monomer; therefore, incomplete polymerization of the composite results in markedly increased solubility. Additional leachable molecules include degradation products of various composite components and may include formaldehyde, benzoic acid, and methacrylic acid. The largest part of the dissolution occurs within the first few hours of placement.

Elements from filler particles dissolve in water to varying degrees and are detected in quantities as high as 180 umol/g. Boron and silicon are the main elements, but barium, strontium, and lead, other additives to glass particles, also leach out. The presence of silicon in solution may indicate degradation of the surface treatment of the filler.

Alcohol is a solvent of bis-GMA and acidulated fluoride gels increase the rate of dissolution of filler particles. Therefore, alcohol-free rinses and neutral fluoride products should be used.

**Color stability**

Darkening and a color shift to yellow or gray has often been noted in self-curing systems and has been attributed to the presence of the tertiary amine accelerator, which produces colored products upon oxidation. Photo-initiated systems do not contain a tertiary amine and have shown considerably improved color stability over long periods of time.
Under accelerated aging conditions in a weathering chamber, erosion of the resin matrix and exposure of filler particles of microfilled composites resulted in a lightening of the color of the material. Color stability of microfilled composites, however, was affected by erosion only to a small degree.

Radiopacity

A degree of radiopacity slightly exceeding that of enamel may be useful in diagnosis. Radiopacity may be conferred by incorporating elements of high atomic number, such as barium, strontium, and zirconium, into the filler. The relative number of these atoms is still small, however, and the materials are much less radiopaque than is amalgam. Many composites currently available have some degree of radiopacity, and several for posterior use have radiopacities greater than that of enamel.

Mechanical properties

The higher compressive and tensile strengths of fine-particle and hybrid composites, as compared to microfilled composites, reflect the higher volume fraction of the high-strength filler component. Note that for all materials compressive strengths are several times higher than tensile strengths, reflecting the somewhat brittle behavior of composites. More highly filled composites have tensile strengths near that of dentin and compressive strengths similar to or higher than that of dentin (Table 8-3). Some highly filled composites have compressive strengths greater than that of enamel.

<table>
<thead>
<tr>
<th>Table 8-3 Mechanical properties of enamel and dentin</th>
</tr>
</thead>
<tbody>
<tr>
<td>Compressive strength (MPa)</td>
</tr>
<tr>
<td>---------------------------</td>
</tr>
<tr>
<td>Enamel</td>
</tr>
<tr>
<td>Dentin</td>
</tr>
</tbody>
</table>

The elastic (Young's) modulus is a measure of a material's stiffness. A material with low elastic modulus deflects under stress. As a group, composites have elastic moduli that are only a fraction of that of enamel. Fine-particle materials, however, have moduli similar to that of dentin. Under high loads, such as occur in posterior teeth during mastication, deflection of the restoration strains the tooth-composite bond. Deflection additionally places considerable tensile stresses on adjacent cusps.

The lower filler content of microfilled composites results in elastic moduli of one quarter to one half that of the more highly filled fine-particle composites, and are therefore recommended for cervical (Class 5) restorations since deflection could reduce stresses at the tooth-composite interface.

Microindentation hardness of composites is directly related to volume fraction of the hard, inorganic filler component. Hardness is also related to the degree of polymerization. In laboratory experiments, composites that underwent a secondary heat treatment to increase the degree of polymerization showed higher Knoop hardness values than did composites that were light cured only.
As a group, the hardness of composites is a fraction of that of enamel but is similar to or higher than that of dentin.

**Wear**

Wear in composites is a complex phenomenon that depends on several intrinsic and extrinsic factors. The large amount of data collected for various available composites is confusing, at least in part because measuring techniques have not been standardized. In vivo wear has not been shown to correlate well with any single material property. In addition, the appearance of wear patterns in restorations of long duration is complicated by the presence of erosion, a degradative uniform loss of material across the composite surface; several wear mechanisms are shown in Fig 8-7. Nevertheless, a number of factors that contribute to wear have been identified.

**Fig 8-7** Several possible wear mechanisms for dental composites. (F = filler particle; M = matrix.) (From O'Brien and Yee, 1980. Modified with permission.)

**Filler content, particle size, and hardness**

Increased filler volume results in decreased wear. Laboratory studies demonstrate a greater loss of material volume during abrasive action for microfilled as compared to more highly filled fine-particle composites. A higher filler volume results in a higher fracture toughness, as shown in Fig 8-8.
The effect of filler volume fraction on fracture toughness, $K_{IC}$. (Courtesy of J. Ferracane.)

Keeping volume fraction constant, wear resistance is increased by decreasing the size of the filler particle. Large, hard particles transmit considerable stress to the matrix, possibly resulting in microcracking and subsequent loss of material (Fig 8-9). By contrast, a reduced load per particle results when a large number of small particles is present per unit volume.

It has been reported that wear of composites with filler particles smaller than 1.0 μm occurs at a constant rate with time. Wear of composites with particles larger than 1.0 μm is greatest in the year after placement, declining thereafter.

The incorporation of softer filler particles with hardness characteristics similar to that of enamel appears to result in decreased wear. It is thought that soft particles are more capable than hard particles of absorbing energy generated during the masticatory process, thereby transmitting lower stresses to the matrix. Scanning electron micrographs of a material with hard quartz particles and another with soft glass
particles are shown in Figs 8-9 and 8-10. Note the difference in the shapes of the particles that project from the occlusal surface.

![Worn surface of a composite with a soft glass filler. (SEM, original magnification × 2,500.)](image)

Fig 8-10

Interestingly, the presence of filler particles with hardness values greater than that of enamel has been shown to increase the roughness of opposing enamel over time.

Tooth position in the arch

In general, the more distally located the restoration, the higher the rate of wear.

Porosity

Internal porosity, particularly in stress-bearing areas, increases wear. It has been proposed that porosities concentrate stresses in the matrix and, under loading, contribute to the formation of microcracks. In addition, voids, produced during the spatulation process or during incorporation of filler during manufacture, are air-filled. An air-inhibited layer of incompletely polymerized matrix may exist at the void surface. Light-initiated systems requiring no spatulation have demonstrated higher wear resistance than self-curing systems.

Degree of polymerization

Strength properties of polymers are directly related to molecular size. During polymerization, molecular size increases enormously. The enhanced wear resistance found in some studies of heat-processed composite inlays is thought to be related to their higher degree of polymerization.

Coupling agent

The absence of a silane coupling agent at the matrix-filler interface reduces wear resistance by about half.

Method of finishing
Wear resistance decreases with the use of carbide or diamond finishing burs. Their use is thought to weaken the surface through the formation of microcracks or to degrade the matrix through the generation of heat. It has been reported that treatment of the surface after occlusal adjustment and subsequent curing with a low-viscosity unfilled resin decreases wear by about 50%.

Wear resistance of composites developed for posterior use has improved steadily over recent years.

**Depth of cure**

Polymerization in both chemically and light-activated composites is incomplete. Degrees of conversion are reported to be in the range of 60% to 75%. The number of unreacted double bonds at 24 hours is higher in light-activated than in chemically activated systems and results from a number of factors affecting depth of cure.

Light intensity at the surface and time of exposure are critical. The tip of the light source should be held within 3 to 4 mm of the surface in order to cure a light shade of material to a depth of 2.0 to 2.5 mm using a standard exposure time of 40 seconds. A longer exposure time will increase the degree of polymerization at all depths and is a necessity when using darker shades or more opaque materials. A reduction in thickness of the increment to be cured is a more reliable way of achieving polymerization than an increase in exposure time. In addition, hardness of the top surface of a cured restoration is not a good indication of the extent of polymerization at the bottom surface.

Depth of cure is also influenced by the wavelength of light and the concentration of the activator-initiator system. The refractive indices of the resin and filler, as well as the size, shape, and number of filler particles, are important to the dispersion of the light beam. The small, highly numerous colloidal-sized particles of microfilled composites scatter incident light very efficiently, necessitating a longer exposure time to obtain adequate polymerization.

Chemically activated systems are considered to have an infinite depth of cure.

**Biocompatibility**

Histological studies of the effect of residual monomer molecules on pulp tissue have shown a moderate degree of cytotoxicity, even in low concentrations. Recent in vivo biocompatibility studies, however, show that resin composites, whether completely or incompletely cured, cause little irritation to the pulp if an adequate marginal seal is present. It has been proposed that a significant degree of sensitivity after the placement of a restoration is a consequence of microbial invasion from the oral environment and not of toxicity of the material itself. A second possibility is that postoperative sensitivity may be a consequence of debonding between dentin and the composite at the cavity floor, causing a pumping action of dentinal fluid during chewing (percolation), possibly expressing irritants or bacterial toxins into dentinal tubules.
Until the precise mechanisms involved are understood, pulpal protection is recommended in deep parts of the cavity preparation.

**Manipulation**

**Placement**

Eugenol inhibits the polymerization of resin composites. Therefore, liners, bases, and interim restorations containing eugenol are not recommended. The use of cavity varnish is not recommended under composite restorations, because monomers present in the composite may solubilize and disrupt the integrity of the varnish film. Also, varnish will prevent bonding.

Following cavity preparation and prior to placement of the composite, a sealing procedure of some type is indicated. If any dentin bonding agent is to be used, the use of a rubber dam is indicated because moisture in exhaled air may interfere with bonding. The dentin is first conditioned according to the manufacturer's directions. Deep preparations may require the placement of a glass-ionomer or resin-modified glass-ionomer liner or base over the dentin. Very deep cavities require pulpal protection in the form of a thin layer of a calcium hydroxide product on the dentin over the pulp.

Enamel and dentin are treated strictly according to manufacturer's directions depending on the bonding agent used. Generally, enamel and dentin are etched for 15 seconds using a 35% to 50% phosphoric acid solution or acid gel. High-viscosity gel etchants have the advantage of ease of control of the application to enamel walls. The preparation is thoroughly washed with water for at least 15 seconds to remove all residue. The surface is gently air-dried, at which point the enamel should have an opaque, white appearance. Any contamination by saliva after this step requires re-etching to clean the surface thoroughly.

A dentin bonding agent is applied to the clean enamel and dentin according to the manufacturer's directions. The bonding resin should be air-blown gently to ensure a thin film application. Dentin bonding agents work as well on enamel as do enamel bonding agents. While components of bonding systems should not be interchanged, any composite can be used with most bonding agents.

A transparent matrix band is sometimes placed for the purpose of contouring the restoration.

**Two-paste system**

To ensure uniform distribution of filler particles in the matrix, each paste should be stirred periodically with disposable plastic mixing sticks, taking care to avoid cross-contamination that will cause polymerization of the pastes in the jars. Less particle settling and an increase in shelf life results if the pastes are refrigerated.

Equal amounts of the two pastes are dispensed onto a mixing pad with a disposable two-bladed plastic spatula. One blade of the spatula should be used to dispense one
paste, and the other blade, the second paste. When needed, the two pastes are mixed thoroughly, requiring 20 to 30 seconds. Care should be taken to avoid the incorporation of air during mixing. Metal spatulas are not recommended for mixing, because filler particles are capable of abrading metal and small amounts of metal may be incorporated into the composite, resulting in discoloration.

Two-paste composites have a working time from the start of mixing of 1 to 1.5 minutes and a setting time of 3 to 5 minutes. The mixed material is inserted in one of two ways, depending on the viscosity of the particular product. Viscous material is best placed with plastic instruments that do not adhere to the unset material. A small amount of bonding resin on the tip of the plastic instrument will prevent sticking. If the viscosity is low enough, the material may be injected into the cavity preparation from a syringe. A syringe placement tends to minimize the incorporation of voids into the restoration.

The cavity preparation is slightly overfilled. At 3.5 to 4 minutes after the start of mixing, the matrix band, if used, is removed. After an additional 2 to 6 minutes, the composite surface is sufficiently hard for finishing to begin.

Single-paste system

The shelf life of composites supplied as single pastes and stored in a cool, dry environment is about 1 year.

The composite is best placed in small layers to minimize polymerization shrinkage. Shrinkage per layer placed is less if the tooth-composite bonding area per layer thickness is large. Each layer should be light cured for at least 40 seconds. After curing, a tacky, air-inhibited layer is present, through which the subsequent layer bonds.

Microfilled composites require longer exposure times than do fine-particle composites because their colloidal-sized filler particles scatter blue light more efficiently.

**Finishing**

Composites are finished and polished in order to establish a functional occlusal relationship and a contour that is physiologically in harmony with supporting tissues. In addition, proper contour and high gloss give the restoration the appearance of a natural tooth structure. Early composites had large, hard quartz particles. Polishing preferentially removed the resin matrix, leaving filler particles exposed and giving the surface a dull appearance. In addition, quartz has a hardness about 2.5 times that of enamel and is difficult to polish compared to glasses, which have hardness characteristics similar to that of enamel.

Particles smaller than about 0.05 um cannot be detected visually and allow polishing to a high luster. Fine-particle composites have no microfilled particles, are considered to be only semipolishable, and tend to have a rather opaque appearance. The colloidal-sized filler particles of microfilled materials scatter light efficiently, giving these restorations a pleasing esthetic appearance. Hybrid composites are polishable, but are not as translucent as microfilled composites.
Common prophylaxis pastes are very abrasive to composites and should not be used for polishing.

The composite surface may be contoured with a plastic matrix strip, but some gross reduction is often required. Finishing begins with coarse abrasives, such as 9- and 12-bladed finishing burs and fine diamonds, and progresses to 16- and 30-bladed finishing burs, ultrafine diamonds, and medium-grit abrasive points, disks, and strips. Polishing is accomplished with aluminum oxide polishing paste on a rubber cup at low speed.

A postocclusal adjustment cure of 40 seconds further hardens the surface. It is recommended that this step be followed by re-etching and the application and curing of a low-viscosity unfilled resin. The resin penetrates microcracks on the finished surface, further reducing wear.

Pitting has been observed on the surface of microfilled-particle-reinforced composites after polishing. The pits are thought to occur at the junction of the new resin and the prepolymerized resin/filler, resulting in an increased susceptibility to chipping.

A properly placed anterior composite restoration has an expected clinical life of 7 to 10 years. Posterior restorations are generally serviceable for less than that. The major reasons for replacing any composite include deterioration of esthetics, interfacial staining, wear, microleakage, and secondary caries.

**Material Selection**

*Introduction*

A variety of materials are available for use as direct esthetic dental restorations: resin composites, compomers, hybrid ionomers, and glass ionomers. Each has a resin component, except the traditional glass ionomer. The unique characteristics of each of these materials make them desirable for a range of applications. Choosing the most suitable material requires matching a material's desired properties and manipulation to application for use. The continuum of resin composite to glass-ionomer esthetic restorative materials is illustrated in [Fig 8-11](#) and their uses are presented in [Table 8-4](#).
**Table 8-4** Direct esthetic materials and their uses

<table>
<thead>
<tr>
<th>Restorative Material</th>
<th>Application</th>
</tr>
</thead>
<tbody>
<tr>
<td>Flowable resin composite</td>
<td>Restorations following air abrasion, small Class 3 and 5 restorations, liner/base under packable composite</td>
</tr>
<tr>
<td>Hybrid and microhybrid resin composite</td>
<td>Anterior restorations, posterior restorations</td>
</tr>
<tr>
<td>Microfilled resin composite</td>
<td>Areas of esthetic concern, anterior restorations</td>
</tr>
<tr>
<td>Packable resin composite</td>
<td>Class 1 and 2 restorations, class 3 and 5 restorations</td>
</tr>
<tr>
<td>Polyacid-modified resin composite (compomer)</td>
<td>Cervical erosion or abrasion lesions, class 1 and 2 restorations in pediatric patients, sandwich technique for Class 2 restorations</td>
</tr>
<tr>
<td>Glass ionomer</td>
<td>High caries-risk patients, cervical erosion or abrasion lesions, class 5 restorations where esthetics is not a concern</td>
</tr>
<tr>
<td>Resin-modified glass ionomer (hybrid ionomer)</td>
<td>Class 3 and 5 restorations, cervical erosion or abrasion lesions, class 1 restorations in pediatric patients, sandwich technique or Class 2 restorations, liners, bases, and luting cements</td>
</tr>
</tbody>
</table>

**Resin composites**

Composites are the most commonly used direct esthetic restorative material today. They offer excellent esthetics, strength, and wear resistance but have little or no fluoride release. Microfilled composites are used for anterior applications, and hybrid composites are recommended for posterior applications; however, the difference between posterior and anterior composites has become less distinct due to the development of the microhybrid composites.

**Fig 8-11** Continuum of glass-ionomer and composite restorative materials. (After Albers, 1996).
Composites can also be selected based on their handling characteristics. **Packable composites** have been termed an alternative to amalgam (Fig 8-12).

![Fig 8-12 A packable composite material. An interlocking particle network allows the packable handling properties.](image)

They differ from anterior and posterior composites in that they have higher filler loading (greater than 80% by weight and 66% to 70% by volume) with either fibers, porous filler particles, irregular filler particles, or viscosity modifiers. They are recommended for use in Class 1 and 2 cavity preparations. Because of the high depth of cure and lower polymerization shrinkage of packable composites, a bulk-fill technique may be possible. A low-viscosity (flowable) material (Fig 8-13) can be used as a liner to fill irregular internal surfaces and proximal boxes before placing the packable composite. Flowable composites or compomers will wet and adapt to the tooth surface, sealing it and possibly reducing postoperative sensitivity.

![Fig 8-13 A flowable composite with a filler load of about 60% by weight and average particle size of 0.6 um.](image)
Compomers

Compomers, or polyacid-modified composites (Fig 8-14), contain monomers modified by polyacid groups with fluoride-releasing silicate glasses. The properties and clinical characteristics of compomers are directly related to their composition of composite and glass ionomer. Compomers release fluoride by a mechanism similar to glass and hybrid ionomers, but the amount of release and its duration are lower. Additionally, compomers do not recharge from fluoride treatments or brushing with fluoride dentifrices as much as do glass and hybrid ionomers. Like composites, compomers require a bonding agent to bond to tooth structure. Single-bottle bonding agents that contain acidic primers have been used with some compomers. Acidic primers will bond to both enamel and dentin without the need for phosphoric acid etching. However, studies have shown the bond strength of compomers to teeth is improved with additional etching with phosphoric acid. Compomers set by light-activated polymerization and by an acid-base reaction with water absorbed from the tooth. They are packaged as single-paste formulations in compules and syringes. Compomers are recommended for cervical lesions, Class 3 and 5 restorations in adults, primary teeth, Class 1 and 2 restorations in children, sandwich technique for Class 2 restorations, and with medium-caries-risk patients.

Resin-modified glass ionomers

Resin-modified glass ionomers, or hybrid ionomers, contain fluoride-releasing glass and polyacid monomers. They set by an acid-base reaction and both chemical and light-cured free-radical reactions. They are recommended for cervical lesions, Class 3 and 5 restorations in adults, primary teeth, Class 1 restorations in children, sandwich technique for Class 2 restorations, and with high-caries-risk patients. Because of the resin content, these restorations are more esthetic and stronger than glass ionomers.
Hybrid ionomers are packaged in powder-liquid form or are pre-encapsulated to be mixed in an amalgamator. The use of a bonding agent prior to placement is not recommended because it decreases fluoride release. Hybrid ionomers naturally bond to tooth structure once the tooth is conditioned with polyacrylic acid or a primer. See also chapter 9 for the use of hybrid ionomers as cements. A comparison of the properties of several of these materials is given in Table 8-5.

Table 8-5 Mechanical properties of glass ionomer and resin composite combinations*

<table>
<thead>
<tr>
<th>Materials class</th>
<th>Flexural strength (MPa)</th>
<th>Flexural modulus (GPa)</th>
<th>Compressive strength (MPa)</th>
<th>Diametral tensile strength (MPa)</th>
<th>Shear bond strength (MPa)</th>
<th>Shrinkage (% vol)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Conventional glass ionomer</td>
<td>25</td>
<td>8</td>
<td>180-200</td>
<td>22-25</td>
<td>3-5</td>
<td>3</td>
</tr>
<tr>
<td>Resin-modified glass ionomer</td>
<td>35-70</td>
<td>4</td>
<td>170-200</td>
<td>35-40</td>
<td>8-14</td>
<td>3.5</td>
</tr>
<tr>
<td>Compomer</td>
<td>97</td>
<td>6-8</td>
<td>210-245</td>
<td>45-47</td>
<td>14</td>
<td>4.5</td>
</tr>
<tr>
<td>Fluoride-releasing resin composite</td>
<td>85</td>
<td>8</td>
<td>—</td>
<td>40</td>
<td>24</td>
<td>3</td>
</tr>
<tr>
<td>Flowable resin composite</td>
<td>111-167</td>
<td>4-8</td>
<td>—</td>
<td>—</td>
<td>20-22</td>
<td>5</td>
</tr>
<tr>
<td>Resin composite</td>
<td>140</td>
<td>18</td>
<td>350</td>
<td>65-75</td>
<td>24-28</td>
<td>3</td>
</tr>
</tbody>
</table>

*From Burgess et al (1996). Used with permission from The Compendium of Continuing Education in Dentistry.

Pit and Fissure Sealants

These sealants are mainly fluid resins that are applied to posterior occlusal surfaces of caries-prone teeth to seal pits and fissures from bacterial action. The principal monomer used is bis-GMA, which may be lightly filled with ceramic filler particles to improve wear resistance. Prior to application of the fluid monomer, the tooth enamel is etched with phosphoric acid. Upon application of the fluid resin, it penetrates into the pits and fissures by capillary action. There also is penetration into the microstructure of the etched enamel to form "tags" that lead to micromechanical action. Deep penetration is promoted by a high penetration coefficient (PC) which results from low viscosity, high surface tension, and low contact angle, that is, good wetting (chapter 5). Figure 8-15 shows the penetration of a sealant into a fissure.
Sealants have been found to be effective in several studies. Long-term studies have shown around 50% decay reductions in sealed teeth compared with controls. About 90% of sealants are at least partially present in 5-year studies. Also, partial loss or wear of a sealant does not prevent its preventive action in deep fissures. One disadvantage to using sealants is that they may trap food debris and bacteria that could lead to continued caries action. Studies have shown that although some food debris is usually trapped under sealants, the bacteria become dormant if they are isolated. Fluoride-containing sealants are also available.

### Clinical Decision Scenarios for Restorative Materials

This section presents a formal approach to choosing restorative materials for specific situations. Each scenario uses the same format as that presented in chapter 7. Advantages and Disadvantages of each material are prioritized using the following codes: * = of minor importance, ** = important, *** = very important.

**Situation**  Light-cured vs chemically cured composite materials for anterior Class 3 restorations

**Description**  A patient has a small proximal caries lesion on the mesial surface of a maxillary permanent central incisor. He desires an esthetic restoration. The dentist has both visible light-cured composite material and chemically cured material in his armamentarium.

**Critical Factor**  Esthetics

**Light cure**

**Advantages**

*** Good esthetics

** Better color stability
Less porosity
More working time

_Disadvantages_
Less complete cure
Curing light needed

_Chemical cure_

_Advantages_
*** Good esthetics
* More complete cure
Curing light not needed

_Disadvantages_
More porosity
** Poorer color stability
Less working time

**Analysis/Decision** Although either material would be satisfactory, the greater color stability provides the determining factor in this instance and leads to selection of the light-cured material. The greater working time of light-cured material is convenient but not necessary to a dentist experienced in the use of the chemically cured material. The higher porosity of the chemically cured material—induced by the need to mix the paste and catalyst—as well as the activators and accelerators needed in the chemical cure, probably contribute to the lower color stability.

**Situation** Resin composite vs amalgam for cores

_Description_ A patient comes to the office complaining of a broken tooth. Upon examination, the dentist finds that the lingual cusps of the mandibular right permanent first molar have fractured at the gingival line. The tooth already has a large mesio-occlusodistal (MOD) amalgam in it, and the facial cusps are not sturdy. The patient is leaving for a winter vacation to Florida in 10 days and hopes to have the tooth repaired before then. The treatment plan is to prepare a pin-retained core buildup and then a full gold crown. The dentist has both amalgam and a composite core material available.
Critical Factor Time

Amalgam

Advantages

Greater strength
Long clinical history

Disadvantages

*** 24-hour set
No bond to tooth

Resin composite

Advantages

*** Immediate set
Bonds to tooth
Can be prepared immediately

Disadvantages

Lesser strength

Analysis/Decision Although this may seem like an abbreviated list, these are essentially the main considerations. While the strength of the amalgam is thought to be greater in these applications, composites have served well if properly done. The deciding factor is often that the composite can be placed and prepared for the crown immediately, whereas it is recommended that the amalgam be allowed to set 24 hours before preparation. Thus, the composite may be placed and the tooth prepared in one appointment. If amalgam had been the material of choice, a second appointment would have been necessary for the crown preparation. This was the deciding factor that made the dentist choose the composite in this instance.

Materials Microfilled resin composite vs hybrid composite for direct veneer

Description A recent college graduate requests an improvement in the appearance of her maxillary lateral incisors, which are slightly crowded, and she wants it done before a job interview in 2 weeks. She is advised that orthodontic treatment would be a more permanent solution, but it would take 1 or 2 years. Due to urgency, the dentist agrees to place direct veneers using either a microfilled or hybrid composite material.
**Critical Factors**  Appearance, strong bond to enamel, strength

**Hybrid composite**

*Advantages*

- Ease of placement
  - * Stronger than microfilled composite
  - ** More wear resistance
    - Lower expansion and contraction
  - *** Strong bond to enamel

*Disadvantages*

  - *** Less natural appearance
    - * Less smooth finish

**Microfilled composite**

*Advantages*

- Ease of placement
  - *** More natural appearance
    - * Smoother finish
  - *** Strong bond to enamel

*Disadvantages*

  - ** Less wear resistance
    - * Weaker than hybrid
  - Higher expansion and contraction

**Analysis/Decision**  Given that the microfilled composite material has a smoother and more natural appearance, the dentist chose to use this material. No tooth removal was necessary. The lateral incisors were isolated, cleaned with pumice, and enamel etched, then natural-looking restorations were created. The patient was delighted with the results. The dentist explained to the patient that these restorations may need to be resurfaced/polished on a regular basis and replaced every 4 or 5 years due to wear.
Glossary

**acid etch** Selective etching of portions of the enamel rods with phosphoric acid, resulting in both high surface area and increased surface energy. Resin is able to flow into the enamel substructure and upon polymerization provides a mechanical bond to the enamel.

**compomer (polyacid-modified resin composites)** A blend of resin composite and glass ionomer; requires a bonding agent to bond to tooth.

**coupling agent** A chemical attached to the filler surface for the purpose of creating a bond with the resin matrix upon polymerization. The presence of a coupling agent improves several properties of the composite, including strength and wear resistance.

**cross-linked polymer** A polymer with a three-dimensional network structure.

**dentin bonding agent** Low-viscosity resin used for adhesion to enamel and/or dentin.

**dimethacrylate** A methacrylate monomer with reactive, or polymerizable, groups at each end.

**fine-particle resin composites** Composites containing fine-sized filler particles (0.5 to 3.0 µm). Less polishable than microfilled composites.

**hybrid ionomer** A resin-modified glass ionomer that sets by an acid-base reaction and resin polymerization. The powder is similar to glass ionomer and the liquid contains monomers, polyacids, and water.

**hybrid (blend) resin composites** Composites containing colloidal silica (0.01 to 0.12 µm) in addition to fine-sized particles.

**microfilled resin composites** Composites containing colloidal silica as filler. Most polishable and translucent of the resin composites.

**microhybrid resin composite** Composites containing filler particles in the range of 0.4 to 1.0 µm; a blend of hybrid and microfilled composites.

**monomer** A single molecule with double or triple bonds that are capable of uniting the monomers into oligomers or polymers.

**oligomer** A short polymer made up of two to four monomer units.

**packable composites** A composite with a higher filler content than traditional resin composites with either fibers, porous filler particles, irregular filler particles, or viscosity modifiers.

**polymer** A macromolecule formed by the linkage of monomers or oligomers.

**polymerization** The process by which a polymer is formed from monomers or oligomers.
oligomers.

**resin composite** A physical mixture of silicate glass particles (filler) with an acrylic monomer (matrix phase) that is polymerized during application. A material having two or more distinct components and with properties different from those of the individual components.

**Discussion Questions**

1. Why are these materials called "polymeric restorative materials" when they usually contain mostly ceramic phases?

2. How have composites overcome many of the problems of unfilled resin restorative materials?

3. How well are composite materials competing against dental amalgam?

4. Although microfilled composite materials have inferior mechanical properties, they continue to be widely used. Why?

**Study Questions**

*(See Appendix E for answers.)*

1. What is the general composition of resin composite restorative materials?

2. What are the applications for resin composite restorative materials?

3. How long do composite restorations last?

4. What are the advantages of fine-particle composites?

5. What are the advantages of microfilled composites?

6. What is a hybrid, or blend, composite?

7. What is polymerization shrinkage, and why is it a problem?

8. Which restorative materials release fluoride? List them in order of amount released.

9. What is the material of choice for cervical restorations?

**Recommended Reading**


Armstrong SR, Seol J, Boyer DB (1999). Fracture toughness of direct tooth-colored


Chapter 9. Dental Cements

Introduction

Although dental cements are used only in small quantities, they are perhaps the most important materials in clinical dentistry because of their application as (1) luting agents to bond preformed restorations and orthodontic attachments in or on the tooth, (2) cavity liners and bases to protect the pulp and foundations and anchors for restorations, and (3) restorative materials. This multiplicity of applications requires more than one type of cement because no one material has yet been developed that can fulfill the varying requirements.

Over the last two decades, the emphasis has been on materials for luting in view of the increased use of fixed partial dentures. More recently, with the advent of glass-ionomer cements, interest in restorative applications has revived also. These different applications require different physical properties and appropriate clinical manipulative characteristics, and so, in response to the changing situation, new international standards are being developed (International Standards Organization [ISO]), as are various national standards (American National Standards Institute/American Dental Association [ANSI/ADA]) based on performance criteria rather than specific composition.

For acceptable performance in luting and restorative applications, the cement must have adequate resistance to dissolution in the oral environment. It must also develop an adequately strong bond through mechanical interlocking and adhesion. High strength in tension, shear, and compression is required, as is good fracture toughness to resist stresses at the restoration-tooth interface. Good manipulation properties, such as adequate working and setting times, are essential for successful use. The manipulation, including dispensation of the ingredients, should allow for some margin of error in practice. The material must be biologically acceptable.

Most cements are powder-liquid materials that may be dispensed and mixed manually or predispensed in capsules that are mixed mechanically. Some recent materials are composed of two pastes. Cements set by chemical reaction between the ingredients (often an acid-base reaction) or involve polymerization of a monomeric component.

In the early 20th century zinc oxide-phosphoric acid, zinc oxide-eugenol (clove oil 85%), and silicate glass-phosphoric acid cements were discovered. These zinc phosphate, zinc eugenate, and silicate cements were widely used until the 1970s, when new cements began to be developed.

The introduction of new types of cements was prompted by the emphasis on improved biocompatibility and bonding to the tooth that began to develop 20 years ago. New information on pulpal histopathology resulting from particular clinical techniques and materials, as well as the demonstration of marginal leakage involving penetration of bacteria to the dentin interface and a reduction in retention of restorations, led to the realization that new materials possessing good wetting and bonding to enamel and dentin and low toxicity were needed.

These concepts were the basis of the development of cements based on polyacrylic
acid: first the zinc polyacrylate (polycarboxylate) cements, then the glass-ionomer cements, and more recently the resin cements and hybrid ionomer cements. The newer cements have gradually become established as alternatives to zinc phosphate cement because of their minimal effects on pulp, similar strength and solubility characteristics, and adhesive properties.

The advent of the acrylic resins led to the development of poly(methyl methacrylate) in the mid-1950s. These materials had limitations such as lack of adhesion, leakage, and toxicity that terminated their use for routine cementation. In the last 15 years, polymerizable bis-GMA and other dimethacrylate monomer cements have become available in various forms for attachment of cast restorations and orthodontic brackets to enamel. More recently, similar systems containing (potentially) adhesive monomers have been marketed for fixed partial denture cementation.

The cements based on the reaction between calcium hydroxide and a liquid salicylate also originated 25 years ago. They were primarily fluid, two-paste materials intended for the lining of deep cavities that had actual or potential exposure, thus providing an antibacterial sealing action to facilitate the formation of reparative dentin. The susceptibility to acid erosion of the original formulations, both through marginal leakage of restorations and exposure to phosphoric acid during acid-etch techniques, has resulted in more resistant compositions and, quite recently, to a light-cured, resin-based material.

As a result of the research of the last 15 years, cements of five basic types are available, classified according to the matrix-forming species, as shown in Fig 9-1 and Table 9-1:

1. Phosphate bonded
2. Phenolate bonded
3. Polycarboxylate bonded
4. Dimethylacrylate bonded
5. Polycarboxylate and dimethylacrylate combinations
**Fig 9-1** Classification of dental cements according to bonding mechanism.

**Table 9-1** Classification of dental cements

<table>
<thead>
<tr>
<th>Type (matrix bond)</th>
<th>Class of cement</th>
<th>Formulations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Phosphate</td>
<td>Zinc phosphate</td>
<td>Zinc phosphate&lt;br&gt;Zinc phosphate fluoride&lt;br&gt;Zinc phosphate copper oxide/salts&lt;br&gt;Zinc phosphate silver salts&lt;br&gt;Zinc silicophosphate&lt;br&gt;Zinc silicophosphate mercury salts</td>
</tr>
<tr>
<td>Phenolate</td>
<td>Zinc oxide-eugenol</td>
<td>Zinc oxide-eugenol&lt;br&gt;Zinc oxide-eugenol polymer&lt;br&gt;Zinc oxide-eugenol EBA/alumina&lt;br&gt;Calcium hydroxide salicylate&lt;br&gt;Calcium hydroxide salicylate</td>
</tr>
<tr>
<td>Polycarboxylate</td>
<td>Zinc polycarboxylate</td>
<td>Zinc polycarboxylate&lt;br&gt;Zinc polycarboxylate fluoride&lt;br&gt;Glass ionomer&lt;br&gt;Calcium aluminum polyalkenoate&lt;br&gt;Calcium aluminum polyalkenoate-polymethacrylate</td>
</tr>
<tr>
<td>Resin</td>
<td>Acrylic</td>
<td>Poly(methyl methacrylate)&lt;br&gt;Dimethacrylate&lt;br&gt;Dimethacrylate unfilled&lt;br&gt;Dimethacrylate filled&lt;br&gt;Adhesive&lt;br&gt;I-META</td>
</tr>
<tr>
<td>Resin-modified glass ionomers</td>
<td>Hybrid ionomers</td>
<td>Self cured Light cured</td>
</tr>
</tbody>
</table>

Numerous brands of each type are available, and there is some overlap between properties. Since clinical and in vivo evaluation of cements is still very limited, the predictive value of laboratory data for assessment of clinical performance requires knowledgeable interpretation, especially because generalizations on specific types of
cement cannot be made on the basis of the behavior of one or two brands. The applications of the different types of cements are given in Table 9-2. Examples of current widely used brands of permanent luting cements are given in Table 9-3. Typical properties of luting cements are given in Table 9-4.

### Table 9-2 Selection of dental cements

<table>
<thead>
<tr>
<th>Application</th>
<th>Cement type</th>
</tr>
</thead>
<tbody>
<tr>
<td>Luting inlays, crown posts, multiretainers, fixed partial denture in or on:</td>
<td>Glass-ionomer cement, hybrid ionomers, resin</td>
</tr>
<tr>
<td>Nonvital teeth or teeth with advanced pulpal recession and average retention</td>
<td>Zinc phosphate</td>
</tr>
<tr>
<td>Vital teeth with average retention, average pulpal recession, thin dentin, especially for single units and small-span fixed partial dentures</td>
<td>Zinc polycarboxylate</td>
</tr>
<tr>
<td>Multiretainer splints on vital teeth with above-average retention, minimal dentin thickness; hypersensitive patients</td>
<td>Zinc oxide-eugenol polymer</td>
</tr>
<tr>
<td>Provisional cementation</td>
<td>Zinc oxide-eugenol polymer</td>
</tr>
<tr>
<td>Provisional cementation and stabilization of old, loose restorations; fixation of facings and acid-etched cast restorations</td>
<td>Dimethacrylate resin composite</td>
</tr>
<tr>
<td>Base/liner in:</td>
<td>Glass-ionomer cement, resin ionomer</td>
</tr>
<tr>
<td>Cavity with remaining dentin greater than about 0.5 mm</td>
<td>Zinc polycarboxylate</td>
</tr>
<tr>
<td>Cavity with minimal dentin or exposure</td>
<td>Zinc phosphate (low-acid type)</td>
</tr>
<tr>
<td>Calcium hydroxide salicylate</td>
<td>Zinc oxide-eugenol polymer</td>
</tr>
</tbody>
</table>

### Table 9-3 "Permanent" luting cements

<table>
<thead>
<tr>
<th>Product</th>
<th>Type</th>
<th>Manufacturer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fuji Plus</td>
<td>Hybrid ionomer</td>
<td>GC America</td>
</tr>
<tr>
<td>Vitremer Luting</td>
<td>Hybrid ionomer</td>
<td>3M/Espe</td>
</tr>
<tr>
<td>Fleck's Extraordinary</td>
<td>Zinc phosphate</td>
<td>Mizzy</td>
</tr>
<tr>
<td>Fuji I</td>
<td>Glass ionomer</td>
<td>GC America</td>
</tr>
<tr>
<td>Fynal</td>
<td>Zinc oxide-eugenol</td>
<td>L.D. Caulk</td>
</tr>
<tr>
<td>Hy-Bond Polycarboxylate cement</td>
<td>Zinc carboxylate</td>
<td>Shofu Dental</td>
</tr>
<tr>
<td>Hy-Bond Zinc Phosphate cement</td>
<td>Zinc phosphate</td>
<td>Shofu Dental</td>
</tr>
<tr>
<td>Ketac-Cem</td>
<td>Glass ionomer</td>
<td>3M/Espe</td>
</tr>
<tr>
<td>Advance</td>
<td>Hybrid ionomer</td>
<td>L.D. Caulk</td>
</tr>
<tr>
<td>Modern Tenacin</td>
<td>Zinc phosphate</td>
<td>L.D. Caulk</td>
</tr>
<tr>
<td>Super EBA</td>
<td>Zinc oxide-eugenol</td>
<td>Bosworth</td>
</tr>
<tr>
<td>Tylok Plus</td>
<td>Zinc carboxylate</td>
<td>L.D. Caulk</td>
</tr>
<tr>
<td>Zinc Cement Improved</td>
<td>Zinc phosphate</td>
<td>Mission White Dental</td>
</tr>
</tbody>
</table>
Phosphate-Based Cements

Zinc phosphate cement

Applications

Because of their long history, these materials have the widest range of applications, from the cementation (luting) of fixed cast alloy and porcelain restorations and orthodontic bands (Table 9-3) to their use as a cavity liner or base to protect pulp from mechanical, thermal, or electrical stimuli.

Composition and setting

The powder is mainly zinc oxide with up to 10% magnesium oxide and small amounts of pigments. It is fired at high temperature (> 1,000°C) for several hours to reduce its reactivity. The liquid is an aqueous solution of phosphoric acid containing 45% to 64% $\text{H}_3\text{PO}_4$ and 30% to 55% water. The liquid also contains 2% to 3% aluminum and 0% to 9% zinc. Aluminum is essential to the cement-forming reaction, whereas zinc is a moderator of the reaction between powder and liquid, allowing adequate working time and permitting a sufficient quantity of powder to be added for optimum properties in the cement.

Some zinc phosphate cements have modified compositions. One material, widely used as a cavity liner, has 8% aluminum and only 25% $\text{H}_3\text{PO}_4$ in the liquid and a powder that contains calcium hydroxide. Others may contain fluoride and have as much as 10% stannous fluoride.

The amorphous zinc phosphate formed binds together the unreacted zinc oxide and other components of the cement. The set cement consists of a cored structure of residual zinc oxide particles in a phosphate matrix:

$$\text{zinc oxide} + \text{phosphoric acid} \rightarrow \text{amorphous zinc phosphate}$$

Manipulation

The measurement of components and the timing of mixing are essential to consistent success. The mixing slab must be thoroughly dried before use. The powder is added to the liquid in small portions to achieve the desired consistency. Dissipation of the heat of reaction by mixing over a large area on a cooled slab will allow a greater incorporation of powder in a given amount of liquid. The cement must be undisturbed until the end of the setting time. The cement liquid is kept sealed with a stopper to prevent changes in the water content. Cloudy liquid should be discarded. Increasing the powder/liquid ratio gives a more viscous mix, shorter setting time, higher strength, lower solubility, and less free acidity. Use of a chilled (5°C) thick glass slab slows the initial reaction and allows incorporation of more powder, giving superior properties in the set cement.

Properties
The long persistence of zinc phosphates in clinical practice indicates that reasonable performance is obtained. Although the properties are far from ideal, they are usually regarded as a standard against which to compare newer cements (see Table 9-4). The principal reasons for their satisfactory performance under routine conditions are that they can be easily manipulated and that they set sharply to a relatively strong mass from a fluid consistency.

**Table 9-4** Properties of dental luting cements

<table>
<thead>
<tr>
<th>Material</th>
<th>Film thickness (mm)</th>
<th>Setting time (min)</th>
<th>Solubility (wt%)</th>
<th>Compressive Strength (MPa)</th>
<th>Tensile Modulus of elasticity (GPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zinc phosphate</td>
<td>25-35</td>
<td>5-14</td>
<td>0.2 max</td>
<td>80-100</td>
<td>5-7</td>
</tr>
<tr>
<td>Zinc oxide-eugenol Unmodified</td>
<td>25-35</td>
<td>2-10</td>
<td>1.5</td>
<td>2-25</td>
<td>1-2</td>
</tr>
<tr>
<td>Polymer reinforced EBA-alumina</td>
<td>40-60</td>
<td>7-13</td>
<td>1</td>
<td>55-70</td>
<td>3-6</td>
</tr>
<tr>
<td>Zinc polycarboxylate</td>
<td>20-25</td>
<td>6-9</td>
<td>0.06</td>
<td>55-85</td>
<td>8-12</td>
</tr>
<tr>
<td>Glass ionomer</td>
<td>25-35</td>
<td>6-9</td>
<td>1</td>
<td>90-140</td>
<td>6-7</td>
</tr>
<tr>
<td>Polymer based</td>
<td>20-60</td>
<td>3-7</td>
<td>0.05</td>
<td>70-200</td>
<td>25-40</td>
</tr>
</tbody>
</table>

For a given brand of material, the properties are a function of the powder/liquid ratio. For a given cementing consistency, the higher the powder/liquid ratio, the better the strength properties and the lower the solubility and free acidity.

At room temperature (21°C to 23°C) the working time for most brands at luting consistency is 3 to 6 minutes; the setting time is 5 to 14 minutes. Extended working times and shorter setting times can be achieved by use of a cold mixing slab, which permits up to an approximate 50% increase in the amount of powder, improving both strength and resistance to dissolution.

The cement must have the ability to wet the tooth and restoration, flow into the irregularities on the surfaces it is joining, and fill in and seal the gaps between the restoration and the tooth. The minimum value of film thickness is a function of powder particle size, powder/liquid ratio, and mix viscosity. As measured by ISO and ANSI/ADA specifications, acceptable cements give film thicknesses of less than 25 µm. In practice, the cement fills in the inaccuracies between the restoration and the tooth and allows most castings to seat satisfactorily. Unless escapeways or vents are provided with full crowns, separation of powder and liquid may occur, with marginal defects in the cement film.

At the recommended powder/liquid ratio (2.5 to 3.5 g/mL), the compressive strength of the set zinc phosphate cement is 80 to 110 MPa (11,000 to 16,000 psi) after 24
hours. The minimum strength for adequate retention of restorations is about 60 MPa (8,500 psi). The strength is strongly and almost linearly dependent on powder/liquid ratio. The tensile strength is much lower than the compressive strength, 5 to 7 MPa (700 to 900 psi), and the cement shows brittle characteristics. The modulus of elasticity (stiffness) is about 13 GPa ($1.8 \times 10^6$ psi).

According to the standard method, the solubility and disintegration in distilled water after 23 hours may range from 0.04% to 3.3% for inferior material. The standard limit is 0.2%. The fluoride-containing cements give a figure of about 0.7% to 1.0% because of the leaching of fluoride. The solubility in organic acid solutions, such as lactic or citric acid, is 20 to 30 times higher. These data are only a rough guide to solubility under oral conditions. The comparative evaluation of cement solubility under clinical conditions has shown significant loss but conflicting results. Dissolution contributes to marginal leakage around restorations and bacterial penetration. This may be facilitated by dimensional change. The cement has been found to contract about 0.5% linearly, giving rise to slits at the tooth-cement and cement-restoration interfaces.

Biologic effects

The freshly mixed zinc phosphate is highly acidic with a pH of between 1 and 2 after mixing. Even after setting 1 hour, the pH may still be below 4. After 24 hours, the pH is usually 6 to 7. Pain on cementation is due not only to the free acidity of the mix but also to osmotic movement of fluid through the dentinal tubules. Hydraulic pressure developed during seating of the restoration may also contribute to pulpal damage. Prolonged pulpal irritation, especially in deep cavities that necessitate some form of pulpal protection, may be associated with the extended duration of the set material's low pH. This is minimized by a high powder/liquid ratio and rapid setting. One material that has a low acid content and incorporates calcium hydroxide has little effect on pulp when used as a liner. Very thin mixes will also lead to etching of the enamel.

Advantages and Disadvantages

The main advantages of zinc phosphate cements are that they can be mixed easily and that they set sharply to a relatively strong mass from a fluid consistency. Unless the mix is extremely thin (for instance, with a very low powder/liquid ratio), the set cement has a strength that is adequate for clinical service, so manipulation is less critical than with other cements.

However, zinc phosphates' distinct Disadvantages include pulpal irritation, lack of antibacterial action, brittleness, lack of adhesion, and solubility in oral fluids.

**Modified zinc phosphate cements**

Copper and silver cements

Black copper cements contain cupric oxide (CuO); red copper cements contain cuprous oxide (Cu$_2$O). Others may contain cuprous iodide or silicate. Since a much lower powder/liquid ratio is necessary to obtain satisfactory manipulation characteristics with these cements, the mix is highly acidic, resulting in much greater
pulpal irritation. Their solubility is higher and their strength is lower than zinc phosphate cements. Their bacteriostatic or anticariogenic properties seem to be slight. Silver cements generally contain a small percentage of a salt such as silver phosphate. Their advantages over zinc phosphate cement have not been substantiated.

**Fluoride cements**

Stannous fluoride (1% to 3%) is present in some orthodontic cements. These materials have a higher solubility and lower strength than zinc phosphate cement due to dissolution of the fluoride-containing material. Fluoride uptake by enamel from such cements results in reduced enamel solubility and potentially anticariogenic effects.

**Silicophosphate cements**

These materials have been available for many years as a combination of zinc phosphate and silicate cements. The presence of the silicate glass provides a degree of translucency, improved strength, and fluoride release.

Applications

Their principal applications have been for the cementation of fixed restorations and orthodontic bands (Type I), as a provisional posterior restorative material (Type II), and as a dual-purpose material (Type III).

Composition and setting

The powder in these materials consists of a blend of 10% to 20% zinc oxide (zinc phosphate cement powder) and silicate glass (silicate cement powder) mechanically mixed or fused and reground. The silicate glass usually contains 12% to 25% fluoride. Some materials have been labeled "germicidal" because of the presence of small amounts of mercury or silver compounds. The liquid is a concentrated orthophosphoric acid solution containing about 45% water and 2% to 5% aluminum and zinc salts.

The setting reaction has not been fully investigated, but may be represented as follows:

\[
\text{zinc oxide/silicophosphate glass + phosphoric acid} \\
\rightarrow \text{zinc aluminosilicate phosphate gel}
\]

The set cement consists of unreacted glass and zinc oxide particles bonded together by the aluminosilico-phosphate gel matrix.

Manipulation

The mixing is analogous to that for a phosphate cement; a nonabradable spatula and a cooled mixing slab should be used. The filling mix should be glossy, with puttylike consistency.

Properties
At cementing consistency, the setting time is 5 to 7 minutes; working time is about 4 minutes and may be increased by using a cold mixing slab.

These cements generally have shorter working times and a coarser grain size, leading to a higher film thickness than with zinc phosphate cements. One material is improved in these respects, and film thickness is adequate for cementation of cast gold and porcelain restorations.

The compressive strength of the set cement is in the range from 140 to 170 MPa (20,000 to 25,000 psi); the tensile strength is considerably lower at 7 MPa (1,000 psi) (see Table 9-4). The toughness and abrasion resistance are higher than those of phosphate cements.

The solubility in distilled water after 7 days is about 1% by weight. Solubility in organic acids and in the mouth is less than for phosphate cements. Fluoride is leached out and may contribute to anticariogenic action. The durability in bonding orthodontic bands to teeth is greater, and less decalcification is observed.

The glass content gives considerably greater translucency than phosphate cements, making silicophosphate cements useful for cementation of porcelain restorations.

Biologic effects

Because of the acidity of the mix and the prolonged low pH (4 to 5) after setting, pulpal protection is necessary on all vital teeth. Fluoride and other ions are leached out from the set cement by oral fluids, resulting in increased enamel fluoride and probable anticariogenic action.

Advantages and Disadvantages

Silicophosphate cements have better strength, toughness, and abrasion resistance properties than zinc phosphate cements, and show considerable fluoride release, translucency, and, under clinical conditions, lower solubility and better bonding.

Disadvantages include an initial pH and total acidity that are greater than those for zinc phosphate cements. Pulpal sensitivity may be of longer duration, and pulpal protection is essential. Manipulation is more critical than with zinc phosphate cements.

Phenolate-Based Cements

Introduction

There are three main types of cement under this classification:

1. The simple zinc oxide-eugenol combination that may contain setting accelerators
2. The reinforced zinc oxide-eugenol materials

3. The ortho-ethoxybenzoic acid (EBA) cements

Cements have also been formulated using other phenolic liquids, but these have seen little use except for those containing calcium hydroxide and a salicylate.

**Zinc oxide-eugenol cements**

**Applications**

The basic combination of zinc oxide and eugenol finds its principal applications in the provisional cementation of crowns and fixed partial dentures, in the provisional restoration of teeth, and as a cavity liner in deep cavity preparations.

**Composition and setting**

The powder is essentially pure zinc oxide (United States Pharmacopeia [USP] or equivalent, arsenic free). Commercial materials may contain small amounts of fillers, such as silica. About 1% of zinc salts, such as acetate or sulfate, may be present to accelerate the setting. The liquid is purified eugenol or, in some commercial materials, oil of cloves (85% eugenol). One percent or less of alcohol or acetic acid may be present to accelerate setting together with small amounts of water, which is essential to the setting reaction.

A chemical reaction occurs between zinc oxide and eugenol, with the formation of zinc eugenolate (eugenate):

\[
\text{Zinc oxide} + \text{eugenol} \xrightarrow{\text{water}} \text{zinc eugenolate (eugenate)}
\]

The precise mechanism is not fully understood, but the set mass contains residual zinc oxide particles bonded by a matrix of zinc eugenolate and some free eugenol. Water is essential to the reaction, which is accelerated also by zinc ions. The reaction is reversible because the zinc eugenolate is easily hydrolyzed by moisture to eugenol and zinc hydroxide. Thus, the cement disintegrates rapidly when exposed to oral conditions. The rate of reaction between the zinc oxide and the eugenol is dependent on the nature, source, reactivity, and moisture content of the zinc oxide and on the purity and moisture content of the eugenol.

**Manipulation**

The zinc oxide is slowly wetted by the eugenol; therefore, prolonged and vigorous spatulation is required, especially for a thick mix. A powder/liquid ratio of 3:1 or 4:1 must be used for maximum strength.

**Properties**

The working time is long because moisture is required for setting. Variable results are obtained with different samples of zinc oxide, depending on their mode of preparation.
and reactivity. For a given oxide, set time is controlled by moisture availability, accelerators, and the powder/liquid ratio. Mixes of cementing consistency set very slowly unless accelerators are used and/or a drop of water is added. Commercial materials set in the range of 2 to 10 minutes, resulting in adequate strengths at 10 minutes for amalgam restorations to be placed (see Table 9-4).

The particle size of the zinc oxide and the viscosity of the mix govern the film thickness. Use of a fluid mix gives values of about 40 um.

Because of the weak nature of the binding agent, the compressive strength is low, in the range of 7 to 40 MPa (1,000 to 6,000 psi). The tensile strength is very low also.

The solubility is high, about 1.5% by weight in distilled water after 24 hours. Eugenol is extracted from the set cement by the hydrolytic decomposition of the zinc eugenolate/eugenate. The cement disintegrates rapidly when exposed to oral conditions.

Biologic effect

The presence of eugenol in the set cement under clinical conditions appears to lead to an anodyne and obtundent effect on the pulp in deep cavities. When exposed directly to oral conditions, the material maintains good sealing characteristics despite a volumetric shrinkage of 0.9% and a thermal expansion of $35 \times 10^{-6}/^\circ C$. The sealing capacity and antibacterial action appear to facilitate pulpal healing; however, when in direct contact with connective tissue, the material is an irritant. Reparative dentin formation in exposed pulp is variable. Eugenol is a potential allergen.

Advantages and Disadvantages

The main advantage of these materials is their bland and obtundent effect on the pulpal tissues, together with their good sealing ability and resistance to marginal penetration.

Disadvantages include low strength and abrasion resistance, solubility and disintegration in oral fluids, and little anticariogenic action.

Reinforced zinc oxide-eugenol cements

Applications

These materials have been used as cementing agents for crowns and fixed partial dentures, cavity liners and base materials, and provisional restorative materials.

Composition and setting

The powder consists of zinc oxide with 10% to 40% finely divided natural or synthetic resins (eg, colophony [pine resin], poly[methyl methacrylate], polystyrene, or polycarbonate) together with accelerators. The liquid is eugenol, which may also contain dissolved resins as mentioned earlier and accelerators such as acetic acid, as well as antimicrobial agents such as thymol or 8-hydroxyquinoline.
The setting reaction is similar to zinc oxide-eugenol cements. Acidic resins such as colophony (abietic acid) may react with the zinc oxide, strengthening the matrix.

Manipulation

More powder is required for a cementing mix than with other cements. The proper ratio must be adhered to for adequate strength properties. Measures are provided for some commercial materials. The mixing pad or slab should be thoroughly dry. The powder is mixed into the liquid in small portions with vigorous spatulation until the correct amount has been incorporated. Adequate time should be allowed for setting without disturbance of the cement. Both powder and liquid containers should be kept closed and stored under dry conditions.

Properties

These cements may have a long working time because moisture is needed for setting. Some commercial materials contain moisture and, therefore, have working and setting times in the same range as zinc phosphate cements, that is, 7 to 9 minutes in oral conditions. Setting time is also lengthened by reducing the powder/liquid ratio.

At cementing consistency, values of film thickness from 35 to 75 um have been obtained with commercial materials (see Table 9-4). Clinical trials have shown satisfactory performance in seating castings for cements with the lowest values.

These materials have compressive strengths in the range from 35 to 55 MPa (5,000 to 8,000 psi). The tensile strength is 5 to 8 MPa (700 to 1,000 psi) (see Table 9-4). The strength is adequate as a lining material and for luting single restorations and retainers with good retention form. The modulus of elasticity is 2 to 3 GPa (300,000 to 400,000 psi). The mechanical properties of these cements are reduced by immersion in water, which results in loss of eugenol, although this appears to be slower than with simple zinc oxide-eugenol materials. This tendency seems less pronounced with the polymer-reinforced materials.

Because of the presence of the resin, the solubility of these cements appears to be somewhat lower than that of zinc oxide-eugenol materials.

Biologic effects

Polymer-reinforced zinc oxide-eugenol cements have biologic effects similar to basic materials, although there is some variation in inflammatory reaction in connective tissue with the brand of material. There may be softening and discoloration of some resin restorative materials.

Advantages and Disadvantages

The main advantages of these materials are the minimal biologic effects, good initial sealing properties, and adequate strength for final cementation of restorations.

The principal Disadvantages are the lower strength, higher solubility, and higher
disintegration compared to zinc phosphate cements; hydrolytic instability; and the softening and discoloration of some resin restorative materials.

**EBA and other chelate cements**

To further improve on the basic zinc oxide-eugenol system, many researchers have investigated mixtures of zinc and other oxides with various liquid chelating agents. The only system that has received extensive commercial exploitation for luting and lining is that containing ortho-ethoxybenzoic acid.*

*Noneugenol cements have also been developed in which fatty acids or low-odor phenolic derivatives are used to overcome the smell and taste of eugenol.

**Applications**

These materials have been used for the cementation of inlays, crowns, and fixed partial dentures, for provisional restorations, and as base or lining materials.

**Composition and setting**

In EBA materials the powder is mainly zinc oxide containing 20% to 30% aluminum oxide or other mineral fillers. Polymeric reinforcing agents, such as poly(methyl methacrylate), may also be present. The liquid consists of 50% to 66% ethoxybenzoic acid with the remainder eugenol.

The setting mechanism has not been fully elucidated. It appears to involve chelate salt formation between the EBA, eugenol, and zinc oxide. The setting is accelerated by the same factors that are operative for zinc oxide-eugenol cements.

**Manipulation**

In general, the manipulation is similar to that of reinforced zinc oxide-eugenol cements. The cement mixes readily to a very fluid consistency even at a high powder/liquid ratio. In order to obtain optimal properties, it is important to use as high a powder/liquid ratio as possible; this is about 3.5 g/mL for cementation and 5 to 6 g/mL for liners or bases. Vigorous spatulation is required for about 2 minutes to incorporate all the required powder. The correct mix flows readily under pressure because of the long working time. Adequate setting time in the mouth should be allowed. Several days may be required to reach maximum strength.

**Properties**

The working time at room temperature is long because of the dependence on moisture. The setting time ranges between 7 and 13 minutes under oral conditions (see Table 9-4).

The film thickness appears to be in the range of 40 to 70 μm for the different brands and seems adequate for permanent cementation of restorations at the lower level.

At cementing consistency, the compressive strength of these materials is in the range
of 55 to 70 MPa (8,000 to 10,000 psi); higher values, similar to those of zinc phosphate cements, can be obtained by increasing the powder/liquid ratio. The tensile strength is considerably lower, about 3 to 6 MPa (500 to 900 psi). The modulus of elasticity is about 5 GPa (700,000 psi). The EBA cements show viscoelastic properties with very low strength and large plastic deformation at slow (0.1 mm/min) rates of deformation and at oral temperature (37°C). This may explain why the retention values for crowns and orthodontic bands are considerably below those obtained using zinc phosphate cements.

The solubility is similar to that of the polymer-reinforced zinc oxide-eugenol materials in distilled water, although loss of eugenol also occurs. The resistance to solubility in organic acids appears to be greater than that of the zinc phosphate cements. When exposed to moisture, greater oral dissolution occurs than for other cements. However, a clinical survey by Silvey and Myers (1978) of the performance of an EBA-alumina cement over 3 years showed only very slightly worse results than for zinc phosphate and polycarboxylate cements. Oral breakdown may thus depend on the precise brand and manipulation.

Biologic effects

The biologic properties of these materials appear to be similar to those of zinc oxide-eugenol materials.

Advantages and Disadvantages

The principal advantages of EBA cements are their easy mixing, long working time, good flow characteristics, and low irritation to pulp. Strength and film thickness can be comparable to those of zinc phosphate cements (see Table 9-4).

The main Disadvantages are the critical proportioning, hydrolytic breakdown in oral fluids, liability to plastic deformation, and poorer retention than zinc phosphate cements. These materials seem best suited for luting restorations with good fit and retention where there is no undue stress and as cavity bases.

**Calcium hydroxide chelate cements**

The value of calcium hydroxide as a pulp-capping material that facilitates the formation of reparative dentin has long been recognized. This action appears to be largely attributable to its alkaline pH and consequent antibacterial and protein-lyzing effect. Although a number of aqueous paste materials based on calcium hydroxide are available, they are not easy to manipulate and the dried films tend to crack. In the early 1960s, phenolate-type cements based on the setting reaction between calcium hydroxide and other oxides and salicylate esters were introduced.

Applications

These materials are used as liners in deep cavity preparations.

Composition and setting
These materials are usually formulated as two pastes: One paste contains calcium hydroxide, zinc oxide, and zinc salts in ethylene toluene sulphonamide; the other contains calcium sulfate, titanium dioxide, and calcium tungstate (a radiopacifying agent) in a liquid disalicylate ester of butane-1,3-diol. The calcium hydroxide is intended to be in excess to produce an alkaline pH that will effect an antibacterial and remineralization action. There is some variation among the materials in this respect. At least one material contains fluoride.

Calcium and zinc oxide react with the salicylate ester to form a chelate similar to the zinc oxide-eugenol reaction. Likewise the reaction is greatly accelerated by moisture and accelerators.

**Manipulation**

Equal lengths of the two pastes are mixed to a uniform color.

**Properties**

Working time may be 3 to 5 minutes, depending on the availability of moisture. In the mouth, setting is rapid, about 1 or 2 minutes.

The compressive strength at 7 minutes is about 6 MPa (900 psi), and the tensile strength 1.5 MPa (200 psi); at 1 hour the corresponding values are about 10 MPa (1,500 psi) and 1.5 MPa (200 psi); and at 24 hours the values are 14 to 20 MPa (2,000 to 3,000 psi) and 1.7 to 2 MPa (250 to 300 psi). Thin films become resistant to 8 MPa (1,100 psi) penetration force in 90 seconds. At 37°C plastic flow without fracture occurs.

The solubility in 50% phosphoric acid during acid-etching procedures is significant. These cements seem to be subject to hydrolytic breakdown. When continued marginal leakage takes place, complete dissolution of the linings of these materials can occur.

**Biologic effects**

These cements appear to exert a strong antibacterial action when free calcium hydroxide is available and to assist in remineralization of carious dentin. They facilitate the formation of dentin bridges when used for pulp capping on exposures. Their effect on exposed pulp is superior to that of zinc oxide-eugenol materials. These materials can also exert a pulp protective action by neutralizing and preventing the passage of acid and by acting as a barrier to the penetration of other agents such as methyl methacrylate.

**Advantages and Disadvantages**

The advantages of these materials include their easy manipulation, rapid hardening in thin layers, good sealing characteristics, and beneficial effects on carious dentin and exposed pulp.

Their Disadvantages are that they show low strength even when fully set, exhibit plastic deformation, are weakened by exposure to moisture, and will dissolve under
acidic conditions and if marginal leakage occurs. The data on physical properties and clinical experience suggest that further improvements in these materials are required before they can be utilized as the sole liner in deep cavity preparations.

More recently polymerizable resin compositions containing calcium hydroxide have been introduced as alternatives to these materials.

**Polycarboxylate (Carboxylate)- Based Cements**

**Zinc polycarboxylate cements**

The polycarboxylate cements were developed in the late 1960s as adhesive dental cements that would combine the strength properties of the phosphate system with the biologic acceptability of the zinc oxide- eugenol materials.

**Applications**

Zinc polycarboxylates are used for the cementation of cast alloy and porcelain restorations and orthodontic bands, as cavity liners or base materials, and as provisional restorative materials.

**Composition and setting**

The powder in these cements is zinc oxide with, in some cases, 1% to 5% tin or magnesium oxide; 10% to 40% aluminum oxide or other reinforcing filler may be present in some brands. A small percentage of stannous or other fluoride may also be included to improve mechanical properties and provide leachable fluoride. The liquid is approximately a 40% aqueous solution of polyacrylic acid or an acrylic acid copolymer with other organic acids, such as itaconic acid. The molecular weight of the polymer is generally in the range of 30,000 to 50,000, which accounts for the viscous nature of the solution. In some brands of the material the polyacrylic acid component is dried and added to the powder. In a brand that is encapsulated the liquid is a weak solution of $\text{NaH}_2\text{PO}_4$, which both reduces the viscosity of the polyacrylic acid and retards the setting of the cement. In other brands water is simply added to the powdered ingredients.

The zinc oxide reacts with the polyacrylic acid, forming a cross-linked structure of zinc polyacrylate. The set cement consists of the residual zinc oxide particles bonded together by this amorphous gel-like matrix:

\[
\text{zinc oxide} + \text{polyacrylic acid} \rightarrow \text{Zinc polyacrylate}
\]

**Manipulation**

The material should be carefully proportioned and the freshly dispensed components mixed rapidly in 30 to 40 seconds. The mix should be used while it is still glossy, before the onset of cobwebbing. The correct cementing mix is more viscous than a zinc phosphate mix, but because of its different rheology it flows adequately under pressure. The water mix materials are more fluid initially (Fig 9-2). The interior of restorations and tooth surfaces should be clean and free of saliva. The powder and
liquid should be stored under cool conditions and kept sealed with a stopper. Prolonged or cold storage may cause the liquid to gel; to reverse this, it must be warmed to 50°C. Loss of moisture from the liquid will lead to thickening.

**Fig 9-2** Typical consistency for water mix polycarboxylate and glass-ionomer cements. The mix is comparable to zinc phosphate cements.

**Properties**

The rate of setting is affected by the powder/liquid ratio, the reactivity of the zinc oxide, the particle size, the presence of additives, and the molecular weight and concentration of the polyacrylic acid. At luting consistency the recommended powder/liquid ratio for most materials is about 1.5:1 by weight. The working time is 2.5 to 3.5 minutes at room temperature, and the setting time is 6 to 9 minutes at 37°C; the water mix materials tend to give slightly longer setting times. As with other cements, working time can be substantially increased by mixing the material on a cold slab and by refrigerating the powder. The liquid should not be chilled, as this encourages gelation due to hydrogen bonding.

The freshly mixed cement shows shear thinning. Contrary to the subjective impression that the correct mix for a zinc polycarboxylate cement is much thicker than that of a luting zinc phosphate mix, under pressure they flow out to the same degree to a film thickness of 25 to 35 um. In fact, the zinc phosphate mix tends to thicken more quickly than the zinc polycarboxylate mix. One of the most common errors made with the polycarboxylate cements is to make a mix that appears to be as fluid as a zinc phosphate mix; this will result in a low powder/liquid ratio with consequent poor properties in the cement. Measuring devices for these materials will ensure correct proportions.

At cementing consistency, the compressive strength of these materials is in the range
of 55 to 85 MPa (8,000 to 12,000 psi), and the tensile strength is 8 to 12 MPa (1,100 to 1,700 psi) (see Table 9-4). Strength increases with the powder/liquid ratio, reaching a maximum at about 2:1 by weight, and it is increased also by additives such as alumina and stannous fluoride. In general these cements have somewhat lower compressive strengths than zinc phosphate cements but are significantly stronger in tension. The cement gains strength rapidly after the initial setting period; the strength at 1 hour is about 80% of the 24-hour value. The modulus of elasticity is about 6 GPa (850,000 psi).

In distilled water, the solubility ranges from less than 0.1% to 0.6%. The latter high value relates particularly to cements that contain stannous fluoride. However, as in the zinc phosphate system, the solubility is appreciably higher in acids such as lactic and citric acid. In vivo solubility is similar to or less than that for zinc phosphate cements.

Bonding to clean enamel and dentin surfaces can occur through calcium complexation. In practice, adhesion to dentin may be limited because of debris and contamination. The material also sticks to clean stainless steel, amalgam, chromium-cobalt, and other alloys. Bond strength is related to the strength of the cement.

Biologic effects

The effect of zinc polycarboxylate cements on pulp is comparable to or less than that of zinc oxide-eugenol. The formation of reparative dentin in exposed pulp is variable. The generally good biocompatibility appears to be primarily due to the low intrinsic toxicity and also to (1) the rapid rise of the cement pH toward neutrality; (2) localization of the polyacrylic acid and limitation of diffusion by its molecular size and ion binding to dentinal fluid and proteins; and (3) the minimal movement of fluid in the dentinal tubules in response to the cement. The presence of stannous fluoride does not appear to affect the mild response. The fluoride-containing cements release fluoride, which is taken up by neighboring enamel and which presumably will exert anticariogenic effects.

Advantages and Disadvantages

The main advantages of these materials are the low irritation, adhesion to tooth substance and alloys, easy manipulation, strength, solubility, and film thickness properties comparable to those of zinc phosphate cements.

The Disadvantages are the need for accurate proportioning for optimal properties and thus more critical manipulation, the lower compressive strength and greater viscoelasticity than zinc phosphate cements, the short working time of some materials, and the need for clean surfaces to utilize the adhesion potential.

**Polymer-Based Cements**

**Introduction**

The majority of the materials in this group are polymethacrylates of two types: (1) materials based on methyl methacrylate and (2) materials based on aromatic
dimethacrylates of the bis-GMA type. The closely related cyanoacrylate monomers, notably ethyl and isobutyl, have found some limited use for the attachment of facings and for pin cementation. However, the hydrolytic stability and biologic effects in this situation are suspect and little use is made of them.

_Acrylic resin cements_

Applications

Acrylic resin cements are used for the cementation of restorations, facings, and provisional crowns.

Composition and setting

The powder in these materials is a finely divided methyl methacrylate polymer or copolymer containing benzoyl peroxide as the **initiator**. Mineral filler and pigments may also be present. The liquid is a methyl methacrylate monomer containing an amine accelerator.

The monomer dissolves and softens the polymer particles and concurrently polymerizes through the action of free radicals from the peroxide-amine interaction. The set mass consists of the new polymer matrix uniting the undissolved but swollen original polymer granules.

Manipulation

The liquid is added to the powder with minimal spatulation to avoid an incorporation of air. The mix must be used immediately because working time is short. Excess material must be removed at the final set, hard stage and not when the material is rubbery, otherwise marginal deficiencies will be created.

Properties

The properties of these materials are comparable to those of the cold-curing acrylic resin filling materials. They are stronger and less soluble than other types of cement but display low rigidity and viscoelastic properties. They have no effective bond to tooth structure in the presence of moisture; thus they permit marginal leakage, although they may show better bonding than other cements to resin facings and polycarbonate crowns.

Biologic effects

As with acrylic resin filling materials, marked pulpal reaction may occur and pulpal protection is necessary.

Advantages and Disadvantages

The advantages of these materials include relatively high strength and toughness and low solubility.
Disadvantages include a short working time, deleterious effects on pulp, and difficulty in removal of excess cement from margins.

**Adhesive resin cements**

Adhesive acrylic materials have been formulated by adding an adhesion promoter, 4-methyloxy ethyl trimelletic anhydride (4-META), to the methyl methacrylate monomer as well as an additional polymerization initiator, tributyl boron, which is also believed to aid chemical bonding to dentin. Such materials have been developed as cements for metal fixed partial dentures especially of base metal (Superbond, Parkell) and for bonding amalgam to dentin and composites (Amalgambond, Parkell). In vitro tests have shown high bond strengths for the luting cement to oxidized, etched, or silica-coated casting alloy surfaces. Shear bond strength to amalgam is significantly less than the bond strength to dentin, which is comparable to other dentin bonding systems in the region of 20 MPa. Since these materials have only low (<10%) filler content, the physical properties are typical of acrylic resins, that is, moderate strength with high deformation under load. Although the materials have been widely used for cementation of fixed partial dentures, there is little clinical data on longevity, and the cements are said to be technique sensitive.

**Dimethacrylate cements**

Dimethacrylate cements are usually based on the bis-GMA system: They are combinations of an aromatic dimethacrylate with other monomers containing various amounts of ceramic filler. They are basically similar to composite restorative materials.

**Applications**

Dimethacrylate cements are used for bonding crowns (usually porcelain), fixed partial dentures, inlays, veneers, and indirect resin restorations.

**Composition and setting**

These cements are classified according to the following methods of curing:

1. Chemically (or auto-) cured: These are usually paste-paste systems and are used to cement metal and opaque ceramic core (eg, Procera, In-Ceram) restorations (Fig 9-3).
2. Dual cured: These cements start curing with light and continue with chemical curing. The chemical cure will polymerize more thoroughly than light curing alone (Fig 9-4). These are used to cement translucent restorations (eg, porcelain, indirect resin restorations).

![Fig 9-3](image)

**Fig 9-3** Chemically cured resin cement for use under opaque restorations (Panavia 21, Kurarary/J. Morita).

![Fig 9-4](image)

**Fig 9-4** Dual-cured resin cement in syringe form for use when light may not penetrate enough for complete curing (Dual Cement, Vivadent).
3. Light cured/dual cured: These can be used for light curing only or can be dual cured when dual-cure catalysts are added to the light-cure base. These products are used for both light-cure applications (eg, thin porcelain veneers) and dual-cure applications (Fig 9-5).

Fig 9-5 Light-cured/dual-cured cements may be cured with light or with addition of dual-cure catalyst, which will allow setting to continue after light is turned off (Variolink II, Vivadent).

In the powder-liquid materials, the powder is generally a finely divided borosilicate or silica glass together with fine polymer powder and an organic peroxide initiator. The liquid is a mixture of bis-GMA and/or other dimethacrylate monomers containing an amine promoter for polymerization. Some materials contain monomers with potentially adhesive groups, such as phosphate or carboxyl, similar to dentin bonding materials. The two-paste materials are of similar overall composition but with the monomers and fillers combined into two pastes. In light-cured and dual-cured materials, light-sensitive polymerization systems such as diketones (eg, camphorquinone) and amine promoters are present, respectively, in the two cement components in addition to the chemical-initiator systems.

On mixing the components, polymerization of the monomers occurs, leading to a highly cross-linked resin composite structure.

Manipulation

Correct proportioning of powder and liquid components using measures is important. Paste materials are usually proportioned 1:1 (equal lengths). Rapid, thorough mixing, minimizing air inclusion, until uniform is critical.

Properties
As with resin composite restorative materials, monomer conversion is incomplete, even under optimum cure conditions, and thus manipulation is critical to optimum physical properties. For light- and dual-cured materials, the maximum light exposure is desirable. Maximum properties are generally reached about 10 minutes after polymerization; only small changes occur over the ensuing 24 hours.

Since polymerization systems vary and filler contents range between 20% and 80% for the various products, physical properties vary widely (see Table 9-4) and the solubility of a specific material for a particular clinical application should be checked individually. Compressive strengths have been reported to range between 100 and 200 MPa (14,000 and 28,000 psi), and diametral tensile strengths from 20 to 50 MPa (3,000 to 7,000 psi) with corresponding differences in microhardness. These values are considerably higher than traditional cements, and therefore high values can be obtained for retention of well-fitting crowns. However, optimum luting performance is very dependent on fluidity, seating capability, and film thickness. Many resin cements tend to show unacceptably high values for film thickness. Recently, to improve wetting of the tooth, preparation, seating, and bond strength, some resin cements have been used with dentin bonding primers, thus increasing the clinical complexity of the system. Although these materials have been used widely in adhesive techniques, especially for ceramic restorations, there are comparatively few clinical reports of their longevity. Aside from failures induced by material and technique shortcomings during the critical clinical manipulation, studies indicate that resin cement bonds will most likely fail through cyclic fatigue stresses. Some studies on etched metal restorations cemented with chemically cured cements have indicated a median survival time of about 8 years.

Biologic effects

The materials themselves appear to pose few problems, although some patients experience objectionable odors. Cases of allergy among dental personnel have occurred, especially where reactive dentin bonding systems have been used. Skin contact should be avoided.

Pulpal pathology may be due to poor seating, polymerization contraction, and consequent microleakage. All systems show some microleakage that may contribute to tooth sensitivity and clinical failure. Microleakage appears to occur least with systems employing dentin bonding agents, but there are no long-term studies on this aspect.

Advantages and Disadvantages

The advantages of these cements include high strength; low oral solubility; and high micromechanical (and possible chemical) bonding to prepared enamel, dentin, alloys, and ceramic surfaces.

Disadvantages include the need for a meticulous and critical technique, more difficult sealing and higher film thickness than traditional cements, possible leakage and pulpal sensitivity, and difficulty in removal of excess cement.
Glass-Ionomer Cements

Introduction

These materials were formulated in the 1970s by bringing together the silicate and polyacrylate systems. The use of an acid-reactive glass powder together with polyacrylic acid solution leads to a translucent, stronger cement that can be used for luting and restorative materials.

Applications

Glass-ionomer cements are used for the cementation of cast-alloy and porcelain restorations and orthodontic bands, as cavity liners or base materials, and as restorative materials, especially for erosion lesions. They are being replaced by hybrid ionomer cements, which allow better handling.

Composition and setting

The powder in these materials is finely ground calcium aluminum fluorosilicate glass with a particle size around 40 um for the filling materials and less than 25 um for the luting materials. One brand (Zionomer Liner, Den-Mat) also contains zinc oxide. Silver powder is fused into the glass in Ketac-Silver (3M/ESPE) for improved physical properties. The liquid is a 50% aqueous solution of a polyacrylic-itaconic acid or other polycarboxylic acid copolymer that contains about 5% tartaric acid. Some other materials contain 10% to 20% added silver, silver alloy, or stainless steel. In some materials the solid copolymer is added to the powder, and the solution contains tartaric acid; in others, all the ingredients are in the powder, and the liquid is water.

On mixing, the polyacrylic and tartaric acids react with the glass, leaching calcium and aluminum ions from the surface, which cross-link the polyacid molecules into a gel. The tartaric acid serves to increase working time and gives a sharp setting by forming metal ion complexes. Differences in composition between brands affect the hardening rate and properties. Some recent evidence suggests that a polysilicate matrix may also form within the polygel over time.

Manipulation

The material should be carefully proportioned and the freshly dispensed components mixed rapidly in 30 to 40 seconds. Some brands are encapsulated, mechanically mixed, and injected. The powder/liquid ratio for luting is about 1.3:1 for the conventional types of glass-ionomer cement. This ratio appears to be critical with these cements to obtain optimal cementation properties. Best results are obtained by mixing the chilled powder with the liquid on a chilled slab. The correct cementing mix is fluid, similar to zinc phosphate. The lining mix is somewhat more viscous, depending on the brand. The restorative mix should have a puttylike consistency and a glossy surface. Tooth surfaces should be clean and free from saliva but not dehydrated. Restoration surfaces should be free from debris and contamination. The cement hardens slowly and should be protected from loss or gain of moisture when set
clinically. Restoration margins or filling surfaces should be protected with a varnish or a light-curing sealant. This is less important with light-cured materials.

Properties

For the luting materials, the setting time is in the range of 6 to 9 minutes. The lining materials set in 4 to 5 minutes, and the restorative materials set in 3 to 4 minutes.

Materials that are light-cured set in approximately 30 seconds when exposed to a visible light source. The acid-base reaction continues slowly and properties improve over time.

Film thickness is in the range of 25 to 35 um, which is adequate to seat castings satisfactorily, although the flow properties are quite dependent on the powder/liquid ratio.

For the luting cements, the compressive strength increases over 24 hours to 90 to 140 MPa (13,000 to 20,000 psi) depending on the brand. The tensile strength increases similarly to 6 to 8 MPa (900 to 1,100 psi). The compressive modulus of elasticity is about 7 GPa (900,000 psi). The lining materials have compressive and tensile strengths in the same range with some light-cured materials at the higher end of the range reaching 150 to 160 MPa (21,000 to 23,000 psi) in compression and 10 to 12 MPa (1,400 to 1,700 psi) in tension. The light-cured materials are significantly tougher in some brands, with a lower modulus. The restorative materials range from 140 to 180 MPa (20,000 to 26,000 psi) in compression and 12 to 15 MPa (1,700 to 2,100 psi) in tension. The light-cured restorative materials may have strengths as high as 200 MPa in compression and 20 MPa in tension. Some silver-containing materials are in this range, and even higher strengths have been achieved in recent materials.

In general, with light-cured materials, properties are dependent on the depth of cure.

The solubility of the cements in water is about 1% for a luting material, and this is higher in lactic acid. Good resistance to dissolution is observed under oral conditions. Resistance to dissolution and disintegration is improved by varnish protection for conventional cements.

Erosion of clinical restorations of conventional cements by acid phosphate fluoride preventive-treatment solutions has been observed, making these solutions contraindicated.

Some studies show that light-cured glass-ionomer materials continue to absorb water over several months, with swelling and reductions in strength and stiffness. The clinical significance of this behavior is not yet clear.

Glass-ionomer cements exhibit bonding to enamel, dentin, and alloys in a manner similar to zinc polycarboxylates. In vitro and in vivo the adhesion is variable and is affected by surface conditions. Slight and variable marginal leakage has been observed. Bonding to dentin for conventional materials is not improved by pretreatment with polyacrylic acid solutions, whereas with light-cured materials it is dependent on the use of dentin primers.
Biologic effects

Pulpal response to the lining and restorative materials appears generally favorable. Variable behavior has been reported for the various luting materials with instances of postoperative sensitivity. This has been attributed to a prolonged initially low pH coupled with the effects of the toxic ions. This may be accentuated by dehydration of dentin and marginal leakage of bacteria. Leaching of fluoride and uptake by adjacent enamel occurs with these cements, and this continues for at least a year with potentially cariostatic effects. Antibacterial action has been attributed to low initial pH, leaching, release of silver and other ions, or a combination of these. Light-cured materials have been observed to show greater cytotoxicity.

Advantages and Disadvantages

The advantages of glass-ionomer cement materials include easy mixing, high strength and stiffness, leachable fluoride, good resistance to acid dissolution, potentially adhesive characteristics, and translucency.

The Disadvantages include initial slow setting and moisture sensitivity, variable adhesive characteristics, radiolucency, and possible pulpal sensitivity.

*Resin-modified glass-ionomer cements*

Applications

A recent addition to the spectrum of materials, these versatile cements, sometimes also called *hybrid ionomers* (see Fig 8-11), have many uses: cavity liners, bases, core buildups, and luting cements. One hybrid ionomer (**Fig 9-6**) is used for permanent cementation of crowns, orthodontic appliances, and core buildups.

**Fig 8-11** Continuum of glass-ionomer and composite restorative materials. (After Albers, 1996).
Composition and setting

In hybrid ionomers, the acid-base setting reaction in glass-ionomer cements has been modified by the introduction of water-soluble polymers and polymerizable monomers into the composition. The use of copolymers of acrylic acid and methacrylate monomers in the liquid leads to materials that undergo the customary acid-base reaction on setting and can also be light cured via the methacrylate groups. This gives improved lining and restorative materials with an immediate command set and thus higher early strength and water resistance. Some commercial materials contain a preponderance of polymeric components with minimal acid-base reaction.

The classification of these materials as glass-ionomer cements is controversial. Some light-cured restorative glass-ionomer cements are used with a dentin primer similar to dentin bonding resin composite systems and thus depend on surface infiltration for bonding in addition to chemical interaction.

Resin-modified glass ionomers are available in hand-mixed and predosed capsules (Fig 9-7). The resin monomers in the liquid depend on the product and include bis-GMA, hydroxyethylmethacrylate, and methacrylate-modified polyacrylates along with photoinitiators.
Manipulation

For hand-mixed compositions, the powder should be fluffed before dispensing. The powder and liquid should be mixed quickly, within 30 seconds, on the pad. These cements have a working time of about 2.5 minutes. For luting, the cement is applied to the undesiccated tooth to avoid possible postoperative sensitivity.

Properties

*Table 9-5* gives a comparison of properties of resin-modified glass ionomers versus glass-ionomer cements. The strength properties of the two types of cements are similar, with considerable variations among brands. There is a major difference in flexibility, with the hybrid ionomers being twice as flexible, as indicated by lower modulus of elasticity values. Also, many of the hybrid ionomers have been found to expand on setting, possibly due to the absorption of water, which is more than for resin cements. Therefore, hybrid ionomers are not recommended for luting all-ceramic crowns, to avoid possible expansion stresses and crown fracture.

*Table 9-5* Mechanical properties of glass-ionomer and hybrid ionomer cements*

<table>
<thead>
<tr>
<th></th>
<th>Glass ionomers</th>
<th>Resin-modified glass ionomers</th>
</tr>
</thead>
<tbody>
<tr>
<td>Flexural strength (MPa)</td>
<td>25</td>
<td>35-70</td>
</tr>
<tr>
<td>Flexural modulus (GPa)</td>
<td>8</td>
<td>4</td>
</tr>
<tr>
<td>Compressive strength (MPa)</td>
<td>180-200</td>
<td>170-200</td>
</tr>
<tr>
<td>Diametral tensile strength (MPa)</td>
<td>22-25</td>
<td>35-40</td>
</tr>
</tbody>
</table>
Biologic effects

Hybrid ionomers release fluoride from the glass component, which is favorable for caries prevention. Long-term, large-scale clinical data are not available for an overall assessment of their biologic effects.

Advantages and Disadvantages

Their advantages include dual cure, fluoride release, higher flexural strength than glass-ionomer cements, and ease of handling. Also, they are capable of bonding to composite materials.

One problem is a possible setting expansion that may lead to cracking of all-ceramic crowns. Therefore, resin cements, zinc phosphate, and glass-ionomer cements are still recommended for metal-free crowns.

Selection and Use of Cements

None of the cements available is free from deficiencies in the required clinical characteristics, such as biocompatibility, ease of manipulation, satisfactory sealing, retentive properties, and long-term stability. A proportion of clinical failures is inevitable, but this can be minimized by proper selection and manipulation of the cement. The following factors should be kept under review:

1. Rapid uniform and reproducible dispensing of the components
2. Rapid, thorough mixing
3. Moisture isolation where practical
4. An undisturbed setting
5. Careful removal of excess
6. Avoidance of excessive drying of dentin

Factors within the clinician's control, such as the design and execution of the preparation, adequate isolation, proper seating of the restoration, and finishing of the margins, are also important determinants of success as is the manipulation of the cement.

These considerations may influence cement selection and use, but a governing factor is the biological state of the tooth tissue. Thus, as indicated in Table 9-2, the selection

| Shrinkage (% vol) | 3 | 3.5-expansion |

*Data from Burgess et al (1996). Used with permission from *The Compendium of Continuing Education in Dentistry.*
of particular cements for specific clinical situations is limited both by the preparation and the properties of the cement.

**Clinical Decision Scenarios for Dental Cements**

This section presents one approach for choosing dental cements for specific situations. Each of the following scenarios uses the same format as that presented in chapter 7. Advantages and Disadvantages of each material are prioritized using the following codes: * = of minor importance, ** = important, and *** = very important.

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**Materials**  Zinc phosphate vs resin cement for crown cementation

**Description**  A dentist has finished preparation of a full gold crown for a maxillary left permanent second molar and is ready to cement the crown. This is a routine crown cementation. The preparation is normal, the tooth is vital, and there is no known pulpal pathology. The dentist may choose between a resin cement and zinc phosphate cement.

**Critical Factors**  Ease of use, cost, effectiveness

**Zinc phosphate**

*Advantages*

*** Low cost
   
   Long clinical history
   
   High rigidity
   
   Long working time

*** Easy to use

*Disadvantages*

*** No bond to tooth

   Slow setting time
   
   Moisture sensitivity during mixing

**Resin**

*Advantages*

*** Bonds to tooth
Fast setting time

Higher strength

*** Easy to use

Disadvantages

*** High cost

Short working time

Film thickness varies widely among brands

Difficult to remove excess

**Analysis/Decision** Comparing the properties of the two cements reveals they are balanced in advantages and Disadvantages for this situation. The choice finally rests in the personal preference of the dentist. Since this dentist had more experience with zinc phosphate, she chose to use that material.

**Materials** Glass ionomers vs zinc phosphate cement for crown cementation

**Description** An older gentlemen with a history of periodontal surgery to correct bony defects has broken the cusps off a previously restored mandibular first molar. The patient's plaque control is only fair, and he is taking medications that could result in some degree of "dry mouth." He has had a slightly increased level of caries activity since the surgery and since beginning the medication. The dentist has elected to restore the tooth with a cast crown, which has now been returned from the laboratory and is ready for cementation. The dentist has both zinc phosphate and glass-ionomer luting cements available and must decide which to use.

**Critical Factors** Caries resistance, good seal

**Zinc phosphate**

**Advantages**

*** Good seal

** Reasonable cost

** Adequate strength

** Little sensitivity

**Disadvantages**
*** No fluoride release

**Glass ionomer**

**Advantages**

*** Good seal

*** Fluoride release

** Adequate strength

** Reasonable cost

**Disadvantages**

** Occasional sensitivity

**Analysis/Decision** Both cements have a good seal, but only the glass ionomer releases fluoride after cementation. Fluoride is generally accepted to reduce caries and seems to do so in individuals such as this patient. For this reason, the glass ionomer was selected. Careful attention to instructions allowed cementation without the sensitivity sometimes experienced with glass-ionomer cements. The dentist was satisfied with this treatment choice.

**Materials** Glass ionomers vs zinc phosphate for bases under amalgam restorations

**Description** A patient has a mandibular molar that exhibits extensive carious destruction. Due to the patient's finances, a large, pin-retained amalgam restoration is selected. Following caries removal and preparation, it is determined that the cavity needs a base prior to placement of the amalgam restoration. The dentist has zinc phosphate and glass-ionomer cements available and must choose between them for a base material.

**Critical Factors** Strength, modulus of elasticity

**Zinc phosphate**

**Advantages**

*** Adequate modulus of elasticity

*** Adequate strength

** Ease of use

** No sensitivity
Disadvantages

** No fluoride release

Glass ionomer

Advantages

*** Adequate modulus of elasticity

*** Adequate strength

** Fluoride release

** Ease of use

Disadvantages

** Occasional sensitivity

Analysis/Decision  Since the advantages and Disadvantages were essentially balanced for this situation, neither cement had a clear advantage. The highly significant modulus of elasticity, essential for a good base, was roughly equivalent. In this case, the dentist chose glass ionomer because of the added advantage of fluoride release, which is thought to reduce recurrent decay. However, because that was not a problem with this patient, either cement would have been an acceptable choice.

Materials  Zinc oxide-eugenol vs calcium hydroxide as liners under amalgam restorations

Description  A patient in need of a slightly deep mesio-occlusodistal amalgam on a mandibular left first molar was seen by the dentist. The patient reported that the last two amalgams the dentist placed had been quite sensitive for several weeks following placement. They had finally lost their sensitivity, but the patient felt they had hurt unusually long for new restorations. Upon preparation, the dentist decided that a base was indicated. Both calcium hydroxide and zinc oxide-eugenol base materials were available.

Critical Factors  Strength, modulus of elasticity, reduction of sensitivity

Zinc oxide-eugenol

Advantages

** Ease of use

** Decreased sensitivity
** Low cost

Disadvantages

*** Low strength

*** Low modulus of elasticity

Calcium hydroxide

Advantages

** Ease of use

** Stimulates secondary dentin

** Low cost

Disadvantages

*** Low strength

*** Low modulus of elasticity

Analysis/Decision  In this situation, again, there is no clear advantage for either base. Both have low strength and low modulus of elasticity, necessitating that they be applied as very thin layers. The zinc oxide-eugenol has the advantage of the anodyne eugenol, which is known to reduce tooth sensitivity. The calcium hydroxide, on the other hand, will stimulate the formation of secondary dentin and thus is good if the restoration is near the pulp. This was true in the situation described, but the patient also had a history of sensitive teeth following placement of amalgam restorations. Therefore, the dentist selected the zinc oxide-eugenol as the liner of choice. Had the cavity preparation been deeper and the pulp visible through the dentin, calcium hydroxide would have been selected due to its ability to stimulate formation of secondary dentin.

Materials  Flowable resin composite vs hybrid ionomer for Class 5 restorations in high-caries-risk patient

Description  An older patient has recurrent caries around the margins of a few old Class 5 amalgam restorations, probably due to poor hygiene and a mild xerostomia resulting from a cholesterol-lowering drug. The patient requested that the replacement restorations be more esthetic. The dentist considers flowable composites and hybrid ionomers.

Critical Factors  Ease of placement, prevention of recurrent decay, reasonable esthetics
Flowable composite

Advantages
*** Very simple placement
*** Excellent esthetics
* Less soluble

Disadvantages
*** Higher incidence of recurrent decay
** Higher sensitivity to moisture contamination during placement

Hybrid ionomer

Advantages
*** Ease of placement
*** Less recurrent decay
*** Reasonable esthetics
** Less sensitivity to moisture during placement

Disadvantages
* More soluble than flowables

Analysis/Decision  Due to the patient's high risk of recurrent decay, the dentist decided to use the hybrid ionomer restorative material. She chose a dual-cure hybrid ionomer for its ease of use and potential to be recharged with fluoride at future visits.

Materials  All-purpose adhesives vs cavity varnish under amalgam restorations

Description  A patient is seen and diagnosed as needing numerous amalgam restorations, some large. Upon preparation, the cavities turn out to be normal in size and depth. There is little remarkable about the patient or the restorations. The dentist has both cavity varnish and new all-purpose adhesives to use under the amalgam restorations. The all-purpose adhesives are reported by the manufacturer to bond the amalgam to the teeth, but the dentist knows that such claims are as yet probably exaggerated.

Critical Factors  Marginal leakage, bond to tooth, cost, time required for application
Varnish

Advantages

*** Low cost

*** Easy placement

Disadvantages

*** No bond to tooth

*** Mediocre seal

Adhesives

Advantages

*** Superior marginal seal

*** Bond to tooth

Disadvantages

** High cost

*** Complex placement

Analysis/Decision  Since there is no evidence of excessively weakened cusps, the possible bonding of the new adhesives is not significant in this case. The superior seal makes them desirable, but their high cost and complex placement process complicates the choice. The dentist had been to a recent lecture on dental materials in which the speaker claimed the new materials should be a standard of practice, so he chose to place the adhesives. The patient reported a complete lack of postoperative sensitivity in the new restorations.

Materials  Dentin adhesives vs glass-ionomer liners under composite restorations

Description  A patient has several Class 5 toothbrush abrasion lesions on his maxillary and mandibular premolars. At first, they were no problem, but over the years they have become sensitive, and the patient wants the discomfort relieved. The lesions are not carious but are somewhat deep. The occlusal, mesial, and distal margins are on enamel, but the cervical margin is on cementum. The dentist decides against glass-ionomer restorations because she feels a better esthetic result can be obtained with a composite material. In addition, she feels the composite will better resist further toothbrush abrasion. She is uncertain whether to use a dentin bonding agent alone or a glass-ionomer lining cement under the cervical margin to prevent leakage. She has heard that composites often leak on margins that are on cementum.
Critical Factors  Marginal seal, bond strength

Dentin adhesives

Advantages

** Good seal
*** Adequate bond strength
** Lower cost

Disadvantages

*** Poorer marginal seal

Glass ionomer

Advantages

*** Better marginal seal
*** Adequate bond strength

Disadvantages

** Additional cost
** Additional procedure

Analysis/Decision  Although there are more advantages for the bonding agent, the most significant advantage—better marginal seal—favors the glass-ionomer liner. It adds to the cost and the time to do the procedure, but it provides a significantly better seal. The better seal results in a superior restoration that justifies the extra cost and time. The glass-ionomer liner was selected.

Materials  Hybrid ionomer vs glass ionomer for treatment of recurrent caries at gold crown margins

Description  During a dental hygiene appointment, patient examination revealed recurrent caries around the margins of a few gold crowns. Replacement of the crowns is not feasible due to the patient's lack of funds. Normally, the dentist would use a glass-ionomer material to restore the marginal areas after the decay has been removed. However, he is considering the use of the newer hybrid ionomers due to their dual-curing ability.

Critical Factors  Prevention of recurrent caries, ease of placement, bond to tooth
structure

**Hybrid ionomer**

*Advantages*

*** Easy to place
*** Tooth colored

* Low cost

*** Fluoride release

** Bonds to tooth

* Smoother finish

** Light/dual cure

*Advantages*

*** Resonably easy to place
*** Tooth colored

* Low cost

*** Fluoride release

*** Bonds to tooth

*Advantages*

** Bonds to tooth

* Isolation is critical

**Disadvantages**

* Isolation is critical

**Disadvantages**

* Isolation is critical

**Chemical cure only

**Disadvantages**

** More plaque retentive

* Isolation is critical

** Chemical cure only
Analysis/Decision  Though the two materials have comparable fluoride release, the dentist chose a resin-modified glass ionomer mainly for its greater ease of use. The hybrid ionomer can be placed and cured in less than 1 minute due to the dual-cure feature, while the traditional chemically cured glass ionomers require protection from oral fluids for 4 to 5 minutes during setting. The dual-curing hybrid ionomer may also release more fluoride because it contains more fluoride glass, which is the source of fluoride ions and aids in the chemical setting reaction.

Glossary

**accelerator (promoter)** Substance that facilitates decomposition of an initiator.

**anodyne** Relieves pain.

**chelating** Ring structure reaction with metal ions.

**cored structure** A material consisting of at least two phases, for example, as residual particles of a component embedded in a matrix of reaction product.

**initiator** Substance capable of decomposing into free radicals that initiate polymerization.

**lute** A cementlike material that also fills and seals gaps.

**obtundent** A material that reduces irritation or has a soothing effect on tissue.

**rheology** Science of the deformation and flow of matter.

**setting time** Time from the beginning of mixing of the cement to the development of a hard and rigid (usually brittle) state in the mouth.

**working time** Time available, measured from the beginning of mixing at room temperature, for clinical manipulation of a cement before viscosity becomes too great for seating of the restoration.

**Discussion Questions**

1. Why is fluoride release so important in a cement?

2. What is the source of fluoride in a glass-ionomer cement?

3. What is the difference in function between a cement liner and a cement base?

4. Why are polymer cements recommended for use with CAD/CAM inlays with poor marginal fits?

**Study Questions**
1. What is the minimum compressive strength required of a dental cement for adequate retention of restorations?

2. What is the structure of set zinc phosphate cement?

3. How does the solubility of phosphate cements in citric or lactic acid compare with their solubility in water?

4. What are adhesive resin cements?

5. What are resin-modified glass ionomers?

6. What agents accelerate the setting of zinc oxide-eugenol cements?

7. Why do zinc oxide-eugenol cements have a high solubility?

8. What materials can be added to zinc oxide-eugenol cements to improve their strength?

9. What effects do zinc oxide-eugenol cements have on resin restorative materials?

10. How does the composition of EBA cements differ from that of zinc oxide-eugenol cements?

11. What factors affect the setting reaction of polycarboxylate cements?

12. Give possible reasons for the minimal effect of polycarboxylate cements on pulp.

13. What are important considerations in manipulating polycarboxylate cements?

14. What are the advantages of light curing glass-ionomer cements?

15. What are the major advantages of glass-ionomer cements?

16. Define the two types of polymer-based cement.

17. What are the principal Disadvantages of polymer-based cement?

18. It has been reported that hybrid ionomers expand on setting, which could cause cracking of all-ceramic crowns. This was discovered years after the cements were marketed. Using the hierarchy of evidence, which type of study would have warned dentists about this problem?

19. Although short-term pulpal studies on glass-ionomer cements gave a satisfactory overall biocompatibility, there were occasional signs of cytotoxicity with certain products. After many years of clinical use, it has been reported that, if the dentin has been dehydrated by excessive drying, there is a definite chance of postoperative sensitivity when these cements are used to cement crowns. Speculate on why this
A serious clinical problem was not clearly identified in the histologies of treated tooth pulps that were used in the biocompatibility tests.

**Recommended Reading**


Chapter 10. Abrasion, Polishing, and Bleaching

Introduction

Surface finish and appearance are important aspects of restorative dentistry. The surfaces of normal, healthy-looking teeth have a high gloss and an unstained white/ivory color. Abrasion is a process of surface roughening that can either wear away tooth structure destructively or be used to correct tooth shape or attain an optimal smooth finish on the surface of a restoration. Polishing and bleaching are two different approaches to whitening teeth by attacking surface stains. Polishing is an abrasive process of smoothing a rough surface and removing stains with very fine particles. Bleaching lightens or eliminates surface stains chemically by application of strong peroxide solutions. This chapter deals with the basic processes of these techniques.

Abrasion

Introduction

Abrasion is the process of wear on the surface of one material by another material by scratching, gouging, chiseling, tumbling, or other mechanical means. The material that causes the wear is called an abrasive; the material being abraded is called the substrate.

Grinding is the gross reduction of the surface of a substrate by the process of abrasion; it is usually performed with large-particle-size abrasives. The surface texture of the substrate after a grinding procedure is usually rough to the touch and gives a diffuse reflection to incident light (Fig 10-1(b)).

Many abrasives used in dentistry are particles that are bonded to a mandrel or disk for use in a rotary handpiece. Some abrasives are used as powders (or pastes) to be applied to the substrate using a cloth wheel, brush, or cup. The basic principles described in this chapter apply whether an abrasive is used in a bonded form or a powdered form.

Fig 10-1 Schematic diagram of diffuse and specular reflection: (a) high gloss; (b) low
Factors affecting rate of abrasion

1. A large difference in hardness between the abrasive and substrate (e.g., tooth enamel, amalgam) allows the most efficient grinding to take place. Brinell and Knoop hardness values are functions of a material's resistance to indentation, whereas Mohs values indicate one material's resistance to scratching by another (Table 10-1).

Table 10-1 Hardness of dental abrasives and substrates

<table>
<thead>
<tr>
<th>Material</th>
<th>Mohs</th>
<th>Brinell</th>
<th>Knoop</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Abrasives</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Talc</td>
<td>1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gypsum</td>
<td>2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Chalk</td>
<td>3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rouge</td>
<td>5-6</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pumice</td>
<td>6</td>
<td>450</td>
<td>560</td>
</tr>
<tr>
<td>Tripoli</td>
<td>6-7</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Garnet</td>
<td>6.5-7</td>
<td>550</td>
<td></td>
</tr>
<tr>
<td>Tin oxide</td>
<td>6-7</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sand</td>
<td>7</td>
<td>650</td>
<td>800</td>
</tr>
<tr>
<td>Cuttle</td>
<td>7</td>
<td>650</td>
<td>800</td>
</tr>
<tr>
<td>Tool steel</td>
<td>—</td>
<td>800</td>
<td></td>
</tr>
<tr>
<td>Zirconium silicate</td>
<td>7-7.5</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tungsten carbide</td>
<td>9</td>
<td>1,200</td>
<td>2,100</td>
</tr>
<tr>
<td>Aluminum oxide</td>
<td>9</td>
<td>1,700</td>
<td>1,900</td>
</tr>
<tr>
<td>Silicon carbide</td>
<td>9-10</td>
<td>3,000</td>
<td>2,500</td>
</tr>
<tr>
<td>Boron carbide</td>
<td>9-10</td>
<td></td>
<td>2,800</td>
</tr>
<tr>
<td>Diamond</td>
<td>10</td>
<td>&gt;3,000</td>
<td>7,000</td>
</tr>
<tr>
<td><strong>Substrates</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Acrylic</td>
<td>2-3</td>
<td></td>
<td>25</td>
</tr>
<tr>
<td>Pure gold</td>
<td>2.5-3</td>
<td></td>
<td>30</td>
</tr>
<tr>
<td>Hard gold alloys</td>
<td>3-4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Amalgam</td>
<td>4-5</td>
<td></td>
<td>90</td>
</tr>
<tr>
<td>Dentin</td>
<td>3-4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Enamel</td>
<td>5-6</td>
<td></td>
<td>270</td>
</tr>
<tr>
<td>Glass</td>
<td>5-6</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Resin composite</td>
<td>5-7</td>
<td></td>
<td>200</td>
</tr>
<tr>
<td>Porcelain</td>
<td>6-7</td>
<td></td>
<td>400</td>
</tr>
</tbody>
</table>
2. *The particle size* of an abrasive may be expressed in micrometers. By convention, particles are classified as fine (0 to 10 μm), medium (10 to 100 μm), and coarse (100 to 500 μm), according to the average particle size of the sample. Larger abrasive particles will abrade a surface more rapidly than will smaller particles; however, they tend to leave coarser scratches in the abraded surface than do fine particles. Equivalent-sized scratches can be produced by different sizes of particles by varying the applied pressure (Fig 10-2(a)).

![Fig 10-2](image.png)

**Fig 10-2** Factors affecting abrasion. (a) Large particle produces similar scratches with lower applied force ($F_1$) than smaller particle with greater applied force ($F_2$). (b) Sharp particle produces deeper abrasion than rounder particle under equal applied force. (c) Deeper and wider scratches are produced by increasing the applied force from $F_1$ to $F_2$.

3. *The particle shape* also has an effect on rate of abrasion. Sharp, irregularly shaped particles will abrade a surface more rapidly than will more rounded particles having duller cutting angles. Also, the former will produce deeper scratches than the latter (Fig 10-2(b)). The abrasion rate of an abrasive decreases during use; this is due partly to rounding of the particles and partly to contamination of the abrasive with some of the substrate material (debris).
4. *The greater the speed* at which the abrasive travels across the surface of the substrate, the greater the rate of abrasion. The greater friction at higher speeds, however, tends to create higher temperatures at the surface of the substrate.

5. *The greater the pressure* applied, the more rapid will be the abrasion for a given abrasive. Greater pressure produces deeper and wider scratches (Fig 10-2(c)) and creates higher temperatures (and patient discomfort).

6. *Lubricants* (eg, silicone grease, water spray, glycerol) are used during abrasion for two purposes: to reduce heat buildup and to wash away debris to prevent clogging, or *blinding*, of the abrasive instrument. Too much lubrication can reduce the abrasion rate because it may prevent some of the abrasive from coming in contact with the substrate.

*Applications of abrasives*

**Dental prophylaxis pastes**

Prophylaxis pastes should be chosen carefully and applied to remove exogenous stains without damaging the underlying tooth structure or adjacent restorative materials. The abrasive selected should be harder than the surface stain being removed and softer than the tooth surface, although this is not always practicable. If the tooth structure is excessively roughened during the procedure, it should be polished with a fine abrasive (such as zirconium silicate); otherwise, plaque and food substances will easily adhere to it.

The most common abrasives used in prophylaxis pastes are pumice, silica, zirconium silicate, and other silicates. They are usually supplied in various particle sizes (coarse, medium, fine), which produce different rates of abrasion and sizes of scratches. Sodium fluoride or stannous fluoride are incorporated into some prophylaxis pastes to help prevent dental caries.

**Dentifrices**

Dentifrice pastes are used for removing debris and minor stains from teeth and for polishing tooth surfaces. The most commonly used abrasives are dibasic calcium phosphate dihydrate, anhydrous dibasic calcium phosphate, tricalcium phosphate, calcium pyrophosphate, and hydrated alumina.

Many dentifrices contain therapeutic agents, such as sodium fluoride, stannous fluoride, or sodium monofluorophosphate, to decrease the acid solubility of tooth enamel, decrease hypersensitivity, and interrupt the mechanisms of plaque attachment and calculus formation on tooth structure. Dentifrice pastes additionally may contain a humectant to reduce evaporation of water, a surface-active detergent, binders, flavoring and sweetening agents, and a preservative.

Abrasive values for dentifrice products have been reported as an *abrasivity index* (AI), which is a measure of the abrasion of dentin.
In selecting a dentifrice for a patient, the following factors should be considered: degree of staining, toothbrushing habits (force, stiffness of brush, method of brushing), presence of relatively soft restorative materials (e.g., acrylic resin veneers, silicates), and the amount of exposed cementum and dentin.

Denture cleaners

Food debris, plaque, calculus, and stains may accumulate on denture base materials in the same way as natural teeth and are best removed on a regular basis by the patient. Daily soaking in a denture cleanser solution or brushing with or without a paste or powder is usually effective.

Chemical cleansers may contain sodium perborate, which releases peroxide. As peroxide decomposes, oxygen is released, resulting in effervescence. Effervescence, along with the oxidizing ability of peroxide, are assumed to be responsible for the cleaning action.

Dentures may also be soaked in a dilute solution of 5% sodium hypochlorite (1:3 water). Other chemical denture cleansers may contain dilute acids or enzymes. Dentures should not be soaked in hot water, which may warp the denture base material.

Brushing may be needed to remove stains and stubborn deposits. Hard-bristle brushes or brushing with much force may abrade the plastic surface of the denture and should be avoided. Dentifrices are generally too abrasive for use with dentures, although some with gentle abrasives (sodium bicarbonate or acrylic resin) can be used. Organic solvents should be avoided, as they may cause crazing and eventual cracking of the denture material. Ultrasonic vibration has not been shown to adequately remove plaque from dentures.

Handpiece instruments

Handpiece instruments include stones, burs, rubber wheels, and disks. Bonded abrasives are available for dental use in various shapes, abrasive sizes, and hardnesses:

1. **Dental stones** are composed of abrasive particles that have been sintered together or bonded with an organic resin to form a cohesive mass. These stones are available in fine, medium, and coarse grades. The color of the stone is an indication of the particular abrasive used; green stones contain silicon carbide, and white stones contain aluminum oxide. Diamond stones generally have a higher cutting efficiency than silicon carbide or aluminum oxide.

2. Dental excavating burs have a cutting action on tooth structure that is similar to grinding and polishing. Low-speed burs are composed of either carbon tool steel or tungsten carbide. High-speed burs are almost exclusively composed of the harder tungsten carbide. Burs with eight blades can be used for gross reduction of tooth structure and removal of old restorations, and they produce a rather rough surface texture on the substrate. Burs with 12, 20, or 40 blades can be used for producing a
smooth surface on tooth structure and restorations. Of significance is the relative hardness values of the burs and the substrates (see Table 10-1). Hard restorative materials (such as quartz-filled composite restorative materials and dental porcelain) will rapidly dull the sharp cutting edges of the high-speed tungsten carbide burs. Cutting efficiency and durability of the burs varies greatly among manufacturers and design types. Crosscut fissure burs have higher cutting efficiency than plain fissure burs, but they produce a rougher surface.

3. **Rubber wheels** are used for fine grinding of restorative materials (removing coarse scratches from rough grinding). They are made by molding fine abrasives (such as aluminum oxide, silicon carbide, and chromium oxide) in an elastomeric matrix.

4. **Disks** and **strips** are made by bonding abrasive particles onto a thin plastic backing. They generally wear out rapidly due to the loss of abrasive particles. They are particularly useful in finishing relatively flat surfaces. Abrasives commonly used on disks and strips are garnet, emery, aluminum oxide, and quartz (cuttle).

**Polishing**

*Introduction*

**Polishing** is the process of making a rough surface smooth to the touch and glossy (mostly specular reflection of incident light) (see Fig 10-1(a)). Polishing is usually performed with very-small-particle-size (submicron-size) abrasives.

To produce a smooth, lustrous surface by abrasion, successively smaller abrasive sizes must be used. Larger abrasive particles remove large amounts of material from the substrate, and smaller particles smooth out the roughness produced by the larger particles. Final polishing of a surface with a very fine abrasive produces a virtually scratch-free surface by creating a thin microcrystalline or amorphous layer on the surface of the substrate.

It is extremely important to remove all debris and abrasive particles from the surface of the substrate before using a finer abrasive during the polishing sequence. Even a single abrasive particle left on the substrate from a previous step will continue to scratch the surface during the subsequent polishing steps.

*Appearance and texture of polished surfaces*

A polished surface is important for esthetic and functional reasons. If the scratches produced by the abrasives are greater in width than the wavelength of visible light (ie, approximately 0.5 um), the surface will appear to have a dull finish. If the scratches are less than about 0.5 um in width, the surface will appear shiny. In addition, it has been found that the tongue can distinguish subtle differences in roughness; scratches more than 20 um deep feel rough, whereas those less than 2 um deep feel smooth.

*Techniques for polishing restorative materials*

Dental amalgam
A polished surface is desirable on dental amalgam to retard the collection of plaque and help retard tarnish as well. Burnishing alone does not create as smooth a surface as does polishing. Although an amalgam restoration can be burnished immediately after carving, most brands should be left undisturbed for at least 24 hours before polishing to allow the amalgam to set completely. Polishing can then be performed with a rotary instrument (cup, brush, or felt) with a fine abrasive mixed with water or alcohol in a slurry or paste; flour of pumice (ground volcanic glass), extra-fine silex (various silicates, such as quartz or tripoli), or tin oxide may also be used for this purpose. Care must be taken to use sufficient water or alcohol to avoid frictional heating of the restoration, which could cause pulpal damage.

Fast-setting high-copper amalgams may be successfully polished 10 to 12 minutes following placement.

Gold alloys

Gold alloys are finished by using coarse, medium, and fine abrasives in sequence. Coarse scratches are removed with fine pumice or coarse abrasive rubber wheels; the surface is finished with a rubber wheel impregnated with a fine abrasive and finally polished with tripoli and rouge on rag wheels. Particular care must be taken to avoid overfinishing of contours and margins.

Acrylic resin denture bases and veneers

Gypsum material left on the denture base following processing and deflasking may be removed with a "shell blaster." Small blemishes and bumps on the resin surface may be scraped off or removed with an acrylic finishing bur. Denture base material is comparatively soft and can be finished easily with a rag wheel and fine pumice followed by tripoli or tin oxide. Care must be taken not to alter the contour of the denture during finishing. Acrylic denture teeth are particularly easily abraded by pumice.

Resin composite restorations

Resin composite restorations present a unique polishing problem because they are composed of a relatively soft polymeric resin and a hard filler. Relief polishing may result due to unequal wear rates of the resin and filler, leaving "valleys" between the filler particles (Fig 10-3).

Fig 10-3 Uneven wear of composite surface produces valleys in the resin matrix between the hard filler particles.
This is particularly apparent in large-particle (conventional type) resin composites, which still look and feel rough after finishing with available dental abrasives (Fig 10-4).

Although relief polishing still takes place during the finishing of microfilled resin composites, the valleys and filler particles are so small that the surface appears glossy (large specular component of reflection) and feels smooth (Fig 10-5). The new types of hybrid and small-particle resin composites also tend to appear shiny and smooth but cannot be polished as easily as the microfilled resin composites.
A typical sequence for polishing resin composites would be as follows: coarse grinding with a diamond stone or a green stone followed by a series of coarse to fine quartz or aluminum oxide abrasive disks or rubber wheels. Carbide burs with 12 or more blades have also been used to polish resin composites. Finishing instruments for resin composites are available commercially. The effects of several polishing sequences on the surface roughness of resin composites are shown in Fig 10-6, which are tracings from the surface analyzer shown in Fig 10-7.
Fig 10-6 Roughness profiles of conventional resin composite surfaces finished with: (1) green to white stone; (2) silicon carbide disks; (3) 12-fluted bur; (4) 40-fluted bur; (5) green to white stone to alumina paste; and (6) diamond stone. For scale: A = 50 um; B = 2 um. (From Tolley et al, 1978. Reprinted with permission.)
Porcelain

The best way to obtain a smooth, glossy surface on dental porcelain is by glazing in a porcelain oven. After minor adjustments of the surface of a porcelain restoration, the porcelain can be polished using a series of coarse to fine abrasive rubber wheels (containing silicon carbide or aluminum oxide), followed by a fine-particle-size diamond paste applied on a felt wheel.

Bleaching

Bleaching has been used to whiten teeth since the 1800s, but with the introduction of home bleaching systems in 1989, its use has widened.

Three types of bleaching treatments are currently used (Table 10-2).

<table>
<thead>
<tr>
<th>Table 10-2 Tooth bleaching options</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Home bleaching</strong></td>
</tr>
<tr>
<td><strong>Patient selection</strong></td>
</tr>
<tr>
<td><strong>Materials used</strong></td>
</tr>
<tr>
<td><strong>Location</strong></td>
</tr>
</tbody>
</table>
### Technique

<table>
<thead>
<tr>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>A custom-fitted tray is made in a dental office. The correct strength of solution and the tray are taken home with the patient, who administers the treatment and returns periodically for follow-up.</td>
</tr>
<tr>
<td>CP is applied in a customized whitening tray and placed in the patient's mouth for at least 30 minutes per treatment. Additional applications are sent home with the patient.</td>
</tr>
<tr>
<td>The solution is applied to the teeth by a dental professional and activated with a special light or heat source.</td>
</tr>
</tbody>
</table>

### Outcome

<table>
<thead>
<tr>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Takes about 3-4 weeks for measurable results to be seen.</td>
</tr>
<tr>
<td>Some results are seen within 30 minutes of treatment.</td>
</tr>
<tr>
<td>In most cases dramatic results are seen immediately.</td>
</tr>
</tbody>
</table>

### Average number of treatments

<table>
<thead>
<tr>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Once per day for 2-3 hours for 4-6 weeks.</td>
</tr>
<tr>
<td>Can be used as a &quot;jump-start&quot; for daily home treatments.</td>
</tr>
<tr>
<td>One visit. An at-home regimen may be suggested depending on the stains being treated.</td>
</tr>
</tbody>
</table>

The first is home treatment, which involves the patient using carbamide peroxide materials in a custom-fitted tray overnight or for a few hours a day. Home bleaching kits are available for daytime or nighttime use (Fig 10-8). The other two types are performed in the dental office. One method involves application of carbamide peroxide for about 30 minutes, while the other, power bleaching, employs a 30% to 50% hydrogen peroxide solution further activated by a strong light or heat. Dental office treatments may also involve microabrasion to remove superficial staining from enamel with hydrochloric acid and a fine abrasive.

**Fig 10-8** Home bleaching kit for nighttime use.
The results of bleaching last about a year. Deep, dark stains are better handled with porcelain veneers. Side effects of bleaching include tooth hypersensitivity, soft tissue lesions, and sore throat and nausea from swallowing the bleach. Due to the possibility of tooth hypersensitivity, anesthesia should not be used during treatment, which would mask irritation of the pulp. About three office appointments are usually required for dark stains. Considerable care is needed to protect the patient's eyes and to discontinue the treatment if painful. Although bleaching can be effective, many patients have an unrealistic view of the natural color of teeth and undergo unnecessary treatment.

**Glossary**

- **abrasion** The mechanical process of wear of the surface of one material by another.
- **abrasive** The material that causes the wear of another material.
- **abrasivity index (AI)** A method of rating the abrasiveness of dentifrices.
- **amorphous** Without crystalline structure; having random arrangement of atoms in space.
- **blinding** Clogging of an abrasive wheel with debris, causing reduction of abrasive action.
- **dental stones** Grinding instruments composed of abrasive particles bound in a hard resin matrix or sintered together into a hard mass.
- **disks** Grinding and polishing rotary instruments composed of abrasive particles cemented to a flexible plastic backing.
- **microcrystalline** Composed of tiny (submicron) crystals.
- **polishing** The process of making a rough surface smooth and glossy.
- **rubber wheels** Grinding and polishing instruments composed of abrasive particles in a flexible rubber matrix.
- **strips** Instruments for grinding and polishing interproximal areas; composed of abrasive particles cemented to a flexible plastic backing.
- **substrate** The material being abraded.
- **whitening** Removal of surface discoloration by polishing or bleaching.

**Discussion Questions**
1. Since many esthetic porcelain restorations are given a final thin layer of stain glaze for color matching, how could a dentifrice cause an undesirable change in color?

2. How could polishing teeth reduce the need for bleaching?

3. Why is trying to obtain snow-white teeth by bleaching unrealistic?

4. What are the dangers of bleaching teeth, and what are the current regulations regarding the use of these products?

**Study Questions**

(See Appendix E for answers.)

1. What is an abrasive?

2. Which factors affect the rate of abrasion?

3. Place the following in order of hardness: cuttle, rouge, silicon carbide, chalk, sand, diamond, aluminum oxide, and pumice.

4. Rouge is used to polish gold alloys. What effect do you think it would have on dental porcelain?

5. Why is it important to obtain polished surfaces on dentition?

6. During grinding and polishing, what would be the effect if coarse abrasive particles from a previous step were present during the final polishing step? How can this be avoided?

7. How sensitive is the tongue to scratches?

8. What are some common abrasives found in (a) prophylaxis pastes; (b) dentifrices; and (c) stones, rubber wheels, and disks?

9. Which criteria would you use in selecting a dentifrice for a patient?

10. When should you polish an amalgam restoration? Which materials would you use?

11. Why are composite restorations difficult to polish? When would a rough surface on a composite restoration be desirable?

12. What evidence is provided regarding the performance of the large number of bleaching products?

**Recommended Reading**

American Dental Association (1983). Dentists' Desk Reference: Materials,
Instruments and Equipment, ed 2. Chicago: ADA.


Chapter 11. Structure and Properties of Metals and Alloys

Introduction

A wide variety of metals is used in dentistry, as indicated in Table 11-1. Each has a melting or solidification temperature that is characteristic of that element. When elements are alloyed together to change their properties, this single melting temperature is changed to a range of temperatures over which the liquid is in equilibrium with solid crystals nucleated in the liquid metal.

<table>
<thead>
<tr>
<th>Element</th>
<th>Unit cell</th>
<th>Melting temperature (°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mercury</td>
<td>Rhombohedral</td>
<td>-39</td>
</tr>
<tr>
<td>Gallium</td>
<td>Orthorhombic</td>
<td>30</td>
</tr>
<tr>
<td>Indium</td>
<td>Tetragonal</td>
<td>156</td>
</tr>
<tr>
<td>Tin</td>
<td>Face-centered cubic</td>
<td>419</td>
</tr>
<tr>
<td>Aluminum</td>
<td>Face-centered cubic</td>
<td>660</td>
</tr>
<tr>
<td>Silver</td>
<td>Face-centered cubic</td>
<td>960</td>
</tr>
<tr>
<td>Gold</td>
<td>Face-centered cubic</td>
<td>1,063</td>
</tr>
<tr>
<td>Copper</td>
<td>Face-centered cubic</td>
<td>1,083</td>
</tr>
<tr>
<td>Manganese</td>
<td>Cubic</td>
<td>1,244</td>
</tr>
<tr>
<td>Beryllium</td>
<td>Hexagonal close pack</td>
<td>1,284</td>
</tr>
<tr>
<td>Nickel</td>
<td>Face-centered cubic</td>
<td>1,452</td>
</tr>
<tr>
<td>Cobalt</td>
<td>Body-centered cubic</td>
<td>1,493</td>
</tr>
<tr>
<td>Iron</td>
<td>Body-centered cubic</td>
<td>1,535</td>
</tr>
<tr>
<td>Palladium</td>
<td>Face-centered cubic</td>
<td>1,552</td>
</tr>
<tr>
<td>Titanium</td>
<td>Hexagonal close pack</td>
<td>1,668</td>
</tr>
<tr>
<td>Platinum</td>
<td>Face-centered cubic</td>
<td>1,769</td>
</tr>
<tr>
<td>Chromium</td>
<td>Body-centered cubic</td>
<td>1,875</td>
</tr>
<tr>
<td>Molybdenum</td>
<td>Body-centered cubic</td>
<td>2,610</td>
</tr>
</tbody>
</table>

The upper temperature for the liquid-solid alloy range is called the liquidus temperature, and the lower temperature limit is called the solidus temperature. When a liquid alloy melt is being cooled, the liquidus temperature is the temperature at which solid crystals start to nucleate. When a mixture of an alloy liquid and crystals is being heated, the liquidus temperature is the temperature at which the crystals dissolve into liquid. The solidus temperature is the temperature at which the last liquid solidifies on cooling or the first liquid is formed on heating.

Complete melting is needed for casting and soldering and at least an additional 100°C superheat is needed for a fluid melt to cast. Note that a gas/air torch cannot be used to cast a metal with a liquidus temperature above 1,000°C because the flame does not get hot enough. Also, a reducing flame rich in gas should be used for casting or
soldering, otherwise the liquid metal will be oxidized by the oxygen in air. Oxidation keeps the solder from wetting the surface and flowing. To provide a margin of safety against melting the castings in a fixed partial denture, the liquidus temperature of the solder should be at least 50°C lower than the solidus temperature of the casting alloy being soldered.

**Unit Cells of Crystal Lattices**

Liquid metals nucleate crystals upon cooling (Fig 11-1). The atoms joining the crystals form a unique packing arrangement in space that is characteristic of that metal or alloy at equilibrium. The smallest division of the crystalline metal that defines the unique packing is called the **unit cell**. When the unit cell is repeated in space, the repeating atomic positions form the **crystal lattice** structure of a crystalline solid (Fig 11-2). The atoms at the corners of the unit cell are shared among the adjacent eight unit cells, as shown for the body-centered unit cell. Therefore, one eighth of the corner atom is associated with the cell; there are eight corner atoms, so they each contribute one atom to the unit cell. The body-centered atom is totally inside the unit cell and is not shared, so it contributes the second atom to the unit cell mass. Using the **lattice parameters** to calculate the volume of the cubic cell, the density of the metal can be calculated by dividing the mass of atoms in the unit cell by its volume. The lattice parameters for metals and alloys range between 2 Å and 10 Å for the different unit cells formed.

![Fig 11-1](image)

**Fig 11-1** Formation of crystal nuclei in liquid metal. O = atoms in liquid state; ● = atoms in solid state.
Fig 11-2 (a) The body-centered cubic unit cell is typical of the crystal lattice of pure iron at room temperature. The lattice parameter for iron is 2.87 Å. (b) A part of a body-centered cubic crystal lattice. It could extend in all directions. In this "hard sphere model," the atoms are visualized as hard spheres of a definite radius in contact. (c) The body-centered cubic space lattice can be visualized as a "point skeleton" of the body-centered cubic crystal lattice.

It has been observed that the position of the neighboring atoms surrounding every atom of a crystal lattice is identical in a pure crystalline metal. When the property of identical periodic points in space was explored mathematically, it was discovered that there are 14 unique ways to arrange points in space (Fig 11-3).
Fig 11-3 Unit cells of the 14 space lattice contain atoms arranged so that each one has identical surroundings. (From Mott, 1967. Reprinted from Scientific American.)

These are called space lattices. A pure metal crystal lattice is similar to one of the space lattices except that each mathematical point is the site of an atom. Complex crystal lattices like amalgam alloy and enamel have the points of their space lattices replaced by the different atoms of the material or by groups of atoms. The unit cell of each crystalline material, no matter how complex, corresponds with one of these 14 space lattice unit cells (see Table 11-1).
Nucleation and Polycrystalline Grain Structure

Introduction

As the melt of metal is cooled, clusters of atoms come together from the liquid to form solid crystal nuclei. These nuclei will be stable and grow into crystallites or grains if the energy of the system is favorable, that is, the energy is lowered by the process. The energy is lowered by an atom bonding to the solid nuclei, thereby giving up its liquid-state kinetic energy of motion. However, when an atom bonds to the nuclei, the energy can also be raised by the creation of more interfacial surface energy as a result of the increased surface area of the nuclei in contact with the liquid. The energy of the system is favorable for stable nuclei and growth when more energy is lost by bonding than is gained by increasing the interfacial surface area (i.e., energy).

Nucleation can occur by two processes. The first, called homogeneous nucleation, is enhanced by rapid cooling so the nuclei are supercooled. The result for the system is that more energy is lost when an atom of the liquid bonds to the solid. With rapid cooling (quenching in water) more nuclei are formed per unit volume. These nuclei grow together to form the irregular polycrystalline grains or crystallites that fit together like a three-dimensional puzzle to form the bulk of the metal shape (Figs 11-4 and 11-5).

Fig 11-4 Irregular polygons called grains or crystallites. An average distance measured across the faces of the crystal grains is called the grain size. It may be less than 1,000 Å or more than 1 cm, depending on the number of nuclei present during solidification. (From Guy et al, 1971. Reprinted with permission.)
Fig 11-5 The grain structure of a metal is revealed by polishing the surface to a mirror finish and etching lightly in acid. To study the grain structure of metals used for dental appliances, a light or scanning electron microscope is needed for magnification because of the small grain sizes. (From Guy et al, 1971. Reprinted with permission.)

The more nuclei that are formed by rapid cooling, the smaller the grain size or crystallite dimensions. Another means of decreasing the grain size (grain refining) is by adding to the melt a foreign solid particle or surface to which the atoms are attracted, such as a very fine high-melting metal or oxide powder. This process of seeding the nuclei is called heterogeneous nucleation.

Grain size and properties

Decreasing the grain size can have a number of beneficial effects on the cast alloy structure of a crown or removable partial denture. The finer grain size can raise the yield stress, increase the ductility (percent elongation), and raise the ultimate strength. The change in these properties with grain size is related to the processes of plastic deformation and fracture, and to how the boundaries between grains relate to these processes. The size of metal grains in different metals may range from less than 1,000 Å to more than 1 cm. Grains contain large numbers of unit cells—even grains of only 1,000 Å across. The lattices of the grains are formed in random directions when they grow from the melt. A boundary is formed where the grains grow into contact, because the atoms in one grain's crystal lattice are not in position to mesh with the repeating atoms in the crystal lattices of adjacent grains. These grain boundaries are layers several atoms thick that are distorted from normal atomic positions in order to bridge the mismatch in the lattice orientations of adjacent grains.

Only metals with simple body-centered or face-centered cubic unit cells have enough densely packed planes of atoms in their lattices to allow plastic deformation at yield stress. These lattice types permit shearing of the densely packed planes of atoms like cards of a microscopic deck sliding over each other. However, the lattice of adjacent
grain can be viewed as a second microscopic card deck at a different angle. To get the metal to deform, it is necessary to force the cards of one deck into other decks at an angle. But the more grains per unit volume, the more difficult it is to get the planes (cards) to slide because the dislocated slipping planes run against the grain boundaries sooner. Thus, a greater resistance to slippage is created by more grain boundaries, and higher yield stress results.

On the other hand, a material will fracture because a crack opens up on a grain boundary. This is more likely to occur in large-grain metals, when the planes cannot be slipped into the adjacent grains. Many smaller grains in various orientations can divide the plastic strain among the grains more easily, with more of them oriented for slipping. Large grains must each accommodate a larger strain and will have fewer properly oriented to slip. The result is lower ductility and lower ultimate strength for large-grain metals, which open cracks more readily at grain boundaries because the plastic deformation cannot be accommodated. For these reasons, grain-refined or "micrograin" alloys produced by heterogeneous nucleation are advantageous for developing fixed partial denture alloys with higher yield stress, better ductility, and improved ultimate strength.

**Alloy Systems**

Most pure metals are miscible in the liquid state when melted together. When two metals form a solution in the liquid state so their atoms mix randomly on the atomic scale, they are said to form an alloy. As the alloy liquid freezes, the atoms may remain randomly distributed on the unit-cell lattice sites in each crystal grain. This random distribution in the solid alloy is called a solid solution. But, if like atoms tend to prefer to bond among themselves, then as the nuclei form from the melt, the atoms of different elements may segregate in different grains. The grains of the two different metal elements are mixed together.

Different grains may be practically pure if their elements are insoluble in each other's lattices in the solid state. Or they can have a limited solubility in the other's crystal lattice if the elements exhibit partial solubility in one another. Metal atoms of two different metal elements are more likely to be soluble in each other's lattices if they (1) have the same atomic lattice type, (2) have similar atomic radii (ie, a difference of less than 10%), (3) have the same valence number, and (4) form bonds to other atoms with strengths similar to those they form among themselves. On the other hand, if these rules are not followed and the unlike atoms have a strong affinity to each other, grains of an intermetallic compound may be formed at definite ratios of the alloying elements (eg, dental amalgam alloy Ag₃Sn).

The energetically stable (ie, at equilibrium) crystal lattice structures and their compositions for an alloy that is preferred by "nature" varies with temperature and ratio of the alloying elements that are melted together. It is not possible to calculate the equilibrium composition and structures and at what temperatures these change. They must be determined experimentally by measuring the temperatures at which the latent heat is liberated when alloy liquid solidifies or solid lattices transform to different crystal lattices. The type of crystal lattice is determined by x-ray diffraction from the crystal atomic planes. The angle and intensity of the x-ray beam reflections (ie, diffraction) are characteristic of the atomic composition, type of crystal lattice,
and position of atoms in their unit cell. Thus, experimental detection of the temperature when heat is liberated indicates when an alloy is changing its structure, and x-ray scattering identifies what lattices are present. This information is portrayed in a **binary phase diagram** of the alloy system (Figs 11-6 and 11-7). The alloy system represents all possible ratios of the elements.

![Copper-gold phase diagram](image)

**Fig 11-6** Copper-gold phase diagram. The disordered solid solution (softer) and ordered solid solutions (harder) are produced by heat treatment. FCC = face-centered cubic.

When two elements are alloyed, the system is called a **binary alloy**; when three elements are alloyed, the system is a **ternary alloy**, and so on. An alloy is named by listing its elements in descending order of percent composition. For example, Ag₃Sn is called silver-tin alloy. If more than two elements are involved, the number of phase changes and their representation becomes complex, but their description follows the same principle.
Fig 11-7 Silver-copper phase diagram showing a eutectic (lowest melting) point at 28.1% copper.

An equilibrium phase is defined as a homogeneous body of matter that is physically distinct and mechanically distinguishable. For a pure material like water, the vapor, liquid, and solid phases are physically distinct because there is a definite boundary between the regions when they exist together. Also, they are mechanically distinguishable in properties like hardness, compressibility, and elastic modulus. However, when the phase definition is applied to an alloy system, it is important to recognize that if two different types of unit cells nucleate from the melt—as, for example, at the eutectic (lowest melting) point in Fig 11-7—a two-phase region is formed. The two phases nucleate as separate grains. They are physically distinct, as indicated by grain boundaries that define their limits. Their mechanical properties differ, as can be measured by a microhardness tester impinging on individual grains. Note that two-phase alloys are not as corrosion resistant as like single-phase alloys, because microscopic galvanic corrosion cells are set up between the grains of the different phases. Also, porcelain bonding to multiphase alloys is considered potentially weaker because of composition differences of the grains.

**Deformation in Metals**

There are three types of deformation that can occur in metals, which arise from different mechanisms. The simplest deformation, elastic strain, is the elastic stretching of lattice in which all the atoms are shifted from their equilibrium positions by a fraction of their atomic spacing. The strain is directly proportional to the applied stress (ie, force/area or force intensity) up to the proportional limit stress. When the stress is removed, the atoms return to equilibrium atomic spacing. Compared with most polymeric materials, metals generally have strong metallic bonds and resist
elastic stretching. This stiffness or resistance to elastic strain is indicated by the high elastic modulus of metals. Stiffness is desirable for removable partial dentures, so forces can be transmitted by the framework across an arch to better distribute the load. It is desirable for the alloy of resin-bonded fixed partial dentures to resist flexing of the bond.

Another type of deformation is plastic deformation, a permanent deformation that begins when the elastic limit stress or its approximation, yield stress, is reached. This mode of deformation requires that atoms be shifted to new atomic sites on the lattice. These lattice sites must be identical to the old sites and not far away, so the energy to shift atoms is not too great. Thus, ductility is associated with face-centered and body-centered cubic metal lattices, which have more identical sites and more closely packed planes, so atoms do not have to slide far to reach the new lattice sites. Intermetallic compounds are usually brittle because the atomic sites are specific to the different atoms of the compound and not interchangeable.

The mechanism of plastic deformation is called **dislocation motion** and produces the slipping of the closely packed planes over each other. A **dislocation** (Fig 11-8a, $D_x$) is the line of atoms that denotes the edge of an additional half-plane of atoms that appears to be wedged into the lattice. The lattice is distorted (strained) by the presence of the dislocation line of atoms.

![Fig 11-8a](image) The distortion (strain) around a dislocation in a simple cubic crystal lattice. The size of the atoms has been reduced so the perspective can be seen. The row of atoms of the dislocation is $D_x$. (From Ziman, 1967. Reprinted from *Scientific American*.)

Dislocation motion shifts atoms from one lattice site to the next, rather than moving all the atoms of the plane at one time, which would take much more energy (Fig 11-8b).
Fig 11-8b The crystal lattice as it would appear after the yield stress has been exceeded and the dislocation has moved through the lattice. The solid has been deformed to this new shape. (From Ziman, 1967. Reprinted from *Scientific American*.)

It is important to understand that any process that impedes dislocation movement tends to harden a metal, raise its yield stress and often lower its ductility. Some processes are reversible, allowing the hardness to be low when making an appliance, for example, after which the metal can be hardened to provide better service if necessary.

One process for hardening metal is called **cold working**, or work hardening—any plastic deformation of metal by hammering, drawing, cold forging, cold rolling, or bending. These processes produce many dislocations in the metal that cannot slip through each other as easily as the lattice becomes more distorted. The yield stress can be raised more than 100% when a drawn orthodontic wire is compared to the as-cast metal. In dentistry, cold working occurs when gold foil is compacted, a denture clasp is bent, an inlay margin is burnished, or a deformed metal layer forms on a crown during finishing and polishing.

A second process for hardening a metal is precipitation hardening. In this process a second phase of finely dispersed clusters of atoms are precipitated from a metastable supersaturated solid solution by reheating an alloy that was quench-cooled to form the metastable supersaturated solid.

Heat treatment can also be used to harden gold-copper alloys by a slightly different process. In this case, because the gold-copper system forms a complete solid solution at all compositions, the atoms can be interchanged on the lattice sites. However, as the atoms are cooled, the copper and gold atoms tend to separate on alternating planes of the lattice in ordered arrangement. This ordering makes dislocation motion more difficult, raising the yield stress. These alloys are soft if quenched but hard if cooled.
slowly on the bench top or held in the furnace at the ordering temperature range (350°C for 30 minutes).

Other heat treatments are used to homogenize the grains of an alloy that have developed composition gradients by rapid cooling and crystallization of the melt. When cooled rapidly, the grains of the alloy cannot maintain an equilibrium or uniform composition in the grain. The first part of the grain to cool will be richer than the average composition of the higher-melting-temperature element, and the last part of the grain to solidify is richer in the low-melting alloy component. This inhomogeneity tends to reduce corrosion resistance of the metal because of the galvanic cell created between the center of the grain and the grain boundary. Heating the solid at a temperature and for a duration that allows the atoms to reach equilibrium will improve corrosion resistance.

**Diffusion in Metals**

Atoms of a crystalline solid vibrate about their fixed lattice positions. As they absorb heat with increasing temperature, they vibrate at greater amplitude. Also, a small fraction of the atom sites are vacant in a metal. As the temperature increases, an atom may momentarily experience an increase in vibrational energy, which will be sufficient to permit it to jump from one lattice site to another. As the temperature and time increase, more atoms will be able to jump or diffuse to new lattice sites. This random jumping during heat treatment allows the rearrangement of the crystal lattice. Atomic diffusion also permits rearrangement of cold-worked metals and permanent deformation of metals at stress levels far below the yield stress if the stress lasts long enough. In general, these rearrangements by diffusion begin to occur at a significant rate as the temperature of metal exceeds one half of its absolute melting temperature. The rate of rearrangement accelerates rapidly as the metal or alloy approaches its solidus temperature.

There are several heat treatment processes for metals that utilize diffusion. Among them is an annealing heat treatment for release of residual stress. It allows the dislocations and atomic vacancies to move and realign to lower the internal residual stress fields in a cold-worked metal at a relatively low temperature and short heating time.

If the temperature and heating times are extended for a cold-worked metal, the metal can experience recrystallization, in which stress-free grains are crystallized out of the deformed grains. A recrystallization temperature is the temperature at which it would take 1 hour for the cold-worked metal to recrystallize. This temperature is between one third and one half of the absolute melting temperature for most metals (Fig 11-9). As the temperature and time are further extended, grain growth occurs, during which the stress-free grains grow larger at the expense of the disappearing small grains.
A microscopic view of the stages that a cold-worked metal, such as an orthodontic wire, goes through when subjected to prolonged heating. The higher the temperature and the greater the cold working, the more quickly the transitions occur. This figure shows recrystallization being completed in 1 hour, which means the temperature of heating was the recrystallization temperature as per the definition. The changes in properties can be compared with the structural changes. (From Jastrzebski, 1959. Reprinted with permission.)

The residual stress heat treatment relaxes the internal stress, which may cause warpage over time. The chance of corrosion related to residual stress differences on the metal surface are reduced. Recrystallization heat treatment lowers the yield stress and increases the ductility. Grain growth further reduces yield stress and increases ductility yield stress. When soldering a cold-worked wire in which a high yield stress is desired (eg, a partial denture wire clasp or an orthodontic appliance), a low-melting solder and a short soldering time are desirable to minimize recrystallization and grain growth.

**Glossary**

*alloy* An atomic mixture of metallic elements.

*annealing* A relatively low-temperature heat treatment for removing residual stress.

*binary alloy* An alloy composed of two elements.
**binary phase diagram** A map with temperature and composition as coordinates, which displays the regions where each stable phase exists.

**cold working (work hardening)** Deforming a metal at temperatures that are low compared with its melting temperature.

**crystal** A solid with periodic arrangement of atoms in space, usually with atomic planes forming facets on the surface.

**crystal lattice** The periodic arrangement of atoms in three-dimensional space.

**crystallite** A crystal with irregularly shaped surfaces instead of facets.

**crystallization** The process of crystal and crystallite formation.

**deformation** The process of changing the shape of a metal by applied stresses.

**dislocation** A row of atoms displaced from their normal positions in the lattice.

**dislocation motion** The movement of a dislocation through a crystal under an applied stress.

**eutectic alloy** An alloy, easily melted with respect to its component elements, that can transform at one temperature from liquid to two phases separated as distinct grains in the solid metal.

**eutectic composition** The alloy composition at which the eutectic transformation occurs.

**eutectic temperature** The temperature at which the eutectic transformation occurs.

**grain** Another name for crystallite.

**grain boundary** The interface or junction of adjacent grains.

**grain growth** Enlargement of grains by heating.

**hardening** A process in which the yield stress and resistance to indentation are increased.

**homogeneous** Having uniform composition throughout.

**intermetallic compound** An alloy phase with a composition usually near a definite fixed atomic ratio of the elements.

**lattice parameters** The distances between corners of the unit cell.

**nuclei** The embryos of the crystallites formed from the liquid.
**phase** A homogeneous body of matter that is physically distinct and mechanically distinguishable.

**polycrystalline** Composed of many crystallites.

**recrystallization** The process of forming new crystallites from existing crystallites by heating.

**residual stress** Internal stress remaining between parts of a solid after the applied stress is removed.

**solid solution** An alloy phase in which one alloying element enters the lattice of the other.

**space lattice** A pattern of points in space that satisfies the condition that each point is surrounded by the same arrangement of points.

**ternary alloy** An alloy composed of three elements.

**ultimate strength** The maximum stress a solid can support based on its original cross-sectional area.

**unit cell** The minimum grouping of atoms of a homogeneous crystalline solid that gives the geometric relationships, composition, and distance between the atoms in space.

**yield stress** The stress at which dislocation motion, permanent deformation, and plastic flow begin.

**Discussion Questions**

1. Gold and copper are completely soluble in each other in the liquid state. What is the connection between this mutual solubility and their lattice structure?

2. Which mechanical properties of a metal are affected if dislocation motion in the lattice is impeded?

3. How can both the strength and ductility of gold castings be raised by alloying and process control?

4. Why does overheating orthodontic wires lead to brittleness?

**Study Questions**

*(See Appendix E for answers.)*

1. How does the melting of a pure metal differ from the melting of an alloy?

2. What effect does incomplete melting have on dental castings and solder joints?
3. What is the relationship between the density of a molten alloy and its crystalline solid? Is it the same, less, or greater? Why?

4. How can the density change from liquid to solid damage a dental casting?

5. How many atoms are contained in a face-centered cubic unit cell?

6. Outline how the density of a face-centered cubic metal can be calculated from knowledge of the composition of the unit cell, the atomic weight of the atoms present, the type of unit cell, and the lattice parameters for the unit cell.

7. Which of the following metals are capable of experiencing a large amount of ductility? (Their unit cell types are provided.)

   Ag₃Sn—orthorhombic

   beryllium—hexagonal

   copper—face-centered cubic

   gold—face-centered cubic

   iron—body-centered cubic

8. List several ways in which the yield stress of a metal or alloy may be increased.

9. How can dislocation motion be impeded so as to raise the yield stress?

10. Is it better to have large or small crystallites in order to have a high yield stress and ultimate strength?

11. The grain boundaries of the crystallites at the surface of a metal can be made visible by polishing to a mirror finish with fine abrasives and etching the surface lightly with an acid.

   a. If 0.5 um is the resolution of a light microscope, how small a crystal grain might one see?

   b. Why does acid etch the grain boundaries more than the interior of the grain? Why are different grains etched at different rates?

12. What are the eutectic temperature and composition for the silver-copper system?

13. At what temperatures does solidification begin and end for an alloy containing 40% by weight copper and 60% by weight silver?

14. The composition of an alloy can be specified by the weight percentages or the atomic percentages of its elements. Given the alloy Ag₃Sn, calculate the atomic percentage of silver and the weight percentage of silver if the gram atomic weight of silver is 107.9 and the gram atomic weight of tin is 118.7.
15. How do a crystal lattice and a space lattice differ?

16. Discuss how the yield strength and percentage elongation change with time during the heating of a drawn wire.

17. If the recrystallization temperature of an iron wire is 450°C, how long would it take a drawn iron wire to recrystallize at that temperature? What fraction of the absolute melting temperature (ie, degrees Kelvin) of iron is 450°C?

**Recommended Reading**


Chapter 12. Dental Amalgams

Introduction

Amalgam has been an accepted part of dental therapeutics for more than 150 years. It is still used for more than 75% of direct posterior restorations today. The reasons for its popularity lie in its ease of manipulation, relatively low cost, and long clinical service life. Some concern has arisen with reference to mercury from both a biological and an environmental viewpoint; however, it is presently believed that dental amalgam presents an acceptable risk-to-benefit ratio when properly used. An exception to this position has been taken in northern Europe, where concerns have been raised regarding amalgam use in populations thought to be susceptible to mercury exposure, such as pregnant women and pedodontic patients.

Amalgam's primary application in dentistry is for the restoration of posterior teeth and, to some degree, for cores for fixed partial denture buildups. Small Class V restorations, although formerly representing a major clinical use for amalgams, are virtually always carried out using composite and hybrid glass-ionomer restorative materials. Posterior lesions are restored primarily by amalgam, although composites have been employed successfully in small cavities as a result of recent improvement in their clinical handling.

Chemical Composition and Microstructure

Introduction

A general classification of dental amalgam alloys is given in Fig 12-1 and Table 12-1. Contemporary amalgams are mainly classified as high-copper amalgams and have existed since the 1960s. In an attempt to circumvent the problems with mercury, a radically new composition containing a gallium-indium-tin liquid that completely replaces mercury was introduced. To understand the significance of these materials and how knowledge of the solid-state reactions occurring in amalgams has made superior clinical materials, it is first necessary to look at the so-called traditional amalgams.
Fig 12-1 A general classification of amalgam alloys.

**Table 12-1** Classification of amalgam alloys

<table>
<thead>
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<th>Composition</th>
<th>Morphology</th>
<th>Example</th>
</tr>
</thead>
<tbody>
<tr>
<td>Traditional</td>
<td>Lathe cut</td>
<td>Aristaloy</td>
</tr>
<tr>
<td>Traditional</td>
<td>Spherical</td>
<td>Spheraloy</td>
</tr>
<tr>
<td>High copper</td>
<td>Lathe cut (single composition)</td>
<td>Epoque 80</td>
</tr>
<tr>
<td>High copper</td>
<td>Spherical (single composition)</td>
<td>Tytin</td>
</tr>
<tr>
<td>High copper</td>
<td>Admixed (traditional + Ag-Cu</td>
<td>Dispersalloy</td>
</tr>
<tr>
<td></td>
<td>eutectic)</td>
<td></td>
</tr>
</tbody>
</table>

**Traditional amalgam alloys**

Lathe cut

Until the 1960s, the chemical composition and microstructure of available amalgam alloys were essentially the same as those of the most successful systems investigated by G.V. Black (Black, 1895). **Traditional alloys** were delivered to the dentist as filings, which were lathe cut from a cast ingot. Milling and sifting produced the ultimate particle size distribution, as well as the final form of the amalgam alloy particles. Figure 12-2 illustrates a typical traditional **lathe-cut** alloy.
A commercial alloy evolved into a blend of different particle sizes rather than a unimodel system, in order to optimize packaging efficiency. The length of particles in a commercial lathe-cut alloy might range from 60 to 120 um, their width from 10 to 70 um, and their thickness from 10 to 35 um. The particle size has become still smaller (< 30 um) due to the introduction of so-called spherical alloys, as discussed later. The traditional alloys contain 66% to 73% silver by weight; tin varies from 25% to 29% by weight, and the amount of copper may be as high as 6% by weight. Zinc may be present up to 2% by weight. Up to 3% mercury by weight may also be present.

The structures of these traditional alloys are essentially phase mixtures of the \textit{gamma phase} of the silver-tin system (Ag$_3$Sn) and the \textit{epsilon phase} of the copper-tin system (Cu$_3$Sn). It has been shown that Ag$_3$Sn produces the best physical properties when reacting alloys of the silver-tin system with mercury (Gruber et al, 1967). Some of these traditional alloys are still available, but they represent only a minor component of the overall amalgam market.

\textbf{Spherical}

The \textit{spherical alloys} were introduced on the market during the 1960s. Generally, their particle shape is created by means of an atomizing process whereby a spray of tiny drops is allowed to solidify in an inert gaseous (ie, argon) or liquid (ie, water) environment. Although all alloys produced in this way are classified as spherical, their particle shape might be irregular (\textit{Fig 12-3}). Generally, the maximum particle size in a spherical alloy powder is 40 to 50 um or less, although there usually is a particle size distribution. Spherical traditional amalgam lowered the necessary \textit{mercury-alloy ratios} and dramatically reduced \textit{condensation} pressures.
High-copper blended and single composition amalgam alloys

During the late 1960s, alloys with a significantly different chemical composition were introduced on the market. All of these alloys could be characterized by their higher copper content. A list of current high-copper alloy products is given in Table 12-1.

The first alloy of this type (Dispersalloy, Johnson & Johnson) (Innes and Youdelis, 1963) was a mechanical mixture of two parts of a traditional lathe-cut alloy with one part of a spherical alloy (Fig 12-4).
The chemical composition of the spherical particle was 72% silver by weight and 28% copper by weight; it corresponds to the **eutectic composition** of the silver-copper system. The overall composition of this alloy contained approximately 13% copper by weight. This was more than twice the maximum amount permitted in the American Dental Association's (ADA's) specifications for dental amalgam alloy at that time. Amalgams made from this alloy, however, were clinically superior to traditional amalgams with respect to marginal integrity (Mahler et al, 1970), and consequently other manufacturers developed similar compositions featuring some with a copper content greater than that found in traditional amalgam. At present, the copper content varies up to approximately 30% by weight in some commercial amalgam alloys (Table 12-2).

### Table 12-2

Typical compositions of amalgam alloys (wt %)

<table>
<thead>
<tr>
<th>Type*</th>
<th>Ag</th>
<th>Sn</th>
<th>Cu</th>
<th>An</th>
<th>Other</th>
</tr>
</thead>
<tbody>
<tr>
<td>TL</td>
<td>70.9</td>
<td>25.8</td>
<td>2.4</td>
<td>1.0</td>
<td></td>
</tr>
<tr>
<td>TS</td>
<td>72.0</td>
<td>26.0</td>
<td>1.5</td>
<td>0.5</td>
<td></td>
</tr>
<tr>
<td>HCS</td>
<td>41.0-61.0</td>
<td>24.0-30.5</td>
<td>13.0-28.3</td>
<td>0-0.5</td>
<td>In 3.4</td>
</tr>
<tr>
<td>HCAd</td>
<td>62.0-69.7</td>
<td>15.1-18.6</td>
<td>12.0-22.7</td>
<td>0-0.9</td>
<td>In 10</td>
</tr>
<tr>
<td>HCL</td>
<td>43.0</td>
<td>29.0</td>
<td>25.0</td>
<td>0.3</td>
<td>Hg 2.7</td>
</tr>
<tr>
<td>GA</td>
<td>50.0</td>
<td>26.0</td>
<td>15.0</td>
<td></td>
<td>Pd9</td>
</tr>
</tbody>
</table>

*TL = traditional lathe cut; TS = traditional spherical; HCS = high-copper spherical; HCAd = high-copper admixed; HCL = high-copper lathe cut; GA = alloy for gallium amalgam.

The structures of several high-copper alloys are similar to that of Dispersalloy. They can be classified as **blended alloys** in which the traditional and high-copper phases are mechanically blended. Other alloys are produced by melting together all components of a high-copper system and creating a single-composition spherical or lathe-cut alloy, rather than a mechanical mixture of two distinct powders. Depending on the number of components involved, these systems are also referred to as **ternary** or **quaternary alloys**, or as a **single-composition alloy**.

Some amalgam alloy producers, in an effort to improve clinical handling properties, supply **admixed alloys**. In these, the chemical compositions and physical forms of the basic powders (lathe cut or spherical) are varied. This system further differs from those using Dispersalloy in that both blended components are representative of copper-enriched alloys. Several typical classifications and compositions of high-copper systems are presented in Table 12-2. It is important to stress that all of these copper-enriched alloys contain > 10% copper by weight in the form of either the silver-copper eutectic or the copper-tin system. The list in Table 12-2 is representative but by no means comprehensive. The dynamics of the current marketplace preclude a comprehensive listing as a great number of alloys appear and disappear worldwide in response to local demand.

Although amalgam alloys containing many other metals have been proposed or investigated on an experimental basis, at present only indium, palladium, and selenium have been used as commercial additives. Because of economic reasons, the alloys intended for mercury amalgamation and containing palladium feature a
relatively low (< 1%) concentration. Selenium has been added in an attempt to improve the biocompatibility of the amalgam (Sato and Kumei, 1982). Indium has been admixed in large concentrations (10% by weight) in metallic form to a high-copper amalgam in order to reduce the mercury vapor released during mastication (Powell et al, 1989; Youdelis, 1992).

**Gallium alloys**

Although called gallium alloys, these systems make use of the fact that the melting temperature of gallium can be suppressed below room temperature with the addition of appropriate amounts of indium and tin. This liquid can then be triturated with a silver-tin-copper alloy powder (spherical) in the same fashion as dental amalgam. Significant amounts of palladium are added to the alloy powder in current commercial compositions to improve corrosion properties. A current composition marketed in Japan known as Gallium alloy GF (Tokuriki Honton) comes as a powder and contains the following elements (by weight): silver, 50%; tin, 25.7%; copper, 15%; palladium, 9%; and traces, 0.3%. It is also available as a liquid containing gallium, 65%; indium, 18.95%; tin, 16%; and traces, 0.5%.

**Setting Reactions and Microstructure**

**Traditional amalgams**

The amalgamation reaction of the traditional alloy with mercury (known as trituration) as well as its microstructure after setting are described on the basis of a reaction of Ag$_3$Sn (gamma) with mercury. Copper and/or zinc are not usually taken into account, but their presence has important effects. During hardening, new reaction products with mercury are formed at the cost of the original alloy particles. The main reaction products formed are the gamma 1 (silver-mercury) and gamma 2 (tin-mercury) phases. Formation of a network is completed before all the original reactant is consumed. This amalgamation reaction can be symbolized as follows:

\[
\text{Ag}_3\text{Sn} + \text{Hg} \rightarrow \text{Ag}_2\text{Hg}_3 + \text{SnHg} + \text{Ag}_5\text{Sn} \\
\gamma \gamma + \text{Hg} \rightarrow \gamma \text{gamma 1} + \gamma \text{gamma 2} + \text{gamma (remnant)}
\]

After completion of the amalgamation reaction, the remnants of the high-melting-point silver-tin particles are embedded in a matrix of reaction products with mercury (Fig 12-5). In the majority of the traditional amalgams both the gamma 1 and gamma 2 phases form a continuous network. The formation of such an interconnecting structure is extremely important because the gamma 2 phase is prone to corrosion and should be considered the weak link in many traditional dental amalgams. The copper contained in the original alloy will react with tin during trituration to form the eta prime phase Cu$_6$Sn$_5$. The presence of copper has long been associated with improving the physical properties of amalgam, particularly its flow or deformation under static load. This effect is magnified in high-copper amalgams. The presence of zinc appears to extend the working time and, hence, the plasticity of the traditional amalgam.
High-copper amalgams

All high-copper amalgams are characterized by the gamma 2 phase being either absent or substantially reduced, because tin preferentially reacts with copper rather than with mercury, preventing the formation of the tin-mercury reaction product. During amalgamation of blended alloys, Cu₆Sn₅ is created from copper and tin. (The same process occurs in traditional amalgams, but to a lesser extent, because the copper concentration is less.) Because most of the reactive copper is present in the silver-copper spheres, the Cu₆Sn₅ phase is formed at the surface of these particles, creating a reaction zone that is easily identified in the microstructure (Fig 12-6). The mechanism to form Cu₆Sn₅ can be described by

\[ 6 \text{Cu} + 5\text{Sn} \rightarrow \text{Cu}_6\text{Sn}_5 \]

In single-composition systems, Cu₆Sn₅ also will be formed during amalgamation reactions. In this case, however, the reaction is thought to be

\[ \text{Cu}_6\text{Sn} + 3\text{Sn} \rightarrow \text{Cu}_6\text{Sn}_5 \]

because the source of copper is the epsilon phase in single-composition alloys. It is obvious that in dental amalgam alloys where equivalent amounts of copper and Cu₃Sn are present, both types of reaction to form Cu₆Sn₅ may play equally important roles. It should be stressed here that some high-copper amalgams may initially contain the gamma 2 phase if the mercury content is higher than a certain critical percentage. In these amalgams, the elimination of gamma 2 may occur over a substantial time period as the reactions are diffusion controlled. In general, in lathe-cut and blended alloys the mercury-alloy ratios are \( \geq 1.0 \), whereas in spherical alloy systems the mercury-alloy ratios are \( < 1.0 \) and may be as low as 0.7. As mentioned earlier, the gamma 2 phase is considered the weak link in a traditional low-copper amalgam. However, in high-
copper amalgams the gamma 2 phase is absent. In that case, attention should be focused on the least resistant phase in the multiphase structure associated with high-copper amalgams. Preferential corrosion of the Cu$_6$Sn$_5$ phase reportedly has been shown to be significant both in vivo (Marshall et al, 1980) and in vitro (Averette et al, 1978).

Evidence has been presented for the presence of an additional tin-mercury phase, delta 2 (Sarkar, 1994a), at the grain boundaries of the resulting gamma 1 network. This phase results from the lower tin concentration in the last mercury to solidify. Since it is located at grain boundaries, it will have significant influence in determining the structure-sensitive properties of amalgam. Since copper and tin will preferentially combine in dental amalgam, the higher copper concentrations will also reduce the formation of delta 2.

As mentioned earlier, admixing of indium has lowered the amount of mercury vapor released from amalgam. This phenomenon has also been verified recently for amalgams prepared from a mercury-indium liquid in which the indium concentration was as high as 30% (Okabe et al, 1994). It is possible that through solid solution, indium may increase the stability of the gamma 1 phase (Sarkar, 1994b).

![Fig 12-6 Scanning electron micrograph of Dispersalloy amalgam. (a) Ag$_3$Sn; (b) silver-copper eutectic; (c) eta prime phase (Cu$_6$Sn$_5$); (d) gamma 1 phase (Ag$_2$Hg$_3$); (e) epsilon phase (Cu$_3$Sn). (Original magnification × 1,000. Courtesy of T. Okabe.)](image)

**Gallium amalgams**

The structure of gallium amalgams has been interpreted in terms of a reaction zone of
CuGa₂ and PdGa₅ surrounding the unreacted alloy particles, which are held together by a matrix of Ag₉In₄ in which islands of Ag₉Ga₃ and beta-tin can be found. The structure of set gallium amalgam (GF alloy) is shown in Fig 12-7.

**Physical Properties**

The physical properties of dental amalgam are usually compared to those specified in the American National Standards Institute/American Dental Association (ANSI/ADA) specifications for dental amalgam. These properties are (1) 1-hour compressive strength, (2) creep (or resistance to static load), and (3) dimensional change. The ANSI/ADA limits are (1) 1-hour compressive strength of at least 80 MPa (11,000 psi), (2) dental creep of no more than 3%, and (3) dimensional change of ± 20 μm/cm. The corresponding properties of several commercial alloys are given in Table 12-3. The rationale for these properties is that high early strength is important to withstand dental finishing procedures and occlusal stresses. Low creep is desirable for maintaining marginal integrity, and dimensional change must be controlled to prevent excessive marginal leakage.

**Table 12-3** Physical properties of amalgam

<table>
<thead>
<tr>
<th>Type*</th>
<th>Compressive strength (MPa) (30 min/1 h/1 day)</th>
<th>Tensile strength (MPa)</th>
<th>Knoop hardness</th>
<th>Creep (%)</th>
<th>Dimensional change (μm/cm)</th>
</tr>
</thead>
</table>

Courtesy of S-Y Lee.)
Continual reaction occurs as a function of time. The 24-hour compressive strengths shown in Table 12-3 are adequate for most occlusal loadings. If the bite force is assumed to be 750 N (170 lb) and the contact area $2 \text{ mm}^2$, the compressive stress offered to the amalgam would be on the order of 380 Mpa (55,000 psi). As can be seen from Table 12-3, this is similar to the compressive strengths of most set amalgams. Little additional hardening occurs beyond 24 hours, although additional phase changes are possible.

The amount of residual mercury is very important in the determination of mechanical properties. In general, the compressive strength will decrease 1% with each 1% increase in mercury above 60%. Low mercury-alloy ratios after condensation are therefore desired. In addition to the effects of residual mercury, compressive strength will also decrease 1% with each 1% of porosity. Adequate condensation of amalgam is, therefore, mandatory in achieving maximum strength.

It should also be emphasized that amalgam, in both traditional and high-copper compositions, is a brittle material. Generally, the tensile strength of a brittle material is much less than its corresponding value in compression. For amalgam, the tensile strength values are about one seventh of the compressive strength values (Table 12-3). This means tensile failure is much more likely to occur than compressive failure. Tensile failure is particularly apt to occur in the margins where the amalgam may be unsupported or the mercury concentration is higher due to the condensation process. Obviously the last bit to condense will have the higher mercury concentration, because mercury expression occurs as the amalgam is packed. Because of the higher mercury concentration at the margin, this area may contain greater amounts of the delta 2 phase, contributing to weakness in this region. Tensile failure may also occur at the isthmus of mesio-occlusodistal (MOD) restorations with too little bulk at the step.

Creep and flow are both deformations produced by constant load. The creep of amalgam is important because amalgam at oral temperatures is at $0.9 T_m$, where $T_m$ is the melting temperature. At these temperatures, atomic diffusion occurs easily, and deformation under static load is possible. As seen in Table 12-3, the creep of high-copper amalgam is at least an order of magnitude lower than the upper limit of 3% for traditional amalgams. This lower creep has been associated with the presence of $\text{Cu}_6\text{Sn}_5$ in the gamma 1 network and the decreased amount of available tin (Okabe et al, 1977). The lower creep of high-copper amalgams may also now be related to the absence of the delta 2 phase. The lower creep of high-copper amalgam has been
suggested as a possible reason for its demonstrably better marginal integrity.

The wear of amalgams is approximately the same magnitude as that of tooth enamel. The wear resistance of amalgams exceeds that of most posterior composite restorative materials; therefore, amalgams are much more likely than most composite restorative materials to maintain occlusal contacts.

The physical properties of gallium amalgam are intermediate compared to traditional and high-copper amalgams (Table 12-3).

**Corrosion**

**Introduction**

Traditional amalgams are susceptible to corrosion, with chlorides attacking the gamma 2 phase. This phase has been shown to corrode according to:

\[
\begin{align*}
8\text{Sn}_7\text{Hg} + 2\text{H}_2\text{O} + 2\text{Cl}^- \rightarrow \\
14\text{Sn}_4(\text{OH})_6\text{Cl}_2 + 8\text{Hg}
\end{align*}
\]

This process then leads to two deteriorating effects: (1) The corrosion of interconnected gamma 2 further weakens the amalgam, particularly the tensile strength; and (2) the mercury liberated by the corrosion process can react with the remaining unreacted gamma in the amalgam to produce additional reaction products (gamma 1 + gamma 2). The formation of these new reaction products could produce an additional dimensional change (mercuroscopic expansion), leading to unsupported amalgam at the margin, which can easily fracture in tension. The entire mechanism has been associated with the phenomenon of amalgam ditching (Fig 12-8), which was quite prevalent in the clinical use of traditional amalgam. The liberation of mercury as a corrosion by-product of amalgam has created additional concerns from a biocompatibility point of view.

![Fig 12-8](image-url) Marginal breakdown of traditional amalgam. (Original magnification × 2.5.)
The advent of high-copper amalgams eliminated or reduced the corrosion associated with gamma 2, because the formation of this phase was prevented or retarded and the formation of eta prime \((\text{Cu}_6\text{Sn}_5)\) occurred instead. As will be discussed in the next section, this immediately led to an improvement in the marginal integrity of high-copper amalgams. However, the eta prime phase has also proven to be susceptible to corrosion in the oral cavity, with the following reaction possible:

\[
4\text{Cu}_6\text{Sn}_5 + 19\text{O}_2 + 18\text{H}_2\text{O} + 12\text{Cl}^- \rightarrow 6[\text{CuCl}_2 \cdot 3\text{Cu(OH)}_2] + 2\text{O}_2 + 5\text{SnO}
\]

This reaction will not substantially affect the strength of the high-copper amalgam in the margin because the \(\text{Cu}_6\text{Sn}_5\) is not an interconnected phase. However, corrosion of \(\text{Cu}_6\text{Sn}_5\) has raised questions as to possible biocompatibility of the copper-containing corrosion products of high-copper amalgams.

The addition of < 1% by weight of palladium to a commercial high-copper single-composition amalgam alloy and 5% by weight to an experimental dispersed-phase high-copper amalgam alloy has produced amalgams that appear to have superior corrosion behavior (Greener and Szurgot, 1982). Palladium may be soluble in gamma 1 with a resultant improvement in the corrosion behavior of that phase.

Recent studies have also shown that mercury is released during free corrosion of amalgam in vitro in various artificial salivas. Over the short term, this mercury burden was found to be in the range of 4 to 20 \(\mu\)g/day, or about the same value as the dietary intake; over longer periods, the mercury released from amalgams was considerably lower than dietary intake (Brune, 1986). The concentration of the dissolved mercury found in such in vitro tests may be unrealistically high compared with that found in vivo, because the natural buffering capacity of saliva, along with the attendant organic proteins, may appreciably lower corrosion kinetics. Porosity will have a significant effect on corrosion of both traditional and high-copper amalgams in effecting increases in surface area and surface energies.

**Corrosion of gallium amalgams**

In vitro corrosion studies have shown that gallium amalgams corrode at rates similar to those of traditional amalgams with the attendant release of cations of gallium, copper, tin, indium, and silver. This is understandable in terms of the high concentrations of beta-tin present in the matrix. In addition, substantial amounts of gallium were leached out into neutral and acidic saline solutions (Hero and Jorgensen, 1993).

**Clinical Performance**

**Introduction**

Amalgam is a successful clinical material when careful technique is applied to all
parts of the restorative process, such as cavity preparation, mixing of the alloy and mercury (trituration), packing of the plastic amalgam mix into the preparation (condensation), and finishing. The most significant factor in amalgam behavior under oral conditions is the choice of amalgam alloy. Estimated median clinical lifetimes vary from values as low as 6 years for some traditional products available two decades ago up to median lifetimes exceeding 20 years for the best contemporary high-copper systems.

Unfortunately, the long-term clinical function has been documented for only a few of the available commercial systems for two reasons: (1) The tradition of clinical research is relatively young, and (2) many of the high-copper systems have only recently been put on the market. However, a comprehensive analysis of laboratory and clinical research studies by Marshall and Marshall (1992) supports the long-term performance superiority of high-copper amalgam alloys.

The choice of the amalgam alloy is the most prominent factor in marginal deterioration. Marginal deterioration is measured in vivo by comparison of intraoral photographs with a graded scale of marginal failure (Fig 12-9).

![Fig 12-9 Categorical ranking scale for clinical evaluation of marginal breakdown. (Courtesy of H. Letzel and M.M.A. Vrijhoef.)](image)

When treated with appropriate statistics, a quantitative measurement of marginal deterioration may be obtained. These techniques have also allowed for measurement of the kinetics of marginal breakdown (Fig 12-10).
High-copper amalgams tend to give less marginal fracture, although the best traditional systems available might be as good as, or even better than, the least effective high-copper amalgams. The admixing of up to 10% by weight indium did not adversely affect the marginal integrity or luster of a high-copper amalgam (Johnson et al, 1992). Other factors, such as cavity preparation, application of a cavity varnish, mode of packing, and type of finishing, reportedly may have an influence on marginal fracture in a statistical sense. However, their influence is moderate when compared with the differences due to alloy selection.

Several investigators have shown that the risk of marginal fracture increases with increasing cavity size (Mahler and Marantz, 1980). An investigation of the influence of the cavosurface angle (CSA) on marginal fracture indicated that a CSA of 90 degrees produced slightly less marginal fracture than a CSA of less than 90 degrees (Akerboom, 1985). This observation is consistent with published theoretical considerations (Jorgensen, 1965). Finishing the margins with tungsten carbide burs or chisels did not have any detectable influence; however, amalgam restorations with an applied layer of cavity varnish have been shown to have slightly more marginal fracture than those without it (Borgmeijer, 1985). It has also been shown that from the standpoint of marginal fracture, careful application of cotton rolls in combination with vacuum ejection was equivalent to restorations made using a rubber dam (Letzel et al, 1979). Similarly, different modes of packing or condensing (hand vs. four distinct commercial mechanically and air-driven vibrators) did not reveal any marginal deterioration differences (Letzel and Vrijhoef, 1982).

Commercial amalgamators are available for the mechanical trituration of the alloy and mercury (Fig 12-11). In the case of modern high-copper amalgams, different types of postcarving burnishing produce some differences in marginal fracture compared with the traditional polishing methods (carried out after one day). These differences are smaller than those between the distinct amalgam systems commercially available (Letzel and Vrijhoef, 1984).
Thus, the differences in marginal fracture due to operative variables are small compared with differences between amalgam systems and patients. The dentist may, therefore, seem to have only a small influence on the results of clinical reports found in the literature. However, caution must also be taken in extrapolating clinical literature results to the general practice situation because neither the dentists nor the patients in these published studies may be representative. In this regard, cross-sectional clinical performance data from general practices show a dramatic reduction in lifetimes as compared to controlled clinical trials in academic centers (Letzel et al, 1989). It is thus necessary to realize that the manufacturer makes the amalgam alloy, but the dentist makes the amalgam restoration. It is of ultimate importance to control the dentist factor, as well as the patient factor, in a general practice situation.

**Clinical behavior of gallium amalgams**

Changes in marginal integrity, surface texture, luster, and color were measured clinically over time periods of up to 2 years by several Japanese clinical groups (eg, Sakai et al, 1993). It was found that significant changes in luster and surface roughness occur within time periods as early as 4 months after placement, with occlusal changes being more severe than buccal changes. In light of the significant corrosion behavior associated with the presence of free tin and the selective attack of gallium in the current commercial formulation, this is probably not too surprising.

**Correlations Between Laboratory Properties and Clinical Data**

*Creep*
In 1970, correlation between creep and the susceptibility of both traditional and high-copper amalgam restorations to marginal fracture was reported with a higher creep corresponding to more marginal breakdown (Mahler et al, 1970). This correlation between creep and marginal breakdown was subsequently verified in a large number of published studies. It attracted, and still attracts, much attention among researchers and in product advertisements. Frequently, statements can be found in contemporary literature expressing, either explicitly or implicitly, that a low-creep amalgam would solve the marginal fracture problem. This statement is not necessarily true, however. Several investigators have shown that no correlation between creep and marginal deterioration is present for the gamma 2-free amalgams. However, there are indications of a positive correlation between creep and marginal fracture for the group of traditional amalgams containing gamma 2 (Fig 12-12). Enough clinical and laboratory studies support this correlation to stress that caution is required not to overestimate creep as a selection criterion for the prediction of marginal fracture. General practitioners should not buy their alloy on the basis of creep data alone but rather as part of a complete picture in which clinical evidence from a controlled clinical study plays a substantial role.

Creep will also be affected by porosity. An increase in creep of 30% for a 1% increase in porosity was noted for traditional amalgams.

**Fig 12-12** Correlation of marginal fracture and creep for traditional and high-copper amalgams. (After Vrijhoef and Letzel, 1986.)
Corrosion

A well-known and frequently recognized phenomenon in the early life of an amalgam restoration is the "galvanic pain" due to galvanic action of amalgam restorations in combination with other metallic restorations in interproximal or occlusal contact (Fig 12-13). Clinical studies have found that this type of pain occurred only in a small percentage of cases and was generally not serious; it usually occurred in the first hours and in no instance did it last longer than a few weeks after the insertion of the material. It is also possible that the galvanic currents produced could have a harmful effect on the soft tissues or on the organism as a whole; however, the frequency of the occurrence of these effects is supposed to be low or absent.

![Fig 12-13 Corroding amalgam (a) in interproximal contact with gold casting alloy (b).](image)

Galvanic corrosion can adversely affect both the amalgam and the dissimilar metal in contact with it. For example, if the amalgam is in interproximal contact with a gold restoration the amalgam will corrode as the anode. It will suffer surface attack, lose luster, and become rough. Mercury, when produced as a corrosion product, will amalgamate with the gold alloy producing a color change as well as an embrittling effect.

Galvanic effects with dissimilar metals in occlusion are often greater, because occlusal forces at points of contact are usually great enough to rupture protective films of corrosion products that would form in the absence of contact. Thus, mixed metals in contact, either permanent or intermittent, should be avoided in clinical practice.

The loss of luster and the discoloration of the restoration's surface have been the subject of many clinical studies. They are the first indications as to the deteriorating surface of the amalgam restoration and the bulk of material underneath. A rough surface of the amalgam restoration is prone to the formation of plaque and can cause irritation to adjacent soft tissues. The aggressive attack of the oral cavity on amalgam is primarily due to the chloride ion contained in saliva. In addition to this attack, the
silver in amalgam can tarnish due to action of the sulfide ion. This damage is confined
to the surface layers and can easily be removed by polishing. Further, the sulfide ion
concentration is controlled by such factors as the patient's diet and health.

Amalgam corrosion products coming out of the restoration also might have a positive
effect. If these products precipitate in the interspace between the cavity wall and the
restoration (Fig 12-14), less saliva can penetrate it and marginal leakage through
percolation may be minimized. Sealing of the cavosurface margin takes place within
the first several days. Both traditional and high-copper amalgams display this effect.
The corrosion products at the cavosurface margin are phosphates, whereas the
corrosion products seen within the bulk of the restoration are oxides and
hydroxychlorides.

**Fig 12-14** Scanning electron micrographs of Dispersalloy, high-copper admixed, and
conventional amalgams, that have been retrieved from failed clinical restorations.
Note buildup of tin, calcium, and phosphorous corrosion products at the cavodontinal
**Sensitivity and amalgam bonding**

Clinical experiences indicate that many of the modern high-copper spherical amalgams produce more postoperative sensitivity than other systems. It is speculated that these materials, which universally contract upon setting, produce large gaps at the cavosurface margins that cannot be sealed by the resulting formation of corrosion products. In addition, the dynamic dimensional changes produced by thermal and mechanical stresses to the margin may potentiate this phenomenon. Obviously, an open margin is a preferred pathway for bacterial invasion. For these reasons, the sealing of amalgam cavity walls with a dentin adhesive or glass ionomer has become popular. Traditional dentin adhesives, newer amalgam/dentin adhesives, and hybrid glass ionomers have all shown the ability to reduce leakage and in some cases reduce caries when used under amalgams (Pashley et al, 1991; Manders et al, 1990). Bond strengths of amalgam to dentin with these systems range from a few MPa for traditional dentin adhesives and ionomers up to 10 MPa for the newer systems. A choice of system should not be made solely on the basis of strength values, however, as some of the systems that are lower in strength have been more effective in sealing the margin (CRA, 1994).

**Biocompatibility**

Dental amalgam has been clinically employed for over 150 years. Yet periodically, concern is expressed about its biocompatibility. Additionally, contemporary society has expressed great concern over the contribution of mercury to the total environmental burden of dental amalgam.

A report submitted to the US Public Health Service (USPHS) summarizes the risk/benefit ratio involved in amalgam usage (USPHS, 1993). From a risk standpoint there is no doubt that the presence of amalgams will increase mercury levels in blood, urine, and tissue. The level of increase, however, is currently thought to be below that associated with clinical symptoms. **Table 12-4**, adapted from the USPHS report, summarizes what is currently known about mercury uptake from a wide variety of sources, and includes as benchmarks the World Health Organization (WHO) total mercury uptake and the Occupational Safety and Health Administration (OSHA) workplace limit. Values for mercury intake from amalgam are around one half to one tenth of these levels. Mercury uptake from amalgam is similar to that absorbed from food associated with normal saliva.

<table>
<thead>
<tr>
<th>Source</th>
<th>Type</th>
<th>Absorption (ng/d)</th>
</tr>
</thead>
<tbody>
<tr>
<td>WHO</td>
<td>Total</td>
<td>43,000</td>
</tr>
<tr>
<td>OSHA (air workplace)</td>
<td>Elemental</td>
<td>429,000</td>
</tr>
<tr>
<td>Amalgam</td>
<td>Elemental</td>
<td>1,240-29,000</td>
</tr>
<tr>
<td>Water</td>
<td>Inorganic</td>
<td>5</td>
</tr>
<tr>
<td>Food</td>
<td>Organic + organic</td>
<td>2,220-5,572</td>
</tr>
<tr>
<td>Saliva</td>
<td>Inorganic</td>
<td>180-1,400</td>
</tr>
<tr>
<td>Air (home)</td>
<td>Elemental</td>
<td>4,160</td>
</tr>
<tr>
<td>Air (ambient)</td>
<td>Elemental</td>
<td>32-96</td>
</tr>
</tbody>
</table>

Biological testing procedures have revealed some interaction within cell cultures that has been associated either with unreacted mercury or with copper leached out during corrosion of high-copper amalgam systems. Implantation studies have demonstrated that traditional and high-copper amalgams are well tolerated by connective tissue and bone.

Inflammatory reactions and formation of secondary dentin have been noted, but proper use of lining materials in deep cavities should minimize possible pulpal and dentinal reactions.

Amalgam may also produce gingival reaction due to corrosion or to "tattoos" caused by the accidental subgingival condensation of amalgam. Use of amalgams of superior corrosion resistance is, in any case, indicated to minimize these effects.

Under normal service conditions, amalgam restorations are covered by a film of saliva. This both reduces the vapor pressure dramatically and largely reroutes the mercury from the respiratory to the esophageal tract. The esophageal route's tolerance to mercury is greater than that of the respiratory route.

Elevated sera mercury levels have been identified in populations of dental practitioners after only 20 years of clinical practice. However, these levels were far below those producing clinical toxicity, and the dentists were symptom free.

Recent research has found that the greatest amount of mercury vapor released from amalgam is during the setting. This probably accounts for increased mercury levels in urine after the placement of restorations (Frykholm, 1957). The presence of palladium during the setting greatly reduces the amount of mercury vapor released (Fig 12-15). Palladium has the strongest bond with mercury of any of the transition metals of the periodic table and reduces the mercury vapor pressure of the amalgam.
Properly handled dental amalgam should be regarded as safe for general use as a direct restorative material. This position has been accepted, based on available scientific evidence, by the leading regulatory, professional, and research organizations in the world (USPHS, 1993). An important step in the use of this material is the awareness of clinicians and support staff in the proper handling and disposal of mercury and dental amalgam.

Clinical Decision Scenarios for Dental Amalgams

This section presents one approach for choosing dental cements for specific situations. Each of the following scenarios uses the same format as that presented in chapter 7. Advantages and Disadvantages of each material are prioritized using the following codes: * = of minor importance, ** = important, and *** = very important.

**Situation** Glass ionomer vs amalgam for cores

**Description** A patient comes to the office complaining of a broken tooth. Upon examination the dentist finds that the lingual cusps of the mandibular right first permanent molar have fractured right at the gum line. The tooth already has a large MOD amalgam in it, and the facial cusps are not sturdy. The treatment plan is to prepare a pin-retained core buildup and then a full gold crown. The dentist has both amalgam and a glass-ionomer core material available.

**Critical Factor** Strength
Amalgam

Advantages

*** High strength

* Long clinical history

Disadvantages

* 24-hour set

* No fluoride release

* No bond to tooth

Glass ionomer

Advantages

* Immediate set

* Fluoride release

* Bonds to tooth

Disadvantages

*** Low strength

Analysis/Decision  Despite the more numerous advantages for the glass ionomer, its low strength makes it unsuitable for a full buildup, and the amalgam was selected. Glass ionomers have been advertised for this use, but time has shown that they do not hold up well if they are the entire support for the crown. They are useful for filling in some small depressions in the preparation but not for the complete core.

Situation  Amalgam vs glass ionomer for crown margin repair in patient with low caries risk

Description  A patient with several all-ceramic crowns comes to the office for a check-up. He has had the crowns for about 15 years and has gingival recession resulting in supragingival margins around these crowns. His oral hygiene is good, but a small caries lesion can be seen on the buccal margin of the crown on his mandibular right first molar. A new crown could be made, but the patient does not want to pay for it. The dentist decides to place a small repair restoration on the margin after removal of the caries lesion. She compares the use of either amalgam or glass-ionomer restorative material.
**Critical Factor**  Low cost, durability of restoration, ease of placement

**Amalgam**

*Advantages*

*** Easy to place

*** Excellent durability

*** Low cost

Long history of success

*Disadvantages*

Not tooth colored

* No fluoride release

* Difficult interproximal placement

** No bond to tooth

**Glass ionomer**

*Advantages*

*** Reasonably easy to place

Tooth colored

*** Low cost

* Fluoride release

** Bonds to tooth

*Disadvantages*

*** Less Durrable

** More plaque retentive

*** Isolation is critical

**Analysis/Decision**  The durability and ease of placement of amalgam, combined with its low cost, made it the best choice for this restoration. The access was easy, and the patient was not too concerned with the appearance of the repair. The low caries risk of
the patient was also significant in making the decision.

Situation  Amalgam vs glass ionomer for repair of a proximal margin on a metal crown in patient with high caries risk

Description  The same patient as in the previous scenario returns to the same dentist 6 years later for a dental hygiene appointment. He is now 72 years old and is starting to show the signs of advancing age. He is more frail, forgetful, and is taking medication for high cholesterol, which has a side effect of slight xerostomia (reduced saliva output). The result of his new circumstances is a lowered hygiene level. There is quite a bit of plaque buildup interproximally, and several all-ceramic crown margins show small areas of decay around them. Cost remains a problem, and the question remains how to treat marginal areas of decay on a budget.

Critical Factor  Caries resistance, low cost

Amalgam

Advantages

** Easy to place

** Excellent durability

*** Low cost

** Long history of success

Disadvantages

** Not tooth colored

*** No fluoride release

** Difficult interproximal placement

No bond to tooth

Glass ionomer

Advantages

** Reasonably easy to place

** Tooth colored

*** Low cost
*** Fluoride release

Bonds to tooth

Disadvantages

** Less Durable

** More plaque retentive

Isolation is critical

** Analysis/Decision ** This time the dentist chose the glass-ionomer material because it is capable of preventing recurrent decay due to its reliable and high fluoride release. New crowns are not feasible due to the patient's poor financial situation and insurance that would pay only a small fraction of the total cost. The dentist explained to the patient the need for supplementation with artificial saliva and the possible need to replace the restorations every 3 years because of erosion.

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** Glossary **

** admixed alloy ** An amalgam alloy containing particles of different composition, that is, silver-tin particles and silver-copper particles.

** alloy for dental amalgam ** A silver-tin alloy containing other metals, usually copper and zinc, that will be mixed with mercury to form dental amalgam.

** amalgamation ** Reaction that occurs between mercury and an amalgam alloy.

** beta (β)-tin ** An allotropic form of tin.

** biocompatibility ** The ability of a material to provide successful service in a host while causing minimal response.

** blended alloy ** An alloy containing both spherical and lathe-cut particles.

** burnishing ** Smoothing the surface of a dental amalgam after initial carving by rubbing with a metal instrument having a broad surface.

** condensation ** Packing dental amalgam into a prepared cavity.

** corrosion ** Degradation due to the electrochemical process.

** creep ** Permanent (plastic) deformation under constant load after the material has set.

** delta 2 (-2) phase ** A tin-mercury compound that is a reaction product in dental amalgam.
**dental amalgam** An alloy that results when mercury is combined with a silver alloy and that is initially a plastic mass that hardens after placement in a prepared cavity.

**epsilon (ε) phase** A copper-tin compound (Cu₃Sn) that occurs in particles in traditional alloys.

**eta prime (η') phase** A copper-tin compound (Cu₆Sn₅) that is a reaction product in dental amalgam.

**eutectic composition** An alloy or solution whose components are proportioned so the melting point is the lowest possible for those components. Upon cooling, the single liquid phase is transformed into two or more solid phases, the number of solid phases being equal to the number of components.

**flow** Permanent (plastic) deformation that occurs while the material is setting and in the process of developing its final strength.

**gallium alloy** A product formed by the reaction of an alloy powder (silver-tin-copper) with a gallium-based liquid alloy (gallium-indium-tin).

**galvanic action** Electric potentials in the mouth caused by the contact of dissimilar metals in interproximal or occlusal contact.

**gamma (γ) phase** A silver-tin compound (Ag₃Sn) that forms a substantial part of the amalgam alloy and is also present in the resulting amalgam structure after reaction with mercury has occurred.

**gamma 1 (γ₁) phase** A silver-mercury compound (Ag₂Hg₃) that is a reaction product in dental amalgam.

**gamma 2 (γ₂) phase** A tin-mercury compound (Sn₇Hg) that is a reaction product in dental amalgam.

**high-copper amalgam** Dental amalgam alloy with a relatively high copper content (10% by weight). This alloy is characterized by the corrosion-prone gamma 2 phase being either absent or substantially reduced.

**ingot** Cast rod of alloy.

**lathe-cut alloy** Amalgam alloy made by machining small, irregularly shaped chips from a large cast bar of alloy.

**mercuroscopic expansion** The expansion that occurs when mercury, released by the corrosion of the gamma 2 phase, reacts with the remaining amalgam alloy particles. This will produce an unsupported wedge at the margin of the restoration.

**mercury-alloy ratio** The ratio of the amount of mercury to be mixed with an amount of amalgam alloy.

**quaternary alloy** An alloy containing four elements.
**single-composition alloy** Each alloy particle contains the same components in the same ratios.

**spherical alloy** An alloy whose particles are created by means of an atomization process whereby a spray of tiny drops is allowed to solidify in an inert gaseous or liquid environment.

**ternary alloy** An alloy containing three elements.

**traditional (or conventional) alloys** An alloy with the following composition: 66% to 73% silver by weight, 25% to 29% tin by weight, 2% to 6% copper by weight, 0% to 2% zinc by weight, and 0% to 3% mercury by weight.

**trituration** Mixing dental amalgam alloy with mercury.

**Discussion Questions**

1. Are dental amalgams a health hazard?

2. Why is dental amalgam so popular, even though it is not esthetic?

3. What are the advantages and Disadvantages of using a bonding agent with amalgam restorations?

4. How does copper affect the set microstructure of amalgam?

**Study Questions**

*(See Appendix E for answers.)*

1. How can the working time of amalgam be controlled?

2. Should a zinc-free amalgam be used?

3. Can some amalgams be finished the same day they are placed?

4. Why are spherical amalgams difficult to condense?

5. Does creep predict marginal fracture?

6. Why are seal/bond amalgam restorations of particular interest?

7. How safe are amalgams?

**Recommended Reading**


Chapter 13. Precious Metal Casting Alloys

Introduction

Since the introduction of investment casting to dentistry by Taggart in 1907, precious metal alloys have traditionally been used for several types of restorations. High-gold-content alloys have been used for inlays and still are because soft alloys that can be burnished are desirable. Full-cast crowns and three-quarter crowns are still cast from gold alloys. However, most crowns and fixed partial dentures for the anterior part of the mouth are of the porcelain-fused-to-metal (PFM) type. Alloys for PFM fixed partial dentures have revolutionized this field and include palladium and nickel alloys, as well as high-gold-content alloys. Another change has been the introduction of lower-gold-content alloys to replace the traditional 18-carat American Dental Association (ADA)-certified alloys developed during the 1930s. The tarnish resistance of these lower-gold-content alloys is sufficient for some oral environments but not others.

Composition and Properties

Introduction

Precious metal casting alloys contain mainly gold, palladium, and platinum (which are classified as noble metals), and silver. They also contain limited amounts of nonprecious alloying elements such as copper, indium, iron, tin, and zinc (Table 13-1). The carat scale expresses the relative amount of gold in an alloy, with 24-carat being pure gold. Twelve- and 18-carat alloys contain 50% and 75% gold, respectively. The fineness of a gold alloy is the percentage gold content multiplied by a factor of 10 (eg, 75% is 750 fine). Fineness is used with dental gold solders, but the carat scale is seldom used in dentistry. Copper, silver, palladium, and platinum generally serve as hardening elements in alloys with high gold content. Iron and tin, at much lower concentrations, are hardening additions in PFM alloys. Indium, iron, and tin also serve to promote bonding of porcelain to PFM alloys by formation of stable, adherent oxides.

Table 13-1 Role of alloying elements in dental gold alloys

<table>
<thead>
<tr>
<th>Property</th>
<th>Gold</th>
<th>Platinum</th>
<th>Palladium</th>
<th>Copper</th>
<th>Silver</th>
<th>Zinc</th>
<th>Iridium</th>
</tr>
</thead>
<tbody>
<tr>
<td>Specific gravity</td>
<td>9.32</td>
<td>11.45</td>
<td>12.0</td>
<td>3.96</td>
<td>0.49</td>
<td>1.31</td>
<td>2.4</td>
</tr>
<tr>
<td>Melting point in °C</td>
<td>1,063</td>
<td>1,769</td>
<td>1,552</td>
<td>1,083</td>
<td>0.61</td>
<td>120</td>
<td>1,443</td>
</tr>
<tr>
<td>Face-centered cubic</td>
<td>1,945</td>
<td>3,224</td>
<td>2,829</td>
<td>1,981</td>
<td>1,761</td>
<td>4,429</td>
<td></td>
</tr>
<tr>
<td>Atomic diameter (A)</td>
<td>1.88</td>
<td>2.77</td>
<td>2.74</td>
<td>2.55</td>
<td>1.88</td>
<td>2.66</td>
<td>3.32</td>
</tr>
<tr>
<td>Close-packed hexagonal cubic</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 13-1 Role of alloying elements in dental gold alloys*
<table>
<thead>
<tr>
<th>Chemical activity</th>
<th>inert</th>
<th>inert</th>
<th>Mild</th>
<th>Very active</th>
<th>Active</th>
<th>Very active</th>
<th>Active</th>
</tr>
</thead>
<tbody>
<tr>
<td>Color</td>
<td>yellow</td>
<td>White</td>
<td>White</td>
<td>Red</td>
<td>White</td>
<td>White</td>
<td>White</td>
</tr>
<tr>
<td>Approximate content</td>
<td>0%-95%</td>
<td>0%-20%</td>
<td>0%-12%</td>
<td>0%-17%</td>
<td>0%-20%</td>
<td>0%-2%</td>
<td>.005%- .1%</td>
</tr>
<tr>
<td>Density (specific gravity)</td>
<td>increases markedly</td>
<td>increases markedly</td>
<td>lowers slightly</td>
<td>lowers</td>
<td>lowers</td>
<td>lowers</td>
<td>increases slightly</td>
</tr>
<tr>
<td>Effect of color on alloy</td>
<td>lends yellow color</td>
<td>whitens slowly; 12% required; not pure white</td>
<td>whitens rapidly; as little as 5%</td>
<td>lends red color; dark plate high in Cu</td>
<td>whitens very slowly, counteracts redness of Cu; creates green gold</td>
<td>percentage too low to have effect</td>
<td></td>
</tr>
<tr>
<td>Melting</td>
<td>raises melting point mildly</td>
<td>raises melting point fairly rapidly</td>
<td>raises melting point rapidly</td>
<td>lowers melting point even below its own</td>
<td>slight effect; may raise or sometimes lower mildly</td>
<td>lowers no melting point rapidly; in most solders</td>
<td></td>
</tr>
<tr>
<td>Tarnish resistance</td>
<td>essential to good tarnish resistance</td>
<td>contribute importantly to tarnish resistance</td>
<td>increases tarnish resistance but less than Au and Pt</td>
<td>contribute to tarnish in flame or with sulfurous food</td>
<td>tarnishes in presence of sulfur</td>
<td>will tarnish, but in low percentage has little effect</td>
<td></td>
</tr>
<tr>
<td>Heat hardening</td>
<td>contribute importantly with Cu</td>
<td>increases with Cu</td>
<td>some increase with Cu</td>
<td>essential if alloy heat hardens</td>
<td>increases with Cu</td>
<td>slight with Cu no effect</td>
<td></td>
</tr>
<tr>
<td>Gas absorption</td>
<td>—</td>
<td>—</td>
<td>rather high for hydrogen</td>
<td>—</td>
<td>rather high for oxygen</td>
<td>A good deoxidizer no effect</td>
<td></td>
</tr>
<tr>
<td>Castability</td>
<td>—</td>
<td>—</td>
<td>Effects not critical</td>
<td>Effects not critical</td>
<td>decreases surface tension and increases fluidity</td>
<td>No effect</td>
<td></td>
</tr>
</tbody>
</table>

*Adapted from Brumfield, 1955.*
High-gold alloys

Traditional dental casting alloys contain 70% by weight or more of gold, palladium, and platinum. American National Standards Institute/American Dental Association Specification no. 5 for Dental Casting Gold Alloy divides these alloys into four types based on mechanical properties:

Type I—Soft (VHN* 60 to 90)

Type II—Medium (VHN 90 to 120)

Type III—Hard (VHN 120 to 150)

Type IV—Extra hard (Quenched VHN minimum 150; Hardened VHN minimum 220)

Table 13-2 identifies currently available alloys and their mechanical properties.

<table>
<thead>
<tr>
<th>Type</th>
<th>Approximate composition (%)</th>
<th>Representative product</th>
<th>Vickers hardness number</th>
<th>Yield strength psi (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>I. Soft: one- and two-surface inlays</td>
<td>32 2 12</td>
<td>Degussa-Oey-Williams</td>
<td>85</td>
<td>20,000</td>
</tr>
<tr>
<td>I. Medium: inlays, MOD, crowns</td>
<td>77 2 12</td>
<td>3-Inlay Medium</td>
<td>120 145 (225)</td>
<td>33,000 (290)</td>
</tr>
<tr>
<td>II. Hard: inlays, crowns, fixed partial dentures</td>
<td>73 2 12</td>
<td>Degullor-C-Oey-Williams</td>
<td>130 200</td>
<td>37,000 (255)</td>
</tr>
<tr>
<td></td>
<td>50 4 25</td>
<td>Special-Oey-Hard</td>
<td>140 240</td>
<td>56,000 (385)</td>
</tr>
</tbody>
</table>

*VHN = Vickers hardness number.
Type I alloys are weak, soft, and highly ductile. They are useful only in areas not subject to occlusal stress and are not widely used. They do not harden by heat treatment. Type IV alloys are relatively strong, hard, and nonductile. They are intended for high-stress applications such as partial dentures. They also are not widely used at present.

The intermediate Type II and Type III alloys are used for most restorations. Type II alloys are used for inlays in which burnishability of margins is more important than high strength. Type III alloys are used in higher-stress applications for inlays, onlays, and three-quarter crowns, and for fixed partial denture retainers and pontics where the restoration design makes burnishability less important than strength.

Typical composition ranges for high-gold alloys are shown in Table 13-2. Iridium in small amounts, around 0.1%, is added as a grain refiner by several manufacturers. Type III and Type IV alloys may contain a high percentage of both palladium and platinum as hardening elements, as shown in the table, while retaining a light gold color. Type III and Type IV alloys are susceptible to heat treatment and may be
hardened or softened by appropriate heating cycles (Table 13-2 for hardness values).

Alloys containing more than 6% palladium are normally white (silver colored). Typical composition ranges of these alloys are given in Table 13-2 also. They are hard, strong, and heat treatable and have mechanical properties characteristic of Type III and Type IV alloys.

*Vickers hardness number. The Vickers hardness test, or the 136-degree diamond pyramid hardness test, is a microindentation method. The indenter produces a square indentation, the diagonals of which are measured. The diamond pyramid hardness is calculated by dividing the applied load by the surface area of the indentation.

**Low-gold alloys**

Low-gold alloys are composed mainly of gold, silver, and copper, with a small percentage of palladium. Gold content ranges from 45% to 60%. The main incentive for the use of these alloys is financial; when the price of gold increases, the use of low-gold alloys increases relative to that of high-gold alloys.

Few alloys are marketed with gold contents between 55% and 70% because the cost savings are not high enough. Few alloys contain less than 45% gold because of tarnish and corrosion problems with use. There is some tendency for alloys in this group to contain additional palladium to partially make up for low gold content (Table 13-2).

The mechanical properties of the low-gold alloys generally correspond to the properties of ADA Type III alloys. Thus, the alloys are strong and hard and have only moderate ductility.

Low-gold alloys are rarely used for inlays but can be quite suitable for full-cast crowns.

**Palladium-silver alloys**

Alloys that have palladium as their main ingredient have several applications. Palladium-silver alloys provide mechanical properties similar to those of Type III gold alloys. An increase in silver content leads to increased ductility and lowered hardness but increases corrosion problems. These alloys are more commonly used for crowns than for inlays. Unless other alloying elements are added, these materials are not heat treatable, because they form only a continuous solid solution at all compositions (Fig 13-1).
High-palladium alloys contain only a small percentage of other precious metals as alloying elements. They have been used as PFM alloys. Such PFM alloys have caused problems with porcelain discoloration due to the formation of silver oxide and volatilization of the silver.

Porcelain-fused-to-metal alloys

Alloys intended for use as bases for porcelain have special requirements because of the need to develop and maintain strength at the temperature involved in porcelain applications and to provide a firm bond to the applied porcelain. In addition, the designs of many PFM restorations emphasize the need to be able to cast thin sections and the need for high yield strength. Three groups of precious alloys are used for PFM application (Table 13-2):

Type I—Alloys containing over 90% gold, platinum, and palladium with small amounts of iron, indium, and tin as hardening and bonding agents

Type II—Alloys containing approximately 80% gold, platinum, and palladium, with
trace additions of iron, indium, tin, and silver making up the balance

Type III—Palladium-silver alloys

Porcelain-fused-to-metal alloys are the most widely used alloys in commercial laboratories. They constitute approximately 70% of all cast units currently in use. Precious PFM alloys are now in direct competition with nonprecious PFM alloys.

Properties of importance for PFM alloys include the ability to produce good casting precision in their cross sections, high yield strength, controlled thermal expansion, and suitable surface characteristics for bonding to dental porcelain. Burnishability is a secondary consideration.

Normally, bonding involves the use of a precious metal alloy containing small quantities of iron, indium, and tin. Controlled oxidation of the castings during a "degassing heat treatment" produces an oxide coating on the alloy surface to which the porcelain adheres.

The properties of PFM alloys are improved by heat treatment. A precipitation reaction during the porcelain firing procedure strengthens and hardens the alloys. Iron-platinum and gold-tin phases are common precipitates.

The high-precious-metal alloys are still widely used. Nickel-chromium and other base metal alloys also used for PFM applications are, in general, less desirable because of problems with casting accuracy, fit, finishing, and porcelain bonding.

Alloys for PFM restorations are discussed in more detail in chapter 14.

**Heat Treatment**

Except for Type I and Type II high-gold alloys, precious metal casting alloys respond to heat treatment by changes in properties and microstructure.

**Homogenizing** (softening) heat treatment consists of heating the alloy to a temperature approximately 75°C below the solidus temperature, holding at that temperature for 10 to 30 minutes, and then quenching to room temperature. Hardening can be produced by one of two methods: slow cooling or a constant-temperature heat treatment. In either case, the important factor is the time spent in a critical temperature range between the softening range and room temperature. This critical temperature varies among alloys; it generally lies about halfway between room temperature and the alloy's softening temperature. Typical cycles for a Type III high-gold alloy are as follows: the softening treatment consists of holding the alloy at 700°C to 750°C for 10 minutes, followed by quenching to room temperature. The hardening treatment consists of 10 minutes at 350°C to 400°C, followed by quenching or low cooling to room temperature.

High-fusing alloys may require more time as well as higher temperatures for both softening and hardening heat treatments.

**Basic Crystal Structure**
All precious metal casting alloys are based on metallic elements having face-centered cubic crystal structures. With the appropriate homogenizing heat treatment, most of them can be converted to a single phase. Hardening heat treatment results in precipitation of phases with other crystal structures, and hardening alloys may contain several different phases. In many alloys, ordering can occur. This is a rearrangement of atoms within the unit cells of the crystal structure. It can cause a characteristic microstructure to appear. This mechanism is important in the hardening of binary gold-copper alloys but is of secondary importance in dental alloys (Figs 13-2 and 13-3).

Fig 13-2 The lattice arrangement in an ordered gold-copper alloy with a regular alternation of unlike atoms in layers.
Fig 13-3 The gold-copper phase diagram. The ordered phases $\text{AuCu}_3$ and $\text{AuCu}$ may be formed by heat treatment. The formation of these phases results in hardening. Quenching from 700°C avoids their formation and hardening.

**Cast Microstructure**

Both the cooling and nucleation rates are quite high for most dental castings. Thus, the typical cast structure consists of fine uniform grains. Large-grain structures are found more often with PFM white alloys unless a grain refiner (eg, iridium) is used (Figs 13-4 and 13-5). Normally, single-phase structures are found in alloys that have received a thorough softening (homogenizing) heat treatment. Hardening produces the appearance of a discontinuous grain boundary precipitate. However, this structure is unrelated to hardening. A submicroscopic continuous precipitate occurring throughout the grains is the primary cause of hardening. This is the result of the separation of silver-rich and copper-rich gold alloy phases. Alloys that undergo ordering during hardening exhibit changes in microstructure.
**Fig 13-4** Dendritic (skeleton- or needle-shaped) grain structure of precious metal alloy as cast. (Original magnification × 100.)

**Fig 13-5** Normal grain structure of cast gold alloy after softening heat treatment. (Original magnification × 300.)

**Glossary**

**burnishability** Ability of a metal to undergo surface deformation (stretching) with a special tool (burnishing instrument).

**carat** A measure of the gold content of an alloy, with 24 carat being 100% gold.

**corrosion** Attack on a metal surface involving loss of material (eg, rusting of iron).

**fineness** The percentage of gold content in an alloy multiplied by a factor of 10 (eg, 75% gold is 750 fine).
grain A single crystal of metal as seen in the microstructure.

grain refiner An ingredient of an alloy (eg, gold alloys) that helps form smaller grains when the alloy is cast.

homogenizing heat treatment A process of heating an alloy to produce a more uniform distribution of elements by diffusion, usually resulting in softening of the alloy.

noble metal A metal that is resistant to oxidation; includes gold, platinum, palladium, and the other platinum group metals.

ordering The regular arrangement of atoms of an element in a lattice structure rather than a random distribution.

precious metal A metal that is relatively high in cost; includes gold, platinum, palladium, and silver.

precipitation The separation of a phase from a solution upon cooling, due to reduced solubility.

tarnish The formation of objectionable reaction products on the surface of an alloy (eg, black oxides or sulfides on silver).

white gold alloy An alloy containing gold and white metals (eg, silver, palladium) that impart a white appearance to the entire mass.

Discussion Questions

1. What is the relationship between carat and platinum metal content and the tarnish of gold restorations?

2. How have the high price of gold and the patients' desire for esthetic restorations affected the use of precious metal restorations?

3. What major effect does iridium have on the microstructure of gold alloys?

4. How would the temperature of quenching a gold casting mold affect its mechanical properties?

Study Questions

(See Appendix E for answers.)

1. List the elements classified as noble.

2. Which elements contribute to the hardening of dental gold alloys?

3. Give the main applications of the ADA Type I, II, III, and IV alloys.

4. Which three elements are added in fractional amounts to harden high-gold-content
alloys to be used with porcelain?

5. What effect does palladium content have on the color of gold alloys?

6. Describe a softening heat treatment for the ADA Type III and IV alloys.

7. Describe a hardening heat treatment for the ADA Type III and IV alloys.

8. Which element is used as a grain refiner in gold casting alloys? Why is it added?

9. Describe the purpose of the heat treatment given to alloys prior to porcelain application.

10. Which five elements are usually present in the white gold alloys used with porcelain (PFM)?

11. What is ordering in the gold-copper system?

12. What process is currently believed to be responsible for the hardening of gold-copper-silver alloys by heat treatment?

13. What is the main risk involved in using low-gold-content (less than 45%) fixed partial denture alloys?

14. Which mechanical property is usually considered a measure of the burnishability of a soft inlay alloy?

15. What has been the main problem with high-silver-content alloys for use with porcelain?

**Recommended Reading**


Wise EM, Eash JT (1933). The role of the platinum metals in dental alloys. III. The influence of platinum and palladium and heat treatment upon the microstructure and constitution of basic alloys. Tr AIME Inst Metals Div 104:276.
Chapter 14. Alloys for Porcelain-Fused-to-Metal Restorations

Introduction

In response to the fluctuating prices of gold and other precious metals, many alternative alloys have been introduced into the dental profession. Although the development of these alternative alloy systems was largely motivated by economics, the resultant properties of the alternatives often make them superior choices even when compared with more costly alternatives. Selection of the optimal alloy to use for crowns and fixed partial dentures should be based on a rational appraisal of the properties relevant to the intended use of the alloy. A classification of current alloy systems is given in Fig 14-1, and commercial alloys are listed in Table 14-1.

Fig 14-1 A classification of alloys for PFM restorations.

The proliferation of alloy systems has complicated the dentist's choice of products for specific restorative situations. As a result, many practitioners rely solely on the advice of dental laboratories for their selections. Many laboratories base their choices on cost factors rather than on a material's properties. Additionally, many less significant criteria for alloy selection are used by dentists, for example: alloy color, gold cement, "precious only," high cost, or "looks, feels, or is cast like gold."

Table 14-1 Examples of commercial PFM alloys

<table>
<thead>
<tr>
<th>Alloys</th>
<th>Representative products</th>
<th>Manufacturer</th>
</tr>
</thead>
<tbody>
<tr>
<td>high gold</td>
<td>Jelenko O</td>
<td>Jelenko</td>
</tr>
<tr>
<td></td>
<td>Jelenko O, Biostar, Y-Ceramic</td>
<td>Generic/Pentron</td>
</tr>
<tr>
<td></td>
<td>Will-Ceram Y, Golden Ceramic</td>
<td>Williams</td>
</tr>
<tr>
<td>gold-palladium (no silver)</td>
<td>Olympia</td>
<td>Jelenko</td>
</tr>
<tr>
<td></td>
<td>SFC, Everest</td>
<td>Generic/Pentron</td>
</tr>
<tr>
<td></td>
<td>Eclipse, Encore</td>
<td>Ney</td>
</tr>
<tr>
<td>gold-palladium-silver</td>
<td>Cameo, Collegiate</td>
<td>Jelenko</td>
</tr>
</tbody>
</table>
Rational selection of a casting alloy for porcelain-fused-to-metal (PFM) restorations should be based on the following:

1. Physical properties
2. Chemical properties
3. Biocompatibility
4. Laboratory workability
5. Porcelain compatibility

Lastly, in a balanced decision, cost should be considered relative to these criteria. Factors such as single-unit or multiple-unit, presence or absence of metal occlusal surfaces, span length, and porcelain brand often indicate different alloy choices. A practitioner using only one alloy is unlikely to make an optimal choice in every clinical situation.

Classifications

Before discussing alloy characteristics, it is worth reviewing the terms noble, precious, semiprecious, and nonprecious. Noble metals are defined on the basis of their chemical properties; that is, they resist oxidation and are not attacked by acids. Seven metals meet this definition, but only three are widely used in dental alloys: gold, palladium, and platinum. These metals give noble metal alloys their inert properties in the mouth.

The term precious refers only to cost, which is controlled by supply and demand.
Many elements in the periodic table, including the seven noble metals, are precious by today's standards. In dental advertisements *precious* usually refers to silver, a metal that is precious but not noble; silver is usually a major ingredient in most alloys advertised as precious.

The term *semiprecious* was originally coined for noble metal alloys that contained significant amounts of silver, and it subsequently has been applied to a variety of alloys, some of which are mixtures of precious and nonprecious ingredients. It is advisable to drop the term *semiprecious* from the dental vocabulary, as it is not well defined and leads to much confusion.

*Nonprecious* alloys are composed of nonprecious ingredients, except for the common inclusion of 1% to 3% beryllium, a precious but ignoble metal. Most nonprecious alloys are based on a combination of nickel and chromium, although cobalt-chromium and iron-based alloys also exist. See Table 14-2 for the compositions of alloys commonly used in PFM restorations.

| Table 14-2 Composition (%) of alloys for PFM restorations* |
|-----------------|-----|-----|-----|-----|-----|-----|-----|---------------------|
| **Alloys**       | Au  | Pt  | Pd  | Ag  | Sn  | In  | Ga  | Other metals        |
| High gold        | 74-88| 0-20| 0-16| 0-15| 0-3 | 0-4 | ___ | Zn < 2; Fe < 0.5; Ta < 1 |
| Gold-palladium   | 45-68| 0-1 | 22-45|___  | 0-5 | 2-10| 0-3 | Zn < 4              |
| (no silver)      |     |     |      |     | 25-40| 5-16| 0-6 | 0-2 Zn 0-3          |
| Gold-palladium-silver | 42-62| ___ | 25-40| 5-16| 0-8 | 0-8 | 3-9 | Cu 4-20; Zn 0-4     |
| Palladium-copper | 0-2 | 0-1 | 66-81|___  | 0-8 | 0-8 | 0-8 | Zn 0-4; Mn 0-4       |
| Palladium-silver | 0-2 | 0-1 | 50-75| 1-40| 0-9 | 0-8 | 0-6 |                     |
| Nickel-chromium  |     |     |      |     |     |     |     | Ni 59-74; Cr 10-22   |
| Nickel-          |     |     |      |     |     |     |     | Ni 70-80; Cr 12-15; Be 0.6-2 |
| chromium-        |     |     |      |     |     |     |     |                     |
| beryllium        |     |     |      |     |     |     |     |                     |
| Cobalt-chromium  |     |     |      |     |     |     |     | Co 54-65; Cr 24-32   |
| Titanium         |     |     |      |     |     |     |     | CP Grades 2 and 4    |
| Titanium alloys  |     |     |      |     |     |     |     | Ti-6Al-4V; Ti-Nb-Al  |

*Courtesy of A. Prasad.

When an alloy is chosen for a particular clinical situation, a number of characteristics have clinical significance and should be considered. Among the most important of these characteristics are physical and chemical properties, casting accuracy, and porcelain-metal compatibility.

**Physical and Chemical Properties**
Color is one of the most obvious physical properties of an alloy. Although the color has no biologic significance, it is equated with quality in the minds of many dentists. Sometimes this color factor seems to matter more to the dentist than to the patient.

When the gold content of an alloy is decreased and metals such as silver and palladium are substituted, yellow color is lost. These less yellow dental alloys are not yet widely accepted. In fact, the profession's desire for gold color is so strong that gold-colored semiprecious and nonprecious alloys are commercially available, even though their other physical and chemical properties fall far short of those of even the cheapest white alloys. In some countries, yellow alloys of copper and nickel are currently quite popular.

If an alloy is gold colored, it must contain copper, gold, or both. However, an alloy can contain substantial amounts of gold or copper without appearing yellow. Good examples of this apparent contradiction are jewelers' white gold and some popular gold alloys for PFM restorations (such as Degudent U, Degussa; SMG-3, Ney). The latter products contain more than 80% gold, yet no yellow color is seen because of the strong whitening effects of palladium and platinum.

Color can be a misleading indicator of composition; dentists should consider other physical and chemical properties as more important than color when a casting alloy is selected.

Some important physical and chemical properties to consider when choosing a cast alloy are:

1. **Noble metal content:** the weight (or better, the atomic) percentage of the seven noble metals contained in an alloy

2. **Hardness:** the Vickers hardness number (VHN), a measure of resistance to indentation

3. **Yield strength:** a measure of the stress required to cause permanent deformation under tension

4. **Elongation:** the amount of permanent deformation a metal undergoes when loaded to its fracture point

5. **Fusion temperature:** the approximate temperature at which an alloy separates under its own weight from partial melting

All of these characteristics have clinical significance. The noble metal content determines, to a large extent, the corrosion resistance and inert properties of the alloy. Hardness is important in relation to occlusal wear resistance and finishing and affects polishing properties. Yield strength is necessary in determining load-bearing ability, especially in fixed partial dentures. Elongation relates to margin-finishing properties, especially important in partial veneer crowns and abutments. It is important to remember that the elongation value for an alloy may be clinically irrelevant if the yield strength is high. To use the potential elongation, stresses exceeding the yield
Strength must be applied to move the metal. Within each group of alloys, yield strength generally increases with increasing hardness. Fusion temperature is important in relation to solder melting ranges and correlates with sag resistance.

**Porcelain-Metal Compatibility**

Thermal expansion, bond strength, and composition are also important properties to consider when choosing among alloys for PFM restorations. These characteristics determine porcelain-metal compatibility.

Thermal expansion is important because a state of zero residual stress is desirable for porcelain in the final restoration. Such a state is achieved when the total expansions and contractions of the porcelain and metal are matched between the porcelain firing temperature and room temperature.

Porcelain-to-metal bond strength ensures retention of porcelain both in the oral environment and during thermal processing, when the induced thermal stresses can be quite high.

Composition is a key factor in porcelain-metal compatibility because some components of an alloy can affect the color of the porcelain, perhaps compromising the esthetics of a restoration. Among the alternative alloys, those containing silver are often associated with porcelain color changes and can cause "greening" of some brands of porcelain. The mechanism for this porcelain discoloration is an exchange between silver from the alloy and sodium from the porcelain. The exchange process requires an oxidizing atmosphere, but a subsequent reducing atmosphere is required to produce the colloidal precipitate responsible for color changes in the porcelain.

**Other Properties**

Because the cross-sectional area of metal used in PFM restorations is usually smaller than that of all-metal restorations, physical properties such as yield strength of the alloy are crucial in design. Stress in turn controls the minimum allowable dimensions of critical areas like connectors. The elastic modulus is equally important because it determines the flexibility of the metal framework. Flexibility is inversely proportional to elastic modulus; an alloy with a high elastic modulus will flex less under load than an alloy of low elastic modulus.

Chemical properties are important because they affect tarnish resistance, corrosion resistance, and thermal stability. Thermal properties are critical in alloys for PFM restorations because the alloy must have a sufficiently elevated melting temperature range to provide dimensional stability during the porcelain firing cycle. Thermal creep results in distortions such as sag in fixed partial denture frameworks and margin opening during the porcelain firing cycles.

Casting accuracy must, of course, be sufficient to provide clinically acceptable castings. In addition to dimensional accuracy (a strong function of technique), the mold-filling ability also contributes to casting accuracy.

Biocompatibility includes a number of factors, among them cytotoxicity and tissue
irritation. Potential biologic hazards from the base metal alloys, particularly nickel and beryllium, are controversial. These potential hazards may affect not only the patient but also the dentist or technician who makes the restoration. The lack of data and long-term clinical experience suggests caution in using base metal alloys, particularly for people with known sensitivity to base metals. To date, however, neither experimental data nor clinical experience unequivocally contraindicate the use of alloys containing these potentially toxic elements, even in patients known to be sensitive to them.

The following discussion of each alloy group is intended to be general and not necessarily specific to the proprietary products. The product examples were chosen based on their status as the historical forerunners of each alloy group. Table 14-3 lists properties of alloys used in dentistry for crowns and copings.

<table>
<thead>
<tr>
<th>Group</th>
<th>Vickers hardness number</th>
<th>Elastic modulus psi × 10^6 (GPa)</th>
<th>Yield strength psi (MPa)</th>
<th>Specific gravity</th>
</tr>
</thead>
<tbody>
<tr>
<td>High gold</td>
<td>182</td>
<td>13 (90)</td>
<td>65,000 (448)</td>
<td>18.3</td>
</tr>
<tr>
<td>Gold-palladium (no silver)</td>
<td>220</td>
<td>18 (124)</td>
<td>83,000 (572)</td>
<td>13.5</td>
</tr>
<tr>
<td>Gold-palladium-silver</td>
<td>218</td>
<td>16 (110)</td>
<td>63,600 (439)</td>
<td>13.8</td>
</tr>
<tr>
<td>Palladium-copper</td>
<td>425</td>
<td>14 (96)</td>
<td>166,000 (1,145)</td>
<td>10.6</td>
</tr>
<tr>
<td>Palladium-silver</td>
<td>242</td>
<td>20 (138)</td>
<td>77,000 (531)</td>
<td>11.1</td>
</tr>
<tr>
<td>Nickel-chromium</td>
<td>257</td>
<td>29 (207)</td>
<td>58,000 (400)</td>
<td>8.7</td>
</tr>
<tr>
<td>Nickel-chromium-beryllium</td>
<td>357</td>
<td>31 (213)</td>
<td>116,000 (800)</td>
<td>7.8</td>
</tr>
</tbody>
</table>

High-Gold Alloys

Porcelain-fused-to-metal technology was introduced to the dental profession with the introduction of Ceramco No. 1 alloy in 1958. The alloy was a forerunner of the improved high-gold alloys that remain on the market today, such as Jelenko O (Jelenko).

The high-gold alloys are composed principally of gold and platinum group metals. Gold content in these alloys varies from 78% to 87% by weight, and total noble metal content is about 97%. Small amounts of tin, indium, and iron are added for strength and to promote a good porcelain bond to metal oxide. Because of their high nobility, these alloys tend to be costly, both in terms of their cost per ounce and their high density, resulting in heavy castings.

High-gold alloys are usually light yellow in color, although some are white. Some are very yellow, apparently in response to the gold mystique previously discussed. The properties of the very yellow alloys are usually inferior to other products in the group, and their low tensile strength in particular makes them a questionable choice for fixed
partial dentures.

The hardness of alloys in this group is considered ideal for working characteristics and ease of finishing, and the tensile strength for all but the very yellow products is good. Corrosion resistance is excellent because of high nobility. Porcelain discoloration is not a problem because the alloys contain little or no silver.

Besides cost, the principle Disadvantages of the high-gold alloys are low elastic modulus and poor sag resistance during the porcelain firing cycle. These factors are also troublesome for fixed partial dentures and suggest the use of alternative alloys for these situations.

**Gold-Palladium-Silver Alloys**

Gold-palladium-silver alloys were the first alternative systems, introduced in 1970 as Will-Ceram W (Williams), and they remain on today's market. The addition of substantial amounts of silver (10% to 15%) and a relatively high palladium content (20% to 30%) may reduce the cost of these alloys as compared with the higher-gold-content group. Elastic modulus is higher, and the alloys are less susceptible than the high-gold group to dimensional changes during the porcelain baking cycle. Corrosion resistance and clinical working characteristics are generally good.

The principle disadvantage of these alloys is their tendency to induce color changes in porcelain because of their silver content. Silver transport into the porcelain results in a yellow-green color change, depending on the brand of porcelain.

The gold-palladium-silver group has been largely superseded by silver-free gold-palladium alloys, which eliminate problems with porcelain color change. Although gold-palladium-silver alloys are successfully used by many practitioners and have had excellent commercial success, they are used less since the introduction of the cost-competitive silver-free alloys.

**Palladium-Silver Alloys**

The first palladium-silver alloy was introduced to the dental profession in the 1970s, but the one that has remained on the market the longest is Will-Ceram W-1 (Williams), introduced in 1975 and at one time the largest-selling alloy in the United States.

Palladium-silver alloys usually include 50% to 60% palladium, with most of the balance being silver. The physical and chemical properties are favorable for PFM restorations and are comparable to other noble metal alloys. The 50% to 60% nobility assures a satisfactory degree of tarnish and corrosion resistance and good clinical working characteristics.

The elastic modulus for this group is the most favorable of all the precious metal alloys and results in the least flexible castings. Only nonprecious alloys have superior elastic moduli. Palladium-silver alloys solder well and have the lowest sag tendency of the precious metal alloys. Porcelain bond strength is also excellent.
The principal disadvantage of this group is a porcelain color change to green—which occurs to a greater degree in this group than in alloys with lower silver content, such as the gold-palladium-silver alloys. Color problems vary considerably, depending on the brand of porcelain; with some brands this disadvantage is eliminated. Will-Ceram and Ivoclar (Ivoclar) porcelains are more resistant to silver discoloration than others.

Some manufacturers recommend the use of metal surface coupling agents to reduce porcelain color problems. Some of these coupling agents are modified porcelains and others are 24-carat gold. The colloidal gold agents are reasonably effective in reducing surface activity of silver in the alloy, thus preventing diffusion into the porcelain. However, if these gold coupling agents are used in excessive amounts, the gold interferes with surface oxidation necessary for a porcelain-metal bond. Selecting a brand of porcelain that does not change color is a more reliable solution to the problem than using coupling agents.

The palladium-silver group can be a good alternative to the gold-containing group. If the porcelain is one that shows minimal (or no) color change in the presence of silver, it is difficult to find fault with these alloys. Mechanical properties are often superior to even the most costly noble metal alloys.

Gold-Palladium Alloys

The gold-palladium silver-free alloys were developed in the mid-1970s to alleviate the color problems caused by silver. The first silver-free alloy was introduced in 1975 as Olympia (Jelenko). These alloys generally contain about 50% gold and 40% palladium. They have had considerable commercial success. Yield strength and hardness are favorable, and elastic modulus is increased significantly compared with high-gold alloys. Cost is comparable to that of the gold-palladium-silver group.

The only recognized disadvantage of the gold-palladium group is thermal expansion incompatibility with some of the higher-expansion porcelains. The silver-free alloys tend to have lower expansion values than the silver-containing group. Some incompatible combinations are well known and are readily acknowledged by the respective manufacturers.

In the absence of thermal expansion incompatibility, there are no Disadvantages and several recognizable advantages to using alloys from this group in preference to the high-gold and gold-palladium-silver alloys. Rigidity is improved for partial dentures, and the porcelain-metal bonds are adequate. Corrosion resistance is excellent because of high nobility. Sag tendencies are about the same as for gold-palladium-silver alloys and, again, much better than for high-gold alloys. Gold-palladium alloys can be an excellent choice when their relatively high cost is not a major consideration.

Recently, small amounts of silver have been added to otherwise silver-free compositions. The resulting alloys are probably superior to the silver-free compositions. Because the silver content is low (usually less than 5%, compared with 10% to 15% in the gold-palladium-silver group previously discussed), no porcelain color problems are evident. However, marketing appeal may be lacking because of the impression created by advertising "silver-free, trouble-free" alloys. Like all new alloys, the best way to test the efficacy of these products is through clinical
experience. The new alloys seem promising because thermal expansion is increased and castability is better than with the silver-free alloys. The increase in expansion tends to eliminate the incompatible porcelain-metal combinations previously mentioned.

**Palladium-Copper Alloys**

The palladium-copper alloys are a relatively recent development, first introduced to the dental profession in 1982 as Option (Ney).

Palladium-copper alloys are usually composed of 70% to 80% palladium and contain little or no gold, up to 15% by weight of copper, and around 9% of gallium. Copper was an unusual addition to porcelain-bonding alloys; such large amounts of copper would cause problems with bonding and porcelain color in gold-based alloys, but apparently do not cause these problems in alloys rich in palladium. Because the alloys have no silver, they cause none of the porcelain color problems associated with silver. Some palladium-copper alloys have a rather heavy oxide that is difficult to cover with opaque porcelain. High hardness values in some of the alloys are offset by a relatively low elastic modulus, resulting in better working characteristics than would be expected with a high hardness value. Strength is good, and some alloys exhibit extremely high yield strengths.

Palladium-copper alloys generally do not melt or cast as easily as palladium-silver alloys, but they are quite acceptable in this regard. Presoldering has been associated with problems for some, but not all, of these alloys. Additionally, the sag resistance of most of them is not as high as in the palladium-silver alloys, again tending to contraindicate their use in large-span fixed partial dentures.

**Palladium-Cobalt Alloys**

Palladium-cobalt alloys, with around 88% palladium and 4% to 5% cobalt by weight, have been in limited use. The main advantages of these alloys is a higher coefficient of thermal expansion that is useful with certain porcelains. However, the main disadvantage is the formation of a dark oxide that may be difficult to mask at thin margins. Also, these alloys may be more susceptible to hot tearing and embrittlement from carbon if no silver is present. Commercial palladium-cobalt alloys on the market are Jelenko PTM (Jelenko) and Jeneric/Pentron APF (Jeneric/Pentron).

**Base Metal Alloys**

Developed in the early 1970s, most of the base metal alloys are based on nickel and chromium, but a few cobalt-chromium and iron-based alloys are also available. Because they are not noble metals, their corrosion resistance depends on other chemical properties. A thin, invisible chromium oxide layer provides a complete and impervious film that passivates the surface of the alloy. The passive layer is so thin, it does not dull the surface finish. A similar passive oxide layer limits surface corrosion in ordinary stainless steel.

In addition to noticeable differences in chemical properties, the nonprecious alloys have different physical properties than the noble metal alloy groups. The most
significant of these are high hardness, high yield strength, and high elastic modulus. Elongation is about the same as for gold alloys but is negated by the high yield strength, which makes it difficult or impossible to work the metal.

When used for metal occlusal restorations, the nonprecious alloys have only a few recognizable advantages. They are low in cost, and some have high hardness values, which can be important when wear resistance is needed. Some nickel-chromium alloys in this group, especially those containing beryllium, have mold-filling abilities that are superior to all other groups. This mold-filling ability permits easier casting of thin sections and produces sharp margins on castings.

However, base metal alloys have many Disadvantages when used for metal occlusal restorations. Their hardness makes occlusal adjustments, polishing, crown removal, and endodontic opening very difficult. Laboratory labor costs are often higher for crowns made from nonprecious alloys because their hardness increases working time. Increased labor costs offset the slight savings in the cost of material. Although casting accuracy can be excellent, the high casting shrinkage (approximately 2.3%) must be accommodated. This usually requires modification of the casting techniques used for gold alloys, which have a lower casting shrinkage (1.4%). Soldering is unreliable in areas where stresses are involved, although soldering of contacts and minor repairs presents no problem. For the latter, white palladium-based solders work well.

Properties that are considered Disadvantages for metal occlusal applications can be used to advantage in porcelain occlusal restorations. Examples include high tensile strength (up to 120,000 psi, or 830 MPa) and high elastic modulus (about 30 million psi, or 200,000 MPa). The high tensile strength permits use of thinner metal sections than would be possible if noble metal alloys were used (with the possible exception of some high-palladium alloys). Nickel-chromium alloys have the highest elastic moduli of all dental alloys, which decreases flexibility to a significant degree. The flexibility of a fixed partial denture framework constructed of nickel-chromium is less than half that of a framework of the same dimensions made from a high-gold alloy. Unlike the relatively thick metal crowns, PFM crowns can be easily removed by penetrating the porcelain with rotary diamond instruments, followed by separating the thin metal with proprietary carbide burs made for this purpose.

The addition of beryllium to some nickel-chromium alloys results in more favorable properties. Beryllium increases fluidity and improves casting performance. Beryllium also controls surface oxidation and results in more reliable, less technique-sensitive porcelain-metal bonds. Generally, these bonds are satisfactory when the alloy contains beryllium but are often questionable when beryllium is lacking. Beryllium-containing alloys require strict control of grinding dust in dental laboratories according to the Occupational Safety and Health Administration (OSHA).

Nickel-chromium alloys show sag resistance that is uniformly superior to all noble metal alloys. This characteristic, along with increased stiffness and high tensile strength, indicates use of these alloys in fixed partial dentures. The problems with presoldering, often necessary for fixed partial dentures, can be easily overcome by using the cast-joining techniques described by Weiss and Munyon (1980).

Some nickel-chromium alloys have been chemically modified to overcome certain
objectionable properties of this group as compared with noble metal alloys. Examples include products that are advertised to "feel like gold," "cast like gold," or "process like gold." In general, such modified alloys fall short in mechanical or physical properties, or in casting behavior, when compared with the better nickel-chromium-beryllium alloys in this group. The latter alloys are simply different from gold or other noble metal alloys, and their differences, when taken into account, can be used to advantage.

The allergenic and carcinogenic properties of the base metals, especially nickel, are controversial. Some investigators report no allergic response to nickel, even in known allergic patients, whereas others report a high frequency of allergic reactions in similar patients. A high incidence of respiratory cancer has been well documented in persons who have occupational exposure to nickel. Nickel is also known to induce tumors in rat muscle tissue, whereas other metals, such as manganese, chromium, copper, and aluminum, do not. Until the potential danger from dental alloys is better understood, care should be taken to avoid inhalation of base metal dust, and caution should be exercised when nickel alloy restorations are placed in patients known to be sensitive to nickel.

In summary, base metal alloys are a useful alternative in PFM restorations. Although the properties of alloys in this group are quite different from those of noble metal alloys, these differences can be used to advantage in many PFM situations. Disadvantages associated with metal occlusal restorations are largely overcome when these alloys are used for thin copings under porcelain. Until more long-term data are available, the practitioner should keep in mind the potential biologic hazards associated with base metals and should always follow recommended safety precautions when using these materials. Such precautions include strict control of grinding dust (with suction, masks, etc) and screening patients for allergy to nickel (eg. pierced earring posts and other jewelry).

**Titanium Alloys**

Although titanium is not a noble or precious metal, it is often not classified with the base metals in dentistry due to its high biocompatibility. The main problem with the use of titanium for PFM restorations is difficult processing. Casting of titanium alloys is difficult due to a high casting temperature (2,000°C), rapid oxidation, and reactions with investments. Titanium melting is best done in specially designed furnaces with an argon atmosphere. Investments for use with titanium are described in chapter 19. A titanium alloy, Ti-6Al-4V, has been used for PFM restorations with special low expansion porcelains (coefficient of expansion of $9 \times 10^{-6}/°C$). Pure titanium is used and formed by machining and spark erosion with a process developed by NobelBiocare for their Procera porcelain. Jeneric/Pentron markets a Ti-Al-V alloy (R/2) with an ultimate tensile strength (UTS) of 1,000 MPa (145,000 psi) and elongation of 9%.

**Criteria for Selecting Alloys**

Rational selection of a specific alloy should be based on a balanced consideration of cost and the properties relevant to the intended use of the alloy. For single crowns, properties such as strength and sag resistance are less important than they are for fixed
partial dentures. Castability, biocompatibility, tarnish and corrosion resistance, porcelain color, and hardness are usually equally important for both alloy uses. For fixed partial dentures, solder and joining behavior, sag resistance, strength, and elastic modulus become increasingly important as the span increases. Porcelain thermal expansion compatibility also increases in importance as the span width increases, because of the complexity of geometry and consequent stress fields due to porcelain and alloy mismatch.

When cost is not a major factor, the clinician has a wide spectrum of alloys from which to choose for PFM restorations. Selection of the best alloy for a particular case depends on a number of factors, including the brand of porcelain selected. Whereas it is difficult to rationalize the use of high-gold alloys because of their Disadvantages, gold-palladium alloys are considered ideal noble metal alloys by many clinicians. If the clinician were forced to use only one alloy for all PFM restorations, the gold-palladium alloys would probably be the most logical selection. Mechanical and physical properties are good, and there are no biologic objections. Porcelain compatibility and castability are quite good with minor (less than 5%) silver additions to the gold-palladium alloys.

Gold-palladium-silver alloys are comparable in cost to gold-palladium alloys, but unfortunately they have porcelain color problems due to the substantial silver content (10% to 15%). The silver-free alloys and the very low-silver palladium-gold alloys appear to be better choices in most cases.

When cost is a major factor, the nonprecious nickel-chromium-beryllium alloys are alternative candidates. Considering all factors, including labor, cost may not be significantly different among the alloy groups. The nickel-chromium-beryllium alloys are often the alloys of choice where large-span fixed partial dentures are involved, high castability is needed, or esthetic considerations are important. The question of biocompatibility with base metal alloys has yet to be resolved.

Palladium-silver alloys have excellent clinical working characteristics and—provided the porcelain is not one susceptible to discoloration in the presence of silver—have no real Disadvantages. Long-term clinical success is well-known with palladium-silver.

The newer palladium-copper alloys appear to have many of the advantages of palladium-silver alloys without the porcelain color problems. Limited experience indicates slightly more difficult melting and casting than with palladium-silver but generally good working characteristics and excellent strength. These palladium-copper alloys may replace the palladium-silver alloys as more clinical evidence is accumulated and soldering techniques are developed.

Some clinicians have found that the very yellow high-gold alloys lead to better porcelain color because their oxides are more readily opaqued with porcelain, allowing thinner opaque porcelain layers and, consequently, better esthetics. In such situations, the lack of strength and poor sag resistance are probably of minor importance and the high-gold yellow alloys should be considered a rational choice for single crowns.

When metal occlusal restorations are present, or when partial veneer abutments are
cast in the same alloy, metal hardness and ductility can become important. In these situations, rational alloy choices may be more restricted.

Clinical indications for PFM alloys

Long-span FPDs

Rational selections

1. Nickel-chromium-beryllium
2. Palladium-silver
3. Gold-palladium, perhaps with minor silver additions

Irrational selections

1. High gold, especially the very yellow ones, due to their high cost, poor sag resistance, and, for the very yellow examples, poor strength
2. Palladium-copper, due to soldering or joining problems
3. Gold-palladium-silver for the combination of cost and porcelain color problems
4. Nickel-chromium without beryllium, because the addition of beryllium greatly enhances its properties without increasing biologic concern to patients

Short-span FPDs and single crowns

Rational selections

1. Palladium-copper
2. Palladium-silver
3. Nickel-chromium-beryllium
4. Gold-palladium, perhaps with minor silver additions

Irrational selections

1. Gold-palladium-silver, due to the combination of cost and porcelain color problems
2. High gold, due to cost and lack of desirable properties
3. Nickel-chromium without beryllium, because the addition of beryllium greatly enhances its properties
Glossary

**alloy** A mixture of two or more metals.

**base metal** A metal that oxidizes readily.

**coefficient of thermal expansion** A measure of the dimensional change upon heating or cooling, expressed as length change per degree of temperature change.

**noble metal** A metal that is resistant to oxidation; includes gold, platinum, palladium, and other platinum group metals.

**nonprecious metal** Relatively inexpensive base metal, such as nickel, chromium, and cobalt.

**precious metal** An expensive metal; includes gold, platinum group metals, and silver.

**sag resistance** Amount a bar of the material will distort at high temperatures.

**semiprecious** Alloys containing silver as well as precious metals.

**Discussion Questions**

1. Why are the melting temperatures of these alloys so high?

2. What is the role of minor alloying elements in the bonding of these alloys with porcelain?

3. Why does the silver content of these alloys cause a major problem with the color of applied porcelains? What has been done about this?

4. What is the main biological concern with the use of nickel alloys in the mouth?

**Study Questions**

*(See Appendix E for answers.)*

1. What are precious and semiprecious alloys?

2. What are nonprecious alloys?

3. What are the rational considerations for the selection of an alloy?

4. What is the advantage of palladium-silver alloys?

5. What are the advantages and Disadvantages of palladium-copper alloys?

6. Which base metal alloys may be used with porcelain?
7. How do nonprecious alloys compare with precious metal alloys?

8. What are gold-palladium alloys, and what are their advantages and Disadvantages?

9. What are the necessary precautions for using base metal alloys in place of precious metal alloys?

**Recommended Reading**


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Chapter 15. Dental Porcelain

Introduction

Porcelain has been used for denture teeth since 1790. Today the major use is for porcelain individual jacket crowns and for veneering metal crowns and fixed partial dentures. Porcelain has also been used to fabricate inlays, but this has fallen into disuse.

The main advantages of porcelain, responsible for its wide acceptance, are its excellent esthetic properties, durability, and biocompatibility. Most attention today is centered on comparisons of the porcelains used to fabricate porcelain-fused-to-metal (PFM) fixed partial dentures and the new, stronger ceramics used for all-ceramic jacket crowns. There is also increasing use of porcelain veneers bonded to teeth to cover unsightly areas.

Porcelain is defined as a white, translucent ceramic that is fired to a glazed state. Dental porcelains are classified according to fusion temperature as follows:

- **High fusing** 1,260°C to 1,371°C (2,350°F to 2,500°F)
- **Medium fusing** 1,093°C to 1,260°C (2,000°F to 2,300°F)
- **Low fusing** 660°C to 1,066°C (1,220°F to 1,950°F)

Denture Teeth

Introduction

The raw materials for porcelain denture teeth are mainly feldspar, about 15% quartz, and 4% Kaolin clay to improve moldability. A plastic mass made from this mixture and additional pigments is formed into metal molds and fired under vacuum to reduce porosity. During firing the porcelain teeth are glazed by the glass produced from the feldspar. Metal pins or holes are placed in the teeth during manufacture for mechanical attachment to the denture base. Acrylic denture teeth have been steadily improved, and their use has grown as an alternative to porcelain.

Advantages

1. Excellent biocompatibility
2. Natural appearance
3. High resistance to wear and distortion

Disadvantages

1. Brittle
2. No bond to acrylic denture bases; requires mechanical attachments

3. Produces clicking sound on contact

4. Cannot be polished easily after grinding

5. Higher density increases weight of teeth

6. Mismatch in coefficient of thermal expansion produces stresses in acrylic denture base

**Porcelain Enamels Used with Metals**

*Composition*

Dental porcelains are used to bond with metals for an outer layer of natural-appearing porcelain, as shown in [Fig 15-1](#).

![Fig 15-1 Cross section of a ceramic metal crown with full coverage.](#)

These porcelains were developed during the 1950s by raising the coefficient of thermal expansion of feldspar porcelain to match the values of gold alloys, which are 13 to 14 × 10⁻⁶ C. This was accomplished by heating orthoclase feldspar with alkali metal carbonates (e.g., K₂CO₃, Li₂CO₃) to approximately 1,093°C (2,000°F) to form a glass and a high-expansion ceramic phase identified as leucite (K₂O · Al₂O₃ · 4SiO₂) (O'Brien and Ryge, 1964). An analysis of several of these high-expansion porcelain enamels is given in **Table 15-1**.
Table 15-1 Chemical analysis of dental porcelains* (%)  

<table>
<thead>
<tr>
<th>Compound</th>
<th>Biodent opaque</th>
<th>Ceramco opaque</th>
<th>VMK opaque</th>
<th>Biodent dentin</th>
<th>Ceramco dentin</th>
<th>VMK dentin</th>
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</thead>
<tbody>
<tr>
<td>SiO₂</td>
<td>52.0</td>
<td>55.0</td>
<td>52.4</td>
<td>56.9</td>
<td>62.2</td>
<td>56.8</td>
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<tr>
<td>Al₂O₃</td>
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<td>11.65</td>
<td>15.15</td>
<td>11.80</td>
<td>13.40</td>
<td>16.30</td>
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<tr>
<td>CaO</td>
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<td>---</td>
<td>---</td>
<td>0.61</td>
<td>0.98</td>
<td>2.01</td>
</tr>
<tr>
<td>K₂O</td>
<td>11.05</td>
<td>9.6</td>
<td>9.9</td>
<td>10.0</td>
<td>11.3</td>
<td>10.25</td>
</tr>
<tr>
<td>Na₂O</td>
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<td>4.75</td>
<td>6.58</td>
<td>5.42</td>
<td>5.37</td>
<td>8.63</td>
</tr>
<tr>
<td>TiO₂</td>
<td>3.01</td>
<td>---</td>
<td>2.59</td>
<td>0.61</td>
<td>---</td>
<td>0.27</td>
</tr>
<tr>
<td>ZrO₂</td>
<td>3.22</td>
<td>0.16</td>
<td>5.16</td>
<td>1.46</td>
<td>0.34</td>
<td>1.22</td>
</tr>
<tr>
<td>SnO₂</td>
<td>6.4</td>
<td>15.0</td>
<td>4.9</td>
<td>---</td>
<td>0.5</td>
<td>---</td>
</tr>
<tr>
<td>Rb₂O</td>
<td>0.09</td>
<td>0.04</td>
<td>0.08</td>
<td>0.10</td>
<td>0.06</td>
<td>0.10</td>
</tr>
<tr>
<td>BaO</td>
<td>1.09</td>
<td>---</td>
<td>---</td>
<td>3.52</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>ZnO</td>
<td>---</td>
<td>0.26</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>UO₃</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>0.67</td>
</tr>
<tr>
<td>B₂O₃, CO₂, H₂O</td>
<td>4.31</td>
<td>3.54</td>
<td>3.24</td>
<td>9.58</td>
<td>5.85</td>
<td>3.75</td>
</tr>
</tbody>
</table>

*Data from Nally and Meyer, 1970.

During manufacture, the fused glass containing the leucite is quenched or "fritted" in water, freezing it in an amorphous state. The frit is then ground up by ball milling and pigmented with colored ceramic compound such as iron oxide to shade the porcelain. Each kit of porcelain supplied to dental technicians contains about a dozen shades of porcelain in at least three translucency levels for forming layers in building up the crown. The opaque porcelain contains approximately 15% tin oxide, zirconium oxide, or titanium dioxide. These opaque porcelains screen out the underlying metal oxide surface color, even in very thin layers. The main layer above the opaque layer is known as the body or dentin layer. Finally, a highly translucent porcelain called incisal or enamel gives the crowns a natural translucent appearance at the incisal edge.

Condensation and sintering

The anatomy of a porcelain crown is built up by hand by applying a paste of porcelain powder (applied to a metal casting or a platinum foil matrix with a small brush). Generally, distilled water or special liquids are used to form the paste with the porcelain powder on a glass slab. As each layer of paste is added, most of the water is removed by vibrator and contact with an absorbent tissue paper. This gives the wet crown more strength and increases the density of the compact.

As each layer of the crown is built up, it is fired in a porcelain furnace. The wet crown is first dried in front of the furnace to remove the residual water and then fired under vacuum. As the porcelain is heated, adjacent particles bond together in a process called sintering. Although there is no meeting of the porcelain powder particles, they join together by flow on contact as a result of surface energy (Fig 15-2).
Firing in a vacuum furnace greatly reduces the porosity of the final product, as shown in Figs 15-3a and 15-3b. The first firing of porcelain is called the **bisque or biscuit bake**. After the incisal layer is added, the porcelain is brought to the final stage, called the **glaze bake**. Upon reaching the glazing temperature of the porcelain, a layer of glass is formed on the surface. After glazing, the crown is removed from the furnace and cooled under an inverted glass or beaker. An alternative approach is to add a thin layer of a low-fusing glass or glaze to the surface and fire to the flow temperature of the glaze.
**Fig 15-3a** Fracture surface of gingival porcelain fired under vacuum. (Scanning electron micrograph, original magnification $\times$ 300. From Meyer et al, 1976. Reprinted with permission.)

**Fig 15-3b** Fracture surface of gingival porcelain fired under normal atmospheric pressure. (Scanning electron micrograph, original magnification $\times$ 300. From Meyer et al, 1976. Reprinted with permission.)
Properties

Porcelain enamels have a vitreous structure consisting of an irregular network of silica produced by the presence of large alkali metal ions, such as sodium, potassium, and lithium (Fig 15-4a).

![Diagram of glass structure](image)

**Fig 15-4a** Irregularity of glass structure due to the presence of large alkali cations.
(After Warren, 1941.)

This amorphous structure produces physical properties typical of a glass, including brittleness and lack of a definite melting temperature. Glasses are brittle due to their irregular structure and the absence of slip planes present in a true crystalline material. The strength of glasses and brittle materials is actually governed by the presence of small flaws or cracks. When stressed in tension, according to the crack propagation theory, small flaws tend to open up and propagate, resulting in a low tensile strength. This is less of a factor with ductile metals because stress concentration around the tip of flaws is reduced by elongation of the metal, as illustrated in **Fig 15-4b**. However, glasses are much stronger in compression, because compressive stresses tend to close up flaws. Therefore, the tensile strengths of vitreous dental porcelains are around 5,000 psi (35 MPa) as compared with compressive strengths of 75,000 psi (517 MPa).
Fig 15-4b Effects of tensile forces on crack propagation in ceramics and metals, resulting in brittleness and ductility, respectively.

The strength of dental porcelains is traditionally tested in flexure as a beam and reported as **modulus of rupture**. The modulus of rupture of a vitreous body or enamel porcelain is about 13,000 psi (90 MPa). The strengths of vacuum-fired porcelains are higher due to fewer flaws.

Vitreous dental porcelains do not have a definite melting temperature but undergo a gradual decrease in viscosity when heated. A sharp decrease in viscosity occurs around the **glass-transition temperature**, $T_g$, as shown in Fig 15-5. Below $T_g$, the glass has the properties of a solid. Above $T_g$, glass flows more readily, and vitreous sintering takes place.
**Fig 15-5** Viscosity (rigidity) increasing rapidly below glass-transition temperature. (From Jones, 1956.)

A typical thermal expansion (TE) curve of a porcelain bar is shown in **Fig 15-6**. The TE is linear up to around $T_g$. Above $T_g$ there is a rapid increase in the rate of expansion when the glass has a more liquid structure. If heating is continued, the bar will reach the softening temperature and collapse. The TE of dental porcelains for bonding to metals is especially important in relation to the TE of the metal involved. Generally, the metal and porcelain should be matched in coefficients of TE values. If the TE curves of the metal and porcelain are too far apart, undesirable thermal stresses will result in fracture of the porcelain, which is the weaker material. The porcelain and metal are therefore said to be incompatible.

**Fig 15-6** Thermal expansion curve for glass with a glass-transition temperature ($T_g$) and softening temperature ($T_s$).
Adhesion to metals

Several factors have been identified as promoting good adhesion or bonding of a porcelain enamel to a metal, including wetting, adherent oxide, and mechanical retention.

Wetting

Good wetting of the porcelain on the metal is indicated by a low contact angle of a drop of the porcelain when fired on the solid, as shown in Fig 15-7. Good wetting promotes penetration of the glass into surface irregularities and, therefore, a greater area of contact. Good wetting also indicates chemical compatibility between the porcelain and the metal.

![Fig 15-7 Good wetting of molten porcelain on alloy.](image)

Adherent oxide

The presence of an adherent oxide on the metal surface that is wet by the porcelain provides a beneficial transition layer. Diffusion of atoms from the metal and porcelain into this oxide usually can be detected and is cited as evidence of a chemical bond. A nonadherent oxide can lead to a weak boundary failure.

Mechanical retention

The presence of surface roughness on the metal oxide surface can result in mechanical retention on a microscopic level, especially if undercuts are present.

Bond failure classification

The several types of failure possible are shown in Fig 15-8.
**Fig 15-8** Classification of porcelain enamel failures according to interfaces formed. Type III represents cohesive failure indicative of a proper bond. (From O'Brien, 1977.)

A Type I failure, shown in **Fig 15-9**, was the result of using a thick, pure gold coating agent on the alloy surface. Such a coating blocks formation of the trace metal oxide layer necessary for strong bonding.
**Fig 15-9** Type I failure in gold casting coated with pure gold.

**Figure 15-10** shows a clinical failure found in a nickel-chromium alloy, which is classified as Type V. If the oxide layer on nickel alloys becomes too thick, a weak boundary layer is formed. **Figure 15-11** shows a nickel alloy surface with microscopic cohesive attachment sites where the bond was stronger than the porcelain.

**Fig 15-10** Example of Type V failure with fracture through oxide layer of nickel-chromium alloy.
**Fig 15-11** Microscopic cohesive attachment sites on fracture interface between nickel alloy and porcelain; a mixed failure.

The relation between the density of these sites to bond strength is shown in **Fig 15-12**.

**Fig 15-12** Relation between density of cohesive attachment sites and observed bond strengths.
When density is low, mixed types of bond failure are observed. However, when the cohesive plateau is reached, the bond strength is equal to the strength of the porcelain, $S_p$, and cohesive failure is observed. Because the bond strength in tension has been found to be about 5,000 psi (35 MPa) with properly oxidized gold alloys, and the tensile strength of the porcelain has been measured to be about the same, higher bond strengths lack practical significance. Shear bond strength tests indicate values of 16,000 psi (111 MPa) to 21,000 psi (147 MPa), as these represent the shear strengths of the porcelains.

**Advantages**

1. High strength
2. Potential for fixed partial dentures
3. Excellent fit

**Disadvantages**

1. Appearance of metal margins
2. Discoloration by metal
3. Difficulty producing an appearance of translucency
4. Bond failure with metals
5. Possible Disadvantages of alloy used

**All-Ceramic Crowns**

**Introduction**

A classification of porcelain crowns according to composition is shown in Fig 15-13. The **jacket crown** is the traditional, accepted term for all-ceramic crowns used for restoring the entire clinical crown portion of a tooth.
The most significant developments in dental ceramics within past years have been in new materials and processes for fabricating ceramic jacket crowns. Porcelain jacket crowns have been used widely in dentistry since Land developed the platinum foil technique in 1903. They were fabricated with high-fusing feldspathic porcelains and were known for natural esthetics resulting mainly from high translucency and the specialized laboratory skills used. However, they have not been used extensively for decades. Failures of these porcelain crowns include breakage or fracture. It has been suggested that failures are caused by the low strength of the porcelain or possibly poor adaptation to the tooth, resulting in high-stress areas; however, this has not been documented.

In 1965, alumina-reinforced porcelain crowns were introduced. These crowns are constructed of a coping or core of a ceramic material containing 40% to 50% alumina with an outer layer of translucent porcelain. The alumina ceramic core material has a flexural strength of approximately 131 MPa (19,000 psi), which is twice that of feldspathic porcelain. The clinical failure rate for anterior crowns made with alumina ceramic cores has recently been established to be below 2%, which is considered an acceptable risk. However, the failure rate for molar crowns was 15.2%, which is unacceptable. Therefore, core materials for anterior crowns should have flexure strengths not significantly below 131 MPa (19,000 psi); the strength required for posterior crowns is still to be determined.

Interest in all-ceramic crowns in the United States has developed recently for many reasons. First, the all-ceramic crown has greater potential for more esthetic anterior restorations. In the PFM crown, the alloy structure produces an opaque appearance and the metal margins are often visible. Also, the selection of alloys for PFM restorations is a confusing issue for many dentists. High-gold-content alloys are relatively expensive, and the alternatives may have Disadvantages such as risk of metal allergy, bond failure, or porcelain discoloration. In addition, the number of available alloys is extensive, and noble metal content alone is not indicative of clinical performance. Currently, alloys that are classified as acceptable or provisionally acceptable by the American National Standards Institute/American Dental Association (ANSI/ADA) range from high-noble alloys to nickel alloys.

**Fig 15-13** A classification of porcelain crowns according to composition.
**Castable and pressed-glass ceramics**

Castable ceramic systems are used to cast crowns by the lost wax process. Indication for use is in single anterior and posterior crowns. Tooth preparation is either a 90-degree shoulder with a rounded internal line angle or a 120-degree chamfer with adequate tooth reduction—from 1 mm minimum on the gingivo-axial surfaces to 1.5 to 2 mm incisally and occlusally.

Impressions, models, and dies are made in the usual manner. The restoration is waxed on the die, and the wax pattern of the crown is invested in a phosphate-bonded investment following the same procedure used for some metal crowns. An ingot of the ceramic material is placed in a special crucible and melted and cast with a motor-driven centrifugal casting machine at 1,380°C (2,500°F) (Fig 15-14). The cast crown is a clear glass that must be heat treated to form a crystalline ceramic, which is essentially a fluorine mica silicate. The crystallization procedure takes several hours in a heat-treating or "ceramming" furnace, with a final temperature of 1,075°C (1,967°F).

![Fig 15-14 Centrifugal casting machine for castable ceramic crowns (Dicor system).](image)

The fired ceramic crown has a "universal" white shade with a translucency of around 50%. It has a flexure strength of 152 MPa (22,000 psi) and a coefficient of TE of 7.2 × 10^-6/°C. Final shading is achieved using a series of light coats of colored surface porcelains. Shaded zinc phosphate cements are suggested by the manufacturer. Because the entire crown is translucent, these colored cements may also be used to achieve the final shade. Before seating the crown, tight contacts may be adjusted with an abrasive stone or wheel and then polished with rubber wheels.

A second cast ceramic (Cerapearl, Kyocera America) based on a calcium-phosphate glass has been introduced. This system also involves casting by the lost wax process and heat treatment to convert the cast glass into a ceramic. Transverse strength values between 17,000 and 43,500 psi (116 to 300 MPa) are reported. However, experience and documentation of the properties of this new system are limited at this time.
compared with the fluorine-mica system.

A newer glass ceramic system is Empress II (Ivoclar). It forms a lithium disilicate phase when heated, after the crown is formed by injection molding while in the glass state (O'Brien and Boenke, 2000). A microstructure of Empress II showing the large amount of crystalline phase is illustrated in Fig 15-15.

![Fig 15-15 Scanning electron micrograph of etched Empress II core showing lithium disilicate crystals.](image)

**Magnesia core material**

Magnesia core material is compatible with the high-expansion porcelains normally bonded to metals. This is a major advantage because technicians are already familiar with handling these porcelains. Also, shade matching with adjacent PFM bridges is simpler.

Magnesia rather than alumina is used as the basis of the high-expansion core material because it has a coefficient of TE of $13.5 \times 10^{-6}/\text{°C}$ for magnesia. This higher TE is explained on the basis that magnesia has a face-centered cubic structure, whereas alumina has a hexagonal, close-packed structure. Strengthening is achieved by dispersion of the magnesia crystals in a vitreous matrix, and also by crystallization within the matrix. These mechanisms are both necessary to achieve the required strength. The microstructure is shown in Fig 15-16 and crown design in Fig 15-17.
The magnesia core material has a modulus of rupture strength of 19,000 psi (131 MPa) after firing and a coefficient of TE value of $14.5 \times 10^{-6}/^\circ C$. Its strength can be doubled to a value of 39,000 psi (269 MPa) by applying a glaze. The glaze strengthens by two mechanisms: (1) It penetrates into surface pores to fill in these points of high stress; and (2) it places the surface layer in compression.
Advantages

1. Adequate strength for most anterior crowns
2. Esthetics superior to PFM for a given shade and technician (eg, no metal margins, discoloration, etc)
3. No risk in choosing alloy

Disadvantages

1. Not used for fixed partial dentures
2. Requires learning to do good shoulder preparation (special instruments available)

Injection-Molded High-Leucite Porcelain

In this process high-leucite porcelain ingot cylinders are heated to 115°C to produce a plastic state. Then, the ingots are pressure-injected into investment molds formed by the lost wax process for crowns, inlays, onlays, and veneers (Fig 15-18). Due to the relatively high leucite content and pressure forming process, the flexural strength of porcelain formed by this process is around 200 MPa. The advantages of the process are good fit and higher strength for the resulting restorations.

Fig 15-18 Injection-molding oven for high-leucite porcelain.
Alumina Slip Casting

Introduction

This process for forming alumina cores gives very high strength values (around 500 MPa). To form the alumina cores, a slurry of fine alumina particles is painted on plaster dies, which absorb the water to form a green state core. The alumina is then sintered at 1,120°C to form a dense mass. Glass is then applied and fused at 1,100°C for 4 hours to allow glass infiltration (Figs 15-19 and 15-20). The rest of the crown is then formed by firing a body porcelain over the cores by traditional firing. Although the strength of the glass-infiltrated alumina cores is high, the alumina cannot be etched and silane treated for resin bonding.

Fig 15-19 Application of alumina-liquids slip to plaster cast that absorbs water by capillary action. (Courtesy of Vident.)

Fig 15-20 Glass infiltration of alumina copings fired on platinum foil.
**Advantages**

1. High strength
2. Good fit

**Disadvantages**

1. High initial cost
2. Long processing time
3. Lack of bonding to the tooth structure

**Milled Ceramic Restorations**

**CAD/CAM**

A number of systems for machining ceramics to produce inlays, onlays, veneers, and crowns have been introduced. One system uses computer-assisted designs and computer-assisted manufacturer (CAD/CAM) technology and comes from the manufacturing industry. In making a restoration with the Cerec CAD/CAM chairside system (Sirona), the following sequence is carried out. First, a powder is applied to the patient's prepared tooth to provide contrast for the optical scanner. Next, the prepared tooth is scanned with an optical probe, and the image is stored in a computer (Fig 15-21). The restoration is designed on a monitor screen with computer assistance. After the restoration is designed, a bloc of a machinable glass-ceramic is selected by shade. With information from the computer, the restoration is milled in a few minutes in a compartment of the chairside unit. The restoration is then acid etched, and a silane agent is applied in preparation for bonding to the tooth preparation. After cementing with a resin cement, the main surfaces are contoured with a diamond contour instrument and polished.
Although the newest model can fabricate crowns as well as inlays, onlays, and veneers, the expensive equipment, extensive training, and longer chairside time make the CAD/CAM system useful for special practices only. Dental laboratory-fabricated ceramic restorations still compete well.

**Advantages**

1. Work can be done chairside
2. Requires only one visit

**Disadvantages**

1. High initial equipment costs
2. Lack of marginal accuracy
3. Labor intensive

**Porcelain Repair**

Since dental porcelains are brittle, fractures occasionally occur. Fortunately they can be repaired, and the entire restoration does not need to be replaced. A number of repair systems have been developed for chairside use (Fig 15-22).
The general method for repair is as follows:

1. Establish a dry field.

2. Remove the surface of adjacent remaining porcelain with an abrasive bur.

3. Treat the area to be repaired with etching gel, and clean off.

4. Silanize the ceramic surface with a silane component.

5. Apply metal bonding component over exposed metal surfaces.

6. Apply bonding resin to the entire area.

7. Repair the restoration with composite component and cure.

The shear bond strengths of repair systems usually range from 10 to 15 MPa after storage in water for 1 day and thermocycling.

**Surface Treatments**

**Polishing**

Surface treatments to increase strength are appealing because they do not require major investments in new technology. The simplest method of increasing strength with a surface treatment is polishing. Polishing reduces the sizes of surface flaws and thus can dramatically increase strength. Dental manufacturers and dental materials researchers have found through strength testing that highly polished ceramic specimens can have strength values 50% to 100% greater than unpolished specimens. Clinically, the occlusal and internal surfaces of all-ceramic crowns are roughened with coarse burs during placement to improve occlusion and fit; this introduces deep
scratches. A study of crown fractures has shown that failure was often initiated from scratches on the internal surface of the crown. Porcelain can be polished and flaws removed with Sof-Lex (3M) or Shofu finishing disks. Reglazing is another option, but is not as convenient in the typical practice. It is foolish to place an expensive high-strength ceramic material in the mouth in a roughened state. Metals in PFM restorations are much less sensitive to these flaws.

**Ion-exchange treatment**

This process (Tuf-Coat, GC), carried out in the dental laboratory, consists of heating the porcelain restoration, which has been coated with a potassium salt, in a low-temperature oven. As a result, sodium ions from the porcelain surface are exchanged for potassium ions. Since potassium ions are about 35% larger in diameter than sodium ions, the surface layer develops a residual compressive stress (Fig 15-23). Consequently, the flexural strength of the porcelain is increased as long as the surface is not damaged by grinding.

![Fig 15-23 Strengthening of porcelain by replacing sodium ions with larger potassium ions. (From Dunn, 1977.)](image)

This procedure is a simplification of the original process of immersing the porcelain restoration in a bath of molten potassium salt. It is especially useful for the internal surface of an all-ceramic porcelain crown, but is not recommended for use with the new high-strength porcelains (ie, aluminous porcelain, Procera, In-Ceram). An anterior porcelain jacket crown made from a feldspathic porcelain without a core may be the only major application.
Hydrothermal porcelains

Hydrothermal porcelains (e.g., Ducera LFC) have a special nonfeldspathic composition that forms a plasticized surface layer when hydrated. This is unusual in that the surface hardness of the porcelain is significantly reduced and flexural strength is significantly increased. The increase in strength is due to the plastic nature of the hydrated surface, which allows for deformation of surface flaws and prevents them from propagating through the bulk. Raman spectroscopy and scanning electron microscopy verifies hydration of the glass structure (Fig 15-24). The reaction occurs as an ion exchange between the alkali ion and a proton:

\[ \text{Si-O-Na}^+ + \text{H}_2\text{O} \rightarrow \text{Si-OH} + \text{Na}^+\text{OH}^- \]

Molecular water has also been detected in the glass structure.

![Hydroxy groups](image)

**Fig 15-24** Structure of hydrated glass. (From O'Brien, 2000. Adapted with permission from *The Compendium of Continuing Education in Dentistry*.)

**Selection Criteria**

The gold standard for selection of a new all-ceramic crown system is the 5-year clinical failure rate of a system for its intended application, not the reported strength of the ceramic. The clinical failure rate is about 2% for all-ceramic anterior crowns; it is higher for posterior crowns. The clinical failure rate for posterior PFM restorations is also about 2%. Therefore, any of the new ceramic systems (Empress, Procera, Magnesia, Dicor, In Ceram, etc) are acceptable for anterior crowns, but are more risky for posterior restorations.

In addition to porcelain strength, other factors that are important for clinical survival of posterior crowns include adhesive bonding of porcelain to teeth and bonding of the porcelain layers to each other. Bonding to dentin is significant because it reduces
stress concentrations within the ceramic crown; this is much more important for ceramics than for metals. Also, bonding of the weaker outer body and incisal porcelains to a high-strength core is important to distribute stress and prevent fracture of the outer layers. Therefore, a well-documented clinical failure rate of no more than about 2% is the standard for replacing posterior PFM restorations with all-ceramic restorations.

Finally, new ceramic systems do not have magical properties that can overcome the common mistakes of dentists and technicians. The fit of a crown is directly related to the accuracy of the impression taken by the dentist; the esthetics require precise shade-taking and an accurate diagram/photograph for the technician and ultimately rely on the artistry of the technician; and the strength of a ceramic restoration is significantly affected by the amount of surface roughness present when the crown is placed.

**Clinical Decision Scenarios for Dental Porcelain**

This section presents an approach for choosing materials and a system for a specific situation. It uses the same format as that presented in chapter 7. Advantages and Disadvantages of each material/system are prioritized using the following codes: * = of minor importance; ** = important; and *** = very important.

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**Materials** IPS Empress vs Vita In-Ceram

**Description** Dentists in a large practice have decided to buy an all-ceramic crown system for their in-house dental laboratory. They are choosing between the IPS Empress and Vita In-Ceram systems.

**Critical Factors** Etchability, cost, and flexural strength

**IPS Empress**

**Advantages**

*** Ability to be etched and bonded to dentin

** Marginal fidelity of < 50 um

*** High flexural tensile strength

** Versatility (anterior crowns, veneers, inlays, and onlays)

** Relatively simple technique using lost wax technique modified for this system

** Biocompatibility

**Disadvantages**
** Learning period required for technicians

** Vita In-Ceram

Advantages

*** High flexural tensile strength

** Marginal fidelity of 20 to 25 um

*** Versatility (single crown; three anterior fixed partial dentures; and inlays, onlays, and veneers with Spinel In-Ceram)

** No thermal sensitivity

** Biocompatibility

** Flexibility of provisional cementation

Disadvantages

*** Long learning period required for technicians, because of the complexity of slip-cast fabrication, a relatively new concept

*** Cannot be etched or bonded to dentin

*** Time and labor intensive

Analysis/Decision  The dentists chose the IPS Empress equipment. Although both systems have merit, the IPS Empress system requires less time and labor and has a shorter learning period for technicians. In addition, IPS Empress allows bonding to dentin, which is permanent and lowers stress concentrations.

*Courtesy of Waletha Wasson, DDS, MS.

Materials CAD/CAM inlays vs Empress inlays

Description  A patient has had a number of routine restorations placed over the years. They consist basically of Class 1 and 2 amalgam restorations and some anterior composites. The patient has been very satisfied with the results of these restorations, but is now seeking more esthetic, but still long-lasting, alternatives to replace the amalgams. She also indicates that she is leaving on vacation in 2 weeks and would like to have all the dental work done before that.

Critical Factor  Good esthetics, speed of completion

CAD/CAM inlays
Advantages

*** Good esthetics
*** Can be completed in one appointment
* No impressions required
* No temporization required
** Not dependent on lab availability

Disadvantages

* More expensive equipment
** Variability in marginal adaptation

Empress inlays

Advantages

*** Excellent esthetics
** Good marginal adaptation

Disadvantages

*** At least two appointments required for completion
* Impressions required
* Temporization required
** Dependent on lab availability

Analysis/Decision  The dentist chose the CAD/CAM porcelain inlays because they can be made chairside, which is much faster than Empress inlays, which must be sent to an outside dental laboratory.

aCourtesy of Gisele de Faria Neiva.

Glossary

aluminous porcelain  A porcelain-containing alumina (Al₂O₃) as an opacifier and strengthener.

bisque (biscuit bake)  The first firing of a porcelain.
cohesive plateau The apparent bond strength between porcelain and alloy equal to the strength of the porcelain, attained when the bond is stronger than the porcelain.

crack propagation theory The theory that glasses and other brittle materials fail by the propagation of minute flaws or cracks when under stress.

feldspar A crystalline mineral of the general formula $X_2O \cdot Al_2O_3 \cdot 6SiO_2$, where $X$ may be sodium or potassium.

frit A powdered glass.

glass-transition temperature The temperature below which a glass behaves like a solid.

glaze (glaze bake) The shiny layer of surface glass produced on a porcelain by firing. The glass may either come from within the porcelain or be added to the surface before the final firing.

jacket crown An all-porcelain crown.

modulus of rupture The flexural strength of a material determined by loading a beam-shaped specimen.

porcelain A white, ceramic-containing glass with a glazed surface.

residual compressive stress Stress frozen in a material that is independent of an applied force.

sintering The densification of a powdered material, usually by heating.

vitreous Glasslike in properties.

Discussion Questions

1. What is the biocompatibility of porcelain to soft and hard tissues that it contacts?
2. Compare the advantages of porcelain-fused-to-metal and all-ceramic crowns.
3. What is the nature of the bond between porcelain and metal?

Study Questions

(See Appendix E for answers.)

1. What is the difference between vitreous and crystalline ceramics?
2. Which ions form glasses by their incorporation into silicate systems?
3. What is the nature of the coloring agents used in dental porcelains?
4. How is a frit made?
5. How does sintering differ from complete fusion?
6. What is the role of surface tension in sintering?
7. Why are porcelains fired under vacuum?
8. Why is glazing not produced during the biscuit bake?
9. How much shrinkage occurs during the firing of porcelain?
10. Define glass-transition temperature.
11. Why are glasses considerably weaker under tensile stresses than under compressive stresses?
12. Why can residual stresses in ceramics be either beneficial or harmful?
13. How does the bond strength of porcelains to alloys compare with the strength of the feldspar glass alone?
14. What effect does the addition of oxide-forming elements to gold have on the wetting and bonding of porcelain enamels?
15. Using the hierarchy of evidence discussed in the book's Introduction, identify the common belief that all-ceramic crowns will last longer if the core materials of the crown have a very high flexural strength.

**Recommended Reading**


Columbus, OH: American Ceramic Society, Jan-Feb.


Chapter 16. Base Metal Casting Alloys

Introduction

Chromium-containing base metal casting alloys have been used in dentistry for almost 70 years. The attractiveness of these materials stems from their corrosion resistance, high strength and modulus of elasticity, low density, and low cost.

Chromium-type alloys are the principal materials used in the fabrication of removable partial denture frameworks, and they enjoy wide use in fixed prosthodontic procedures as well. Alloys of similar composition to those used in removable and fixed dental restorations are available for use in dental, maxillofacial, and orthopedic implants. Figure 16-1 indicates which chromium-containing alloys are recommended for these procedures.

Fig 16-1 A classification of base metal casting alloys.

Alloys for Removable Partial Dentures

Introduction

Alloys based on cobalt or nickel and containing a substantial amount of chromium are suitable for construction of removable partial denture (RPD) frameworks, full denture bases, and temporary toothborne surgical and periodontal splints. Table 16-1 shows the base metal RPD alloys that are currently available.
Table 16-1 Chromium-type removable partial denture alloys

<table>
<thead>
<tr>
<th>Alloy</th>
<th>Manufacturer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vitallium</td>
<td>Austenal Dental</td>
</tr>
<tr>
<td>Vitallium 2</td>
<td>Austenal Dental</td>
</tr>
<tr>
<td>Dentorium</td>
<td>Dentorium Products</td>
</tr>
<tr>
<td>Regalloy</td>
<td>Dentsply International</td>
</tr>
<tr>
<td>JD Alloy: LG Alloy</td>
<td>J.F. Jelenko</td>
</tr>
<tr>
<td>Neoloy &quot;N&quot; Partial</td>
<td>Neoloy Products</td>
</tr>
<tr>
<td>Alloy-Regular</td>
<td></td>
</tr>
<tr>
<td>Nobilium Alloy</td>
<td>Nobilium/American Gold</td>
</tr>
<tr>
<td>Ticonium Premium 100*</td>
<td>Ticonium</td>
</tr>
<tr>
<td>Premium Hard</td>
<td>Ticonium</td>
</tr>
<tr>
<td>Master Tech</td>
<td>Williams Dental</td>
</tr>
</tbody>
</table>

*Nickel-chromium based alloy.

Cobalt-chromium

The major constituents of most available materials are about 60% cobalt and 25% to 30% chromium, which impart corrosion resistance. They may also contain minor quantities of molybdenum, aluminum, tungsten, iron, gallium, copper, silicon, carbon, and platinum. Manganese and silicon enhance fluidity of the molten alloys; molybdenum, tungsten, and carbon are the principal hardening and strengthening elements.

Cobalt-chromium-nickel

The base of one variant of the cobalt-chromium system consists of about 50% cobalt, 25% chromium, 19% nickel, and minor components that are found in other cobalt-based products. However, the variant's molybdenum level of about 3.7% and carbon level of about 0.2% are significantly lower than those of the more conventional cobalt-chromium alloys.

Nickel-chromium

One representative proprietary nickel-chromium alloy contains about 70% nickel and 16% chromium. Important minor components are about 2% aluminum and 0.5% beryllium. Aluminum and nickel form an intermetallic compound (Ni₃Al) that contributes to strength and hardness, and beryllium lowers the melting range, enhances fluidity, and improves grain structure. Other minor elements include molybdenum, tungsten, manganese, cobalt, silicon, and carbon.

Physical properties

Melting temperatures of the base metal RPD alloys are significantly higher than those of dental golds; a fusion temperature range of 1,399°C to 1,454°C (2,550°F to 2,650°F) is common. Polished cobalt-chromium and nickel-chromium prostheses are
lustrious and silvery white. Chromium-type casting alloys are lighter than their gold alloy counterparts. Densities of the cobalt-chromium and nickel-chromium materials lie between 8 and 9 g/cm³. Lightweight materials are especially useful for the construction of large and bulky maxillary removable appliances. Linear casting shrinkage is relatively high, 2.05% to 2.33%.

**Mechanical properties**

Chromium-containing RPD alloys are about 30% harder than Type IV golds. Usually, indentation hardness is measured on the Rockwell superficial hardness scale (R-30N), and a Vickers (diamond pyramid) number of 370 is typical. Appliances cast from alloys exhibiting such hardness values must be finished and polished with special laboratory equipment.

Ultimate tensile strength values range from 90,000 to 120,000 psi (621 to 828 MPa). Values for yield strength fall between 60,000 and 90,000 psi (414 and 621 MPa) and are comparable to the yield strengths of Type IV golds. When comparing yield strengths of base metals with those of golds, the amount of offset (0.1% or 0.2%) used in analysis of stress-strain diagrams must be the same for both types of materials. Use of a 0.2% offset in the evaluation of base metals is common; resulting values may be as much as 10% higher than those obtained with a 0.1% offset.

The modulus of elasticity (stiffness) of cast base metal alloys is approximately twice that of cast dental gold alloys. Thus, under a given load within the elastic limit, a structure cast from a chromium-type alloy will be deflected only half as much as a structure of comparable thickness cast from a gold alloy. Modulus of elasticity values of the cobalt-chromium and nickel-chromium RPD alloys approach 30 million psi (207 GPa).

Chromium-type alloys are quite brittle. Elongation values are dependent on casting temperature and mold conditions. Available cobalt-chromium alloys exhibit elongation values of 1% to 2%, whereas cobalt-chromium-nickel alloy, which contains lesser amounts of molybdenum and carbon than the other cobalt-based materials, shows an elongation of 10%.

The mechanical properties of cobalt-chromium-based RPD alloys can be neither improved nor controlled by heat treatment. On the other hand, the strength and ductility of some nickel-chromium alloys can be altered markedly by high-temperature heat treatment. A softening treatment (15 minutes at 982°C [1,800°F] followed by water quenching) may be used to improve workability, and subsequent rehardening (15 minutes at 704°C [1,300°F] followed by water quenching) increases the toughness of dental castings.

**Chemical properties**

Clinical experience indicates that RPD alloys containing a total of no less than 85% by weight chromium, cobalt, and nickel exhibit a reasonable degree of intra-oral corrosion resistance. The surfaces of these alloys are made passive in air by the spontaneous development of a thin, transparent, and contiguous chromium oxide film. This protective film reduces the corrosion rate to a relatively low level.
Recently, it was shown that thin coatings of electrolytic zirconium oxide (ZrO$_2$) deposited on cobalt-chromium alloys reduce chromium-release levels in artificial saliva, as compared to uncoated cobalt-chromium alloys. The coated alloys also exhibit better corrosion resistance than their uncoated counterparts.

Modification or repair of a cobalt-chromium-molybdenum RPD framework should not be accomplished with a different alloy (cobalt-chromium-nickel, for instance). If a gold braze were used to join these dissimilar alloys, the least noble component (cobalt-chromium-nickel) would undergo corrosion in a galvanic couple with the gold brazing alloy.

All chromium-type alloys are attacked vigorously by chlorine; household bleaches should not be used for cleaning appliances made from chromium alloys.

**Manipulation**

Alloys melting above 1,300°C (2,372°F) should not be cast in gypsum investments. High-fusing alloys require the use of ethyl silicate- or phosphate-bonded investment materials. These investments preclude the possibility of harmful sulfonation of the cast alloy, which could occur upon breakdown of a conventional gypsum (calcium sulfate dihydrate) investment. The thermal expansion of ethyl silicate- and phosphate-bonded investments compensates, in part, for the relatively high linear casting shrinkage of cobalt-chromium alloys. The manufacturer of one high-fusing nickel-chromium RPD alloy suggests that this product be cast into molds made from a special oxalate-protected gypsum-bonded investment.

Entrapped gases can produce voids in large castings. Care must be exercised to ensure adequate spruing and mold venting, complete wax elimination, and proper melting and casting practices to facilitate the escape of mold gases.

High-temperature equipment (oxygen/acetylene, oxygen/natural gas, or electric induction) is required. An induction melting unit equipped with an optical pyrometer provides the most reliable means for attaining proper melting and casting temperatures. Oxidation of the metal and the formation of brittle nitrides must be avoided.

Casting temperatures affect the microstructure and mechanical properties of chromium-type alloys. Excessive temperatures and overheating can lead to production of casting porosity and interaction between the alloy and constituents of the investment.

Simple "broken-arm" machines are not recommended for the centrifugal casting of lightweight, base metal RPD frameworks. The most satisfactory results are obtained with more complex equipment that allows for adjustment and control of acceleration, centrifugal force, and speed.

Cast molds should be set aside and bench cooled to room temperature before further handling. Investment molds can be cleaved with a small pneumatic mallet for retrieval of the castings. Oxide coatings and remnants of investment should be removed by
liquid honing or abrasive blasting, rather than by "pickling" in mineral acids.

High hardness and strength make necessary the use of high-speed laboratory equipment for sprue removal, gross grinding, and finishing operations. Special stones and abrasive wheels are available.

**Disadvantages**

Allergic responses to the constituents of base metal alloys, especially nickel, are observed occasionally. Most adverse tissue reactions attributed to the wearing of a base metal removable prosthesis, however, are manifestations of improper design or poor fit.

Although certain physical and mechanical features of the chromium-type alloys are superior to those of RPD golds, clinical application of the chromium-containing materials may be burdened by the following occurrences:

1. Clasps cast from relatively nonductile base metal alloys can break in service; some break within a short period of time. Apparently, fatigue causes clasp failure. When tested in the dry state, cobalt-chromium-molybdenum specimen bars can sustain 78,000 loading cycles. Similar tests performed in artificial saliva or water show resistance to fatigue up to 59,000 and 36,000 loading cycles, respectively. These reductions of resistance to fatigue are attributable to corrosion of the alloy (Co-Cr-Mo) in aqueous mediums.

2. Minor but necessary adjustments required on delivery of a base metal RPD can be made difficult by the alloy's high hardness, strength, and accompanying low elongation. Such adjustments consume inordinate amounts of the dentist's valuable chair time.

3. High hardness of the alloy can cause excessive wear of restorations or natural teeth that contact the cast framework.

**Alloys for Fixed Partial Dentures**

**Introduction**

Castings of chromium-containing alloys are used as substructures for porcelain-veneered fixed restorations and, to a lesser extent, as all-metal restorations. Numerous varieties of chromium-type alloys are available for fixed partial denture (FPD) applications. They are listed in Table 16-2.
Table 16-2 Chromium-type fixed partial denture alloys

<table>
<thead>
<tr>
<th>Alloy</th>
<th>Manufacturer</th>
<th>Major elements</th>
<th>Contains beryllium</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vera Bond</td>
<td>Aalba-Dent</td>
<td>Ni-Cr</td>
<td>Yes</td>
</tr>
<tr>
<td>Vera Soft</td>
<td>Aalba-Dent</td>
<td>Ni-Cr</td>
<td>Yes</td>
</tr>
<tr>
<td>Beta</td>
<td>Amp-Sterngold</td>
<td>Ni-Cr-Mo</td>
<td>Yes</td>
</tr>
<tr>
<td>Microbond NP2*</td>
<td>Austenal Dental</td>
<td>Ni-Cr-Ga</td>
<td>No</td>
</tr>
<tr>
<td>Vi-Comp</td>
<td>Austenal Dental</td>
<td>Co-Cr-Mo</td>
<td>No</td>
</tr>
<tr>
<td>Calloy</td>
<td>California Dental Products</td>
<td>Co-Cr</td>
<td>No</td>
</tr>
<tr>
<td>Calloy 1210</td>
<td>California Dental Products</td>
<td>Ni-Cr</td>
<td>No</td>
</tr>
<tr>
<td>Biobond II</td>
<td>Dentsply International</td>
<td>Ni-Cr-V</td>
<td>Yes</td>
</tr>
<tr>
<td>Neptune</td>
<td>Jeneric Industries</td>
<td>Ni-Cr-Mo</td>
<td>No</td>
</tr>
<tr>
<td>Novarex</td>
<td>Jeneric Industries</td>
<td>Co-Cr-W</td>
<td>No</td>
</tr>
<tr>
<td>Rex V</td>
<td>Jeneric Industries</td>
<td>Ni-Cr-Mo</td>
<td>Yes</td>
</tr>
<tr>
<td>Rexalloy</td>
<td>Jeneric Industries</td>
<td>Ni-Cr-Mo</td>
<td>No</td>
</tr>
<tr>
<td>Rexillium III</td>
<td>Jeneric Industries</td>
<td>Ni-Cr-Mo</td>
<td>Yes</td>
</tr>
<tr>
<td>Crown Cast*</td>
<td>Jeneric Industries</td>
<td>Co-Cr-Si</td>
<td>No</td>
</tr>
<tr>
<td>Albond</td>
<td>Jensen Industries</td>
<td>Ni-Cr-Mo</td>
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<tr>
<td>Unitbond</td>
<td>Jensen Industries</td>
<td>Ni-Cr-Mo</td>
<td>Yes</td>
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<tr>
<td>Neobond II</td>
<td>Neoloy Products</td>
<td>Co-Cr-W</td>
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<td>Neobond II Special</td>
<td>Neoloy Products</td>
<td>Co-Cr-W</td>
<td>No</td>
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<tr>
<td>NPX III*</td>
<td>Nobilium/American Gold</td>
<td>Ni-Cr</td>
<td>Yes</td>
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<tr>
<td>NuPent</td>
<td>Pentron</td>
<td>Co-Cr-W</td>
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<tr>
<td>Pent V</td>
<td>Pentron</td>
<td>Ni-Cr-Mo</td>
<td>Yes</td>
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<tr>
<td>Pentiullium</td>
<td>Pentron</td>
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<td>PNP</td>
<td>Pentron</td>
<td>Ni-Cr-Mo</td>
<td>No</td>
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<tr>
<td>Safety Bond</td>
<td>Pentron</td>
<td>Ni-Cr-Mo</td>
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<tr>
<td>Wonder White*</td>
<td>Pentron</td>
<td>Co-Cr-Si</td>
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<td>T-3*</td>
<td>Ticonium</td>
<td>Ni-Cr</td>
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<tr>
<td>Ticon*</td>
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<td>Servalloy</td>
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<td>Ultra NP</td>
<td>Unitek</td>
<td>Ni-Cr</td>
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<tr>
<td>Will-Ceram Litecast B</td>
<td>Williams Dental Gold</td>
<td>Ni-Cr-Mo</td>
<td>Yes</td>
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<tr>
<td>Will-Ceram Litecast</td>
<td>Williams Dental Gold</td>
<td>Ni-Cr-Mo</td>
<td>No</td>
</tr>
</tbody>
</table>

*Not used with fused porcelain.

Minor compositional differences can produce significant variations in the microstructures and properties of chromium-containing FPD alloys. So previous experience with one commercial product cannot be used to predict the handling characteristics and clinical behavior of another.
Nickel-chromium alloys

Available nickel-chromium products contain 62% to 82% nickel and 11% to 22% chromium. Common minor constituents are molybdenum, aluminum, manganese, silicon, cobalt, gallium, iron, niobium, titanium, and zirconium. Beryllium, in amounts ranging from 0.5% to 2% by weight, is a constituent of several commercial alloys.

The ease with which certain nickel-chromium alloys can be etched electrochemically has led to the wide clinical use of resin-bonded fixed prostheses, commonly known as "Maryland bridges" (Fig 16-2).

Fig 16-2 A resin-bonded prosthesis with abutments bonded to enamel. (Courtesy of Dr Van P. Thompson.)

Electrochemical etching creates a large bonding area for a resin cement and thereby precludes the concentration of occlusal stresses in narrow tags of luting medium that protrude from perforated frameworks. Also, etching minimizes exposure of the resin luting medium to untoward events that may lead to abrasion or microleakage between the resin and the metal framework. The etched surface of a chromium-type FPD alloy is shown in Fig 16-3.
**Cobalt-chromium alloys**

Typically, these products contain about 53% to 65% cobalt and about 27% to 32% chromium. Some members of the cobalt-chromium alloy family contain 2% to 6% molybdenum. Other minor components include tungsten, iron, copper, silicon, tin, manganese, and ruthenium, a platinum group metal.

**Properties**

Melting ranges of the nickel-chromium and cobalt-chromium alloys are between 1,232°C and 1,454°C (2,250°F and 2,650°F). When polished, the surfaces of nickel-chromium and cobalt-chromium castings are lustrous and silvery white. These alloys are light, with densities slightly greater than 8 g/cm$^3$.

Available chromium-type alloys for casting single- and multiple-unit fixed restorations offer broad ranges of hardness and strength. Most, however, are harder and stronger than their noble metal FPD counterparts. Typical Rockwell and Vickers hardness values are in the vicinity of 50 and 3,000, respectively. Ultimate tensile strength ranges from 80,000 to 150,000 psi (552 to 1,034 MPa), and yield strength is between 32,000 and 110,000 psi (221 and 779 MPa).

Modulus of elasticity values are close to 30 million psi (207 GPa). High stiffness coupled with relatively high yield strength suggests the usefulness of chromium-type alloys for the fabrication of conventional long-span fixed prostheses and thin resin-bonded cast restorations. The elongation of most chromium-containing FPD alloys is low (2% to 3%).

Bond strengths of many base metal-porcelain combinations are comparable to those of noble alloy-porcelain couples. Nevertheless, nickel-chromium-beryllium alloys produce significantly better bonding to porcelain than beryllium-free nickel-chromium alloys. Bonding of porcelain to chromium-type alloys without beryllium is inhibited by thick oxides that accrue on the underlying cast framework. Usually bond failure occurs within the oxide layer.
Nickel-chromium and cobalt-chromium alloys have higher in vitro corrosion rates than dental golds. The clinical significance of this finding, however, is not known. Overall, nickel-chromium and cobalt-chromium FPD alloys are sensitive to routine dental laboratory procedures. Castings of these alloys are not as accurate as those cast from Type III and Type IV golds and low gold alloys. However, when castings are oversized, discrepancies involving margins are relatively small. Thus, it is difficult to produce a cast chromium-type crown with acceptable retention and proper margins.

**Manipulation**

The alloys' high melting and casting temperatures mandate the use of phosphate-bonded investments. Burnout temperatures of 732°C to 927°C (1,350°F to 1,700°F) are employed, with 815°C (1,500°F) being the most popular temperature.

The thermal expansion of some available "high-heat" investments does not compensate adequately for the casting shrinkage of chromium-type FPD alloys. Often, copings and crowns tend to be undersized. Many of the problems encountered in the adjustment and seating of undersized, tight-fitting copings and crowns, however, can be obviated by the judicious use of a so-called die spacer. Researchers and clinicians agree that the castability of nickel-chromium-beryllium alloys is better than that of beryllium-free chromium-type alloys.

High-temperature equipment (acetylene/oxygen, natural gas/oxygen, or electric induction) is required for melting. Oxides that form during heating of the alloy prevent coalescing of the molten ingots. After casting, the lightweight oxides formed upon melting the alloy will remain in the crucible as slag.

High hardness and strength necessitate the use of high-speed laboratory engines and special abrasive disks and stones for sprue removal and finishing. The use of inappropriate equipment and metal removing instruments is time-consuming and ineffective.

The refractory oxides that form on chromium-type alloys, especially those containing beryllium, make soldering a difficult task. Solder joints exposed to high masticatory forces are unreliable. Whenever possible, multiple-component fixed frameworks should be cast in one piece.

Surface preparation of substrate castings is critical. The porcelain-bearing surface of the cast framework must be ground and finished with successively finer ceramic-bonded aluminum oxide stones. The use of resin-bonded disks and other resin-bonded metal removal devices may deposit a resin-rich layer of debris on the metal's bonding surface, which may preclude the future development of the proper surface oxide.

The ground surfaces of the framework should be pressure blasted with clean, unrecycled fine-grit aluminum oxide until all scratches made by grinding are obliterated. Then the framework must be cleaned ultrasonically in an industrial detergent. The use of household detergents for the cleaning of cast frameworks should be avoided because they tend to bleach tenacious surface debris rather than facilitate its removal.
After cleaning, the frameworks should be transferred with a hemostat to a beaker containing deionized water and rinsed ultrasonically. All subsequent handling of the cleaned and rinsed framework must be accomplished with a clean metallic instrument to prevent contamination of the bonding surface by organic materials.

Manufacturers’ instructions regarding thermal conditioning ("desorbing" or "degassing"), choice of porcelain, and fusion procedure should be followed explicitly.

**Biologic effects**

The biocompatibility of chromium-type alloys, especially those that contain nickel, is controversial. Whereas some researchers report no allergic responses to nickel-containing FPD alloys, even in known nickel-sensitive patients, others report a high incidence of adverse response in similar patients.

The high incidence of malignant neoplastic disease of the lung and nasal sinuses found among workers engaged in the production of nickel alloys is well documented. Similarly, nickel is known to produce sarcoma when implanted subdermally in Wistar rats.

To date, however, neither substantial data nor clinical experience unequivocally contraindicates the use of nickel-based FPD alloys. Conversely, the lack of sufficient, appropriate, and reliable long-term biologic data has precluded definite demonstration of the safety of these materials.

Nonetheless, with respect to biologic safety, it must be understood that corrosion is the most relevant behavior of base metal dental alloys. Elements released in the mouth during corrosion can cause local toxicity, allergy, and carcinogenesis.

Worries concerning systemic effects of casting alloys are based on meager evidence. On the other hand, local toxic effects are not well documented, but involve high risk, because local tissues are exposed to high concentrations of released metal ions. Nickel and cobalt are potent allergens. Beryllium, a component of several chromium-type base metal casting alloys, is a mutagen and, in different forms, is a known carcinogen.

Until the potential risks associated with nickel-containing alloys are better understood, care should be taken to avoid inhalation of nickel-containing dust, and caution should be exercised in using nickel- or cobalt-chromium alloys in restorations for nickel- and cobalt-sensitive patients.

Auger electron microscopy and electrochemical tests concerning four nickel-chromium FPD alloys demonstrated that beryllium-free alloys are more resistant to accelerated corrosion than beryllium-containing alloys. After cold sterilization, all four alloys showed lower corrosion rates in cell culture solutions.

Corrosion products of nickel-based alloys released to cell culture media do not alter cellular morphology or viability of human gingival fibroblasts, but they reduce cellular proliferation. However, metal ions released from nickel-based dental casting alloys interfere with cellular energy metabolism. Ion release stops glucose-6-
phosphate dehydrogenase activity and reduces cellular adenosinetriphosphate (ATP).

Dusts from grinding beryllium-containing alloys should be avoided. Work areas must be equipped with adequate air-exchange systems and kept free from dust. Additionally, laboratory personnel should be equipped with individual respirators.

Advantages and disadvantages

The nickel-chromium alternatives to noble FPD alloys offer high strength, stiffness, and hardness at a seemingly low cost. Nonetheless, some significant disadvantages accompany their selection and clinical application. High hardness complicates occlusal adjustment, polishing, restoration removal, and endodontic opening. Usually, laboratory labor is more costly for base metal fixed restorations than for noble alloy restorations because the former's properties and technique sensitivities increase working time. Often, material cost savings are negated by increased labor costs.

Surgical Alloys

Introduction

Presently, most permanent dental implants are machined from highly biocompatible, commercially pure titanium. Nonetheless, chromium-type casting alloys exhibiting a sufficient degree of electropassivity are available for use in bone surgery as plates, screws, intermedullary bars and trays, and posts for anchorage of fixed and removable dental prostheses.

Two chromium-type alloy systems are available. One is based on about 60% cobalt and 32% chromium (Vitallium, Austenal Dental); the other contains about 54% nickel, 25% chromium, and 15% cobalt (Surgical Ticonium, Ticonium).

Chromium-type surgical alloy systems employ about 4% molybdenum, 0.5% silicon, and 0.6% iron. Both contain manganese and carbon. For the cobalt-chromium-based alloy, however, the contents of manganese (about 0.7%) and carbon (about 0.4%) are 24 and 30 times greater, respectively, than in the nickel-chromium-cobalt alloy.

Properties

Liquidus temperatures of chromium-type surgical alloys are in the vicinity of 1,554°C (2,650°F). When highly polished, the surfaces of cast surgical devices are lustrous and silvery white. Most implant devices, however, are left with a dull matte finish. The surgical alloys are light; their densities (about 8 g/cm³) are comparable to those of chromium-type RPD alloys.

Commercially available chromium-type surgical alloys offer a wide choice of mechanical properties, as outlined in Table 16-3. Chemically, these alloys are not inert. However, in vivo and in vitro testing has shown that these materials are more resistant to corrosion than many stainless steels used as implants. Placement of cast screws and pins requires the use of instruments made from the same alloy to prevent corrosion by the interaction of dissimilar metals.
### Table 16-3 Mechanical properties of surgical alloys*

<table>
<thead>
<tr>
<th>Property</th>
<th>Cobalt-chromium</th>
<th>Nickel-chromium-cobalt</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tensile strength (psi)</td>
<td>130,000</td>
<td>68,000</td>
</tr>
<tr>
<td>Yield strength (^a) (psi)</td>
<td>100,000</td>
<td>48,000</td>
</tr>
<tr>
<td>Proportional limit (psi)</td>
<td>69,000</td>
<td>38,000</td>
</tr>
<tr>
<td>Modulus of elasticity (× 10^6 psi)</td>
<td>36</td>
<td>29</td>
</tr>
<tr>
<td>Elongation (%)</td>
<td>2</td>
<td>20</td>
</tr>
<tr>
<td>Hardness (R-30N)</td>
<td>54</td>
<td>19</td>
</tr>
</tbody>
</table>


\(^a\)0.2% offset.

### Manipulation

The same factors that are important to the proper handling of chromium-containing RPD alloys must be considered with respect to the selection of investment materials, melting and casting techniques, and finishing procedures for chromium-type surgical alloys.

### Biologic effects

Implanted metallic bodies can stimulate both generic and specific local responses. A **generic tissue response** is evidenced by fibrous encapsulation of the implant. This reaction may be a manifestation of rejection or merely a reparative response to surgical trauma. **Specific tissue responses** are induced by identifiable chemical, physical, and mechanical factors.

Chemical responses are traceable to corrosion of the implant. Regardless of alleged corrosion resistance, any metal or alloy placed in contact with tissue will exhibit some degree of ionization or solubility. The severity of a chemical response is related directly to the concentration of metal ions released into the tissue. Ions present in local tissues can be transported to various distant organs, particularly the lungs, liver, and spleen, regardless of the implant site.

Mass, size, and configuration of the implant can influence biologic tolerance. The availability of casting alloys suitable for implantation has encouraged the use of massive forms with greater surface areas. Large surface areas create potential spaces at implant-tissue interfaces, which can be transformed into **adventitious bursae**. Large metallic implants are more likely to elicit signs of clinical failure than are smaller ones. Sharp edges and protuberances can injure intervening soft tissue. Movement of the implant resulting from improper stabilization may produce painful bursae and **decubitus ulcers**.

The formation of an adventitious bursa is an early sign of implant rejection. Continued irritation by corrosion or movement can cause pain, swelling, and necrosis, thereby necessitating removal of the implant. Complete rejection is manifested by the formation of fistulae or by frank extrusion.

Nonetheless, bioluminescent assays reveal that American Society for Testing of
Materials (ASTM) F75 Co-Cr-Mo is more toxic than FL38 316L wrought stainless steel. The most toxic individual alloying elements are hexavalent chromium > nickel > cobalt.

Solutions containing nickel ions are known to elicit expression of inflammatory mediators. Also, it is known that exposing keratinocytes, monocytes, or endothelial cells to solutions containing nickel ions causes expression of inflammatory mediators, such as interleukin-1 beta, tumor necrosis factor-alpha, and intercellular adhesion molecules.

**Advantages and disadvantages**

The availability of surgical casting alloys makes possible the on-site preoperative construction of customized replacements for various parts of the mandible and the facial bones. Cobalt-chromium-molybdenum, the most commonly used chromium-type surgical alloy, is especially suited for the management of situations with high strength requirements. On the other hand, the nickel-chromium-cobalt-based material offers good ductility and lower hardness and yield strength. These features may allow adjustment or shaping often required during the placement of cast implant devices.

Over the years, problems associated with surgical alloy-tissue interaction have catalyzed the search for alloplasts exhibiting a greater degree of inertness. Today, the need for this desirable feature in a dental endosteal implant is being met by the remarkable metal titanium.

**Commercially Pure Titanium and Its Alloys**

**Commercially pure titanium**

The following features make commercially pure (cp) titanium an attractive alternative to chromium-type surgical, RPD, and FPD alloys: low cost, low density, nonstrategic geopolitical status, mechanical properties that resemble those of hard and extra hard casting golds, high corrosion resistance, and remarkable biocompatibility.

"Commercially pure" disclaims 100% purity and acknowledges that small amounts of oxygen (0.18% to 0.40% by weight) and iron (0.20% to 0.50% by weight) are combined with titanium. On the basis of varying oxygen and iron contents, cp titanium is obtainable in four different grades.

Titanium crowns, multiple-unit fixed restorations, and RPD frameworks are being cast in research laboratories and in a growing number of commercial dental laboratories. Irrefutable is the proposition that continued improvements in melting and casting technologies and mold fabrication procedures make feasible the production of titanium dental devices.

Fortunately, the nucleus of trained, titanium technicians capable of assuming control and training others continues to grow. The titanium learning curve is steep and comparable to the curve involving the substitution of chromium-type alloys for conventional golds. Clearly, continuous progress is being made in developing solutions to problems encountered in casting a low-density (4.5 g/cm³) metal that
melts at temperatures in the vicinity of 1,700°C (3,092°F); possesses a strong affinity for oxygen, hydrogen, and nitrogen; and reacts with the components of most investment materials. Thus, attempts to minimize the effects of these untoward factors are starting to improve casting efficiency by favorably affecting the filling of molds with molten metal. Complete "mold fills" ensure the production of complete castings.

Attainment of a small marginal gap width between the restoration and tooth structure requires a complete mold fill. In a clinical study of margin quality, the marginal discrepancies for titanium and gold restorations were 72 ± 18 um and 64 ± 18 um for titanium and gold, respectively. Although gap widths in titanium restorations did not match the gold standard, the data justify the use of titanium as an alternative to gold for inlay and onlay restorations.

However, it appears that little or no progress is being made toward improvement of the soundness of castings. Micro- and macroporosity remain consistent features of dental restorations cast from titanium. One study reports that the use of a double sprue technique for titanium copings produces smoother cast surfaces and less internal porosity than a single sprue design. Also remaining to be resolved definitively is the inordinate amount of difficulty experienced in making relatively thin castings.

To make a suitable titanium coping for a porcelain-metal restoration, the pattern is usually waxed to a thickness of about 0.6 mm. This practice makes necessary the machining of the resultant casting to the desired thickness. When heated in air at temperatures in the vicinity of 750°C (1,382°F), titanium becomes embrittled through the absorption of oxygen, hydrogen, and nitrogen. Such embrittlement may cause thin margins of restorations to fracture during burnishing.

When heated at temperatures below 800°C (1,472°F) for short periods, titanium of high purity or commercial purity forms a compact, adherent oxide scale. Moreover, at higher temperatures and extended periods of heat treatment, titanium forms a porous and poorly adherent scale.

During a few successive firings of a low-fusing (750°C), low-expansion (7.1 × 10⁻⁶/°C on cooling) porcelain to titanium, the smooth adherent oxide produced on degassing at 750°C thickens and becomes flaky. An oxide of this type precludes attainment of a reliable porcelain-to-metal bond.

A study comparing titanium and titanium alloy (Ti-6Al-4V) with cobalt-chromium RPD clasps placed in 0.75-mm undercut specimens produced the following results: (1) Over 3 years of simulated clinical use, there was less loss of retention for clasps made from commercially pure titanium and titanium alloy than for cobalt-chromium clasps. (2) Porosity in pure titanium and titanium alloy clasps was greater than for cobalt-chromium clasps. (3) The observed porosity in the titanium clasps did not provide proof of fracture or permanent deformation.

In another study, fatigue resistance of cast clasps made from an experimental titanium alloy (Ti-50.8Ni) was compared with clasps cast from commercially pure titanium, cobalt-chromium alloy, and a gold-silver-palladium-copper alloy. The clasp tips were engaged in 0.25- and 0.50-mm undercut areas of abutments. No significant changes in retentive force were observed for titanium-nickel clasps in 1,010 repeated cycles.
During testing, the other three clasp types showed a significant decrease in the force required for removal from the retentive undercuts.

Properties

At room temperature, the lattice configuration of virgin, cp titanium is hexagonal close packed. This atomic arrangement, known as alpha (\(\alpha\)) phase, persists until the metal is exposed to high temperatures. Upon reaching its transus temperature of 833°C (1,531°F), titanium's entire habit becomes body-centered cubic. This crystal lattice is known as beta (\(\beta\)) phase. Fabricated titanium structures consisting mainly of beta phase are stronger but less ductile than comparable structures with a dominant alpha (\(\alpha\)) phase. So, to obtain consistently reproducible mechanical properties, the solidified metal's cooling rate and the time and temperature of subsequent heat treatments must be controlled.

The tensile strength and elongation of pure titanium are about 36,250 psi (250 MPa) and 50%, respectively. It is known, however, that seemingly slight variations in oxygen and iron content exert substantial and lasting effects on titanium's properties. Generally, as oxygen or iron content increases, strength increases and ductility decreases.

Furthermore, the consequences of titanium-oxygen interactions seem to vary according to melting and casting practices. When melted and cast with the use of an argon arc-centrifugal casting machine, yield strength (0.2% offset), ultimate tensile strength, and elongation for grade 1 cp titanium are about 84,000 psi (579 MPa), 101,600 psi (701 MPa), and 18%, respectively. Comparatively, castings made in an argon-tungsten arc vacuum pressure machine exhibit greater ductility (elongation = 31%), but yield strength and tensile strength drop to 11,300 psi (285 MPa) and 53,000 psi (365 MPa), respectively.

Titanium's elastic modulus of about 17 million psi (117 GPa) is higher than those of Type III and Type IV casting golds (~100 GPa), but lower than those of most chromium-type alloys (171 to 218 GPa). The Vickers hardness of cast cp titanium is 210.

Titanium alloys

Presently, only a few titanium alloys are being used in the United States for the commercial production of fixed or removable dental prostheses. Nevertheless, a high level of interest in such alloys exists within the dental research community. Titanium alloys of interest include Ti-30Pd, Ti-20Cu, Ti-15V, and Ti-6Al-4V.

Alloying other metals with titanium enables attainment of the benefits of lower melting and casting temperatures. Also, it provides a means to stabilize, or expand, either the alpha phase field or the beta phase field. In Ti-13Nb-13Zr, niobium and zirconium expand the beta phase field by decreasing the (alpha + beta) transus temperature. Conversely, in Ti-6Al-4V, aluminum is an alpha stabilizer because it increases the (alpha + beta) to beta transformation temperature and thereby expands the alpha phase field. Accordingly, Ti-13Nb-13Zr is a near-beta alloy with a substantially lower beta transus (735°C) than Ti-6Al-4V (1,000°C), which has an
(alpha + beta) structure.

The effects of interactions with atmospheric gases during melting, casting, and other high-temperature laboratory procedures are as devastating to titanium alloys as they are to unalloyed titanium. Thus, the problems encountered in producing cast dental restorations from titanium alloys are not unlike those experienced with titanium of high or commercial purity.

Properties

The ultimate tensile strength, 0.2% offset yield strength, and modulus of elasticity of various titanium alloys range from 101,500 to 142,800 psi (700 to 985 MPa); 81,200 to 124,700 psi (560 to 860 MPa); and 16 million to 17 million psi (110 to 117 GPa), respectively. Except for modulus of elasticity, the tensile properties of titanium alloys are quite similar to those of chromium-type alloys.

Biologic effects

The corrosion resistance and biocompatibility of titanium at room, oral, and body temperatures are attributed to the formation of a stable oxide film with a thickness of less than 1 nm (10^-9 meter). If the film is scratched or abraded, the involved area repassivates instantaneously. At high temperatures, the oxide film is not protective because it thickens and becomes nonadherent.

Numerous reports document the superior biocompatibility of titanium. The reaction of tissue that contacts titanium or its alloys, Ti-6Al-4V and Ti-13Nb-13Zr, is extremely mild, and direct bone ingrowth or osseointegration does occur. See chapter 23 for further information on osseointegration with titanium.

Titanium's remarkable biocompatibility perpetuates the insatiable desire to develop simple, reliable, and reasonably priced technology that would make possible the routine production of titanium dental restorations in any dental laboratory. Perhaps this desire will be fulfilled not by further modification of melting and casting-art practices, but by exploitation of computer-aided design (CAD) and computer-aided manufacturing (CAM). These technologies are being used routinely to produce customized titanium valvular and titanium hip prostheses.

Glossary

**adenosinetriphosphate** Adenyl-pyrophosphoric acid, a nucleotide compound occurring in all cells but chiefly in striated muscle tissue; it represents the energy reserve of the muscle (myosin).

**adventitious bursa** Fluid-filled cyst formed between two parts as a result of friction.

**base metal alloy** An alloy composed of metals that are neither precious nor noble.

**decubitus ulcer** Superficial loss of tissue on the surface of skin or mucosa, usually with inflammation, caused by pressure.
**ethyl silicate-bonded investment** A silica refractory bonded by hydrolysis of ethyl silicate in the presence of hydrochloric acid.

**generic tissue response** A reaction that cannot be attributed to a specific characteristic of a foreign body or substance.

**intercellular adhesion molecules** Chemokines that regulate lymphocyte and leukocyte migration, roles of integrins involving interactions of lymphocytes with intestinal mucosa, lymphocyte movement through the central nervous system, and tumor immunotherapy.

**interleukin-1 beta** A cytokine produced especially by monocytes and macrophages that regulates immune responses by activating lymphocytes and mediates other biological processes such as inflammation and infection.

**intermedullary** Between bone marrow spaces.

**intermetallic compound** A definite combination by weight between metals. These compounds have specific, reproducible physical characteristics.

**optical pyrometer** A device to measure temperature by matching the color of an electrically heated wire to the color of the surface being tested. The current required is calibrated against temperature.

**phosphate-bonded investment** Used in casting high-melting alloys. Contains a mixture of silica with a metallic oxide and a phosphate that combine to bind the silica together. On heating, forms complex silicophosphates that increase the strength of the mold.

**Rockwell superficial hardness scale (R-30N)** An indication of resistance to penetration of balls or metal cones of differing diameters under different loads.

**specific tissue response** A reaction caused by a definite characteristic of a foreign body or substance.

**tumor necrosis factor-alpha** A cytokine (one of a class of immunoregulatory substances secreted by cells of the immune system); produced mainly by monocytes and macrophages. It is found in synovial cells and macrophages in the tissues.

**Discussion Questions**

1. What are the biocompatibility problems associated with high nickel and cobalt alloys?

2. What are the advantages and disadvantages of Ni-Cr alloys for porcelain-fused-to-metal restorations?

3. Compare the relative clinical advantages and disadvantages of gold and Co-Cr alloys for removable partial dentures.
4. What are the advantages and disadvantages of titanium as porcelain-fused-to-metal fixed partial denture alloys?

**Study Questions**

**Introduction**

*(See Appendix E for answers.)*

1. What are the three major types of chromium-containing dental casting alloys?

2. How do the mechanical properties of chromium-type fixed partial denture alloys differ from those of dental gold alloys?

3. Which properties of the chromium-type alloys make these materials suitable for the fabrication of long-span fixed partial dentures?

4. What effect do excessive surface oxides have on the bonding of porcelain to chromium-type alloys?

5. How do the tensile properties of cast titanium alloys compare with those of chromium-type alloys?

**Removable partial denture alloys: true/false statements**


2. The performance potential of cobalt-chromium castings is enhanced by nitride inclusions and excessive metallic carbides.

3. Low density is a useful characteristic of chromium-type RPD alloys.

4. The hardness and strength of chromium-type RPD alloys are essentially the same as those of Type IV gold alloys.

5. In function, a chromium-type removable partial denture framework is more likely to flex than one cast from a Type IV gold.

6. Chromium-containing removable prostheses should be cleaned regularly in household bleaches.

7. Gypsum investments are not recommended for the casting of cobalt-chromium alloys.

8. Casting temperatures for cobalt-chromium alloys must be controlled carefully.

9. Conventional laboratory equipment and techniques can be used to finish cobalt-chromium and nickel-chromium castings.

10. Sensitivity to one or more alloy components accounts for a significant incidence
of adverse tissue response among wearers of chromium-type removable partial dentures.

**Fixed partial denture alloys: true/false statements**

1. Chromium-containing FPD alloys are based entirely on the nickel-chromium system.

2. The mechanical properties of the chromium-type FPD alloys are slightly inferior to the properties of dental gold alloys.

3. Certain properties of chromium-type alloys make them suitable for fabrication of long-span fixed restorations.

4. Except for modulus of elasticity, the tensile properties of some titanium alloys are comparable to those of chromium-type alloys.

5. The ability of chromium-type alloys to form surface oxides when exposed to temperatures employed in the porcelain firing cycle ensures the development of a strong and reliable porcelain-to-metal bond.

6. The biocompatibility of cp titanium, certain titanium alloys, and chromium-type FPD alloys is well documented.

**Surgical casting alloys: true/false statements**

1. The tensile properties and hardness of nickel-chromium-cobalt and cobalt-chromium surgical alloys are comparable.

2. Chemical factors alone influence tissue response to alloy implants.

3. Metallic ions liberated from an alloy implant can be transported to distant organs.

4. The surface area of a metallic implant plays a prominent role in biologic acceptance.

5. Certain clinical signs signal complete rejection of an implant device.

**Recommended Reading**

**Introduction**

**Chromium-type removable partial denture alloys**


Chromium-type fixed partial denture alloys


**Chromium-type surgical alloys**


**Titanium and titanium alloys**


Chapter 17. Casting

Introduction

The lost wax casting process, though one of the oldest existing technologies, is still the preferred and most commonly used method for casting dental restorations. This mode of casting is favored for dental applications because asymmetrical castings incorporating extremely fine detail can be fabricated conveniently and inexpensively compared with other modes of casting.

In brief, a metal casting is made using a refractory mold made from a wax replica or pattern. The procedure includes the following steps:

1. Prepare the tooth or teeth to receive a cast restoration.

2. Make an impression of the prepared tooth.

3. Pour gypsum slurry into the impression to make a positive cast, which is an exact replica of the dental arch from which the individual die(s) representing the prepared tooth or teeth is sectioned.

4. Make a wax pattern that will be representative of the lost tooth structure.

5. Sprue the wax pattern (fix it in space) (Fig 17-1).

![Fig 17-1](image)

Arrangement of a pattern in a ring prior to pouring the investment material. (After Craig and Peyton, 1975.)

6. Invest the wax pattern.

7. Eliminate the wax pattern by burning the wax out of the investment in a furnace, thus making the mold.
8. Force molten metal into the mold using one of a variety of means.

9. Clean the cast.

10. Remove the sprue from the casting.

11. Finish and polish the casting on the die.

12. Cement the finished cast restoration on the prepared tooth.

When the direct technique—in which the pattern is made on the tooth rather than on the die—is used, steps 2 and 3 are omitted. The desired accuracy of the casting is about 0.1%; therefore, the lost wax procedure requires specially developed materials that compensate for the dimensional changes indicated by the following equation:

\[
\text{Shrinkage (wax + alloy)} = \text{Investment expansion}
\]

**Thermal Shrinkage**

**Wax**

A wax pattern prepared directly in a patient's mouth will shrink about 0.4% when cooled from oral temperature. In the indirect method of preparing the wax pattern on a die, the wax shrinkage is about 0.2%.

**Gold alloy**

The casting shrinkage takes place as the solidified metal cools to room temperature. The values for this shrinkage depend on the geometry of the casting. For example, the gold shrinkage ranges from 1.25% for a thin three-quarter crown to 1.75% for full crowns and 2% for Class 5 restorations. Table 17-1 summarizes the average amount of mold expansion required for typical types of restorations to compensate for wax and metal shrinkages.

<table>
<thead>
<tr>
<th>Table 17-1 Expansion requirements of casting</th>
</tr>
</thead>
<tbody>
<tr>
<td>Restoration</td>
</tr>
<tr>
<td>Thin three-quarter crown</td>
</tr>
<tr>
<td>Classes 1 and 2, small MOD</td>
</tr>
<tr>
<td>Large MOD, three-quarter crown</td>
</tr>
<tr>
<td>Overlay, pin pontic</td>
</tr>
<tr>
<td>Bulky three-quarter crown</td>
</tr>
<tr>
<td>Small Class 5, full crown</td>
</tr>
<tr>
<td>Large Class 5</td>
</tr>
</tbody>
</table>

MOD = mesio-occlusodistal.
Thermal Expansion

High-heat technique

This method employs cristobalite (a high-expansion form of silica) investment materials (Fig 17-2).

Fig 17-2 Typical thermal expansion curves for cristobalite and quartz.

After the investment has been mixed according to the manufacturer's instructions and allowed to set for at least 45 minutes and no longer than 60 minutes, the mold is placed in a 200°C oven for 20 to 30 minutes to burn out the wax pattern. The temperature of the mold is further elevated by transferring the mold to a second oven and holding at 700°C for no longer than 20 to 30 minutes to obtain the maximum thermal expansion of 1.25% (Table 17-2). Because this type of investment is weak by nature, a metal ring must be used. To increase the setting expansion (SE) of 0.35%, the inside of the ring should be lined with a dampened liner strip that also acts as a cushion against which expansion can take place. This greater expansion caused by the uptake of water from the liner, referred to as hygroscopic expansion (HE), is double the normal SE (Table 17-2). The HE cited in Table 17-2 includes the SE. Adding the 0.70% HE to the 1.25% thermal expansion (TE) gives the maximum expansion one can expect—about 1.95%. It should be pointed out that most of the compensation is TE that takes place after the wax pattern is eliminated from the mold.
### Table 17-2  Mold expansion

<table>
<thead>
<tr>
<th>Technique</th>
<th>Setting expansion (%)</th>
<th>Hygroscopic expansion (%)</th>
<th>Thermal expansion (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>High heat (cristobalite)</td>
<td>0.35</td>
<td>0.70&lt;sup&gt;a&lt;/sup&gt;</td>
<td>1.25 (700°C)</td>
</tr>
<tr>
<td>Hygroscopic immersion (Beauty Cast*)</td>
<td>0.30</td>
<td>1.50</td>
<td>0.55 (480°C)</td>
</tr>
<tr>
<td>Hygroscopic water added (Hygrotrol*)</td>
<td>0.75</td>
<td>2.00</td>
<td>0.55 (480°C)</td>
</tr>
<tr>
<td>Phosphate-high heat (Ceramigold*)</td>
<td>0.23-0.50</td>
<td>0.35-1.20</td>
<td>1.33-1.58 (700°C)</td>
</tr>
</tbody>
</table>

<sup>a</sup>Whip Mix.
<sup>a</sup>Wet liner.

### Water-immersion technique (low heat-hygroscopic)

Investments made for water immersion are much stronger than the high-heat types; therefore, a metal ring is not necessary. Instead, a rubber ring is used to contain the mixed investment. Maximum hygroscopic expansion is obtained by immersing the invested pattern and rubber ring, allowing the investment to set under water (hence the name **hygroscopic investment**). In this case, most of the compensatory expansion is HE (1.50%), which again includes the normal SE of 0.3% (Table 17-2). This expansion takes place with the pattern present in the mold, which may cause distortion in certain pattern configurations (eg, mesio-occlusodistal [MOD]). After the investment has set, the rubber ring is removed from the mold, and the mold is placed directly into a 480°C oven for 30 to 45 minutes to eliminate the wax and acquire the additional necessary TE of 0.55% (Table 17-2). With a water-immersion investment, one can expect an overall expansion of 2.10% (Table 17-2).

### Phosphate-bonded investment

This type of investment is supplied as a powder containing silica, primary ammonium phosphate (NH₄H₂PO₄), and magnesium oxide (MgO). The setting reaction in aqueous solution is:

\[ \text{NH}_4\text{H}_2\text{PO}_4 + \text{MgO} \rightarrow \text{NH}_4\text{MgPO}_4 + \text{H}_2\text{O} \]

A phosphate investment such as Ceramigold (Whip Mix) may be mixed with a liquid containing a silica sol in water, which increases setting and hygroscopic expansions. The expansion of a phosphate investment is adjusted by varying the amount of silica sol liquid used in mixing and by employing hygroscopic and high-heat expansion methods. These investments are employed for high-fusing gold alloys used with porcelain.

### Other investment systems

With the advent of alloy systems that do not incorporate any of the noble metals, a very different investment material is required because of the higher fusion or melting
temperatures of the alloys. To overcome the decomposition of the gypsum- and phosphate-type investment systems, silicate investments, zircate investments, and various magnesia investments may be used.

**Spruing**

*Introduction*

The purpose of spruing the wax pattern is fourfold:

1. To form a mount for the wax pattern and fix the pattern in space so a mold can be made
2. To create a channel for elimination of wax during burnout
3. To form a channel for the ingress of molten alloy during casting
4. To compensate for alloy shrinkage during solidification

*Sprue size and design*

The sprue must be large enough so that it remains open until the casting solidifies and short enough to allow rapid filling of the mold cavity. Large and small inlays require sprues that are 14 gauge (4 to 5 mm long) and 16 gauge (3 to 4 mm long), respectively. Large and small crowns require 10- and 12-gauge sprues, respectively, with an average sprue length of 4 to 5 mm.

*Point of attachment*

Sprue attachment must always be made at the bulkiest portion of the pattern (Fig 17-3). If two bulky portions are separated by a thin cross section (eg, MOD) a Y-shaped sprue (as shown in Fig 17-3) must be used. Turbulence of the molten gold as it enters the mold causes porosity, which is due to entrapped gases and an inappropriate angle of sprue attachment. All attachments, both sprue-pattern and sprue-crucible-former, must therefore be "trumpeted" or "filleted" to eliminate all sharp corners, angles, and instrument marks.
Sprue selection

The wax sprue is the most commonly used. A hollow metal sprue pin is preferable to a solid metal pin because of its stronger attachment. Sticky wax must be used to fill the hollow sprue core before use. Plastic sprues are not recommended because their higher flow temperatures and TE characteristics make it difficult to eliminate the sprue. Since the wax melts at a much lower temperature than the plastic sprue and the TE of the wax is 5 times that of plastic, excessive wax pressure may build up in the mold during burnout before the plastic sprue softens.

Orientation in mold

The wax pattern is mounted on the sprue pin, which in turn is mounted on a clean sprue-crucible-former base, as indicated in Fig 17-1. It is essential that, when the
investment ring is placed over the pattern-sprue-crucible-former assembly, the pattern be 6 mm from the end of the ring. If the pattern is less than 6 mm from the end, there is not enough thickness of investment to keep the molten gold from breaking through. If there is more than 6 mm of space, the gold will solidify before the entrapped air can escape, resulting in rounded margins, incomplete casting, or mold fracture.

**Liner**

A liner is placed inside the ring to allow lateral expansion of the investment. Three millimeters of clearance is allowed at each end of the ring so the mold is sealed and anchored in place. After the liner is placed in the ring, it is dipped in water until saturated, and the excess water is shaken off.

**Melting**

**New metal**

Since gold alloys and other alloys change composition during casting, at least one third new gold by weight must be used for each melt.

**Contamination**

Clean melting crucibles are essential to prevent alloy contamination. Copper-containing gold alloys and noncopper alloys for use with porcelain should not be melted in the same crucible. Previously cast metal must be thoroughly cleaned using appropriate fluxes to remove all gases, oxides, and investment before remelting.

**Methods of melting**

The various modes available for melting alloys may be grouped into two categories: (1) those employing a torch, and (2) those employing some form of electrical assistance, as shown in Table 17-3. Each mode has unique advantages and Disadvantages, and in general the modes requiring some form of electrical energy are more costly than those requiring a torch.

<table>
<thead>
<tr>
<th>Torch</th>
<th>Electrical</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gas/air</td>
<td>Resistance</td>
</tr>
<tr>
<td>Gas/oxygen</td>
<td>Induction</td>
</tr>
<tr>
<td>Air/acetylene</td>
<td>Electric arc</td>
</tr>
<tr>
<td>Oxygen/acetylene</td>
<td></td>
</tr>
</tbody>
</table>

**Torch melting**

The natural gas/air method is the most common for melting gold alloys intended for inlays and fixed partial denture purposes. Such alloys require the maximum heat
production that a gas/air torch is capable of generating (Fig 17-4). For gold alloys with higher fusion temperatures, such as those intended for porcelain-fused-to-metal (PFM) applications, palladium alloys and some base metal alloys can be melted by the higher temperatures provided by the natural gas/oxygen torch.

![Fig 17-4](image)

**Fig 17-4** Zones of torch flame. The innermost cone (A) consists of unburnt gases and is not hot enough to melt gold. The second cone (B) is blue with burning gases and forms a reducing atmosphere. Point X is best for melting or soldering, because it is the hottest zone of the flame. The outer cone (C), which contains burnt gases, forms an oxidizing atmosphere and should be avoided.

Air/acetylene and oxygen/acetylene generate much higher temperatures, with the latter providing the hottest flame. Although air/acetylene and oxygen/acetylene modes have the advantage of melting the alloys faster than gas/air or gas/oxygen modes, the Disadvantages outweigh their advantages for the following reasons:

1. Excessive heat may distill lower melting components, thus changing the composition of the alloys.

2. Overheating some alloys allows environmental gases to be dissolved in the melt, resulting in porous castings.

3. Some alloys, particularly base metal, may be excessively hardened and made too brittle due to the uptake of carbon from the acetylene flame.
4. Since the environment cannot be controlled, the extremely high temperature may cause excessive oxidation of the alloy. In fact, the newer titanium alloys cannot be melted by torch. The best practice is to leave the use of the air/acetylene and oxygen/acetylene modes to a highly skilled technician.

Electric melting

The electric mode of melting includes electric resistance melting, which is suitable for all gold alloys, as well as induction melting and electric arc melting, which are capable of melting cobalt-chromium and titanium alloys. Gold and palladium alloys require a reducing atmosphere as supplied by the use of the middle blue cone of the torch (see Fig 17-4), with flux being added to the melt. A greater convenience is afforded by the electric resistance furnace, which uses a carbon (rather than ceramic) crucible, thus providing a reducing environment throughout the melting regime.

The electric resistance furnace also provides the best means of temperature control. The torch adjustment is always variable with no means of monitoring the temperature, and the induction and electric arc modes are less controllable than the resistance mode because melting is achieved so rapidly. Attempts to control the melt temperature when using induction or electric arc require a pyrometer to be focused on the melt. However, since each alloy has its own unique emissivity, the pyrometer is only reliable when calibrated to a given alloy; when a different alloy is used, the pyrometer must be recalibrated.

Casting Machines

Introduction

Casting machines provide the means for transferring the molten alloy from the melting crucible to the mold. They vary greatly, from the most elementary to the complex, with corresponding variations in cost. The casting machines can be divided into two broad categories: (1) those employing pneumatic forces, and (2) those that provide primarily centrifugal forces to transfer the molten alloy to the heated mold.

Pneumatic forces

A very old method still used to a limited extent throughout the world today uses steam pressure to drive the melt into the mold. The alloy is melted in the sprue-crucible-former part of the mold. A simple handheld device fitted with a wet asbestos plug is quickly placed over the melt, making a seal with the mold ring. When the device is pressed down firmly, the wet asbestos makes contact with the molten alloy, generating sufficient steam pressure to complete the casting.

The pressure/vacuum casting machine produces pressure over the molten alloy. A vacuum is applied to the bottom of the mold; thus, the molten alloy is "pushed and sucked" simultaneously into the mold. In variance, the vacuum/pressure types of casting machines first evacuate the melting chamber to reduce oxidation and then apply air pressure uniformly about the casting ring, forcing the alloy into the mold. An innovative variation of the pneumatic mode is the jet casting machine, which
depends on combustion gas to provide the casting force.

Casting results from these three machines are comparable. One factor must be taken into consideration when using the pneumatic-type casting machines: They all provide good castings when a porous gypsum investment is used, but when a denser investment—such as a phosphate investment or an investment that tends to sinter on heating—is used, castings may be incomplete. Since the melting procedure is usually done directly in the sprue-crucible-former part of the investment mold, which may be subject to thermal decomposition, thin sprues must be used to prevent premature entrance of the alloy into the sprue opening of the mold during melting.

**Centrifugal forces**

Two considerations are fundamental to any casting procedure: (1) the casting force, be it pneumatic pressure or centrifugal (speed of the rotating arm), and (2) the rate of time required to fill the mold. In most cases the two are inversely related—the greater the casting force used the less time required to fill the mold.

The **castability** of an alloy is determined by a number of variables: its melting range, the mold temperature, and the initial acceleration of the casting machine or applied force, among them. Although the rotation speed of the spring-driven casting machines can be increased by increasing the number of windings of the casting arm, and the vertical rotating casting machines reach their maximum revolutions per minute before the horizontal rotating casting machines, the time required to fill the mold is significantly greater than when pneumatic modes are employed. Even more time is required to fill the mold with the motor-driven casting machine because of the longer time needed to achieve maximum rotation. A large posterior full crown, whose occlusal surface cross section is thick with large adjoining cusps and whose margins are feathered, can hinder the maximizing of the casting force. As the molten metal under a given centrifugal force enters the thickest cavernous area of the mold, a pressure drop is realized before the molten alloy is forced into the feathered margin area. The greatest difficulty in the overall casting regime is in the initial acceleration of the casting machine's arm. To improve the castability of an alloy, one must increase the initial acceleration of the rotation so the mold is filled quickly and the ultimate rotation speed of the casting machine becomes relatively unimportant compared with the initial acceleration.

The traditional centrifugal casting machine has been modified (Airspin Pneumatic Caster, Airspin Manufacturing) to enhance or improve the initial acceleration. It is a vertical-motion centrifugal casting machine that is rapidly accelerated by activating a pneumatic cylinder and piston. Compressed gas is applied to the piston to move it within the cylinder, and a connected drive chain engages a drive wheel coupled to the casting arm. Movement of the piston through the drive stroke is thereby coupled to rotate the casting arm and provide such rapid acceleration that the mold is filled with molten metal before any significant cooling can take place.

With the advantage of increased acceleration, casting of finer detail and thinner cross sections can be made. This has clinical significance because the ability to cast thin sections is critical for maximum marginal integrity and is especially important with alloys intended for porcelain bonding. When casting alloys with a high modulus of
elasticity are used, a coping thickness of only 0.1 mm will provide sufficient strength to support the porcelain and allow increased thickness of body porcelain to cover the opaque porcelain, thus greatly improving the esthetics.

**Cleaning and Finishing the Casting**

After the casting has been made, the mold is removed from the cradle of the casting machine as soon as the arm stops rotating. It is immersed in cold water when the button (the excess gold left in the crucible above the sprue) turns a dark red. This softens the gold casting, which can then be hardened in a controlled manner by heat treatment. As the mold is plunged into the water, the water in contact with the mold boils, breaking most of the investment away from the casting. The rest can be removed easily by the careful use of any suitable hand instrument or brush.

The cleaned casting may have a dark or tarnished appearance due to oxide or sulfide deposits on the surface. The deposits are removed by immersing the casting in a pickling solution. Such a solution is usually an acid or a combination of acids with a HCl base. It is highly recommended that the pickling solution be acquired from the alloy supplier, because their formulations are usually nonfuming and emanate no corrosive vapors.

The casting is placed in a small porcelain casserole and just enough solution is added to cover the casting. The solution may be heated carefully until the characteristic gold luster develops. The pickling solution is flushed from the casserole with copious quantities of tap water. Plastic or quartz tongs are not required for this method, eliminating the chance of dropping the casting; also, since only a small quantity of solution is needed (10 mL), new pickling solution can be used each time.

Pickling solution should never be used if the color has changed even slightly, nor should metal tongs be employed. The gold casting should never be heated directly or indirectly in a flame prior to immersing it in pickling solution, because the casting may be dropped or become distorted, or the margins may become rounded from overheating. Furthermore, grain growth is possible, making the casting weak and difficult or impossible to polish. Ultrasonic cleaners and abrasive spray devices (sandblasting) can also play a useful role in the overall cleaning procedure.

Polishing is the final step after precision of fit and marginal integrity have been established on the die. Rubber, rag, or felt wheels impregnated with abrasives are used in the initial stages of finishing. Final polishing is accomplished with various oxides of tin and aluminum used in conjunction with a small rag or chamois buffing wheel, followed with an iron oxide rouge. Since these oxides are often supplied in stick form for convenience of handling and confining the abrasive to the wheel, residual traces of the rosin or waxlike matrix must be removed with a suitable solvent or "polishing compound remover" followed by a hot, soapy water rinse.

**Other Casting Alloys**

The metals used for partial denture frameworks are usually alloys of cobalt, with chromium being the primary additive. Lesser amounts of nickel, molybdenum, and carbon are also added in varying amounts. Gold alloys have been used for this
purpose; however, since they possess a relatively low modulus of elasticity and proportional limit, are relatively soft, and have a density about twice that of the cobalt alloy, the gold alloys are not preferred for partial denture fabrication. Both alloys, however, possess excellent corrosion resistance properties.

The casting technique for cobalt alloys is very similar in most respects to that described for gold alloys. Although the demands for accuracy may not be as stringent, the principles set forth for gold casting are still valid. Only the primary differences between the gold and cobalt alloy techniques will be presented here. Because the general practitioner is not likely to have the appropriate casting facilities, further details of the actual casting technique should be obtained by reading technical manuals supplied by the manufacturers.

The investment used must be of the phosphate- or silica-bonded type. Cobalt allows melt in the temperature range of 1,250°C to 1,450°C (2,280°F to 2,640°F), well above the decomposition temperature of calcium sulfate; gypsum investment therefore cannot be used.

Induction is the preferred method for melting cobalt alloys. An oxygen/acetylene torch may be used, but the control of the flame is critical. If the flame is acetylene-rich, carbon is picked up by the alloy, embrittling it; if the flame is oxygen-rich, the metal is oxidized. The melting range of these alloys is well above the upper limits of an air/gas or an oxygen/gas flame, and therefore these mixtures cannot be used. Because of the relatively great hardness of the cobalt alloys, sandblasting and electrolytic polishing techniques are used. Rouges can often be used for final polishing.

Nickel alloys used for fixed partial dentures are melted with gas/oxygen or acetylene/oxygen torches or by electrical induction. Phosphate-bonded investments are required, since these alloys are cast at about 1,260°C (2,300°F). They are handled similarly to the cobalt-chromium alloys. Titanium-based alloys must be melted in a highly controlled inert environment (ie, argon or nitrogen) using either the induction or electric arc mode. Because titanium alloys have such extremely high fusion temperatures, only zircate or magnesia investments can be used.

Titanium and titanium casting are discussed in chapter 16.

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**Gold Casting Troubleshooting**

**General problems**

1. **Problems with accuracy**

   *Water/powder ratio* Higher values reduce setting, thermal, and hygroscopic expansions, giving smaller casting.

   *Spatulation* Increased spatulation increases setting expansion, giving larger castings.
Burnout temperatures Lower temperatures result in less thermal expansion and smaller castings.

Immersion time Delay results in decreased hygroscopic expansion and smaller castings.

Water-added technique Decreased amounts of water added to investment result in decreased hygroscopic expansion and smaller castings.

Water bath Temperatures below 37°C (100°F) reduce hygroscopic expansion and result in smaller castings.

2. Problems with distortion

Wax too hot Excessive shrinkage results on cooling.

Wax too cool The pattern undergoes stress release with a change in shape.

Insufficient pressure during waxing The pattern distorts because of thermal shrinkage.

Delayed investment The sooner the investment is complete, the less distortion there will be.

Heating pattern during spruing This may create distortion.

Overheating casting during soldering procedure This warps or melts the margins.

3. Problems with bubbles

Inadequate vacuum or ineffective painting procedure The vacuum must have at least 26 in of mercury for vacuum investing.

Water/powder ratio If the investment is too thick, it will not cover the pattern completely.

Excessive vibration of the ring This produces small nodules.

4. Problems with surface roughness

Water/powder ratio A high ratio increases the roughness of the mold.

Excess wetting agent or salivary contamination This may form a film on the pattern surface and be reproduced on the casting surface. Generally the roughness is localized on the exterior and is masslike in appearance.

Prolonged heating or overheating of the mold Prolonged heating may cause investment disintegration, whereas overheating may cause a reaction between the alloy and casting investment. Roughness appears general and feels sharp.

Premature heating of casting investment Wait a minimum of 45 minutes for burnout.
5. Problems with fins on the surface or margins

*Prolonged heating* This may produce cracks in the investment that radiate out from the surface of the pattern.

*Heating rate is too rapid* Cracks may appear in the investment, caused by nonuniform heating of investment.

*Water/powder ratio* A high ratio produces a weak investment that may crack.

*Excessive casting pressure* Metal impact may cause investment fracture.

*Cooling of the investment prior to casting* This may produce cracks in the investment.

6. Problems with short, rounded margins

*Investment or alloy is too cold* In this case, solidification is completed prior to completion of the mold.

*Casting pressure* The casting pressure is inadequate to fill the mold because of trapped air. The situation depends on the pattern position and the porosity of the investment.

*Incomplete burnout* This may result in residual carbon at peripheral portions of the pattern.

7. Problems with miscasting

*Casting is nearly or entirely missing* The pattern detached from the sprue pin, due to excessive vibration.

*Pattern fractured during investing*

*Gold alloy was too cold during casting*

*Incomplete burnout*

*Sprue pin was too small* If the sprue freezes before the alloy fills the mold completely, incomplete casting results.

8. Problems with pits

*Inclusions of investment, asbestos, or other debris are carried to the margin by molten alloy* Pits generally exhibit angular edges. They may be rounded if they are the result of flux inclusion.

*Problems with internal porosity*

9. Problems with localized shrinkage porosity
Sprue pin diameter is too small The sprue channel solidifies simultaneously with or before the casting. Additional molten alloy is prevented from entering the mold to compensate for solidification shrinkage.

Alloy or mold temperature is too low Rapid solidification of the alloy results here.

10. Problems with subsurface porosity

Short, thick sprue pin Rapid entry of the alloy causes skin formation; the bulk of alloy pulls away, forming subsurface porosity.

Alloy or mold temperature is too high The first portion of gold to contact the investment will solidify and form a thin skin. The alloy behind it shrinks during solidification and pulls away, forming small porosities.

11. Problems with microporosity

Alloy or mold temperatures are too low If solidification occurs more rapidly than normal, shrinkage may develop throughout the casting.

Problems with external porosity

12. Problems with back pressure porosity

Insufficient alloy mass Air is entrapped in the solidifying alloy.

Insufficient turns on the casting machine The denser the investment, the greater the force needed to eliminate the gas within the mold chamber.

Pattern is too far away from the end of the ring The situation is aggravated by dense investments and lower burnout temperatures.

Glossary

castability The ability of an alloy to completely fill a mold.

casting A process for forming objects by pouring molten metals in molds that are cooled to cause solidification.

die A replica of a tooth or prepared tooth onto which a wax pattern is formed. The die is usually made of a gypsum material.

gypsum The hydrated product formed when plaster is mixed with water: CaSO$_4$·2H$_2$O.

investment A molding material that surrounds the pattern and subsequently hardens and forms the mold after the wax pattern is eliminated.
**lost wax casting process** A method of casting metals that uses a wax model of the object to form the mold and, after wax elimination, the mold cavity.

**refractory** Any material that has an extremely high melting point. In gold casting, some form of silica (usually cristobalite) would be the refractory.

**sprue** Part of a casting that acts as a channel for the molten metal to flow into the mold cavity.

**Discussion Questions**

1. For an inlay casting, how close is the tolerance at the margins?
2. If an inlay or crown casting is too small, what factors in the process may be responsible?
3. How can porosity, which is a problem in polishing, be reduced?
4. Compare the advantages and Disadvantages of the high-heat and hygroscopic mold expansion techniques.
5. Why are the properties of the wax pattern so important in the accuracy of the lost wax process?

**Study Questions**

*(See Appendix E for answers.)*

1. Describe the lost wax technique of casting.
2. What equation must be satisfied to make a casting that will fit?
3. What are the two primary techniques used in fabricating a wax pattern?
4. What primary factor determines the amount of gold shrinkage?
5. What is cristobalite?
6. Why is it necessary to use an asbestos liner when making a cristobalite investment?
7. Name four commonly used investing techniques.
8. Which investment is designed for the higher-fusing casting golds, as used in porcelain-fused-to-gold techniques?
9. How can the expansion of phosphate investment best be controlled?
10. What are the four primary functions of the sprue?
11. Why must gold alloys be cleaned by melting on a charcoal block before being used for the second time?

12. Name three types of casting machines commonly used in gold casting.

13. What is the nature of pickling solutions used for cleaning gold alloys?

14. What are the primary components of the base metals used for partial denture frameworks?

15. Why is gold not the metal of choice for fabricating a partial denture framework?

16. Name two types of investment used for casting cobalt-chromium alloys.

**Recommended Reading**


Eames WB, MacNamara JF (1978). Evaluation of casting machines for ability to cast
sharp margins. Oper Dent 3:137.


Chapter 18. Soldering, Welding, and Electroplating

Soldering

Introduction

Soldering is often used in the construction of dental appliances. Larger fixed partial dentures are frequently cast in parts that are soldered together after careful fitting to the master cast. This procedure improves dimensional accuracy. Wrought-wire clasp arms can be soldered in place for partial dentures using investment soldering techniques. Orthodontic wires and bands are often soldered together. Soldering can be used to build up certain regions of crowns and inlays where the dimensions should be increased, such as for missing contact points. Some casting defects can be corrected with soldering.

Clean oxide-free metal surfaces brought into intimate contact will bond together as a consequence of metallic bonding forces. However, this does not occur in most practical situations due to surface contamination and/or oxidation. Additionally, metallic bonding forces are of a very short range compared with the surface roughness of even polished flat surfaces. Only small areas will be sufficiently close for bonding to occur, even when clean and oxide free.

The basic technique of soldering is first to obtain metal surfaces that are free of contamination. This is accomplished by cleaning and by using fluxes, which also prevent oxidation during the soldering process. Second, a metal or alloy with a lower melting point than the parts to be joined must be chosen. Third, the parts and solder should be brought to the solder's melting temperature. If the molten metal wets the solid metal, it will spread between the flux and the metal part, providing intimate contact. Upon solidification, the solder will still be in contact and metallic bonding will be established. Solder penetrates joints by capillary action.

Solder compositions

Usually the fineness of a solder is less than that of the alloy being soldered. The proportion of pure gold in gold solders is specified by its fineness. Some manufacturers give a carat designation indicative of the gold alloy for which the solder is to be used (e.g., an 18-carat solder is intended for soldering an 18-carat alloy); the proportion of gold in the solder may be less than 18 carat. The compositions of several solder alloys are given in Table 18-1.

Copper is added to a solder to lower the fusion temperature, improve its strength, and make it amenable to age hardening. Silver in a larger proportion than copper improves the wetting (spreading and penetration) of gold solders. Silver also whitens the alloy. Tin and zinc are present in relatively fixed amounts (2% to 4%) to lower the fusion temperature. Nickel may be added instead of copper if a white alloy is desired.

Preceramic solders for soldering of porcelain-fused-to-metal (PFM) appliances must withstand the high sintering temperatures of porcelain and so contain more noble metals and less tin and zinc. Copper is not used because it colors the porcelain green, as does silver.
Table 18-1 Composition of gold solders*

<table>
<thead>
<tr>
<th>Solder no.</th>
<th>Composition (wt %)</th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Gold</td>
<td>Silver</td>
<td>Copper</td>
<td>Zinc</td>
</tr>
<tr>
<td>A</td>
<td>65.0</td>
<td>16.3</td>
<td>13.4</td>
<td>3.9</td>
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<td>B</td>
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<tr>
<td>C</td>
<td>66.1</td>
<td>12.4</td>
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<tr>
<td>D</td>
<td>72.9</td>
<td>12.1</td>
<td>10.0</td>
<td>3.0</td>
</tr>
<tr>
<td>E</td>
<td>80.9</td>
<td>8.1</td>
<td>6.8</td>
<td>2.1</td>
</tr>
</tbody>
</table>

*From Coleman (1928).

Silver solders contain 10% to 80% silver, 15% to 50% copper, 4% to 35% zinc, and small amounts of cadmium, tin, and phosphorus to lower the fusion temperature.

**Flux and antiflux compositions**

Borax flux for soldering gold can be made from dehydrated borax (Na$_2$B$_4$O$_7$), boric acid (H$_3$BO$_3$), and silica (SiO$_2$). The fused borax flux produces oxide-free surfaces over which the solder will flow easily. The flux can be applied as a powder, a liquid (flux mixed with alcohol), or a paste. A paste gives the most control in application and is formed by mixing the powdered flux with petrolatum or a similar inert base.

**Fluoride fluxes** are used for soldering alloys containing chromium because fluorides dissolve the chromium oxide. A flux consisting of potassium fluoride, boric acid, borax glass, and sodium carbonate or silica is typical. The ingredients are fused and ground to fine powder, which is used either directly or as a liquid in alcohol or a paste in petrolatum.

An **antiflux** prevents flow of solder and is used to confine the solder to the work area. Graphite from a lead pencil is a convenient antiflux; however, it is removed by oxidation at higher temperatures. An effective antiflux for prolonged heating or higher temperatures can be made from a suspension of rouge (ferric oxide) or chalk (calcium carbonate) in alcohol.

**Investment materials**

**Soldering investments** should not expand as much as for casting investments because soldering requires lower temperatures than casting. Consequently, quartz-based, rather than cristobalite-based, gypsum investments are used for soldering with typical low-fusing gold solders. Phosphate-bonded soldering investments are used for preceramic soldering.

**Manipulation of solder**

Selection is made on the basis of the solder's corrosion resistance, strength, fusion temperature, and color (to give an inconspicuous joint). The fusion range of the solder must be at least 100°C (212°F) below that of the parts to be soldered. Additionally, selection of solder is based on how permanent the appliance will be and if the appliance can be removed for cleaning. A solder of 580 fineness has been suggested as the lowest fineness to be used in permanent restorations; higher-fineness solders
are generally recommended. Removable appliances can be removed for cleaning and polishing so a lower-fineness solder may be used.

In soldering appliances, such as orthodontic wires and bands, recrystallization or softening of the wires must be avoided. Lower-melting-point solders such as silver solders or low-fineness gold solders, which also improve the mechanical properties of the joint, are preferable. Silver solders are used for stainless steel and other base metal wires. Gold solders used with nickel-chromium and cobalt-chromium-nickel wires are generally about 450-fine and seldom above 650-fine. Higher-fineness solders have inferior mechanical properties.

Cleaning

Surface films of inorganic gases, organic material, and metallic oxides separate the two surfaces and prevent the solder from wetting the metal's surfaces. These must be removed by cleaning and using fluxes, which also prevent oxidation during the soldering process.

Parts must be thoroughly clean in order to obtain a good joint. Casting oxides are removed by pickling in acids. If the parts are polished prior to soldering, they must be thoroughly washed with soap and water then pickled to remove residual polishing materials. Rubber-bonded finishing wheels and points are useful for removing oxides and contamination from the region to be soldered.

Investment soldering

Investment soldering is recommended for precise arrangement of parts for fixed partial dentures or partial dentures with wrought-wire clasp arms. The general procedure is as follows:

1. To prevent warping or porosity, the parts are placed on the master cast so that the gap distance between them is at least 0.1 mm. Excessive gap distance can lead to distortion and pitting. A typical business card, which ranges in thickness from 0.20 to 0.34 mm, can be used as a gauge for maximum gap distance. Sheets of paper should not be used as a gauge because they are often too thin, ranging from 0.05 to 0.1 mm in thickness.

2. The parts are securely fastened together with sticky wax before removal from the master cast and subsequent placement in the soldering investment.

3. Investment should cover metal parts not to be soldered, but no investment should be at the joint.

4. Antiflux may be applied to confine the flow of solder.

5. To obtain dimensional accuracy, the investment is preheated to eliminate moisture and to provide enough thermal expansion to compensate for the thermal expansion of the crowns. Overheating may cause sulfur contamination.

6. Flux can be applied to the joint area before or after preheating.
7. Soldering is accomplished with a reducing flame when the parts are at 750°C to 870°C (1,382°F to 1,598°F), giving them a yellowish red color. The solder should immediately flow smoothly into the joint area if the surfaces are clean.

8. The investment and appliance are allowed to cool for about 5 minutes before quenching. This allows some age hardening to occur and prevents warping that could result if the investment were quenched immediately. Total bench cooling would make the solder too brittle through age hardening.

9. The flux cools to a glass that can be removed through pickling. Generally, finishing instruments should not be used to remove the flux because the glass is harder than the metal, and surrounding metal will be removed along with the flux. Small amounts of flux can, however, be chipped or crushed off with a hard instrument (#7 spatula or knife).

Free-hand soldering

Soldering of orthodontic appliances is generally accomplished without the use of an investment. Orthodontic torches can be placed on the bench so that both hands are free to hold the parts in position. Solder is generally melted onto one of the parts; then they are held together and the joint is heated.

Thin wire parts must be held in contact to avoid narrowing of the joint caused by the surface tension of the solder. This decrease in diameter concentrates stress and is much more significant than for larger appliances.

The wires can easily be overheated, with deterioration of mechanical properties resulting. Orthodontic torches, which develop small needlelike flames, are used to limit the heating to a small area around the joint. Quenching prevents embrittlement of the solder by age hardening. Overheating of stainless steel wires can lead to carbide precipitation, which may soften the wire.

Defective soldering

Overheating of wrought wires during soldering can lead to diffusion between solder and wire, and to recrystallization and grain growth. Microstructural changes, surface pitting, and internal porosity all result in a weak joint. Microstructural changes result from prolonged heating or overheating.

When the solder does not flow properly, resulting in an incomplete joint, it is usually due to one or more of the following:

1. The parts were too cool when the solder was applied.

2. If the parts were not at similar temperatures, the solder would flow over the hotter part leaving an incomplete joint.

3. Flux was insufficient to cover the joint.

4. Contamination was present due to improper cleaning, poor placement of flux, sulfur released from overheated investment, oxidation from an improperly adjusted
torch, or oxidation caused by removing the reducing portion of the flame from the joint before the solder flowed.

5. The gap distance was too small (< 0.1 mm) for the solder to penetrate between the parts. The joint may appear complete, but there will be a void at the center of the soldered joint that could result in premature failure.

6. If the gap distance was too large, the solder may not bridge the gap or, if it does, the diameter of the joint will be too small for adequate strength. Parts should be shaped to fit together with a proper gap distance separating them.

Properties of solder

Fusion temperature

The melting ranges of several gold solders are given in Table 18-2. The difference in melting range between high- and low-fineness solders is not great. Silver solders begin to melt between 600°C and 700°C (1,112°F and 1,292°F) with a range of about 10°C to 40°C.

<table>
<thead>
<tr>
<th>Solder no.</th>
<th>°C</th>
<th>°F</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>765-800</td>
<td>1,410-1,470</td>
</tr>
<tr>
<td>B</td>
<td>745-785</td>
<td>1,375-1,445</td>
</tr>
<tr>
<td>C</td>
<td>750-805</td>
<td>1,385-1,480</td>
</tr>
<tr>
<td>D</td>
<td>755-835</td>
<td>1,390-1,534</td>
</tr>
<tr>
<td>F</td>
<td>745-870</td>
<td>1,375-1,597</td>
</tr>
</tbody>
</table>

Table 18-3 Tensile properties of gold solders*

<table>
<thead>
<tr>
<th>Solder no.</th>
<th>Heat treatment</th>
<th>Proportional limit kg/cm² psi</th>
<th>Tensile strength MPa psi Elongation (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Soft</td>
<td>2,100 30,000</td>
<td>302 44,000 9</td>
</tr>
<tr>
<td></td>
<td>Hard</td>
<td>5,400 77,000</td>
<td>632 92,000 &lt; 1</td>
</tr>
<tr>
<td>B</td>
<td>Soft</td>
<td>1,890 27,000</td>
<td>283 42,500 14</td>
</tr>
<tr>
<td></td>
<td>Hard</td>
<td>3,850 55,000</td>
<td>431 63,000 1</td>
</tr>
<tr>
<td>C</td>
<td>Soft</td>
<td>2,060 29,500</td>
<td>306 44,500 12</td>
</tr>
<tr>
<td></td>
<td>Hard</td>
<td>5,420 77,500</td>
<td>573 83,500 &lt; 1</td>
</tr>
<tr>
<td>D</td>
<td>Soft</td>
<td>1,680 24,000</td>
<td>247 36,000 7</td>
</tr>
<tr>
<td></td>
<td>Hard</td>
<td>4,300 61,500</td>
<td>480 70,000 &lt; 1</td>
</tr>
<tr>
<td>E</td>
<td>Soft</td>
<td>1,440 20,500</td>
<td>257 37,500 18</td>
</tr>
</tbody>
</table>
*From Coleman (1928).
\[a\] Composition given in Table 18-1.
\[b\] No appreciable age hardening.

Silver solders have mechanical properties comparable to gold solders in the soft condition.

Corrosion resistance

Since there is a composition difference between the solder and the parts, the joint is susceptible to galvanic corrosion.

Interfacial properties

The relationship of the interfacial energies between the metal parts, solder, and flux determines how well the solder will flow. The fluxes used in dentistry will remove surface oxides and protect the parts from further oxidation. Solders penetrate the flux and wet the metal surfaces beneath.

*Soldering of metal-ceramic fixed partial dentures*

Metal-ceramic fixed partial dentures can be soldered together prior to placement and firing of the ceramic (preceramic soldering) or after firing of the ceramic (postceramic soldering). The two methods require different procedures and materials.

In preceramic soldering, a high-fusing precious metal solder must be used to avoid melting or distortion during firing of the porcelain. Usually a single-orifice gas/oxygen torch is used for soldering. Except for use of a phosphate-bonded soldering investment, the procedure is similar to the previously described investment soldering.

A lower-fusing gold solder is used for postceramic soldering so as to avoid thermal damage to the porcelain or its shape. To avoid overheating the porcelain, the parts are oven soldered. Except for the oven heating, the procedure is the same as for investment soldering. Following oven drying and fluxing, the correct amount of solder is placed on the joint and the dried invested pieces are placed in a furnace heated to the upper fusion range of the solder.

Preceramic and postceramic soldering have been successfully used with base metal-ceramic fixed partial dentures. The main difference between soldering of gold and base metal alloys is that special fluoride fluxes must be used for the latter to dissolve the chromium oxides. However, not all fluoride fluxes work equally well. It is important to use the manufacturer's recommended fluxes for these alloys.

*Commercial soldering materials*

A number of companies supply gold ranging from 450 to 800 in fineness (eg, Degussa, Ivoclar Williams, Jelenko, Ney). In addition, most companies carry white
gold solders for soldering white gold. Some companies also have special gold solders (eg, Balance Line Solder, Ney) made to match the colors of their popular gold alloys.

High-Fusing Ceramic Gold Solder (Ivoclar Williams); SMG 1, 2, 3, or YW (Ney), Jelenko O and Olympia Presolder (Jelenko), and WPG Solder (Degussa) are examples of preceramic solders. While manufacturers of preceramic solders also make postceramic solders, the lower-fusing gold solders of the previous paragraph can be used for postceramic soldering.

Soldering Flux (Jelenko), which comes in a paste or powder, and Solder Flux (Ney) are examples of fluxes for soldering of gold alloys. A liquid flux, "T" Flux (Degussa), is recommended for both postceramic and conventional soldering of precious metal alloys.

Some soldering investments (ie, Soldering Investment, Whip Mix) can be used for both high-temperature alloys and conventional low-temperature gold alloys. Chromium-containing orthodontic wires are soldered with special silver wires and with fluxes containing potassium fluoride, which dissolves chromium oxide. These are available from orthodontic wire suppliers.

**Welding**

**Introduction**

Bonding will occur between two metallic surfaces placed in contact if they are free of surface films (including oxides and films of adsorbed gases) and surface roughness. The three methods of welding used in dentistry achieve metal-to-metal contact differently.

**Spot welding**

In spot welding, the two clean metal surfaces to be welded are placed together under pressure (Fig 18-1). Metal-to-metal contact is obtained by passing a current through the joint to cause interfacial melting. If a pulse of sufficient voltage and duration is applied by means of copper electrodes, melting will begin at the interface between the parts and spread outward to form a weld.
Adjustable parameters are the magnitude and duration of the pulse, electrode size and shape, and pressure applied at the electrodes. Typical pulse parameters are 2 to 6 V for 1/25 to 1/50 of a second, which generally results in an applied current of 250 to 750 A, depending on the metal, size and shape of the work, electrodes, and pressure.

The equation

$$\text{Heat} = I^2R$$

gives the amount of heat in watts generated per second by a current of $I$ amperes passing through a structure having a resistance of $R$ ohms. The resistance of the joint is much higher than that of the rest of the parts because of the small contact area at the joint and the surface films present. Because the current is constant, more heat will be generated at the contact areas than in the interior parts. Therefore, the metal will melt first at the contact points.

Liquid contact is established because of the applied pressure and the expansion upon melting. Molten areas spread because the resistance of the metal in its liquid state is greater than in its solid state. Similar conditions exist at the electrode interface, but a higher-energy pulse would be required to cause melting because of the low resistance and high thermal conductivity of copper.
Small welds are generally considered better, because bonding is achieved with a minimum of change in the original grain structure.

Spot (resistance) welding is used to join flat structures, such as orthodontic bands and brackets, and to join orthodontic wires. The work is pressed together between two copper electrodes and an electrical pulse is applied. The magnitude of the pulse depends on the metals and their size and shape at the welding point, and on the size of the electrodes. Typical values for the pulse are 2 to 6 V for 1/25 to 1/50 seconds at 250 to 750 A.

**Pressure welding**

If two metal parts are placed together and a sufficiently large force is applied perpendicular to the surface, pressure welding occurs. Pure gold is extremely malleable and, in the form of thin foils, can be pressure welded by hand. Pure gold has no surface oxides but adsorbed gases prevent metal-to-metal contact. The applied force must be sufficiently large to produce permanent distortions parallel to the surface so as to expose film-free metal. The force must be applied rapidly so the exposed surfaces can be compressed together before surface gases adsorb. In pressure welding, the problems of surface roughness are overcome by compressive forces.

Gold foil, which is extremely thin and clean, may be welded under the pressure of dental instruments known as gold foil condensers to form gold foil restorations. Although rarely used today, gold foil restorations were popular during the nineteenth century. The absence of an oxide or contamination on the surface of pure gold allows room temperature welding.

**Laser welding**

A laser generates a coherent, high-intensity pulse of light that can be focused. By selecting the duration and intensity of the pulse, metals can be melted in a small region without extensive microstructural damage to surrounding areas. In laser welding, the beam is focused at the joint to melt the opposing surfaces. Due to expansion from the locally high temperature and the change of state, the two liquid surfaces contact and form a weld on solidification.

The intensity and duration of the laser pulse is such that a weld can be achieved before much heat is conducted away. This means that there is very little heating of the total appliance, except at the point of application. Consequently, the procedures can be performed on the master cast.

**Properties of welds**

**Strength**

In engineering applications, spot and pressure welds have strengths comparable to other forms of joining metals, such as soldering or arc welding. Laser welds are comparable to soldered joints between cast structures. Spot welding of work-hardened structures, such as bands and wires, destroys the grain structure and softens the metal
at and around the weld.

Corrosion resistance

In general, welds are more susceptible to corrosion than are the metals surrounding them. Spot welding in dentistry has been confined to provisional appliances, where the results have been satisfactory. Pressure-welded gold foil restorations are not subject to corrosion in the oral environment.

Electroplating

Introduction

Dies formed by electroplating are used in the construction of ceramic and PFM restorations particularly for full-mouth reconstruction. The chances for abrasion of the die surface are higher for construction of appliances involving ceramics than for all-metal ones. Consequently, a more durable die is desired when ceramic restorations are involved. Metal-plated dies are more abrasion resistant than stone dies and are therefore often used for making PFM restorations. Metal-plated dies are also utilized in the construction of platinum-bonded alumina crowns.

There is a relationship between electroplating and galvanic corrosion. In galvanic corrosion, the anode is corroded by oxidation and ionic dissolution. Under proper conditions—dependent on electrolyte, composition, potential difference between electrodes, and the metals involved—the dissolved ions from the anode can plate onto the cathode.

Plating solutions vary in their ability to plate concave surfaces, such as tooth surfaces of impressions. This plating ability is referred to as throwing power. Considering the size, depth, and shape of impressions of teeth, solutions with considerable throwing power are required for dental applications.

Plating manipulation

The basic procedure for construction of electroformed dies, dies formed by electroplating, is to plate an impression with an appropriate metal, which is then poured in stone or acrylic. The impression is removed, leaving the metal firmly attached to the die material. Impressions from compound or elastomeric materials can be electroplated. Generally, compound impressions are plated with copper, whereas silver is used for elastomeric impressions. Nickel is also used as plating metal for impressions.

A copper wire is threaded through nonvital regions of the impression to establish electrical contact with a direct current power supply and the inner surface of the impression. The impression is made electrically conductive by a process called metallizing, whereby a camel's-hair brush is used to burnish a fine powder of metal onto the impression. Wax is applied over areas that need not be plated.

The proper plating solution containing the same metal ion as the plating electrode is chosen. The impression is made cathodic to the plating anode by connection to a
direct current power supply. At the proper voltage—dependent on the size of the impression and the materials involved—the process requires about 10 to 15 hours. Approximately 1 mm of metal is deposited on the impression. The impression is rinsed thoroughly in running water and other appropriate solutions to neutralize the plating electrolytes. The impression is poured in stone or acrylic, then removed as usual, leaving the metal attached to the stone or acrylic.

There are potential inaccuracies associated with electroformed dies that do not occur for properly made stone dies. The plating process is of sufficient duration that the impression can distort. The plated metal layer can also slightly distort the impression's surface. Potential inaccuracies may occur if self-curing acrylic is used instead of stone. Curing shrinkage of the acrylic may distort the metal.

Platinum-bonded alumina crowns

Jacket crowns are generally made by firing porcelain to a thin platinum coping that has been adapted to a die. This coping is not bonded to the porcelain and must be removed before cementation, which results in a thick cement layer between crown and tooth. Electroplating the platinum coping with tin has been found to produce a surface conducive to bonding. The electroplated coping is oxidized in a furnace, producing a tin oxide layer to which the alumina core will adhere. This process allows the platinum coping to be left in place during cementation and likely improves the strength of the crown.

Electroformed gold copings

Nearly pure gold copings for PFM restorations can be made by an electroplating process similar to that used to make electroplated dies. In this case, however, an accurate die is metallized and plated with gold to a thickness of 0.2 to 0.3 mm. The die material is then removed from the gold by grinding and dissolving the remainder in an acid solution. The gold is trimmed and fitted to the master die to give an accurate coping. Gold particles and organic binders are then applied to the outer surface of the coping before it is electroplated with tin so as to obtain a good bond with dental porcelains that have fusion temperatures below 927°C. Thin, accurately-fitting gold copings can be made by this process. These copings are too weak to support porcelain fixed partial dentures.

Commercial products

Silver plating solutions are difficult to obtain. They contain silver cyanide and, therefore, should only be used under a hood. Silver plating has been largely discontinued, but copper plating is still used. Apparatus and materials for making electroformed gold copings are available (Captek).

Glossary

antiflux Material placed on the work area before the flux to confine the flow of solder.

electroplating The formation of a metallic surface coating on another by an
electrochemical solution.

**fluoride flux** Fluoride-containing flux for chromium alloys.

**flux** A substance that promotes the flow of solder over the metal parts by cleaning the surfaces and removing oxides.

**gap distance** Space between parts to be soldered, which will be filled with solder.

**laser welding** A form of welding in which the heat for melting the metal is supplied by a focused beam of light generated by a laser.

**metallizing** Coating an impression material with a powdered metal to make it electrically conductive.

**pressure welding** A form of welding in which the weld is made by pressure.

**solder** Metal or alloy melted to unite adjacent, less fusible metal parts.

**soldering investments** Similar in composition to quartz casting investments, which may be used for soldering.

**spot welding** A form of welding in which the heat for melting the metal is generated by the flow of electricity through the parts to be welded.

**sticky wax** Brittle wax-containing resin used to hold metal parts during investment soldering. It fractures rather than deforms under stress.

**throwing power** A measure of the uniformity in plating thickness of irregular surfaces.

**welding** The joining of metal surfaces directly by the application of heat or high compressive forces.

**Discussion Questions**

1. Why do fixed partial dentures often fail clinically at solder junctions?

2. How do the processes of soldering and welding differ on a basic level?

3. What precautions need to be taken when soldering or welding wrought metals?

4. What problems might be expected in the clinical use of pure electroplated gold PFM dentures? What evidence might be needed?

5. What are the consequences of using too much solder or making too large a spot weld when joining orthodontic wire?

**Study Questions**
1. What are the two main components of gold solder, besides gold, and why are they added?

2. What would be expected to happen if a porcelain-gold appliance were presoldered with an ordinary gold solder?

3. Why is a special fluoride flux required to solder alloys containing chromium?

4. What is an antiflux?

5. Give two examples of an antiflux.

6. To avoid melting gold parts, what fineness does the solder usually have relative to the parts?

7. What is the recommended minimum fineness for soldering permanent restorations?

8. Why is silver solder recommended for soldering stainless steel wires?

9. Why is cleaning the parts important to successful soldering?

10. When is investment soldering recommended?

11. What would happen with investment soldering if the sticky wax fractured but the soldering operations were continued?

12. What is the purpose of preheating the investment, and what will happen if it is overheated?

13. How does one determine if all parts are at the proper temperature for soldering?

14. Why is it important to have uniform heating at the joint?

15. What are some of the major differences between investment soldering of relatively large parts and free-hand soldering of wires?

16. What microstructural changes are expected if wrought wires are overheated?

17. What difficulties might occur if soldering were performed in a dark room?

18. A soldered joint of an appliance that failed in service had the following defects: surface porosity, flux incorporated into the solder, and a large pore at the center of the joint. What were the likely causes?

19. If a joint becomes oxidized during soldering, what will happen?

20. What are causes of oxidation?
21. What errors in soldering procedure could lead to a poorly fitting appliance?

22. Why is flux important to successful soldering?

23. Referring to Table 18-3, what property of solders is seriously reduced by age hardening, and how is this avoided in the investment soldering procedure?

24. Why would it be impractical to spot weld flat copper structures together using orthodontic spot welders?

25. Why is it unlikely that thin foils of copper could be pressure welded by gold foil condensers?

26. Why can laser welding be accomplished on the master cast?

27. Can pressure welding be applied to nongold alloys?

28. Why might an organic film prevent welding during spot welding of orthodontic appliances?

29. What is the main application of electroplating in dentistry?

30. What are the advantages and Disadvantages of electroplated dies?

31. Why is the throwing power of a plating solution important?

**Recommended Reading**


Milner DR, Apps RI (1968). Introduction in Welding and Brazing. New York:


Chapter 19. High-Temperature Investments

Introduction

The main use of high-temperature investments is for casting dental alloys that need to be heated to casting temperatures in excess of about 1,200°C to 1,300°C (2,192°F to 2,372°F).* Alloys that fit this description include many high-gold and palladium-based alloys used for the fabrication of porcelain-veneered fixed partial dentures and those dental alloys based on nickel, cobalt, or titanium. These investments are also used for creating fixtures that hold sections of prostheses in proper apposition while joining them together during soldering, brazing, or welding operations.

These investments are not gypsum based, as are those used for the casting of alloys that have casting temperatures less than approximately 1,200°C (2,192°F), but instead have a variety of basic formulations that are better suited to withstand exposure to high temperatures. The predominate compositions, used for the casting of high-temperature alloys and for creating veneering dies, are based on the use of a phosphate binder. Silicate-bound systems are also used (Kondic, 1960), but because of their difficult handling, their use tends to be restricted to the casting of cobalt-chromium alloys at temperatures above 1,425°C (2,600°F).

Either entirely new systems or modifications of the aforementioned ones have been needed for the casting of titanium-based alloys. These alloys require special investments because they are cast from higher temperatures than the others and are also highly reactive. They can easily degrade the more conventional molds at elevated temperatures and, as a result, produce poor surface finishes. Further, they can be contaminated by degradation of the investment components, such as silica and carbon, and by oxygen and nitrogen present in the atmosphere. These contaminations serve to harden and embrittle the surface of titanium castings. Because of the developments of investments for the casting of titanium alloys for dental applications, a separate section is devoted to this subject.

*Casting temperature is approximately 65°C to 150°C (150°F to 300°F) above the top of the melting range for most dental alloys.

Phosphate-Bonded Investments

Applications

Casting of alloys

Phosphate-bonded investments have been used for many years in dentistry to make molds into which high-melting-temperature dental alloys were cast. There are two types of casting investments. Type I traditionally has been employed for the casting of inlays, crowns, and other restorations, especially for alloys based on gold, platinum, palladium, cobalt-chromium, and nickel-chromium, to which porcelain is fused in the construction of esthetic fixed restorations. Type II is used for the casting of removable partial dentures. These applications are still in use, but the number of alloy applications has grown. Some variations are used for cast titanium alloys, discussed later in this chapter.
For the burnout process of the conventional Type I investment for inlays and crowns, it is recommended that the mold be set in a furnace at room temperature, 1 hour after trituration, heated to 800°C (1,471°F) over 1 to 2 hours (depending on the size of the mold) and held at 800°C for 30 minutes before casting. Recently, newer investments that allow insertion of the set investment directly into a furnace at the burnout temperature (800°C) have been developed. These investments are inserted 30 minutes after the powder has been mixed with the liquid.

Soldering and porcelain veneering

Another traditional use of phosphate-bonded investments is to make "soldering" fixtures that hold prosthetic components in alignment while they are being joined with solders, brazing alloys, or welding alloys. Since around 1980 modifications have been used to make refractory dies for the fabrication of custom veneer facings from dental porcelains.

Composition

These investments are available as two-component systems that react to form a solid when mixed together (Takahashi et al, 1990). One component consists of a powder, and the other is an aqueous solution stabilized with colloidal silica. The powder has a variety of ingredients. There are particles of refractory materials such as quartz, cristobalite, or a mixture of the two, in a mass fraction of about 80%, to control thermal expansion and thermal stresses related to thermal phase transformations of cristobalite, and, along with glasses and other metal oxides, to provide bulk and help control the surface finish of a casting. There is also powdered ammonium dihydrogen phosphate, NH₄H₂PO₄ (provided in excess), which, in reaction with water in the presence of calcined magnesium oxide (MgO) powder (another ingredient), provides for the binding of the particles at ambient temperatures. The liquid provides the water needed for the room-temperature setting reaction. The setting reaction of the NH₄H₂PO₄ with calcined magnesium oxide in the presence of water is:

\[
\text{NH}_4\text{H}_2\text{PO}_4 + \text{MgO} + 5\text{H}_2\text{O} \rightarrow \text{NH}_4\text{MgP}_4\cdot 6\text{H}_2\text{O}
\]

After a mixed slurry sets to form a mold, it is fired to burn out consumable patterns (mostly waxes, but some polymers are used also).

Properties

Important properties of phosphate-bonded investments are given in Table 19-1.
Table 19-1 Properties of phosphate-bonded investments used for high-temperature operations with dental materials (except titanium alloys)

**Casting investments**

<table>
<thead>
<tr>
<th>Property</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Compressive strength</td>
<td>2.5 MPa minimum, Type I (for inlays, crowns, etc)* 3 MPa minimum, Type II (for removable appliances)*</td>
</tr>
<tr>
<td>Setting expansion (linear)</td>
<td>Within 15% of manufacturer's stated value* With use of full-strength liquid about 0.4% can be attained with some investments; when a hygroscopic technique is used, about another 0.6% to 0.8% can be realized*</td>
</tr>
<tr>
<td>Thermal expansion</td>
<td>Within 15% of manufacturer's stated value* About 0.8% can be attained with a 50:50 mixture of liquid and water; about 1% to 1.2% can be attained with the use of undiluted liquid</td>
</tr>
<tr>
<td>Modulus of rupture</td>
<td>0.1 to 0.5 MPa (14.4 to 72.5 psi); green, about 0.8 MPa (116 psi), as fired</td>
</tr>
</tbody>
</table>

**Refractory die stones**

<table>
<thead>
<tr>
<th>Property</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Compressive strength</td>
<td>3 MPa minimum</td>
</tr>
<tr>
<td>Setting expansion (linear)</td>
<td>Within 30% of manufacturer's stated value*</td>
</tr>
<tr>
<td>Thermal expansion</td>
<td>Within 15% of manufacturer's stated value* Manufacturers state that their materials are designed to be compatible with their porcelain at temperatures in the vicinity of the glass-transition temperature. Values of about 12 to 13 \times 10^{-6}/^\circ\text{C} would be expected.</td>
</tr>
</tbody>
</table>

*According to ISO #9694 (December 19, 1996).

*Approximate values attainable with some of the commercial brands.

High-temperature mold strength is achieved by the formation of complex silicophosphates from the reaction of some of the silica with the excess dihydrogen phosphate. Liquid that is supplied with the investment may be used either full strength or diluted with water to provide some degree of control over the setting and thermal expansions (Fig 19-1).
Although the basic binding reaction is the same for all of the phosphate-bonded investments, there are some important differences in properties due to composition. Those used for the casting of high-temperature alloys and for making dies used in the fabrication of porcelain veneers contain quartz and cristobalite to achieve expansion, which compensates for the shrinkage (contraction) of the cast alloy during cooling from elevated temperatures. Soldering investments, however, do not require especially fine powders and are designed without high-expansion fillers. The latter is to keep parts that are to be joined from shifting while they and the surrounding investment are heated to the joining temperature. Graphite is found in some investments to render them more permeable after burnout and/or to provide a reducing atmosphere.

*Advantages and Disadvantages*

There are several advantages associated with phosphate-bonded investments. They have both high **green strength** and **fired strength**. This makes them easy to handle without breaking before they are placed in a furnace for the wax burnout process and strong enough afterward to withstand the impact and pressure of centrifugally cast molten alloy. They can also provide setting and thermal expansions high enough to compensate for the thermal contraction of cast-metal prostheses or porcelain veneers during cooling. Finally, they have the ability to withstand the burnout process with temperatures that reach 900°C (1,650°F), and they can withstand temperatures up to 1,000°C (1,831°F) for short periods of time (useful for fabricating porcelain veneers or performing metal-joining operations).

These investments are at a disadvantage when used with higher-melting alloys, those with casting temperatures higher than about 1,375°C (2,500°F). These temperatures, coupled with high mold temperatures, result in mold breakdown and rougher surfaces.
on castings. The high strength of these investments, although an advantage during casting, can make divesting (removal of the casting from the investment) a difficult and tedious task without the use of a simple tool, such as a press, to force the investment out of the metal casting ring (when a casting ring is used). Further, when higher expansion is required, more of the silica liquid is used with the result that a more dense and less porous mold is produced. This can result in incomplete castings if a release for trapped gases is not provided.

A disadvantage when the powder is supplied in bulk form—rather than in sealed, premeasured packages—is that it can react over time with moisture in the air and result in either a lower expansion during setting or a loss of the ability to set to a strong mass. The temperature at which the triturated mixture is held markedly affects setting expansion.

**Ethyl Silicate-Bonded Investments**

**Applications**

Ethyl silicate-bonded investments comprise the second type of investment used in dentistry. These investments have been used since the early 1930s to make molds for the casting of removable partial dentures of cobalt-chromium alloys. This continues to be their primary use, although they are occasionally used for the casting of nickel-based alloys.

**Composition**

Ethyl silicate-bonded investments (ESBIs) rely on binding by the formation of a gel from silicic acid, which is derived from the hydrolysis of tetraethyl orthosilicate, Si(OC₂H₅)₄. It is from this derivation that these investments received their name. As with the phosphate-bonded investments, ESBIs are supplied as a powder that requires mixing with a liquid to bind the mixed mass via a setting reaction at room temperature. The powder consists of refractory particles of silicas and glasses in various forms along with calcined magnesium oxide and some other refractory oxides in minor amounts.

The liquid used for the setting reaction may be supplied as a stabilized alcohol solution of silica gel, or it may be formed from two liquids that are supplied. When the system uses two liquids, one is ethyl silicate and the other may be an acidified solution of denatured ethyl alcohol. Three component systems also exist and may consist of stabilized colloidal silica, ethyl silicate, and denatured ethyl alcohol. Regardless of the number of liquids provided, the active species is the hydrated silica that is formed by the reaction of ethyl silicate with water in the presence of the acid (or a base) according to the reaction:

\[ \text{Si(OC}_2\text{H}_5)_4 + 4\text{H}_2\text{O acid/base} \rightarrow \text{SiO}_4 + 4\text{C}_2\text{H}_5\text{OH} \]

Binding is by the reaction of the hydrated silica in the presence of MgO to form a gel according to the reaction:

\[ n\text{Si(OH)}_4 + \text{MgO} \rightarrow \text{MgO[Si(OH)}_4]_n \]
A wetting agent is added to at least one of the liquid components to reduce the formation of gaseous bubbles on the surface of a pattern.

Properties

The most important property of the ESBIs is their ability to withstand higher temperatures than the phosphate-bonded investments and to achieve sufficient expansion to compensate for the cooling shrinkage from the higher solidification temperatures of the higher-casting-temperature alloys with which they are used (ie, cobalt-chromium alloys). During the vibration of the investment slurry into a casting ring around a pattern, a separation occurs between the finer and coarser particles. The finer particles of the investment rise to the top of the mold where the slurry acquires more liquid. As a result, the top of the mold (near the bottom of the pattern, opposite the sprue end) is prone to cracking due to greater drying shrinkage from the evaporation of the ethyl alcohol, which results from the formation of the silica gel. These cracks must be removed prior to the firing process. Otherwise, when the mold is heated to burn out a pattern and achieve thermally induced expansion, the cracks will grow and result in faulty castings. To overcome this problem, a sufficient header of investment is provided to allow for removal of the cracked portion by grinding. The remaining investment itself undergoes virtually no solidification shrinkage and is free of critical cracks. The expansion of the investment is all due to thermal expansion. Thus, distortion of patterns (such as those that can occur with setting expansion) is minimized, and these investments are well suited for producing large, precise castings.

Because of the low dimensional changes during setting (Table 19-2), models made from these investments may be articulated directly against gypsum models of opposing dentition. However, the green strength of these investments is low, and refractory models are best handled by reinforcing them with a resin dip. A fine-grained surface coat is sometimes applied to the reinforced model and to the pattern on the model in order to achieve a superior surface finish on the cast appliance.

Table 19-2 Properties of ethyl silicate-bonded investments used for high-temperature operations with dental materials

<table>
<thead>
<tr>
<th>Casting investments</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Compressive strength</td>
<td>1.5 MPa, minimum*</td>
</tr>
<tr>
<td>Setting expansion (linear)</td>
<td>There is no requirement for setting expansion. Setting contractions of 0% to 0.4% have been reported. This value will depend on the method of measurement. One manufacturer reports virtually no (0% to 0.05%) contraction.</td>
</tr>
<tr>
<td>Thermal expansion (linear)</td>
<td>Within 15% of manufacturer's stated value*. About 1.5% to 1.8% can be attained between room temperature and 1,000°C to 1,177°C (1,800°F to 2,150°F).</td>
</tr>
</tbody>
</table>

* According to ISO #11246 (June 13, 1996).

a Approximately values attainable with some of the commercial brands.
Advantages and Disadvantages

Ethyl silicate-bonded investments offer the ability to cast high-temperature cobalt-chromium and nickel-chromium alloys, and attain good surface finishes, low distortion, and high thermal expansion (good fit). They are less dense (more permeable) than the phosphate-bonded investments, and thin sections with fine detail can be reproduced. Their low fired strength makes divesture easier than with phosphate-bonded investments.

Their Disadvantages lie primarily in the added processing attention (resin-model reinforced dies, pattern coats) and the extra precaution needed in handling the low-strength fired molds. The low strength and high thermal expansion require a more precise burnout process and firing schedule to avoid cracking and, hence, destruction of a mold.

Other Systems

Applications

Newer investments have been aimed at the casting of titanium and titanium-based alloys, although the possibility of applications for the casting of other alloys or for refractory pattern coats becomes apparent from some of the properties shown in the following section.

Efforts in Japan, Europe, and the United States to cast dental appliances from titanium or titanium-based alloys have shown that the conventional phosphate-bonded investments and ethyl silicate-based investments are deficient for that purpose. This is because molten titanium is highly reactive with oxygen and is capable of reducing some of the oxides commonly found in those investments. Titanium can also dissolve residual oxygen, nitrogen, and carbon from the investment; these elements can harden and embrittle titanium in the solid state. As a result, either modifications of existing refractory formulations and binders or new refractory formulations and binder systems are required.

Composition

A variety of investment formulations for the casting of titanium have been developed over the past several years (Table 19-3) (Togaya, 1993; Miyazaki and Tamaki, 1993). These investments might be classified as phosphate-bonded, ethyl silicate-bonded, and "cemented" according to the source of the binder. Many kinds of refractories such as silica (SiO₂), alumina (Al₂O₃), magnesia (MgO), and zirconia (ZrO₂) have been used.
Table 19-3: Titanium investments available in Japan and Germany

<table>
<thead>
<tr>
<th>Mold-casting temperature (°C)</th>
<th>Binder</th>
<th>Refractory</th>
<th>Burnout temperature (°C)</th>
<th>Country</th>
</tr>
</thead>
<tbody>
<tr>
<td>RT*</td>
<td>MgO, phosphate</td>
<td>Al₂O₃, ZrO₂</td>
<td>900</td>
<td>Japan</td>
</tr>
<tr>
<td>RT*</td>
<td>Al₂O₃, ZrO₂</td>
<td>1,100</td>
<td>Japan</td>
<td></td>
</tr>
<tr>
<td>800</td>
<td>Al₂O₃, ZrO₂</td>
<td>1,150-1,200</td>
<td>Japan</td>
<td></td>
</tr>
<tr>
<td>RT*</td>
<td>Al₂O₃, SiO₂</td>
<td>900-1,000</td>
<td>Japan</td>
<td></td>
</tr>
<tr>
<td>200</td>
<td>Al₂O₃, SiO₂</td>
<td>1,100</td>
<td>Both</td>
<td></td>
</tr>
<tr>
<td>500</td>
<td>Al₂O₃, SiO₂</td>
<td>950</td>
<td>Both</td>
<td></td>
</tr>
<tr>
<td>100</td>
<td>ZrSiO₄, SiO₂</td>
<td>1,100</td>
<td>Japan</td>
<td></td>
</tr>
<tr>
<td>RT*</td>
<td>Al₂O₃, LiAlSi₂O₆</td>
<td>1,050</td>
<td>Japan</td>
<td></td>
</tr>
<tr>
<td>600-700 Ethyl silicate</td>
<td>MgO, Al₂O₃</td>
<td>900</td>
<td>Japan</td>
<td></td>
</tr>
<tr>
<td>600-700 Ethyl silicate</td>
<td>MgO, SiO₂, Al₂O₃</td>
<td>900</td>
<td>Japan</td>
<td></td>
</tr>
<tr>
<td>100 Aluminous cement</td>
<td>MgO, Zr²</td>
<td>850</td>
<td>Japan</td>
<td></td>
</tr>
<tr>
<td>100 Aluminous cement</td>
<td>MgO, Zr²</td>
<td>850</td>
<td>Japan</td>
<td></td>
</tr>
<tr>
<td>600-700 Ethyl silicate</td>
<td>MgO, Al₂O₃</td>
<td>900</td>
<td>Japan</td>
<td></td>
</tr>
</tbody>
</table>

*RT = room temperature.
²Addition for attaining expansion by volume change through oxidation.

Properties

Standards do not exist, nor are they in development, for investments for the casting of titanium and titanium alloys in dentistry. It is reasonable to expect that those mechanical properties being contemplated for the conventional phosphate and ethyl silicate-based binders would apply here as well.

New refractory compositions and binders have been investigated for the casting of titanium prostheses. The objectives have been to reduce breakdown of the investment and the contamination of titanium. One approach to reducing the reaction with investment is to employ molds that have been expanded by the burnout process and then cooled back to near ambient temperatures prior to the casting process. This reduces the time that the alloy is in contact with the mold at elevated temperatures, and the overall reactivity is reduced. However, lowering of the mold temperature requires that either nonreversible expanders, such as metals that expand by oxidation at the elevated temperatures, be used or that the temperature of the mold be kept just above the temperature where a reversal of expansion due to crystalline phase changes takes place.

To avoid contamination of titanium by oxygen through the reduction of refractory oxides of the investment, refractory materials that are less easily reduced by titanium should be used. The Gibbs free energy of formation (FEF) per mole of oxygen (Chase et al, 1985) for titanium oxide (TiO) at 1,727°C is -716 kJ/mol of oxygen, and
that for titanium dioxide (TiO$_2$) at 1,727°C (3,138°F) is -580 kJ/mol of oxygen. The corresponding FEFs of SiO$_2$ in the forms of cristobalite and quartz are -550 and -549 kJ, respectively. From this, it is clear that titanium may be expected to be oxidized by SiO$_2$, which is reduced.

Some modifications of phosphate-bonded investments have been explored for the purpose of rendering them more compatible with molten titanium metals. One investment consisting of a phosphate binder, magnesia, and quartz was developed under the hypothesis that quartz would not be as reactive as silica (Takahashi, 1993). On the basis of FEFs, there is little advantage in the use of one form of SiO$_2$ over another, and any advantages of one form over another would have to lie elsewhere (perhaps their decomposition kinetics are sufficiently different). This investment was recommended for use as a room-temperature mold, ostensibly to reduce reaction with titanium. However, contamination of castings by reaction with the investment was still encountered, as would be expected from the previous discussion on FEFs.

To make use of the setting expansion of a phosphate binder, alumina (FEF = -687 kJ/mol of oxygen at 1,727°C) and magnesia (FEF = -640 kJ/mol of oxygen at 1,727°C), both of which have good heat resistance, can be used as refractories; however, the thermal expansions of these oxides is low. If either of their powders is mixed with colloidal silica to raise the expansion, some contamination from the silica again becomes a problem.

To achieve expansion without the use of reactive powders, a phosphate investment that contains both magnesia and alumina as refractories was developed. This investment can attain large expansion by the spinel reaction of alumina and magnesia (MgO + Al$_2$O$_3$ → MgO - Al$_2$O$_3$) when it is burned out at 1,150°C to 1,200°C. The spinel is, of course, also highly refractory. The spinel-forming temperature can also be reduced by mixing with magnesia acetate. Another approach to obtaining the needed expansion is through the use of spodumen (Li$_2$O - Al$_2$O$_3$ - SiO$_2$) (Okuda et al, 1991). Spodumen expands irreversibly upon heating through the temperature range of 900°C to 1100°C. Figure 19-2 shows the thermal expansion curves of these titanium instruments.
Reaction of ethyl silicate-bonded investments with liquid titanium has been reported to be somewhat less than that of phosphate-bonded investments; this is most likely due to the use of highly refractory oxides in the powder (see Table 19-3). Regardless, these investments require a more complex procedure for their use.

A more recent development is an investment using magnesia bonded by an aluminous cement (CaO - Al₂O₃), which contains a mass fraction of 5% zirconium powder (Togaya et al, 1985). The aluminous cement serves as a binder for the magnesia refractory, and it sets by mixing with water. Oxidation of the zirconium powder to zirconia during the burnout process provides irreversible expansion to compensate for shrinkage of the casting during cooling from the solidification temperature. The zirconia formed is highly stable; it has an FEF of -728 kJ/mol of oxygen, and it should not contaminate the titanium. Titanium castings from this investment were reported to have smooth surfaces free of contamination from mold reaction.

**Glossary**

**articulate** To join the maxillary and mandibular dentition or a replica of it.

**calcined** Rendered a powdery, friable, dry substance by the application of heat.

**colloidal** The state of finely divided particles that are in suspension in a liquid. The particles acquire special properties (electrical, gelatinous, etc) as a result of their fine size and interaction with each other and the liquid. Colloidal silica consists of fine particles of hydrated silicas that can form a gel and bind larger particles of
noncolloidal nature.

**fired strength** The strength of an investment following a burnout process. This strength is often measured by resistance to breakage under three-point loading or bending.

**fixture** A device that is firmly fastened in place in a mechanical sense, often used to secure other devices.

**Gibbs free energy of formation (FEF)** Gibbs free energy is the thermodynamic function $\Delta G = \Delta H - T\Delta S$, where $H$ is enthalpy, $T$ is absolute temperature, and $S$ is entropy. Also called *free energy*. Free energy of formation (FEF) is the free energy change for formation of a particular compound.

**green strength** The prefired strength of an investment, acquired through some chemical reaction at or near room temperature. (See *fired strength*.)

**hydrolysis** Decomposition of a compound by water, or the interaction of positive and negative ions of a salt with hydroxyl and hydrogen ions to form an acid and a base.

**hygroscopic** Pertaining to the absorption of water.

**investment** A heat-resistant material used to form a mold around a wax pattern.

**mold** A cavity into which molten metal is cast.

**refractory** A heat-resistant material.

**ring** A thin-walled structure used to contain an investment slurry before it sets; also provides support for set or fired investments that have poor strength.

**setting expansion** The volumetric or linear increase in physical dimensions of an investment, caused by the chemical reactions that occur during hardening to a rigid structure.

**slurry** A thin mixture of liquid and powder.

**surface coat** The first coating that is applied to a wax pattern or the surface of the mold that comes into direct contact with a cast metal.

**thermal expansion** The increase in dimension of a set investment due to temperature increase during burnout.

**Discussion Questions**

1. Which developments in clinical dentistry have led to a switch from the use of gypsum to more refractory materials?

2. What are the matrix compositions of the two major high-heat investments?
3. How can additional expansion be obtained with magnesia and ethyl silicate investments?

4. Which of the two high-heat investments gives more accurate models?

**Study Questions**

*(See Appendix E for answers.)*

1. Given a multicomponent system that includes ethyl silicate as a reacting component, how can the reaction time for hydrolysis be altered?

2. Cooling hydrated ethyl silicate to 4.4°C (40°F) extends its shelf life. *(a)* Would the shelf life be extended by refrigeration to lower temperatures? *(b)* To what extent may the liquid be cooled without encountering adverse effects?

3. Why is denatured ethyl alcohol used rather than 95% ethyl alcohol as a component of an ethyl silicate system?

4. List advantages and Disadvantages of phosphate-bonded investments.

5. List advantages and Disadvantages of ethyl silicate-bonded investments.

6. For the casting of nonprecious fixed partial denture alloys, which technique would be recommended with a phosphate-bonded investment?

7. What is the investment of choice for casting *(a)* ceramic gold alloy; *(b)* cobalt-chromium partial denture alloy; *(c)* nickel-based fixed partial denture alloy?

8. List the six approaches to altering the fit of a casting.

9. What might be an advantage of a mixed colloidal silica-ethyl silicate system?

10. A carbon-filled and a noncarbon-filled phosphate investment are available to you for casting ceramic alloys. Would you use a gold alloy or a nonprecious alloy for casting? Why?

**Recommended Reading**


Chapter 20. Waxes

Introduction

Waxes have several applications in dentistry. They are used as patterns for inlays, crowns, pontics, and partial and full dentures. Waxes are very useful for bite registration and can be used to obtain impressions of edentulous areas. In addition, they have many applications for processing in all areas of restorative dentistry.

Composition

Waxes are organic polymers consisting of hydrocarbons and their derivatives (eg, esters and alcohols). The average molecular weight of a wax blend is about 400 to 4,000, which is low compared with structural acrylic polymers. Dental waxes are blends of ingredients, including natural waxes, synthetic waxes, natural resins, oils, fats, gums, and coloring agents.

Natural waxes are of mineral (petroleum oil), plant, insect, or animal origin. Paraffin wax is relatively soft with a low melting range (50°C to 70°C). It is a mineral wax obtained from refined crude oil and used in inlay and modeling waxes. Beeswax is brittle with an intermediate melting range (60°C [122°F] to 70°C [158°F]). It is an insect wax obtained from honeycombs and is added to many waxes because of its desirable flow properties at oral temperature. Carnauba wax is a plant wax obtained from carnauba palm trees and is hard and tough with a high melting range (65°C [149°F] to 90°C [194°F]). It is added to toughen paraffin wax and raise its melting range. Microcrystalline waxes have a high melting range (65°C to 90°C) and are added to modify the softening and melting ranges of wax blends. They also serve to reduce stresses that occur on cooling. Microcrystalline waxes are obtained from petroleum.

Synthetic waxes have specific melting points and are blended with natural waxes. Low-molecular-weight polyethylene is an example of a synthetic wax. Natural waxes vary more depending on their sources and need to be monitored more for properties than synthetic waxes, which are more uniform in composition.

Classification

Introduction

Dental waxes are classified according to their applications into the categories of pattern, processing, and impression waxes (Fig 20-1).
Pattern waxes

Pattern waxes include inlay, resin, casting, and base plate waxes. Inlay waxes (Kerr) are used to make inlay, crown, and pontic replicas for the lost wax casting technique. Type I inlay waxes are hard and used for the direct inlay technique. Type II inlay waxes are soft and used for preparing replicas on dies and models. In addition, inlay waxes are sometimes used for the attachment of miscellaneous parts.

Pattern resins are characterized by higher strength and resistance to flow than waxes, good dimensional stability, and burnout without residue. Full-crown patterns fabricated from pattern resins and inlay waxes have similar marginal discrepancies.

A pattern is fabricated by applying the resin in 3- to 5-mm layers and curing in a light chamber or with a handheld light-curing unit. Resin is completely eliminated from the mold before casting by heating at 690°C (1,273°F) for 45 minutes.

Casting waxes are used for thin sections of certain removable and fixed partial denture patterns. They are particularly convenient in the preparation of copings or clasps requiring uniformly thin regions.

Base plate wax is used in the construction of full denture patterns and for occlusal rims, although an occlusal rim wax is also available. Setup wax may be used instead of base plate wax to set denture teeth.

Fig 20-1 A classification of dental waxes.
The American National Standards Institute/American Dental Association has established a specification that includes three types of base plate wax: Type I is a soft base plate wax for veneers and contours. Type II is a medium-hardness base plate wax designed for temperate climates for patterns to be tried in the mouth. Type III is the hardest base plate wax and is for patterns to be tried in the mouth in tropical climates. The hardness is based on the amount of flow the wax shows at 45°C (113°F).

Base plate wax is also used as a mold for the construction of provisional fixed partial dentures and as a bite registration wax. It has some applications in orthodontics.

**Processing waxes**

Processing waxes include boxing, sticky, carding, blockout, white, and utility waxes. Boxing wax is used to form containers for pouring casts. It is also used to fabricate replacement pontics for provisional fixed partial dentures. Sticky Wax (Kerr) is used to join materials temporarily. Carding wax is used for attaching parts and in some soldering techniques. Blockout wax is used to fill voids and undercuts for removable partial denture fabrication. White wax is used for making patterns to simulate a veneer facing. Utility wax (Modern Materials) has miscellaneous applications for various laboratory procedures.

**Impression waxes**

Impression waxes (Bite Wax, Mizzy) exhibit high flow and distort on withdrawal from undercuts. Waxes used for denture impressions are limited to use in edentulous regions of the mouth. Corrective waxes are used as wax washes to record detail and displace selected regions of soft tissue in edentulous impressions. Bite waxes are used in certain prosthetic techniques; a typical use would be bite registration.

**Properties**

Waxes may consist of both crystalline and amorphous components, each with a distribution of molecular weights. Therefore, waxes melt over a range of 5°C to 30°C (41°F to 86°F) rather than at one temperature. Waxes have the highest coefficients of thermal expansion of any dental material. Resultant dimensional changes may produce poor-fitting castings if not balanced by compensating factors of mold expansion. The total wax shrinkage on cooling from liquid to solid at room temperature may be as great as 0.4% and consists of solidification shrinkages plus contraction on cooling to room temperature after solidification.

Flow is a measure of a wax's ability to deform under light forces and is analogous to creep. Flow increases with increasing temperature and force. At a temperature close to its softening range, a wax may flow under its own weight. In liquids, flow is measured by viscosity. In solids, flow is measured by the degree of plastic deformation over a fixed period of time. Type I direct inlay technique waxes* need to flow well to reproduce details of the cavity preparation. However, when the wax is cooled to oral temperature, flow must be minimized to reduce distortion when the pattern is removed.
American National Standards Institute/American Dental Association Specification no. 4 for dental inlay casting wax.

**Wax Distortion**

Waxes are partly elastic in behavior and tend to return to their original shape after deformation. A straight bar of wax bent into a horseshoe shape will slowly straighten out at room temperature (Fig 20-2). This is called the "memory effect." Residual stresses as a result of nonuniform heating also contribute to later distortion. There are four ways of minimizing pattern distortion. First of all, wax for the direct technique should be heated uniformly at 50°C (122°F) for 15 minutes before use. Next, the pattern should be invested quickly. The rigid walls of the set investment constrain the pattern and reduce distortion due to recovery and residual stress. Also, the pattern should be stored at a low temperature if not invested right away.

![Fig 20-2](image.png) The rod on the right was originally bent to the shape of the rod on the left. After floating in room-temperature water for 24 hours, the rod opened as a result of the memory effect.

Elastic recovery is slower at low temperatures; therefore, if immediate investing is impractical, storage in a refrigerator is preferred. If a pattern is refrigerated, however, it should be allowed to warm to room temperature before investing. Finally, it is essential that no wax residues are left in the mold after burnout in the lost wax process. Residues will result in poor castings because of inclusions or incomplete margins.

**Commercial Materials**

Inlay waxes are supplied in geometric and anatomic forms as well as in bulk. Casting waxes are supplied in sheets and rods and in bulk. Preformed patterns are also available for partial denture applications. Bite waxes are supplied in a variety of forms and shapes. Base plate wax is supplied in sheet form.
**Glossary**

**fats** Substances similar to wax but characterized as being soft and greasy to the touch. An example of a fat used in dental waxes is stearic acid.

**flow** Continued deformation resulting from application of a static force.

**gums** Viscous substances from plant or animal sources that harden in air. Gums combine with water to form sticky, viscous liquids. An example of a gum is gum arabic.

**natural resins** Mixtures of high-molecular-weight organic substances obtained directly from plants or trees as exudates. An example of a natural resin used in dental waxes is dammar.

**natural wax** A hydrocarbon or hydrocarbon-derivative polymer with an approximate molecular weight of 400 to 4,000; of mineral, plant, insect, or animal origin. Examples of natural wax are paraffin, beeswax, carnauba, spermaceti, and ceresin.

**pattern** A form used to make a mold.

**recovery** Change in shape resulting from the release of internal stresses.

**residual stress** Internal stress independent of applied force.

**synthetic wax** A man-made wax synthesized from appropriate monomers.

**Discussion Questions**

1. Waxes have the highest thermal expansion values of any dental material. What problems can result from this, especially in casting accuracy?

2. Which type of inlay wax (I or II) should be used for the indirect die technique? How do the different flow properties of the two types make them suitable for their application?

3. If a wax inlay pattern cannot be invested right away, it is recommended that it be stored in a refrigerator. Why and what phenomena will this storage affect?

4. Why are polymer cements recommended for use with CAD/CAM inlays with poor marginal fits?

**Study Questions**

(See Appendix E for answers.)

1. List several common natural waxes used in dentistry.

2. Why must natural waxes be carefully monitored for properties?
3. Name three types of pattern waxes and their applications.

4. Wax pattern contraction arises from two sources. What are they?

5. List two means of increasing wax flow.

6. Describe three steps to minimize wax pattern distortion.

7. Type I pattern wax must have two types of flow behavior. Why?

8. What is the memory or recovery effect in waxes, and what does it lead to?

9. Why is complete burnout of wax patterns critical in the lost wax casting technique?

**Recommended Reading**


Chapter 21. Orthodontic Wires

Introduction

There is a bewildering array of metallic wires available for orthodontic use, including the older stainless steel and cobalt-chromium-nickel wires, the subsequently developed nickel-titanium and beta-titanium wires, and the more recently marketed shape-memory nickel-titanium wires and the ion-implanted nickel-titanium and beta-titanium wires. Esthetic nonmetallic orthodontic wires are under investigation.

The objective of this chapter is to provide information about this wide variety of orthodontic wires, in order to assist the clinician in making a rational selection for patient treatment. A general classification of the alloy systems used for the metallic orthodontic wires is provided in Fig 21-1.

**Fig 21-1** Classification of the alloy systems used for metallic orthodontic wires.

Mechanical Properties

Manufacturing of orthodontic wires

Metallic orthodontic wires are manufactured by a series of proprietary steps, typically involving more than one company. Initially, the wire alloy is cast in the form of an ingot, which must be subjected to successive deformation stages, until the cross section becomes sufficiently small for wire drawing. Several deformation stages and intermediate heat treatments are required because considerable work hardening of the alloy occurs during wire manufacturing. Important proprietary details include the rate of drawing, the amount of cross-section reduction per pass, the nature of intermediate heat treatments, the die material and lubricant in contact with the wires, and the ambient atmosphere, which would be important for the reactive titanium-containing wire alloys. In general, the casting of the starting ingot and the initial mechanical deformation stages are not performed by the orthodontic materials companies that market the wires.

Whereas round orthodontic wires are manufactured by drawing through dies, rectangular cross-section wires are fabricated from round wires by a rolling process.
using a Turk's head, which contains pairs of rolls. The resulting rectangular or square cross-section wires will necessarily have some degree of rounding at the corners, which varies with the specific wire type and the manufacturer. This edge bevel can be of clinical significance for the actual torque delivered by the archwire-bracket combination. Moreover, the surface roughness of the wire, which has a clinically significant effect on the archwire-bracket sliding friction, varies considerably among the various products and is generally greater for the beta-titanium and nickel-titanium wires.

As a result of the wire-drawing sequence, orthodontic wires have a characteristic wrought microstructure. The original equiaxed grain structure of the starting cast ingot is completely eliminated, and when a polished and etched wire specimen is viewed through the optical microscope or the scanning electron microscope, the wrought grain structure appears as a series of closely spaced lines parallel to the original direction of drawing. It is well-known that this microstructure is essential for an orthodontic wire to maintain the desired temper (springiness) or optimal mechanical properties for clinical use. For example, heat treatment of stainless steel wires at temperatures of 700°C (1,300°F) and higher causes rapid softening and loss of the wrought microstructure as a result of recrystallization. Optimum heat treatments would involve only the recovery stage of annealing. In practice, clinicians perform heat treatments using electrical resistance (spot) welding apparatus, and the stainless steel, cobalt-chromium-nickel, and some nickel-titanium wires may be advantageously heat treated.

The properties of a metallic orthodontic wire are thus derived from two principal origins. First, there are four general wire alloys in significant current use: austenitic stainless steel, cobalt-chromium-nickel, beta-titanium, and nickel-titanium. The basic alloy composition will determine the broad range of inherent general properties for each metal type. Second, the particular nature of the drawing, rolling, and other special processes, including heat treatments by the manufacturer and clinician, will have further significant effects on the specific wire properties and archwire-bracket torque delivery characteristics.

In general, an orthodontist should consider the following aspects in the selection of wires: force delivery characteristics, elastic working range, ease of joining individual segments to fabricate more complex appliances, corrosion resistance and biocompatibility in the oral environment, and cost. In a practical sense, wire costs represent a significant concern for the clinician, and there is a considerable difference between stainless steel archwires and beta-titanium and nickel-titanium archwires. However, these more expensive alloys offer unique properties that should be carefully considered when selecting orthodontic wires.

**Bending tests and mechanics principles**

To understand the mechanical properties of primary importance when comparing different wire types and sizes, it is first necessary to review the underlying principles and terminology. Typically, the mechanical properties of orthodontic wires are determined from some type of bending test, because this mode of deformation is considered more representative of clinical conditions than the tension test conventionally used for metals. The cantilever bending test in the original form of
American National Standards Institute/American Dental Association (ANSI/ADA) Specification no. 32 for orthodontic wires not containing precious metals was strongly criticized. A revised specification has been approved; it contains a three-point bending test that better simulates clinical interbracket distances and is more suitable for the very low elastic modulus nickel-titanium wires than the previous test, which used the Olsen stiffness tester.

All bending tests involve the measurement of angular or linear deflection of an archwire segment resulting from a bending moment or an applied force. The bending moment or force and the deflection are represented on the vertical and horizontal axes, respectively, of a graphic plot that is generally a straight line at the force levels for elastic deformation of clinical interest. The exception is when very short test spans are used. Although orthodontists typically activate wires somewhat into the permanent deformation range, the bending properties are based on elastic deformation. Although clinical interest is obviously in the unloading characteristics of activated archwires, investigators have generally determined mechanical properties during the initial loading stage. The elastic loading and unloading plots differ for the nickel-titanium wires, although they do not differ for the other alloy types.

There are several basic mechanical properties of orthodontic wires that are determined from the bending test plot:

1. The **force delivery** is the slope of the initial straight line and is the amount of force or bending moment required for unit **activation**. This property is termed the stiffness of the wire and is the inverse of the property of flexibility.

2. The moment at yielding is the bending moment that corresponds to a designated small amount of permanent deformation, eg, 2.9 degrees in the bending test for the current ANSI/ADA specification. This property is analogous to the yield strength \((Y_S)\) for the tension test, and the analogous value of stress in bending is termed the flexural yield strength.

3. The maximum bending moment occurs during the permanent deformation portion of the bending test. This property is less relevant for the clinician than the moment at yielding, because only elastic deformation is desired for tooth movement.

4. The maximum amount of elastic activation before the onset of permanent deformation is the value of linear or angular deflection corresponding to the maximum elastic force or moment. This property is known as the elastic range or **working range** of the wire.

Wire stiffness or elastic force delivery is dependent on two fundamental factors: \(1)\) the composition and structure of the wire alloy, reflecting both the basic metallurgy and the manufacturing sequence, and \(2)\) the wire segment geometry, that is, the cross-section shape and size and the segment length. The basic metallurgy contribution of the wire alloy is given by the modulus of elasticity \((E)\), or Young’s modulus, which relates tensile and compressive elastic stress and strain independent of the specimen cross-section area and length. The elastic modulus values determined from bending and tension tests should be the same, provided that the bending deformation is properly analyzed.
The resistance of a cross-section shape to elastic bending is given by the **moment of inertia** \((I)\). For a round wire of diameter \((d)\), the moment of inertia is given by

\[
I = \frac{\pi d^4}{64}
\]

whereas for a rectangular wire of width \((w)\) and thickness \((t)\) in the plane of bending

\[
I = \frac{wt^3}{12}
\]

The stiffness of an archwire is inversely proportional to the segment length \((l)\), so that if the length is doubled the wire flexibility or elastic deflection will be doubled for the same applied force or bending moment. Summarizing these contributions, the elastic stiffness or force delivery characteristics are given by the following expressions:

**Round wire:** \(\frac{Ea^4}{l}\)

**Rectangular wire:** \(\frac{Emt^3}{l}\)

There are two additional very useful mechanical properties for orthodontic wires, which are obtained by combining the basic mechanical properties previously discussed (Table 21-1):

1. The modulus of resilience or resilience is the area under the elastic force-activation plot and represents the total biomechanical energy per unit volume available for tooth movement when the wire is loaded to the maximum elastic stress or bending moment or, equivalently, unloaded from this level. The modulus of resilience for an orthodontic wire is usually written as \((YS)^2 / 2E\), as the yield strength is generally used to represent the onset of permanent deformation because of the difficulty in precisely locating the proportional limit \((PL)\) on a bending test plot. The formal expression in materials science for the modulus of resilience is \((PL)^2 / 2E\). It follows that the resilience is much more strongly affected by changes in yield strength or proportional limit than elastic modulus, which is important for the heat treatment response of cobalt-chromium-nickel and stainless steel wires.

2. The **springback** for an archwire after unloading is given by the expression \(YS / E\), which is approximately equal to the maximum elastic strain or working range of the wire. (The formal expression from materials science for springback would be \(PL / E\).) Since the unloading curve from the permanent deformation range for well-behaved orthodontic wire alloys (i.e., other than nickel-titanium wires) is parallel to the elastic loading curve, the value of \(YS / E\) represents the approximate amount of elastic strain released by the archwire on unloading.

Numerous articles listed in the recommended readings have discussed the mechanics of bending tests for orthodontic wires.
### Table 21-1 Summary of conventional terminology for important mechanical properties of orthodontic wires*

<table>
<thead>
<tr>
<th>Basic property</th>
<th>Location on bending plot</th>
<th>Equivalent terms</th>
</tr>
</thead>
<tbody>
<tr>
<td>Force delivery (stiffness)</td>
<td>Slope of elastic loading curve or unloading curve</td>
<td>$EI/l$</td>
</tr>
<tr>
<td>Moment at yielding or flexural yield strength</td>
<td>Bending moment for designated small amount of permanent deformation (vertical axis of graph)</td>
<td>$YS^a$</td>
</tr>
<tr>
<td>Working range</td>
<td>Maximum value of purely elastic deformation (horizontal axis)</td>
<td>—</td>
</tr>
<tr>
<td>Modulus of resilience (resilience)</td>
<td>Area under elastic loading curve or unloading curve</td>
<td>$(YS)^2/2E^a$</td>
</tr>
<tr>
<td>Springback</td>
<td>Elastic strain recovered on unloading from permanent deformation range</td>
<td>$YS^a/E$</td>
</tr>
</tbody>
</table>

*The expressions are applicable for tension tests and for bending tests with well-behaved wire alloys and sufficiently long specimens. In these cases, the elastic loading curve of clinical interest is essentially the same as the initial linear plot for elastic loading. For short loading spans and the nonsuperelastic nickel-titanium alloys, the unloading curves are nonlinear and it is difficult to define a modulus of elasticity. (Terms from Thurow, 1982; Popov, 1968.)

$a$For the idealized definitions, the yield strength ($YS$) should be replaced in the expressions for modulus of resilience (resilience) and springback by the proportional limit ($PL$). Other symbols have their usual meanings: $E =$ modulus of elasticity, $I =$ moment of inertia (proportional to $d^4$ for round wire and $wt^3$ for rectangular wire), $l =$ segment length, $d =$ diameter, $w =$ width, $t =$ thickness in plane of bending.

### Orthodontic Wire Alloys

**Stainless steel**

Stainless steel continues to be the most popular wire alloy for clinical orthodontics because of an outstanding combination of mechanical properties, corrosion resistance in the oral environment, and cost. The wires used in orthodontics are generally American Iron and Steel Institute (AISI) types 302 and 304 austenitic stainless steels, with similar nominal compositions, although the use of 17-7 precipitation-hardening stainless steel has been explored. Type 302 is composed of 17% to 19% chromium, 8% to 10% nickel, and 0.15% maximum carbon. Type 304 contains 18% to 20% chromium, 8% to 12% nickel, and 0.08% maximum carbon. The balance of the alloy composition (Table 21-2) is essentially iron (approximately 70%). These are the well-known "18-8" stainless steels, so designated because of the percentages of chromium and nickel in the alloys.
Table 21-2 General compositions for four major classes of orthodontic wire alloys not containing precious metals

<table>
<thead>
<tr>
<th>Wire alloy</th>
<th>Weight percentage of elements</th>
</tr>
</thead>
<tbody>
<tr>
<td>Austenitic stainless steel*</td>
<td>17%-20% Cr, 8%-12% Ni, 0.15% C maximum, balance principally Fe (approx. 70%)</td>
</tr>
<tr>
<td>Co-based chromium-nickel a (Elgiloy)</td>
<td>10% Co, 20% Cr, 15% Ni, 15.8% Fe, 7% Mo, 2% Mn, 0.16% C, 0.04% Be</td>
</tr>
<tr>
<td>Beta-titanium b</td>
<td>77.8% Ti, 11.3% Mo, 6.6% Zr, 4.3% Sn</td>
</tr>
<tr>
<td>Nickel-titanium c (Nitinol)</td>
<td>55% Ni, 45% Ti (approx. and may contain small amounts of Cu or other elements)</td>
</tr>
</tbody>
</table>

*Data from ASM (1961).
a Data from Anusavice (1996).
b Data from Goldberg and Burstone (1979).
c Data from Civjan et al (1975), Khier (1988), and Quo et al (1994).

Research has shown that the modulus of elasticity in tension for stainless steel orthodontic wires, where values are more reliable than for bending tests, ranges from about 160 to 180 GPa. These values depend on the manufacturer and temper, and are indicative of differences in alloy compositions, wire drawing procedures, and heat treatment conditions. For many years, it was not appreciated that the elastic modulus for stainless steel orthodontic wires can be significantly decreased below the 190 to 210 GPa range given in standard physical metallurgy textbooks for annealed stainless steel, although this reduction in elastic modulus was well-known over five decades ago for heavily cold-worked industrial austenitic stainless steel alloys.

X-ray diffraction has shown that austenitic stainless steel archwires do not necessarily have the single-phase austenitic structure in the as-received condition from the manufacturers. The microstructural phases in these stainless steel wires depend on the manufacturer, temper, and cross-section size; the fundamental factors are the AISI type (particularly carbon content) and thermomechanical processing during manufacturing.

The yield strength for the stainless steel archwires shows a much wider variation than the elastic modulus and has been found to range from approximately 1,100 to 1,500 MPa for a resilient temper from a single manufacturer and several cross-section sizes. The yield strength was increased to about 1,700 MPa for several wire sizes after heat treatment. The range in values of mechanical properties in tension for as-received stainless steel wires of clinically important sizes is summarized in Table 21-3. Heat treatment of these wires also causes significant decreases in residual stress and modest increases (~10%) in resilience. Springback (YS/E) was found to range from 0.0060 to 0.0094 for eight different sizes of as-received stainless steel wires and from 0.0065 to 0.0099 after heat treatment.
Table 21-3  Range of mechanical properties in tension of principal clinical importance for four major orthodontic alloys and as-received wires*

<table>
<thead>
<tr>
<th>Wire alloy</th>
<th>Modulus of elasticity (GPa)</th>
<th>Yield strength (MPa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stainless steel (resilient temper)</td>
<td>160-180</td>
<td>1,100-1,500</td>
</tr>
<tr>
<td>Cobalt-chromium-nickel (Elgiloy Blue)</td>
<td>160-190</td>
<td>830-1,000</td>
</tr>
<tr>
<td>Beta-titanium (TMA)</td>
<td>62-69</td>
<td>690-970</td>
</tr>
<tr>
<td>Nickel-titanium (Nitinol)</td>
<td>34</td>
<td>210-410</td>
</tr>
</tbody>
</table>

*The data for modulus of elasticity and yield strength are for round wires with diameters from 0.016 to 0.020 inch and for rectangular wires with cross-section dimensions from 0.017 inch × 0.025 inch to 0.019 inch × 0.025 inch. (Data from Asgharnia and Brantley, 1986, and Drake et al, 1982.)

a1 MPa = 145 psi; 1 GPa = 145,000 psi.

The use of heat treatment to eliminate residual stresses that might cause fracture during manipulation of stainless steel appliances can be important under clinical conditions. However, austenitic stainless steel alloys can be rendered susceptible to intergranular corrosion when heated to temperatures between 400°C and 900°C, due to the formation of chromium carbides at the grain boundaries. These precipitates deplete the amount of chromium near the grain boundaries in the bulk stainless steel below that needed for corrosion resistance. Since the stainless steel alloys must be heated within this temperature range for soldering, clinicians are cautioned to minimize the time required for this process.

**Cobalt-chromium-nickel**

Cobalt-chromium-nickel orthodontic wires are very similar to stainless steel wires in appearance, mechanical properties (Table 21-3), and joining characteristics, but have a much different composition and considerably greater heat treatment response. Table 21-2 shows that the most commonly used alloy, Elgiloy (Rocky Mountain Orthodontics) has a complex composition of 40% cobalt, 20% chromium, 15% nickel, 15.8% iron, 7% molybdenum, 2% manganese, 0.16% carbon, and 0.04% beryllium, which is somewhat similar to that for some base metal casting alloys for removable partial dentures.

The Elgiloy wires are available in four tempers, soft, ductile, semiresilient, and resilient, which are color-coded for the convenience of the clinician. The differences in mechanical properties arise from proprietary variations in the wire manufacturing process. The soft-temper wires (Elgiloy Blue) are popular because they are easily deformed and shaped into appliances, then heat treated to provide substantially increased values of yield strength and resilience. Increases of 20% to 30% in the yield strength of Elgiloy Blue wires after heat treatment have been reported, and similar heat treatment responses appear to occur for the soft, ductile, and semiresilient tempers. The large increases in modulus of resilience principally arise from the dependence on \((\sigma_S)^2\) (see Table 21-1). The effect of heat treatment on mechanical properties has been attributed to complex precipitation processes. Springback for the
Elgiloy Blue alloy was found to range from 0.0045 to 0.0065 for five different sizes of as-received wires and from 0.0054 to 0.0074 after heat treatment.

The other tempers are less popular than the soft temper because wires made from them have lower formability and are somewhat higher in cost than stainless steel. For Elgiloy Blue, the elastic modulus in tension ranges from about 160 to 190 GPa for as-received wires (see Table 21-3), and from about 180 to 210 GPa after heat treatment. The corresponding ranges in yield strength are approximately 830 to 1,000 MPa in the as-received condition, and 1,100 to 1,400 MPa after heat treatment. It is important to emphasize that the elastic force delivery is very nearly the same for stainless steel and Elgiloy Blue archwire segments of the same size and length, as indicated in Table 21-3. A common misconception is that the elastic force delivery is much less for Elgiloy Blue wires compared to stainless steel wires because of the "feel" of the former. In reality, the yield strength and elastic range are the properties that are diminished relative to the more resilient stainless steel wires.

**Beta-titanium**

A beta-titanium orthodontic alloy, TMA (Ormco/Sybron), was introduced to the orthodontic profession about 20 years ago. The nominal composition of TMA, which is derived from the two major component elements (titanium-molybdenum alloy), is similar to the 77.8% titanium, 11.3% molybdenum, 6.6% zirconium, and 4.3% tin composition of an industrial beta-titanium alloy (see Table 21-2). The presence of molybdenum causes the elevated temperature body-centered cubic beta polymorphic phase of titanium, rather than the hexagonal close-packed alpha phase, to be metastable at room temperature. This results in excellent formability or capability for permanent deformation. Zirconium and tin contribute increased strength and hardness and prevent formation of an embrittling omega phase during wire processing at elevated temperatures. The TMA alloy has somewhat less than half the elastic force delivery ($E$ ranging from about 62 to 69 GPa) of stainless steel wires, with the yield strength ranging from approximately 690 to 970 MPa (see Table 21-3). Springback for four different sizes of as-received TMA wires was found to range from 0.0094 to 0.011.

Another noteworthy characteristic is that the TMA alloy possesses true **weldability**. (Welded joints that are fabricated from stainless steel and cobalt-chromium-nickel alloys must be built up with the use of solders to maintain adequate strength.) Optimum conditions for welding TMA with commercial apparatus have been published. Heat treatment by the clinician is not recommended for TMA, although this alloy does respond to a precipitation-hardening procedure. Solution heat treatment between approximately 700°C and 730°C, followed by water quenching, then aging at approximately 480°C results in precipitation of the alpha-titanium phase and a peak value for the $YS/E$ ratio.

Research has shown that TMA wires have high surface roughness, which leads to high values of archwire-bracket friction. Ion-implanted TMA wires that have substantially reduced archwire-bracket friction are available from Ormco/Sybron.

Recently, a new beta-titanium orthodontic wire (Resolve) was introduced by GAC International. The composition of Resolve (79% titanium, 11% molybdenum, 6%
zirconium, and 4% tin) is very similar to that of TMA, but transmission electron microscopic observations of microstructural precipitates suggest that different processing procedures (wire drawing parameters and heat treatment conditions) are used by the two manufacturers. These differences may have practical significance for the clinically important mechanical properties of the two beta-titanium wire products.

The physical metallurgy of the beta-titanium alloys is complex, and systems other than the titanium-molybdenum-zirconium-tin alloy may also be suitable for orthodontic wire applications.

**Nickel-titanium**

The fourth wire alloy, nickel-titanium, has remained a strong focus of materials research, as well as considerable marketing activity by manufacturers. The name nitinol, which is a generic name for all of these nickel-titanium alloys, is derived from the nickel and titanium composition, along with the Naval Ordnance Laboratory where these alloys were originally developed.

Nickel-titanium orthodontic alloys are based on the intermetallic compound NiTi, which has weight percentages of 55% nickel and 45% titanium (see Table 21-2). While earlier chemical analyses of two nickel-titanium wire products revealed similar nickel-rich compositions (59 to 60 weight percent nickel), recent x-ray energy-dispersive spectroscopic analyses of several nickel-titanium wire products indicated that the compositions are slightly titanium rich (between 50 and 51 atomic percent titanium).

Although the shape-memory effect (SME) associated with these nickel-titanium alloys was not available in the original Nitinol wire (3M Unitek), there were two features of considerable importance for clinical orthodontics:

1. The very low elastic modulus ($E$ about 34 GPa in tension) for Nitinol corresponds to about one fifth of the force delivery for stainless steel archwires and half the force delivery for TMA wires having the same cross-section dimensions and length.

2. Because of the extremely wide elastic working range, 12.5-mm segments in the clinically important size ranges retain a permanent set of not more than 5 degrees, after 90-degree cantilever bending by the original ANSI/ADA specification test procedure and release of the applied moment.

The yield strength for the Nitinol wires generally ranges from about 210 to 410 MPa. Springback for six different sizes of as-received wires was found to range from 0.0058 to 0.016.

Chinese NiTi orthodontic wire (marketed as Ni-Ti by Ormco/Sybron) was subsequently introduced and compared with Nitinol and stainless steel wires, using a cantilever bending test and 5-mm span lengths appropriate to clinical interbracket distances. The nonlinear activation characteristics of clinical interest are quite evident. However, whereas the average unloading stiffness (ratio of bending moment to amount of deactivation in degrees) is the same for all activations of Nitinol, the unloading curves for Chinese NiTi wires are dependent on the level of activation
(amount of bending deflection). These latter unloading curves have relatively high values of unloading stiffness for the initial and final stages (under 10 degrees) of deactivation; through the middle range of deactivation, the unloading stiffness is much smaller and approaches constant force delivery. The average stiffness of Chinese NiTi wire from these unloading curves was found to increase considerably on going from the largest to smallest activations; with the 5-mm test spans, the springback of Chinese NiTi wire was over 4 times that for stainless steel and more than 50% greater than that for Nitinol wire.

Japanese NiTi wire (marketed as Sentalloy by GAC International) was introduced at about the same times as Chinese NiTi wire. The stress-strain curves in tension and the load-deflection curves for three-point bending of 14-mm test spans were compared for 0.016-inch diameter wires of this alloy, Nitinol, stainless steel, and Elgiloy. The tension and bending tests confirmed the existence of superelastic behavior in the Japanese NiTi wires. After a certain level of elastic tensile strain or bending deflection, further deformation (up to about 10% tensile strain or 2 mm deflection for a 0.016-inch diameter wire) takes place at nearly constant tensile stress or bending force. The unloading characteristics exhibit initial and final regions of relatively steep slope, along with an extensive intermediate region where there is little change in stress or force.

When Japanese NiTi wire was heat treated at 500°C from 5 minutes to 2 hours, the bending force remaining during the superelastic region of the deactivation curve could be varied over a wide range. This behavior has been exploited (GAC International) to develop commercial nickel-titanium wires that can deliver low, medium, or high levels of force. A highly convenient electric resistance method has been developed for the heat treatment of Japanese NiTi wires, and a commercial apparatus (GAC International) is available that enables the clinician to heat treat superelastic nickel-titanium archwires as desired for the treatment of individual patients.

The bending properties of several nickel-titanium wire products with both round and rectangular cross-sections have been compared, using cantilever specimens with 6-mm test spans. The bending test plots indicated that the Chinese NiTi and Nitinol SE (3M Unitek) wires possessed superelastic behavior (Fig 21-2) very similar to that for Japanese NiTi wire and that the original Nitinol wire (Fig 21-3) should be classified as nonsuperelastic. The three superelastic wire products evaluated exhibited responses to vacuum heat treatment for 10 minutes and 2 hours at 500°C and 600°C (Fig 21-4) that were similar to those reported for Japanese NiTi wire. The constant bending moment in the superelastic deactivation range was decreased after heat treatment at 500°C, and loss of superelastic behavior occurred after heat treatment at 600°C. In contrast, the bending properties of the three nonsuperelastic wire products evaluated were essentially unaffected by heat treatment.
Fig 21-2 Cantilever bending plots for 6-mm test spans of four different sizes of as-received Nitinol SE (3M Unitek), a superelastic wire. (From Khier et al, 1991. Reprinted with permission from the American Journal of Orthodontics and Dentofacial Orthopedics.)
Fig 21-3 Cantilever bending plots for 6-mm test spans of four different sizes of as-received Nitinol (3M Unitek), a nonsuperelastic wire. (From Khier et al, 1991. Reprinted with permission from the *American Journal of Orthodontics and Dentofacial Orthopedics*.)

Fig 21-4 Cantilever bending plots for 6-mm test spans of 0.016-inch diameter Nitinol SE wires subjected to heat treatments of 10 minutes and 2 hours at 500°C and 600°C. (From Khier et al, 1991. Reprinted with permission from the *American Journal of Orthodontics and Dentofacial Orthopedics*.)
The shape-memory characteristics of the nickel-titanium alloys are associated with a reversible transformation between the austenitic and martensitic NiTi phases, which occurs by a twinning process. The **martensitic phase** (which has a distorted monoclinic, triclinic, or hexagonal structure) forms from the **austenitic phase** (which has an ordered body-centered cubic CsCl structure) over a certain transformation temperature range (TTR) or when the stress is increased above some appropriate level. The austenitic phase forms from the martensitic phase over a different TTR, or when the stress is decreased below the appropriate level. The difference in the temperature ranges for the forward transformation from the martensitic phase to the austenitic phase, and for the reverse transformation from the austenitic phase to the martensitic phase, is termed **hysteresis**. Frequently, an intermediate phase, termed R-phase because of its rhombohedral structure, forms during the forward and reverse transformations between martensitic NiTi and austenitic NiTi. In order for a nickel-titanium archwire to possess shape memory in vivo, the transformation from the martensitic NiTi structure to the austenitic NiTi structure must be completed at the temperature of the oral environment. The shape-memory characteristics of several commercial nickel-titanium wires have been evaluated by measuring the final length of segments that were permanently deformed approximately 7% in tension below the TTR reported by the manufacturers for the transformation to austenite and then heated above the TTR to 150°C. The deformed alloys evaluated exhibited excellent shape (length) recovery ranging from 89% to 94%.

Nickel-titanium alloys with shape-memory behavior activated at body temperature have recently been introduced by many manufacturers. A notable example is the three variants of the Copper Ni-Ti (Ormco/Sybron) wires that achieve shape memory at temperatures of 27°C, 35°C, and 40°C. The 27°C Copper Ni-Ti wire would be useful for patients who are mouth-breathers, whereas the 40°C Copper Ni-Ti wire achieves shape memory only when exposed to hot fluids in the oral environment. It has been reported that Copper Ni-Ti contains nominally 5% to 6% by weight copper and 0.2% to 0.5% by weight chromium. The 27°C variant contains 0.5% chromium to compensate for the effect of copper in raising the temperature for completion of the transformation to austenitic NiTi above that of the oral environment, and the 40°C variant contains 0.2% chromium.

Decreases in the mechanical properties of nickel-titanium wires after long-term deflection have been reported, and time-dependent relaxation of bending deformation in Nitinol was also reported. The clinical significance of these results and their relationship to the metallurgical phases in the different wire products require further study.

Nickel-titanium archwires with ion-implanted surfaces to obtain reduced bracket friction have been introduced (GAC International). The clinical efficacy of these wires requires further investigation. One study has reported no clinical advantage for the initial alignment compared with nickel-titanium archwires without ion-implanted surfaces.
**Considerations for Selection**

Although the stainless steel archwires are still the most widely used in orthodontics and cobalt-chromium-nickel wires are selected by many clinicians, the nickel-titanium and beta-titanium wires have become very popular. The gold alloy wires, while still available commercially, are relatively expensive and have minimal clinical use. Each of the four major nonprecious orthodontic wire alloys has distinct advantages for selection by the clinician, as well as some areas or potential areas of concern, as summarized in Table 21-4.

The stainless steel and cobalt-chromium-nickel wires offer the advantages of relatively low cost, known biocompatibility from extensive clinical usage, and excellent formability for the fabrication of orthodontic appliances. However, these wires have much higher force delivery (elastic modulus) than the beta-titanium and nickel-titanium wires, and the force delivery is increased after heat treatment by the clinician to increase resilience (particularly Elgiloy Blue) and relieve residual stresses.

The relatively high cost of the beta-titanium orthodontic wires is offset by the intermediate force delivery, excellent formability, and true weldability of this alloy. The clinician may select beta-titanium archwires because they more nearly fill the bracket slots due to their lower elastic modulus, compared with the wire sizes that would be selected for a stainless steel alloy. The formability or ductility of beta-titanium provides the orthodontist with the capability of fabricating arches or segments with complicated loop configurations not possible with the nickel-titanium alloys. Clinical examples include the alignment of teeth in an arch during finishing, the rotation and changing of the axial orientation of teeth, the use of specialized springs or auxiliaries such as an intrusive arch or a canine root spring, and the fabrication of closing loops that may contain helices. The direct welding of auxiliaries to archwires enables hooks, tie backs, and finger springs to be easily prepared with commercial welding apparatus.

A clinical study has reported that the rate of orthodontic sliding space closure was not significantly different for ion-implanted and conventional (not ion-implanted) TMA wires, and that the rate of closure was similar to that reported for stainless steel wires. Manufacturing of the beta-titanium wires is difficult because of the reactivity of titanium, and anecdotal reports have described the susceptibility of some batches of TMA wires to undergo fracture during clinical manipulation. Research is in progress to compare the formability and manipulative characteristics of TMA and Resolve archwires.

It is well-known from histologic studies on tooth movement that light, continuous, nearly constant forces represent optimum biomechanical conditions. The Chinese and Japanese NiTi wires, along with numerous more recently introduced superelastic and shape-memory nickel-titanium wires, have excellent springback (range of elastic action), low stiffness, and nearly constant force delivery during deactivation, and provide these desirable mechanical properties better than the stainless steel, cobalt-chromium-nickel, beta-titanium, and original Nitinol wire alloys. However, the original nonsuperelastic Nitinol remains popular for clinical use, and no studies have demonstrated clinical superiority of the superelastic or shape-memory wires compared to the nonsuperelastic nickel-titanium wires. It is possible that under some clinical
conditions the superelastic nickel-titanium wires are not sufficiently deformed for this property to be manifested during initial alignment of the teeth. A recent clinical study has found that there was no difference between multistrand stainless steel, superelastic nickel-titanium, and ion-implanted nickel-titanium archwires for tooth alignment.

Another interesting feature is the increase in force delivery that occurs when an appliance is removed and then retied (ligated) with Chinese NiTi wire. These results would be anticipated with the other superelastic alloys. Superelastic nickel-titanium alloy coil springs have also been developed for orthodontics and have found wide applicability as optimal appliances for tooth movement under certain clinical conditions.

Corrosion resistance of the orthodontic wires is provided by a chromium oxide film for the stainless steel and cobalt-chromium-nickel alloys, and a titanium oxide film for the beta-titanium and nickel-titanium alloys. Clinical experience is that corrosion of orthodontic wires is of minimal concern, and immersion of wires in a 1% sodium chloride solution for periods up to 11 months caused no significant changes in the bending mechanical properties of Nitinol wire segments. When the cyclic potentiodynamic polarization behavior (-500 mV to 300 mV) of stainless steel, cobalt-chromium-nickel, beta-titanium, and Nitinol wires in a 1% sodium chloride solution was compared, breakdown of the surface oxide and pitting were observed only for Nitinol and appeared to be accompanied by the selective dissolution of nickel. Potentiodynamic polarization plots for several superelastic and nonsuperelastic nickel-titanium wires have been published, showing the breakdown potentials for the passivating surface oxide films. A recent in vitro study indicated that the amounts of chromium and copper used to change the superelastic characteristics did not affect the corrosion of nickel-titanium wires immersed in simulated physiologic media.

Research has shown that nickel dissolution does occur at sites of surface damage in nickel-titanium wires. Nickel hypersensitivity reactions to nickel-titanium wires have been observed in orthodontic patients who are nickel-sensitive, although such cases are rare. In vitro studies using human epithelial (He La) cells have shown noncytotoxic responses from nickel-titanium alloys. Moreover, a survey of orthodontic practices did not yield any significant relationship between the use of nickel-containing wires (stainless steel, cobalt-chromium-nickel, and nickel-titanium) and the development of skin lesions. The greater cost of the nickel-titanium archwires and the considerable recent interest in infection control procedures have stimulated several studies that suggest that sterilization and clinical recycling only have small effects on the mechanical properties of the nickel-titanium wires. A recent investigation comparing the surface topography and chemical compositions of as-received and clinically used nickel-titanium archwires concluded that surface scratches and cracks arising from the wire manufacturing process were evidently not sufficiently large to serve as sites for localized corrosion.

A transparent nonmetallic orthodontic archwire (Optiflex, Ormco/Sybron) with a silica core, a silicon resin middle layer, and a stain-resistant outer layer has been developed and used in patient treatment. While sharp bends cannot be placed because of the brittle core, this ceramic-polymer wire is highly resilient and undergoes minimal permanent deformation under clinical conditions. Two different groups have recently described the compositions and mechanical properties of new polymeric
Table 21-4 Advantages and disadvantages of each of the four major orthodontic wire alloys

<table>
<thead>
<tr>
<th>Wire Alloy</th>
<th>Advantages</th>
<th>Disadvantages</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stainless steel</td>
<td>Lowest cost of the wire alloys</td>
<td>High force delivery</td>
</tr>
<tr>
<td></td>
<td>Proven biocompatibility from extensive clinical use</td>
<td>Relatively low springback in bending compared to beta-titanium and nickel-titanium alloys</td>
</tr>
<tr>
<td></td>
<td>Excellent formability for fabrication into orthodontic appliances</td>
<td>Can be susceptible to intergranular corrosion after heating to temperatures required for joining</td>
</tr>
<tr>
<td></td>
<td>Can be soldered and welded, although welded joints may require solder reinforcement</td>
<td></td>
</tr>
<tr>
<td>Elgiloy (soft temper)</td>
<td>Relatively low cost, although higher than stainless steel</td>
<td>High elastic force delivery, similar to that for stainless steel</td>
</tr>
<tr>
<td></td>
<td>Proven biocompatibility from extensive clinical use</td>
<td>Lower springback than stainless steel (using $YS/E$ values from tension test)</td>
</tr>
<tr>
<td></td>
<td>Outstanding formability in as-received condition (heat treated to increase $YS$ and resilience)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Can be soldered and welded, with joining characteristics similar to stainless steel</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Excellent in vivo corrosion resistance</td>
<td></td>
</tr>
<tr>
<td>Beta-titanium</td>
<td>Intermediate force delivery between stainless steel or Elgiloy and nickel-titanium</td>
<td>Expensive</td>
</tr>
<tr>
<td></td>
<td>Excellent formability; the only orthodontic wire alloy with true weldability</td>
<td>High archwire-bracket friction with original TMA (decreased for ion-implanted TMA)</td>
</tr>
<tr>
<td></td>
<td>Excellent springback characteristics (using $YS/E$ values from tension test)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Excellent biocompatibility from high titanium content demonstrated by clinical use</td>
<td></td>
</tr>
<tr>
<td>Nickel-titanium</td>
<td>Lowest force delivery of the four orthodontic wire alloys</td>
<td>Expensive, particularly for newest products</td>
</tr>
<tr>
<td></td>
<td>Excellent springback in bending, particularly for superelastic and shape-memory alloys</td>
<td>Second highest archwire-bracket friction after TMA (decreased for new ion-implanted TMA)</td>
</tr>
</tbody>
</table>
Implanted product

- Difficult to place permanent bends and cannot bend wire over sharp edge or into complete loop
- Wires cannot be soldered and must be joined by mechanical crimping process
- Lowest in vitro corrosion resistance of wire alloys (may be concern about in vivo nickel release

Archwires that are reinforced with either glass or ceramic fibers. Interesting future developments are anticipated in composite nonmetallic archwires and ligature wires, and in polymeric retainer materials.

**Clinical Scenario**

This section presents an approach for choosing materials and a system for a specific situation. It uses the same format as that presented in chapter 7. Advantages and Disadvantages of each material/system are prioritized using the following codes: * = of minor importance, ** = important, and *** = very important.

---

**Materials**  
Superelastic NiTi wire vs stainless steel wires

**Critical Factors**  
Good esthetics, speed of completion

**Superelastic nickel-titanium wires**

**Advantages**

*** Produce lower moments and forces

*** Constant force over wide range of deflection

*** Low stiffness

** High springback

** High stored energy

* More effective in initial tooth alignment

Shape memory

*** Less patient discomfort

**Disadvantages**

* Cannot be soldered or welded
** Cannot be easily formed or bent

*** Tendency for dentoalveolar expansion

*** High cost

    Poor torque control

*** Patient discomfort with larger activation

**Stainless steel wires**

*Advantages*

*** Less expensive

    Good formability

    High resilience

    * Can be soldered or welded

** Good in final detailing

** Great for arch coordination

*** Good torque control

    Relatively predictable biomechanical properties

*Disadvantages***

** More chair time

    Resilience depends on diameter

    No shape memory or superelasticity

** Poor deflection range

*Prepared by Dr David E. Wacker.*

---

**Glossary**

activation Bending a wire that will produce an elastic force for tooth movement.

austenitic phase The face-centered cubic solid solution structure of iron, chromium, nickel, and carbon for stainless steels, or the body-centered cubic structure of NiTi for
the nickel-titanium alloys.

**force delivery** The force produced by an orthodontic wire against a tooth.

**martensitic phase** A body-centered cubic phase in austenitic stainless steels or a phase (reported as monoclinic, triclinic, or hexagonal) in nickel-titanium alloys that forms at low temperatures or as a result of cold working the austenitic phase.

**moment of inertia** A geometric parameter, dependent on cross-section shape and dimensions, which expresses the relative resistance of an orthodontic wire to bending. The stiffness in bending is directly proportional to the moment of inertia of the cross section and the elastic modulus of the wire alloy, and inversely proportional to the length.

**shape-memory effect** A property of certain nickel-titanium wires that will permit shaping at a higher temperature, followed by deformation at a lower temperature, and a return to the original shape by reheating.

**springback** The elastic deformation or strain recovered when an orthodontic wire is unloaded, typically in bending deformation. For a wire that exhibits linear elasticity, the expression for springback in the orthodontic materials literature \((Y/S/E)\) is equivalent to the elastic strain at the yield strength.

**superelasticity** A property of certain nickel-titanium wires that is characterized by an extensive region of elastic activation and deactivation at nearly constant bending moment or force, arising from reversible transformation between the austenitic and martensitic NiTi phases.

**temper** The spring character of an orthodontic wire, which is related to the mechanical properties of yield strength and elastic modulus.

**weldability** Having the capability of being joined by the passage of a strong electric current.

**working range** The maximum deflection of a wire within the elastic range.

**Discussion Questions**

1. What phenomenon is responsible for the force delivery to teeth by deformed wires?

2. How is it possible to obtain different mechanical properties from the same wire?

3. Which type of orthodontic wire is most easily joined by welding?

4. What is shape memory, and which wires have it?

5. What factors would be important for the development and clinical use of composite ceramic-polymer orthodontic wires?

**Study Questions**
1. What are the types of orthodontic wires in current use?
2. What are the main criteria for the selection of an orthodontic wire?
3. What determines wire stiffness or elastic force delivery?
4. What are the advantages of stainless steel wires?
5. What is Elgiloy?
6. What is the composition of beta-titanium wires?
7. What are the advantages of beta-titanium orthodontic wires?
8. What are the Disadvantages of nickel-titanium wires?
9. How do manufacturers obtain shape memory or superelastic characteristics for some nickel-titanium wires?

**Recommended Reading**


Chapter 22. Endodontic Materials

Introduction

Endodontics is the branch of dentistry that is concerned with the morphology, physiology, and pathology of human dental pulp and periradicular tissues. Its study and practice encompass the biology of normal pulp and the etiology, diagnosis, prevention, and treatment of diseases and injuries of the pulp and associated periradicular conditions.

The scope of endodontics includes, but is not limited to, the differential diagnosis and treatment of oral pain of pulpal and/or periradicular origin; vital pulp therapy, including pulp capping and pulpotomy; nonsurgical treatment of root canal systems and the obturation of these systems; selective surgical removal of pathological tissues resulting from pulpal pathosis; repair procedures related to surgical removal of pathological tissues; intentional replantation and replantation of avulsed teeth; root-end resection, hemisection, and root resection; root-end obturation; bleaching of discolored teeth; retreatment of teeth previously treated endodontically; and treatment with posts and/or cores for coronal restorations.

The success of basic nonsurgical endodontic treatment is highly dependent on the triad of access cavity preparation, proper cleaning and shaping of the root canals, and the quality of the obturation of the root canal system. The long-term prognosis is determined by the quality and integrity of the coronal seal, ie, post-core-permanent restoration. It is therefore imperative that further ingress of oral fluids (microleakage), through restorative and endodontic materials, be kept to a minimum.

Endodontic Access

Rubber dam materials

First and foremost in endodontic treatment, a rubber dam must be placed around the tooth or teeth to be treated. Use of rubber dam is mandatory as a standard of care in nonsurgical root canal treatment. It facilitates treatment by isolating the tooth, preventing aspiration of small endodontic instruments and endodontic irrigants, reducing salivary and bacterial contamination during treatment, protecting and retracting the oral soft tissues, and improving the visibility and efficiency of the clinician. The rubber dam system consists of a thin, flat sheet of latex in various colors and thicknesses (thin, medium, heavy, extra heavy). The medium thickness is generally preferred because it resists tear, retracts soft tissue better than the thin materials, and is easier to place than the heavy dams. Dark-colored dams allow for better visual contrast, thus reducing eyestrain, but light-colored dams naturally illuminate the operating field and provide easier film placement underneath the dam. Rubber dams are typically 5 × 5 or 6 × 6 inches, and they are available in nonlatex (eg, nitrile, synthetic) for persons with latex allergies.

For application, a rubber dam punch is used to place a hole in the dam material so that it will fit over the tooth. A rubber dam clamp, which is a plastic or metal retainer, is available in a number of anatomical shapes. A rubber dam forceps is used to secure the clamp around the cervical portion of the tooth, thus stabilizing the rubber dam and
Aiding in retraction of soft tissue adjacent to the isolated tooth. A pronged rubber dam frame, made of plastic or metal, is then placed on the dam extraorally to retract and stabilize it over the patient's mouth.

**Access cavity preparation**

Access to the pulp chamber and canal system is achieved through the use of rotary high-speed burs in a dental handpiece to bore an opening in the affected tooth, typically on the lingual surface of anterior teeth and the occlusal surface of posterior teeth. A variety of bur types can be used depending on the preference of the operator and the status of the clinical crown. Long-shanked tungsten-carbide burs and size 2, 4, or 6 round burs can be used to make the access cavity. After the pulp chamber is located with a sharp, stainless steel endodontic explorer, safe-ended diamond burs or Endo-Z (Dentsply) burs are used to unroof the chamber and refine the axial walls of the cavity preparation. Enhanced illumination and magnification with head lamps, loupes, or a surgical operating endodontic microscope, with magnifications up to 26×, can aid the clinician in locating calcified canals and identifying fractures.

**Biomechanical Instrumentation**

The second component of the endodontic triad involves cleaning and shaping of the canals. Removal of the coronal portion of the pulp is usually performed with a metallic spoon excavator or a rotary bur on a slow-speed handpiece. For the initial debridement of the canals, root canal broaches or rotary orifice-shaping instruments can be used. Manufactured in a number of sizes, the root canal broach is a very narrow, flexible, round stainless steel instrument with barbs along its shaft. Prone to breakage, the broach must be used passively in the canal and should not engage dentin. It is designed to removed gross amounts of pulpal tissue in large canals by locking remnants of pulp with its sharp barbs. Broaches can also be used to remove cotton products that have been placed in the chamber between appointments.

Newer to the marketplace are rotary orifice shapers, which are made of flexible nickel-titanium. These variably tapered instruments are used in special controlled-speed, high-torque handpieces. Their use in the canal is typically limited to the coronal half of the canal. Because of their larger tapers, they facilitate straight-line access to the root apex by removing restrictive coronal dentin early in the cleaning and shaping process.

Gates-Glidden instruments are commonly used to enlarge canal orifices and shape the coronal portion of the canal. Used in a slow-speed handpiece, they consist of flexible, stainless steel, noncutting shafts with flame-shaped burs at their tips. They are available in a variety of lengths and sizes (Fig 22-1).
Specially designed hand instruments known as files and reamers are necessary for biomechanical instrumentation of the various anatomical forms of root canals (Fig 22-2). K-files are tapered metallic instruments made from rectangular, triangular, or rhomboidal cross-sectional wires. They are available in stainless steel or nickel-titanium and are manufactured by twisting or grinding the metal blank. Various cutting angles on the wires are created to plane or scrape the walls of the canal. Depending on the file design and canal size and curvature, these files may be used in a push-pull, twisting, watch-winding, or circumferential motion. The nature of the metal (nickel-titanium being five times more flexible than stainless steel), the size of the instrument, and the cross-sectional configuration determine the relative flexibility of each instrument (Fig 22-3). In order to accommodate different root lengths, these instruments are manufactured in lengths ranging from 21 to 31 mm. Moveable silicone rubber stops on the instruments can be adjusted to correspond to the exact length of each canal.
Many newer file designs incorporate a noncutting tip, which can be used instead of a cutting tip to help guide the file. Cutting tips can create deviations in canal anatomy during instrumentation. Examples of K-type files include Flex-R files (Miltex) (triangular cross section), FlexoFiles (Dentsply) (triangular cross section), and K-Flex files (Kerr Analytic) (rhomboidal cross section).

The size of the file is determined by the diameter of the shaft 1 mm from the tip and is recorded in millimeters. For example, a No. 25 K-file measures 0.25 mm in diameter 1 mm from its tip. The standard taper for these instruments is a 0.02-mm increase for every 1 mm up the shaft from its tip (the length of the working blades). For instance, the width of a No. 25 K-file at 16 mm above its tip is 0.57 mm (0.25 mm + 16 × 0.02 mm). Most traditional instrument designs adhere to this standard of sizing and tapering.

Another type of file is the Hedstrom file. These files are long, thin, and tapered like K-files but are made from round cross-sectional wires. Generally made from stainless steel or nickel-titanium, these aggressive files have cutting angles that are ground into the shaft and can only be used in a scraping or rasping motion as the file moves out of the canal. Rotation of Hedstrom files is contraindicated because they tend to self-thread and have a strong predilection toward fracture when used in this manner. Like the K-files, Hedstroms are manufactured in uniform sizes, tapers, and lengths. They are used primarily for removal of bulk amounts of dentin.

Classical hand root canal reamers are long, tapered, stainless steel instruments made from rectangular, triangular, or rhomboidal cross-sectional wires. They cut only during twisting and have fewer cutting edges than a typical K-file. Their use has diminished over time due to their lack of efficiency and their tendency to deviate from normal root canal anatomy during use. More tapered rotary reamers made of nickel-titanium have essentially replaced the stainless steel hand reamer. These instruments
have radial land-cutting regions along their shafts and thus have been termed "U"-bladed in cross section. Like the orifice openers previously described, rotary reamers are used in a slow-speed, high-torque, gear-reduction, air-driven or electric handpiece. Their tapers range from the standard 0.02-mm increase up to a 0.12-mm increase per millimeter. Examples include Profiles (Dentsply), GT Rotaries (Dentsply), Quanteck 2000 (Kerr Analytic), and Lightspeed (Lightspeed) (Fig 22-4).

![Fig 22-4](image)

**Fig 22-4** Specially designed nickel-titanium hand and rotary instruments with varying tapers.

### Irrigants and Chelating Agents

Endodontic **irrigants** dissolve tissue and necrotic debris as well as destroy bacteria and disinfect the canal spaces. They also serve as lubricants for the metallic instruments used in the canal. Typically, they are delivered to the canal via an irrigating syringe with a nonbinding, blunt-tipped needle. The ideal irrigating solution in contemporary endodontics is sodium hypochlorite (NaOCl), used in percentages ranging from 1% to 5.25% (household bleach). Hydrogen peroxide has been used, but it is significantly less effective than NaOCl due to its inability to dissolve necrotic tissue.

Many practitioners use a chelating agent such as ethylenediaminetetraacetic acid (EDTA) in conjunction with sodium hypochlorite. EDTA has been shown to assist in smear layer removal and may facilitate the loosening of calcific obstructions in the root canal. It is often used in paste form with special additives, or it can be used in pure liquid form as an aqueous base of 17% EDTA. The paste form, such as RC-Prep (Premier) or Glyde (Dentsply), enhances lubrication for rotary nickel-titanium instrumentation as well as traditional instrumentation techniques.
Materials and Instruments for Root Canal Obturation

Introduction

The goal of obturation is to seal off the root canal and its ramifications from oral fluids and bacteria. Although there is no ideal filling material, gutta-percha and sealer cements have proved to be the materials of choice in contemporary endodontics because they exhibit minimal toxicity and tissue irritability when confined to the root canal system.

Gutta-percha

Originating from special trees in Africa and South America, pure gutta-percha is considered to be an isomer of natural rubber known as trans-polyisoprene and is less elastic, more brittle, and harder than natural rubber. It can exist in both alpha and beta crystalline forms; these forms are interchangeable depending on the temperature of the material. The alpha form is the natural state, is less subject to shrinkage, and is often used in obturating systems that use thermoplasticized, or heat-softened, gutta-percha. The beta form, which is typically found in gutta-percha cones or points, is used in cold compaction techniques such as lateral condensation. Gutta-percha cones or pellets used in various obturation techniques contain approximately 19% to 22% gutta-percha, 59% to 75% zinc oxide, and a series of other additives, including waxes, coloring agents, antioxidants, and metallic salts.

The gutta-percha cone or pellet, in conjunction with a root canal sealer, must be compacted in the canal to conform to the prepared root canal system. Gutta-percha cones are available in standardized and nonstandardized forms (Fig 22-5). The standardized forms conform to the same dimensions and uniformity as those used for endodontic files; thus, a No. 40 gutta-percha cone should reasonably fit a canal that has been properly prepared with a No. 40 file. The nonstandardized forms, classified as medium, medium-fine, or fine-fine, have greater tapers than standardized cones and are often used in techniques that involve vertical compaction of heat-softened gutta-percha or filling of the coronal two thirds of a canal after a standardized cone has been compacted in the apical third.

![Gutta-percha cones or points. (Top) Standardized cone; (bottom) nonstandardized cone.](image-url)
A couple of obturating techniques and instruments can be used to compact gutta-percha into the root canal system (Fig 22-6). In lateral condensation, a long (17- to 30-mm), tapered, metallic instrument with a pointed tip known as a spreader is used to compact the gutta-percha cones and sealer laterally against the canal walls. Spreaders are available in both hand and finger forms and are made of stainless steel or nickel-titanium for greater flexibility. Root canal pluggers are used in the vertical condensation method. These are long, slightly tapered, metallic instruments with flattened or blunt tips. Available in both hand and finger types, pluggers are designed to compact gutta-percha and sealer vertically after the gutta-percha has been thermosoftened with a heating device.

**Fig 22-6** Endodontic hand condensers for compacting gutta-percha. *(Top)* Spreader; *(bottom)* plunger.

*Root canal sealers*

Root canal sealers are used to cement the gutta-percha in place, to fill voids and the intricate ramifications of the canal system, and to lubricate the cones during lateral compaction of the relatively nonrigid gutta-percha points. They should be biocompatible with and well tolerated by periradicular tissues.

The most commonly used sealers are zinc oxide- eugenol (ZOE) and calcium hydroxide-based cements because of their good working properties, sealability, biocompatibility, and ease of removal. Resin-, glass-ionomer-, and silicone-based sealers are also available but are more technique sensitive, difficult to remove, and have variable sealing properties.

When mixed into a thick, creamy consistency (Fig 22-7), sealers are placed inside the root canal via paper points or a lentulo spiral, or are deposited in a light layer through the counter-clockwise rotation of an endodontic file. Examples of endodontic sealers
Adjunct Materials

Calcium hydroxide

Calcium hydroxide has been used in dentistry for many years, both as an intracanal medication and as a pulp-capping agent. It is available in a variety of forms, ranging from pure chemical grade to proprietary compounds (Calasept, JS Dental; Tempcanal, Pulpdent). For intracanal use, calcium hydroxide has been proven to be antibacterial and may aid in the dissolution of necrotic pulp tissue. Its high pH is responsible for the destruction of bacterial cell membranes and protein structures. When pulps are exposed during routine cavity preparation, a pulp-capping agent such as Dycal (calcium hydroxide [Dentsply]) or mineral trioxide aggregate (discussed later in this section) can be placed as a "bandage" over the bleeding tissue in an attempt to promote dentinal bridge formation over time, thus preserving the vitality of the pulp.

Mineral trioxide aggregate

Mineral trioxide aggregate (MTA) (ProRoot, Dentsply) is one of the newest and most promising materials to enter the realm of endodontics in many years. This root canal repair material is a grayish powder consisting of fine, hydrophilic particles that set in the presence of moisture (Fig 22-8). The hydration of the powder, composed of tricalcium silicate, tricalcium phosphate, tricalcium oxide, and others, creates a colloidal gel that solidifies to form a strong impermeable barrier. The material sets within 3 to 4 hours and has a working time of 5 minutes; it has been shown to be biocompatible and its seal is superior to that of amalgam. Although the material is somewhat costly and difficult to work with, primarily due to its naturally sandy consistency when hydrated, the indications for its use include clinical situations that often have no other viable options, such as perforation repair. Indications for ProRoot
include pulp capping, internal repair of perforations (noncommunicative), apexification, and root-end filling in endodontic surgery.

Fig 22-8 Mineral trioxide aggregate (ProRoot). (Courtesy of Dentsply, Tulsa, OK.)

**Glossary**

**chelating agents** Chemicals used for the removal of inorganic ions, usually the disodium salt of ethylenediaminetetraacetic acid, from tooth structure.

**debridement** Elimination of organic and inorganic substances as well as microorganisms from the root canal by mechanical and/or chemical means.

**gutta-percha** The purified coagulated exudate from the mazer wood tree. It is a high-molecular-weight stereoisomer of polyisoprene. Since the 1950s, material compounded in the United States for "gutta-percha" points has been made from balata, a nearly identical latex from a special tree in South America.

**irrigants** Liquids used to dissolve and flush out root canal debris; examples include sodium hypochlorite, saline, and hydrogen peroxide.

**obturation** The complete filling and closing of a cleaned and shaped root canal with a root canal sealer and core filling material.

**pulp capping** The procedure of placing a dental material over an exposed or nearly exposed pulp to encourage the formation of irritation dentin at the site of injury.

**pulpotomy** The surgical removal of the coronal portion of a vital pulp as a means of preserving the vitality of the remaining radicular portion.

**sealer cements** Radiopaque dental cements used, usually in combination with a solid or semisolid core material, to fill voids and seal root canals during obturation.
Discussion Questions

1. List all of the materials necessary for the proper placement of a rubber dam on a tooth requiring endodontic therapy.

2. Discuss the types and properties of the materials used in obturating a cleaned and properly shaped root canal.

3. Discuss the various types of instruments that can be used to biomechanically prepare the root canal.

4. Why are irrigants necessary in endodontic therapy?

5. What are the roles of calcium hydroxide and mineral trioxide aggregate in endodontic therapy?

Study Questions

(See Appendix E for answers.)

1. All of the following endodontic instruments are designed to remove dentin except which one: (a) K-file; (b) reamer; (c) broach; or (d) Hedstrom?

2. The endodontic "triad" consists of which of the following: (a) access, biomechanical instrumentation, and obturation; (b) rubber dam placement, access, and obturation; (c) biomechanical instrumentation, obturation, and restoration; or (d) access, obturation, and restoration?

3. All of the following are true about Hedstrom files except which one: (a) they work best by rotation; (b) they are made from a circular wire; (c) they are manufactured in uniform sizes; or (d) they are made from either stainless steel or nickel titanium metal?

4. Which of the following is a chelating agent, helping to loosen calcific obstructions in the root canal: (a) gutta-percha, (b) Roth's 801, (c) sodium hypochlorite, or (d) EDTA?

5. Which of the following instruments is used in the lateral condensation of gutta-percha: (a) plugger, (b) spreader, (c) explorer, or (d) broach?

Recommended Reading


Chapter 23. Implant and Bone Augmentation Materials

Introduction

A critical problem in dentistry is treating the edentulous patient. A survey by the National Institute for Dental Research (NIDR) has shown that 42% of the population over the age of 65 is totally edentulous, and that a substantial number of other patients are partially edentulous, having an average of 10 missing teeth. Although removable dentures and fixed partial dentures offer effective treatments for many edentulous patients, those who have lost substantial tooth-bearing portions of bone and cannot manage prostheses or masticate properly can improve their oral function through the use of dental implants.

The use of implants as a means of treating these patients has accelerated in the last decade, and there are now over 300,000 dental implants in use in the United States. A 1988 conference on dental implants led to the conclusions that they can be effective in providing long-term total and partial support for restorations. Despite the expanded use of implants, however, they are largely evaluated only on a qualitative level. To better understand and quantify the clinical effectiveness of dental implants, a greater understanding of the parameters governing the long-term success of this complex material/tissue aggregate is needed.

In this chapter, the parameters important to implant dentistry are presented. Following an overview of general concepts and indications for implant use, osseointegration is defined and discussed. Next, methods of achieving osseointegration are presented and parameters important to achieving implant success are reviewed, with a primary focus on biomaterials and biomechanical factors.

Indications for Dental Implant Use

The general requirement for use of dental implants is available bone to support the implant. Available bone means there is adequate bone to support the implant with the physiological parameters of width, height, length, contour, and density. Note that the importance of these parameters varies, depending on the specific implant type (Table 23-1). Despite the "glamor" of implant dentistry, a conservative treatment protocol must be stressed. Unsatisfactory treatment with removable dentures or fixed partial dentures remains an important indication for implant use. In other words, dental implants are not a first-treatment option. A number of other contraindications for implant use also exist (see bellow).
Table 23-1 Summary of dental implant type and indications for each*

<table>
<thead>
<tr>
<th>Implant type</th>
<th>Indications</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Endosseous</strong></td>
<td></td>
</tr>
<tr>
<td>Root form</td>
<td>Adequate bone to support implant—width and height of primary concern</td>
</tr>
<tr>
<td></td>
<td>Maxillary and mandibular arch locations</td>
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<td>Blade (plate) form</td>
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<td>Ramus frame</td>
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<td>Adequate anterior bone to support implant—width and height of primary concern</td>
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*Adapted from NIH, 1988.

Contraindications for dental implant use (Adapted from NIH, 1988.)

- Unattainable prosthodontic reconstruction
- Patient sensitivity to implant component(s)
- Debilitating or uncontrolled disease
- Pregnancy
- Inadequate practitioner training
- Conditions, diseases, or treatments that may compromise healing (ie, radiation therapy)
- Poor patient motivation/hygiene
- Perceived poor patient compliance
- Unrealistic patient expectations
Types of Implants

In general, dental implants are classified into three categories, which are subdivided as shown in Table 23-1 and Fig 23-1

1. **Endosseous implants** are embedded in mandibular or maxillary bone and project through the oral mucosa covering the edentulous ridge.
2. Subperiosteal implants rest on the surface of the bone beneath the periosteum.

3. Transosteal implants penetrate the inferior mandibular border and also project through the oral mucosa covering the edentulous ridge.

Root-form endosseous screw-threaded implants are the most commonly used implants in clinical practice today. This class of implants is the only one for which good long-term (eg, 10- to 15-year) clinical tracking of large patient populations is available. Results indicate success rates for mandibular implants of approximately 96%, 94%, and 86% at 5, 10, and 15 years, respectively. For maxillary implants, success rates are 88%, 82%, and 78% at these same times. Breaking down the data based on oral
location indicates the following 5-year success rates: anterior mandible, 99%; posterior mandible, 94%; anterior maxilla, 92%; posterior maxilla, 75%. The clinician's expertise and surgical technique are more important than the specific implant and are the primary factors dictating clinical outcome.

**Osseointegration**

**Introduction**

Unlike many biomaterials, which strive to replace as much of a tissue's natural structure and function as possible, dental implants do not restore function by mimicking the natural function of the periodontal ligament (Fig 23-2).

![Fig 23-2 Schematic of natural tooth vs implant attachment to bone. (From Taylor, 1990. Reprinted with permission.)]
Instead, **osseointegration**, as originally defined, was intended to provide a direct structural and functional connection between ordered, living bone and the surface of a load-carrying implant. This definition was originally based on retrospective radiographic and light microscopic observations, and has since been modified based on scanning and transmission electron microscopic observations. However, the general working definition of osseointegration is fundamentally the same—the host bone responds, in a safe, predictable, and versatile manner, to surgery and placement of an implant in a sterile wound, with a healing cascade leading to interfacial osteogenesis and mechanical stability of the implant (Fig 23-3).

![Fig 23-3 Schematic of localized sections of interfacial zone, showing (a) osseointegrated and (b) fibrous-integrated tissue adjacent to implant surface. Conditions (a) and (b) might occur with minimized (a) and excess (b) tissue-implant relative motion. (From Branemark et al, 1985. Reprinted with permission.)](image)

In a well-functioning implant, interfacial osteogenesis and clinical stability are achieved (Fig 23-4(a)). In comparison, poorly differentiated connective tissue adjacent to an implant leads to clinical mobility and implant failure (Fig 23-4(b)).
Fig 23-4 Radiographic example of (a) well-functioning and (b) failing dental implants. In (a), a well-osseointegrated interfacial zone led to interfacial stability, whereas in (b), poorly differentiated interfacial connective tissue led to mobility and failure.

There are a multitude of interrelated clinical, biological, and engineering factors that control the oral cavity's response and dictate the success of osseointegration. These factors are discussed in the remainder of this chapter.

Mechanisms for achieving and enhancing implant-tissue attachment

An implant must be capable of carrying occlusal stresses. Additionally, stresses must be transferred to the adjacent bone. Not only must stresses be transferred, but they must be of a "correct" orientation and magnitude so that tissue viability is maintained in as near a physiological state as possible. The ability to transmit stress is largely dependent on attaining interfacial fixation. Thus, two further requirements are that the interface stabilize in as short a time postoperatively as possible and that, once stable, the interface remain stable for as long a time as possible.

In an ideal situation, such as the situation that can be achieved with commercially pure (c.p.) titanium, calcified tissue can be observed within several hundred Angstroms of the implant surface. A layer of proteoglycans, 200 to 400 A thick, lies adjacent to the metal oxide, and collagen filaments can be observed about 200 A from the surface (Fig 23-5). Less than optimal surgical techniques, implant surface chemistry, and relative motion can lead to a thicker zone of proteoglycans, soft connective tissue, and disordered bone.
Fig 23-5 Schematic of interfacial zone, showing constituents: bulk metal, metal oxide, proteoglycans, connective tissue, disordered and ordered bone, and relative proportions of each for good and poor osseointegration. (From Branemark et al, 1985. Reprinted with permission.)

Developing an "optimal" implant that meets all of these objectives requires the integration of material, physical, chemical, mechanical, biological, and economic factors. It should be pointed out that while all of these properties are important, they cannot all be optimized in a given design. In fact, optimization of one property often detracts from another. Thus, in implant design, a ranking of requirements and objectives is necessary.

The approach taken to meet a specific property objective should be based on a materials science approach, in that the synergistic relationships between processing, composition, structure, and properties are characterized. Altering any one of these four entities may alter the other three. For example, it is too simplistic to regard all hydroxyapatite (HA)-coated dental implants as similar.

Because of the need to develop a stable interface prior to loading, it is desirable to accelerate tissue apposition to dental implant surfaces. Materials developments that have been implemented in clinical practice include the use of surface-roughened implants and ceramic coatings. Other, more experimental techniques include electrical stimulation, bone grafting, and growth factors and bone proteins.
A variety of surface configurations have been proposed as means of improving the cohesiveness of the implant-tissue interface, maximizing load transfer, minimizing relative motion between implant and tissue, minimizing fibrous integration, and ultimately minimizing loosening and lengthening the service life of the construct. For metal implants, these surface configurations include those that are smooth, textured, screw threaded, plasma sprayed, or porous coated. By far, the most common surface configuration is the screw-threaded dental implant. Osseointegration around screw-threaded implants occurs through tissue ongrowth, or direct apposition between tissue and the implant surface. Alternative methods of implant-tissue attachment, based on tissue ingrowth into roughened or three-dimensional surface layers, yield higher bone-metal shear strength than other types of fixation. Increased interfacial shear strength results in a better stress transfer from the implant to the surrounding bone, a more uniform stress distribution between the implant and bone, and lower stresses in the implant. In principle, the result of a stronger interfacial bond is decreased implant loosening.

Bioceramics can have four different surface types and tissue attachment mechanisms:

1. Fully dense, inert ceramics that attach to bone by either a press fit or bone ongrowth onto a roughened surface
2. Porous inert ceramics, into which bone ingrowth occurs, creating a mechanical attachment
3. Fully dense surface-active ceramics that attach to bone via a chemical bond
4. Resorbable ceramics that integrate with and are eventually replaced by bone

A progression of surfaces from the lowest implant-tissue shear strength to the highest is as follows: smooth, textured, screw threaded, plasma sprayed, and porous surfaced. Two factors must be stressed. First, different surface structures necessitate different osseointegration times. Second, surface roughening, particularly of titanium-based materials, results in reduced fatigue strength. Thus, improvements in implant-tissue attachment strength are often countered by a loss of structural strength and must be met with design compromises to avoid material failure.

**Criteria for Successful Dental Implants**

**Introduction**

Three aspects of an implant-tissue system are important in determining success: (1) the implant material(s) and adjacent tissue(s), (2) the **interfacial zone** between the implant and tissue, and (3) the effect of the implant and its breakdown products on the local and systemic tissues. Although the interfacial zone is composed of a relatively thin heterogeneous metallic oxide, proteinacious layer, and connective tissue, it has an effect on the maintenance of interfacial integrity. Furthermore, interfacial integrity is dependent on material, mechanical, chemical, surface, biological, and local environmental factors, all of which change as functions of time in vivo. In addition, implant "success" is dependent on the patient's overall medical and dental status, the
surgical techniques employed, and the extent and time course of tissue healing. The focus of this section is on the biomaterial and biomechanical factors, which are summarized in Fig 23-6.

![Fig 23-6 Schematic of interdependent engineering factors affecting the success of dental implants. (From Kohn, 1992. Reprinted with permission.)](image)

**Surgical parameters**

Adequate preparation of bone is critical for bone-cell survival, well-ordered connective tissue apposition close to an implant surface, the establishment of a reliable bone anchor, and long-term implant and tissue viability. Poor surgical technique or premature functional loading may result in an inability to achieve osseointegration, fibrous adaptation, and early implant failure. The standard clinical protocol therefore calls for a two-stage surgical procedure. The first stage involves the careful preparation of the implant bed in a manner that minimizes trauma and optimizes healing and interfacial osteogenesis. Certain thermal limits should not be exceeded during surgery. If these temperatures are exceeded, thermal necrosis can occur, resulting in a thicker layer of soft tissue directly apposing the implant surface and jeopardizing osseointegration. Following the initial surgery, the implants are sealed in situ and remain unloaded for 3 to 6 months. During this period, ordered, living bone, with the potential for ultimately carrying occlusal loads, develops within the interfacial zone.

**Surface chemistry and biological response**

Implant materials may corrode and/or wear, leading to the generation of micron- or submicron-sized debris that may elicit both local and systemic biological responses.
Metals are more susceptible to electrochemical degradation than ceramics. Therefore, a fundamental criterion for choosing a metallic implant material is that the biological response it elicits is minimal. **Titanium**-based materials are well tolerated by the body because of their passive oxide layers. The main elemental constituents, as well as the minor alloying constituents can be tolerated by the body in trace amounts. However, larger amounts of metals usually cannot be tolerated. Therefore, minimizing mechanical and chemical breakdown of implant materials is a primary objective.

Linked to biological response are nine questions that must be considered:

1. Is material released?
2. What material is released?
3. What is the form of the material released?
4. How much material is released?
5. What is the rate of release?
6. In what subsequent reactions are the release products involved?
7. What percentage of release products are excreted/retained?
8. Of the percentage that is retained, where do they accumulate?
9. What biological response(s) result from the retained fraction?

Local accumulation of material around an implant may include membrane-bound ions, particles released due to wear or fatigue processes or insoluble reaction products. Excessive metal ion accumulation can lead to metallosis or tissue discoloration and also to reduced phagocytosis and cytotoxicity.

Understanding implant surface chemistry is important to ensure a twofold requirement. First, implant materials must not adversely affect local tissues, organ systems, and organ functions. Second, the in vivo environment must not degrade the implant and compromise its long-term function. The interfacial zone between an implant and the surrounding tissue is therefore the most important entity in defining the biological response to the implant and the response of the implant to the body.

The success of any implant is dependent on its bulk and surface properties, the site of implantation, tissue trauma during surgery, and motion at the implant-tissue interface. The surface of a material is almost always different in chemical composition and morphology than the bulk material. These differences arise from the molecular arrangement, surface reactions, and contamination. In this regard, the interface chemistry is determined primarily by the properties of the metal oxide and not as much by the metal itself. There is little or no similarity between the properties of the metal and the properties of the oxide, but adsorption and desorption phenomena can still be influenced by the properties of the underlying metal. Therefore,
characterization of surface composition, binding state, and morphology are important in the analysis of implant surfaces and implant-tissue interfaces. Surface analysis aids in material characterization, determining structural and composition changes due to processing, and in identifying biologically induced surface reactions.

Metallic oxides dictate the type of cellular and protein binding at the implant surface. Surface oxides are continually altered by the indiffusion of oxygen, hydroxide formation, and the outdiffusion of metallic ions. Thus, a single oxide stoichiometry does not exist. The surface potential may play an important role in osseointegration, and it has been postulated that oxides with high dielectric constants inhibit the movement of cells to an implant surface. It has also been demonstrated that the type and orientation of cells attaching to metal surfaces is influenced by the microscopic geometry of the substrate surface.

**Mechanical parameters**

Mechanical properties important in designing implant materials include stiffness, yield and ultimate strengths, fracture toughness, and fatigue strength. Stiffness, or modulus of elasticity, dictates, to a large extent, the ability of the implant to transmit stresses to the adjacent tissue and maintain tissue viability over time. Static and fatigue strengths obviously are important in minimizing material failures. Fracture toughness is a gauge of the energy needed to cause failure in the presence of existing damage.

Three points about mechanical properties must be emphasized. First, for many materials, including titanium, optimum strength (tensile and fatigue) and optimum ductility (elongation and toughness) require different microstructural morphologies and therefore different thermal processing. Second, depending on the surface configuration, properties differ in their ranked importance. For example, fatigue of smooth-surfaced implants is governed by the initiation stage, whereas fatigue of surface-roughened implants is governed by propagation. Therefore, different microstructures and thermal-processing techniques are desirable for different surface conditions. Third, if any surface coatings are used, mechanical analysis becomes more difficult, as there are now effectively three materials of interest—the substrate material, the substrate-coating composite, and the substrate-coating interface.

Implants are subjected to axial, shear, bending, and torsional loads, so, in addition to the magnitude of the loading, directionality must also be considered. With the above-mentioned considerations and only a qualitative knowledge of "stability"—the maximum allowable displacement at an implant-tissue interface that will still result in osseointegration and bone maintenance—it must be stressed that the postimplantation time at which an implant can begin to undergo loading is most likely implant- and location-specific and generally unknown.

Although rare, material failure of implants, generally by fatigue, does occur. Failure of implant structures or abutments should be not disregarded or viewed as isolated instances. Fatigue of implant materials is clinically important for several reasons. First, fatigue properties of implant materials should be accurately quantified so implants may be designed intelligently. Second, the stress distribution between an implant and surrounding bone tissue is dependent on the section size of the implant as
well as the elastic moduli of both the implant and tissue. Third, coated implants may undergo local fracture processes that do not necessarily compromise the integrity of the implant, but do compromise its functionality and ability to transmit stress to tissue.

**Implant design**

The design of dental implants is based on many interrelated factors, including the geometry of the implant, how this geometry affects mechanical properties, and the initial and long-term stability of the implant-tissue interface. There is no singularly agreed-upon design criterion. Implants can be designed to maximize strength, interfacial stability, or load transfer, with each of these criteria requiring different material and interface properties. Two goals of any implant design are to maximize initial stability (ie, through implant design and surgical precision, create as tight a fit as possible at the time of surgery and accomplish osseointegration in as short a time as possible following implantation) and minimize loosening (ie, maintain osseointegration for as long a time as possible following achievement of stability).

To ensure osseointegration and achieve the potential benefits of biological fixation, the interface must be stable (ie, relative motion must be minimized) prior to loading and throughout the service life of the device. It should be reiterated, however, that the quantification of stability is unknown. In general, if excessive relative motion at the implant-tissue interface occurs, a positive feedback system is created in which relative motion leads to bone atrophy, the formation of a fibrous tissue layer, and further increased displacements (Fig 23-7).

![Fig 23-7](image)

**Fig 23-7** Schematic of positive feedback mechanisms leading to implant loosening. (From Kohn, 1992. Reprinted with permission.)

Quantifying stresses and strains in implants, tissues, and implant-tissue interfaces is important for understanding mechanically mediated response mechanisms and for implant design. Implant and tissue geometry, elastic properties, loading, boundary conditions, interface conditions, and local stresses and strains are all important.

Since most dental implant clinical failures initiate in the interfacial zone, it is the local properties of this region that are most important. It has been hypothesized that the local material and tissue (including bone and fibrous tissue) microstructure within the implant-tissue interfacial zone is the major factor determining the local stresses and therefore regulates tissue integration and adaptation and the success of implants.
Rationale for this hypothesis is provided by the fact that osseointegration is not uniform.

**Biological parameters and properties of tissue**

As discussed earlier, osseointegration is a direct structural and functional contact between loaded, viable bone and an implant surface. Perhaps more important is the corollary that the creation and maintenance of osseointegration depends on the understanding of the tissue's healing, repair, and remodeling capacities. Dental implant design and function, therefore, are not only based on material considerations, but also on the properties of the surrounding tissue.

The microstructure of the mandible and maxilla is complex. For example, the mandible comprises secondary Haversian bone, regular and irregular primary lamellar bone, and plexiform lamellae of varying orientations. In general, the mandible is composed of basilar and alveolar bone, with no well-defined boundary between the two. The basilar bone forms the body of the mandible. Alveolar bone, formed in conjunction with tooth eruption, is a thin lamella that surrounds the tooth roots, attaches to the periodontal ligament fibers, and is surrounded by another layer of bone that supports the tooth sockets. In both humans and animals, the type and orientation of tissue microstructure are regionally dependent, as are density and mechanical properties. For example, compressive strength changes with tissue organization, morphology, and density and progressively increases with increasing numbers of osteons, numbers of lamellar plates, plate density, plate orientation with respect to the stress axis, amount of ground substance, and mineral content. Thus, even within the mandible of one species, held constant with respect to age, sex, and metabolic state, there are substantive differences in architecture and physical/mechanical properties. Location-dependent properties might imply a structure-function relationship in the mandible.

In general, the material and mechanical properties of craniofacial bone are nonuniform and vary as functions of anatomic location, age, sex, and metabolic state. Variations in these properties are functions of variations in composition and microstructure at a local level. The processing-composition-structure-property synergy and mechanistic understanding of synthetic materials at an atom level, discussed previously, holds true for tissue also. In biological materials, it is necessary to understand mechanisms at the cellular and molecular levels. An understanding of regional properties will provide a better understanding of localized bone regeneration, repair, modeling, remodeling, and disease states and possibly facilitate the design of site-specific dental implants and bone augmentation materials.

**Materials Used in Dental Implants**

**Introduction**

In general, two basic classes of materials—metals and ceramics—are used in dental implants, either alone or in hybrid fashion (Fig 23-8). Metallic implant materials are largely titanium based—either commercially pure titanium or Ti-6Al-4V alloy.
However, as already stated, it is essential to recognize that the synergistic relationship between processing, composition, structure, and properties of both the bulk metals and their surface oxides effectively leaves more than two metals. Casting, forging, and machining of metal implants, densification of ceramics, deposition of ceramic and metal coatings onto metal implants, as well as cleaning and sterilization procedures, can all alter the microstructure, surface chemistry, and properties. Thus, the many material-processing sequences necessary to yield a dental implant strongly influence implant properties and functionality, primarily through temperature and pressure effects.

**Fig 23-8** Classification of dental implant materials.

**Metals**

Although cobalt-based alloys have been used experimentally in dentistry, metallic dental implants are almost exclusively titanium based. A good deal of the knowledge about titanium stems from the extensive aerospace and metallurgy literature. Many requirements of an aerospace component, primarily high strength and corrosion resistance, are characteristic properties needed in a dental implant. Thus, titanium has been called the "material of choice" in dentistry because of its strength and the minimal biological response it elicits. The strength of titanium is due to its hexagonal close-packed crystal lattice and crystallographic orientation, whereas its biocompatibility (corrosion resistance) is attributed to its stable, passive oxide layer.

Titanium-based implants are in their passive state (i.e., their oxide is stable) under typical physiological conditions and breakdown of passivity should not occur. Both cp titanium and Ti-6Al-4V possess excellent corrosion resistance for a full range of oxide states and pH levels. It is the extremely coherent oxide layer and the fact that
titanium repassivates almost instantaneously through surface-controlled oxidation kinetics that renders titanium so corrosion resistant. However, even in its passive condition, titanium is not inert. Titanium ion release that does occur results from chemical dissolution of titanium oxide. However, the low dissolution rate and near chemical inertness of titanium dissolution products allow bone to thrive and therefore osseointegrate with titanium. Surface potential may also play an important role in facilitating osseointegration, and it is postulated that oxides with high dielectric constants inhibit the movement of cells to implant surfaces.

The Ti-6Al-4V alloy has a 60% greater strength than pure titanium, but it is more expensive. Both materials have complex, heterogeneous surface oxides. There may be differences in cell adhesion, and tissues may be in closer proximity to pure titanium surfaces than to alloy surfaces. However, there does not seem to be any difference in implant function, and there is no clear rationale for choosing pure titanium over titanium alloy.

The mechanical properties of titanium-based materials are well established. Microstructures with a small (< 20 um) α-grain size, a well-dispersed second (β)-phase, and a small α-β interface area, such as equiaxed microstructures, resist fatigue crack initiation best and have the best high-cycle fatigue strength (approximately 500 to 700 MPa). Lamellar microstructures, which have a greater α-β interface area and more oriented phase colonies, have lower fatigue strengths (approximately 300 to 500 MPa). Surface roughening, whether through screw threading or deposition of coatings, results in a reduced strength compared to smooth-surfaced implants.

Ceramics

The initial rationale for using ceramics in dentistry was based on the relative biological inertness of ceramics compared to that of metals. Ceramics are fully oxidized materials and therefore chemically stable. Thus, ceramics are less likely to elicit an adverse biological response than metals, which only oxidize at their surface. Two types of "inert" ceramics of interest are carbon and alumina (Al₂O₃). Recently, a greater emphasis has been placed on bioactive and bioreabsorbable ceramics, materials that not only elicit normal tissue formation, but may also form an intimate bond with bone tissue and even be replaced by tissue over time. While inert ceramics elicit a minimal tissue response, bioactive ceramics are partially soluble, enabling ion transfer and the formation of a direct bond between implant and bone. Bioreabsorbable or biodegradable ceramics have a higher degree of solubility than bioactive ceramics, gradually resorb and integrate into the surrounding tissue, and are used as bone augmentation materials. Bioactive ceramics are primarily used as scaffold materials or as coatings on more structurally sound metal substrates.

The concept of bioactivity was originally introduced with respect to bioactive glasses via the following hypothesis: The biocompatibility of an implant material is optimal if the material elicits the formation of normal tissues at its surface, and, in addition, if it establishes a contiguous interface capable of supporting the loads that normally occur at the site of implantation. Important examples of these materials are bioactive glasses, glass ceramics, and calcium phosphate ceramics. Bioactive glasses and glass ceramics include Bioglass, which is a synthesis of several glasses containing mixtures of silica, phosphate, calcia, and soda; Ceravital, which has a different alkali
oxide concentration from that of Bioglass; and glass ceramic A-W, a glass ceramic containing crystalline oxyapatite and fluorapatite \([\text{Ca}_{10}(\text{PO}_4)_6(\text{O,F}_2)]\) and \(\beta\)- wollastonite (\(\text{SiO}_2\)-\(\text{CaO}\)) in a \(\text{MgO-CaO-SiO}_2\) glassy matrix. The calcium phosphate ceramics can have varying calcium-to-phosphate ratios, depending on processing-induced physical and chemical changes. Among them, the apatite ceramics, one of which is hydroxyapatite, have been studied most and are the focus of this section.

The impetus for using synthetic hydroxyapatite as a biomaterial stems from the perceived advantage of using a material similar to the mineral phase in bone and teeth for replacing these materials. As such, better tissue bonding is expected. Additional perceived advantages of bioactive ceramics include low thermal and electrical conductivity, elastic properties similar to those of bone, control of in vivo degradation rates through control of material properties, and the possibility of the ceramic functioning as a barrier to metallic corrosion products when it is coated onto a metal substrate. However, processing-induced phase transformations provoke a change in in vitro dissolution behavior, and the different structures and compositions alter the biological response. Given the range of chemical compositions available in bioactive ceramics and the fact that pure hydroxyapatite is rarely used, the broader term calcium phosphate ceramics (CPC) should be used in lieu of the more specific hydroxyapatite. Each individual CPC is defined by a unique set of chemical and physical properties.

Mixtures of hydroxyapatite, tricalcium phosphate, and tetracalcium phosphate may evolve as a result of plasma-spraying deposition processes onto metals. Physical properties of importance to the functioning of calcium phosphate ceramics include:

1. Powder particle size and shape
2. Pore size, shape, and distribution
3. Specific surface area
4. Phases present
5. Crystal structure and size
6. Grain size
7. Density
8. Coating thickness, hardness, and surface roughness

There are variable and conflicting data with respect to implant-tissue bond strength, solubility, and overall in vivo function. In addition, long-term stabilization and fixation strength do not depend on the ceramic coating. The ceramic-metal bond fails before the ceramic-tissue bond and is the "weak link" in the system. Thus, there is reason for concern about the weak ceramic-metal bond and integrity of that interface over a lengthy service life of functional loading.
Problems and Future Directions

Introduction

Although there is no consensus regarding methods of evaluating dental implants and what parameters are most important, clinical evaluations have generally shown that dental implants are successful in at least 75% of cases 5 years postplacement. Despite advances in materials synthesis and processing, surgical technique, and clinical protocols, clinical failures do occur, at rates of approximately 2% to 5% per year. Causes of failure and current problems with dental implants include:

1. Early loosening, stemming from a lack of initial osseointegration
2. Late loosening, or loss of osseointegration
3. Bone resorption
4. Infection
5. Fracture of the implant and/or abutment
6. Delamination of the coating from the bulk implant

The most common failure mechanism with endosseous implants is alveolar crest resorption, leading to progressive periodontal lesions, decreased areas of supporting tissues, and ultimately implant loosening. Aseptic failures are most often the cumulative result of more than one of the above-mentioned factors.

As a result of these clinical problems, basic and clinical research should focus on the complete characterization of materials, including bulk and surface properties, development of new materials, more engineering-based designs for both existing and new materials, fundamental aspects and quantification of stresses and stress transfer between implant and tissue, mechanical and biological responses of tissues, and host response to these foreign materials.

Future materials

Although titanium and, to a lesser extent, ceramic and ceramic-coated implants have an excellent clinical record in implant dentistry, these materials are not necessarily end-stage materials. Continuing developments in the materials and biomedical fields can be expected in the next decade. Because one of the long-term problems with dental implants is stress shielding, or mechanically mediated bone resorption, which is due in part to the elastic mismatch between metal and bone, polymer and composite implants that offer reduced moduli are being considered. The motivation for using composite materials for implants is based on several concepts. Composite materials can be very strong, because materials in fiber form exhibit strengths near the theoretical values. As a result, advanced composites can be as strong as metals and, in some cases, more flexible. The properties of composites can be more easily tailored than those of metals. A specific example is that of the modulus of composites, which can be tailored to some extent to be near that of bone. The rationale for designing
"Isoelastic" implants, or implants with the same modulus as bone, is based on observations of bone resorption in the presence of stiff metal implants. This rationale is based on the hypothesis that an implant that matches the elastic properties of the surrounding tissue will result in a more physiological stress distribution than can be attained with higher modulus metallic implants.

**Augmentation Materials**

One of the oldest biomaterials problems has been the search for materials that can repair or replace bone defects. The standard materials, historically, have been bone grafts. However, given the morbidity associated with autogenous grafts and recent concern about transmission of live viruses with allogenic grafts, increased research into alternative substitute materials, such as ceramics, polymers, composites, bone derivatives, and natural materials, is underway (Fig 23-9). Examples of dense and porous calcium phosphate ceramics are shown in Figs 23-10 and 23-11. Most recently, combinations of synthetic materials and biological constituents (eg, cells or growth factors) have been suggested.

![Classification of bone augmentation materials](image)

**Fig 23-9** Classification of bone augmentation materials.

The three primary application areas for augmentation materials in dentistry are intramucosal, endodontic, and bone-substitute materials. An ideal bone substitute material should be biocompatible; easy to fabricate, sterilize, and shape...
intraoperatively; inexpensive; osteoinductive; and osteoconductive. It should also offer mechanical integrity over a lengthy service life.

**Fig 23-10** (a) Dense hydroxyapatite ceramic augmentation material with starting powders. (From Denissen et al, 1985. Reprinted with permission.) (b) Porous hydroxyapatite augmentation material, 44 x 18 x 16 mm; porosity = 45%. (From Osborn, 1985. Reprinted with permission.)

**Fig 23-11** Porous coralline hydroxyapatite ceramic augmentation material (Pro Osteon Implant 500, Interpore Orthopaedics).
**Glossary**

**calcium phosphate ceramics** A class of ceramics with varying calcium-to-phosphate ratios, which can form a direct bond with bone.

**hydroxyapatite** A specific form of calcium phosphate with a stoichiometry $\text{Ca}_{10}(\text{PO}_4)_6(\text{OH})_2$ and a Ca/P ratio of 1.67. Bone is a nonstoichiometric form of hydroxyapatite.

**interfacial zone** The thin zone at the surface of an implant, which includes the surface oxides, protein layers, and connective tissue.

**osseointegration** A direct structural and functional connection between ordered, living bone and the surface of a load-carrying implant.

**osteconductive material** A material that acts as a scaffold for new bone formation.

**osteoinductive material** A material that causes the conversion of mesenchymal cells preferentially to bone progenitor cells.

**titanium** The "material of choice" in dentistry, primarily because of its excellent biocompatibility (as a result of its stable oxide layer), mechanical properties, and, in implant dentistry, its proven ability to achieve osseointegration.

**Discussion Questions**

1. What are the properties (bulk and surface) of titanium that make it attractive for use as a dental implant?

2. What is/are the rationale(s) for using bioactive ceramics (eg, hydroxyapatite) as coatings on dental implants?

3. What is the importance of implant-tissue interfacial stability, and what are current methods of accelerating osseointegration such that the time between first- and second-stage implant surgery may be reduced?

4. What are the physical, mechanical, and biological parameters affecting the clinical success of dental implants?

**Study Questions**

*(See Appendix E for answers.)*

1. What is osseointegration?

2. What materials are used for osseointegrated implants?

3. What factors dictate the effectiveness of osseointegration?
4. What parameters influence implant success?

**Recommended Reading**


Adhesion of restorative materials to tooth structures has been studied since 1955. The bond strength is the load required to fracture the bond divided by the cross-sectional area of the bond. Bond strengths are measured either in shear or tension. Many factors, including the dentinal substrate, the storage conditions, and the test method, affect the bond strength values.

### Shear bond strength

<table>
<thead>
<tr>
<th>Substrate</th>
<th>Adhesive</th>
<th>Adherend</th>
<th>psi × 10³</th>
<th>MPa</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dentin</td>
<td>Dentin bonding system</td>
<td>Composite</td>
<td>3.2-5.1</td>
<td>22-35</td>
<td>238</td>
</tr>
<tr>
<td>Dentin</td>
<td>No smear layer</td>
<td>Light-cured hybrid glass ionomer</td>
<td>1.5-1.7</td>
<td>10-12</td>
<td>238</td>
</tr>
<tr>
<td>Enamel</td>
<td>Enamel bonding system</td>
<td>Composite</td>
<td>2.6-3.2</td>
<td>18-22</td>
<td>238</td>
</tr>
<tr>
<td>Enamel</td>
<td>Amalgam bonding system</td>
<td>Composite</td>
<td>1.5-1.7</td>
<td>10-12</td>
<td>238</td>
</tr>
<tr>
<td>Enamel</td>
<td>Amalgam bonding system</td>
<td>Amalgam</td>
<td>0.3-0.9</td>
<td>2-6</td>
<td>238</td>
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<tr>
<td>Enamel</td>
<td>No smear layer</td>
<td>Traditional glass ionomer</td>
<td>1.2-1.7</td>
<td>8-12</td>
<td>238</td>
</tr>
<tr>
<td>Enamel</td>
<td>Enamel bonding system</td>
<td>Orthodontic bracket</td>
<td>2.6-2.9</td>
<td>18-20</td>
<td>238</td>
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</table>

### Tensile bond strength

<table>
<thead>
<tr>
<th>Substrate</th>
<th>Adhesive</th>
<th>Adherend</th>
<th>psi × 10³</th>
<th>MPa</th>
<th>Reference</th>
</tr>
</thead>
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<tr>
<td>Dentin</td>
<td>Polyurethane</td>
<td>Composite</td>
<td>0.1-0.9</td>
<td>1-6</td>
<td>147</td>
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<tr>
<td>Dentin</td>
<td>Polyacrylic acid</td>
<td>Composite</td>
<td>0.3-0.6</td>
<td>2-4</td>
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</tr>
<tr>
<td>Dentin</td>
<td>Organic phosphonates</td>
<td>Composite</td>
<td>0.4-1.4</td>
<td>3-10</td>
<td>147</td>
</tr>
<tr>
<td>Dentin</td>
<td>4-META</td>
<td>Composite</td>
<td>0.4-1.0</td>
<td>3-7</td>
<td>147</td>
</tr>
<tr>
<td>Dentin</td>
<td>HEMA + GA^a</td>
<td>Composite</td>
<td>1.6-2.4</td>
<td>11-17</td>
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<tr>
<td>Dentin</td>
<td>NPG-GMA/PMDM^b</td>
<td>Composite</td>
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<tr>
<td>Dentin</td>
<td>Glass ionomer</td>
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<tr>
<td>Dentin</td>
<td>Zinc polycarboxylate</td>
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<td>0.4-0.6</td>
<td>3-4</td>
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<tr>
<td>Dentin</td>
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<td>Dentin</td>
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<tr>
<td>Etched enamel</td>
<td>Fine composite</td>
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<td>2.4-2.8</td>
<td>17-20</td>
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</tr>
<tr>
<td>Etched enamel</td>
<td>Microfine composite</td>
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<td>10</td>
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<td>0.7</td>
<td>5</td>
<td>147</td>
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<tr>
<td>Etched</td>
<td>Adhesive resin</td>
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<td>2.2</td>
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<td>147</td>
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<td>Enamel</td>
<td>Cement</td>
<td>Conventional resin</td>
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<td>10</td>
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<td>--------------</td>
<td>-----------------</td>
<td>--------------------</td>
<td>-----</td>
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</tr>
</tbody>
</table>

*a* 4-META = 4-methoxyethyl-trimellitic anhydride.

*b* HEMA = hydroxyethyl methacrylate; GA = glutaraldehyde.

*b* NPG-GMA = N-phenyl glycine--glycidyl methacrylate; PMDM = pyromellitic acid diethylmethacrylate.
Appendix B. Biocompatibility Tests

Initial Tests

Cytotoxicity tests One of the primary criterion for biocompatibility is that the material not be toxic to cells. This means that the material must not affect the cell number or growth, the integrity of the cell membranes, the activity (primarily enzymatic) of the cells, the genetic integrity, or the genetic expression. These tests are designed to determine how a material sample affects a particular cell type. The results, however, yield little information on how the sample affects a whole organism.

Mutagenesis tests These tests assess how the sample affects (or mutates) the genetic material of the host cells, either directly or indirectly, through a degradation product of the sample. An example is the Ames test, in which histidine-dependent mutants are exposed to the sample and the rate of reversion to nonhistidine-dependent strains is used as a measure of mutagenic potential.

Immune function These tests detect immune responses to the sample, including proliferation, differentiation, chemotaxis, cytokine production, and bioactivity changes of immune cells such as T-cells, macrophages, and lymphocytes. Immune function tests are not used as often as many of the other biocompatibility tests, possibly in part due to the difficulty in quantifying the results and interpreting their meaning/severity. Some researchers believe that a mild immune response is not a substantial problem and may even be beneficial as a method of programmed biodegradation.

Complement activation assay These tests are primarily concerned with engineered blood vessels and other devices that are in contact with the blood. Specifically, it determines whether the sample activates complement that will initiate thrombosis or generate inflammation. This is less of an issue for dental materials, but is pertinent with respect to a material’s effect on the pulp and gingiva.

Hemolysis assay These tests are concerned with a material's potential to damage red blood cell membranes, either directly by permeating the membrane or by lysis. Such damage may result in malfunction or death of the red blood cells. A material is screened for this effect by searching for cell components in the medium's solution that should be internalized in healthy cells. Conversely, this can be identified by using tracers that should be excluded from the cell but have been allowed to permeate due to membrane damage.

Oral and intraperitoneal LD50 assays These tests determine acute lethal effects administered orally or intraperitoneally to the lab specimen, typically rats. The sample is mixed into a solution or suspension of water, propylene glycol, vegetable oil, etc, and then injected into the stomach or peritoneum. The standard LD50 is the minimum amount that will kill 50% of the animals in 2 weeks. If this amount is less than 1 g sample per 1 kg body weight, then it is considered to have acute systematic toxicity.
Secondary Tests

*Mucous membrane irritation test* This test determines if there is an inflammatory response when in contact with mucous membranes or abraded skin. The skin sample is usually taken from the oral tissue of hamsters or rabbits. After a period of weeks, the test samples are compared to controls to evaluate the inflammatory response. Photographs are taken, followed by biopsies for a more in-depth analysis.

*Skin sensitization tests* For these tests, materials are injected intradermally to test for hypersensitivity. The reaction can be augmented with Freund's adjuvant. Hypersensitivity is subsequently tested via adhesive patches containing the test substance. The frequency and severity of inflammation in the test subjects is a measure of the allergenic sensitization of the material.

*Implantation tests* These tests are required for samples that will contact tissue or bone and evaluate a material's potential to cause chronic inflammation or tumor growth. After inflammation from surgery has subsided, the samples are implanted in plastic tubes at sites that correspond with its intended use. The response is then analyzed using histological, histochemical, and/or immunohistochemical methods. Short-term tests last 1 to 11 weeks. Long-term tests are performed similarly but last 1 to 2 years. Other methods of implantation are being developed due to the complicating tissue reactions caused by some plastic tubes.

Usage Tests

*Dental pulp irritation tests* For this test, the materials are placed into Class 5 cavity preparations in intact, noncarious teeth of monkeys or other suitable test subjects. The cavities are prepared in all types of teeth under sterile conditions with a water-spray coolant to minimize pulpal trauma and should be of uniform size and shape. Zinc oxide-eugenol and silicate cement are used as negative and positive controls, respectively. After a period of 1 to 8 weeks, the teeth are extracted and sectioned for microscopic evaluation. Necrotic and inflammatory responses are analyzed. The thicknesses of the remaining and reparative dentin are quantified. The pulpal response is classified as slight, moderate, or severe based on this data. These tests are presently under development, particularly in attempt to account for bacterial insults to the pulp.

*Dental implants into bone* There are three basic evaluations for the success of implants: the penetration of a periodontal probe along the side of an implant, mobility of the implant, and bone-implant integration. A successful implant should have no mobility, no peri-implant radiolucency, minimal vertical bone loss, minimal fibrous encapsulation (preferably none), and no soft tissue complications. Ideally, the bone will contact the implant with strong, healthy tissue without any fibrous encapsulation.

*Mucosa and gingiva tests* These tests are performed in cavity preparations with subgingival extensions. The effects are typically examined at 7 and 30 days. Based on the degree of inflammation and the reaction of the epithelium, the responses are categorized as slight, moderate, or severe. Complications include pre-existing inflammation; inflammation due to preparation, bacterial effects, or surface properties (ie, roughness) of the implant; and poorly designed contours. Prophylaxis minimizes complications due to plaque, but causes minor inflammatory complications of its own.
# Appendix C. Periodic Chart of the Elements

## Periodic Chart of the Elements

<table>
<thead>
<tr>
<th>GROUP</th>
<th>I</th>
<th>II</th>
<th>3</th>
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<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>1</th>
<th>2</th>
<th>III</th>
<th>IV</th>
<th>V</th>
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<tr>
<td>VALENCES</td>
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<td>+2</td>
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<td></td>
<td></td>
<td></td>
<td>VARIABLE</td>
<td></td>
<td>+3</td>
<td>-4+4</td>
<td>-3+5</td>
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<td>4</td>
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<td>(226)</td>
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<td></td>
<td></td>
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</tr>
</tbody>
</table>

### Lanthanum Series

| | Ce | 140.112 | Pr | 140.9077 | Nd | 144.24 | Pm | 145 | Sm | 150.35 | Eu | 151.964 | Gd | 157.25 | Tb | 158.924 | Dy | 162.50 | Ho | 164.93 | Er | 167.26 | Tm | 168.93 | Yb | 173.04 |
| | | (233) | | | | | | | | | | | | | | | | | | | | | | | | |

### Actinium Series

| | Th | 232.036 | Pa | 231.036 | U | 238.03 | Np | 237 | Pu | 244 | Am | 243 | Cm | 245 | Bk | 247 | Cf | 248 | Es | 249 | Fm | 254 | Md | 258 | Lr | 261 | |
Appendix D. Units and Conversion Factors

**Unit SI equivalent**
- Angstrom (Å) $1 \times 10^{-10}$ meter (m)
- Nanometer (nm) $1 \times 10^{-9}$ meter (m)
- Micrometer (µm) $1 \times 10^{-6}$ meter (m)
- Inch (in) 0.0254 meter (m)
- Centimeter (cm) 2.54 meter (m)
- Pound (lb) 0.4536 kilogram (kg)
- Newton (N) dyne $1 \times 10^{-5}$ newton (N)
- Pound force (lbf) 4.4482 newton (N)
- Joule (J) Calorie (Cal) 4.1868 joule (J)
- Btu 1055.06 joule (J)
- Newton/meter² (N/m²) atmosphere 1.013 $\times 10^{5}$ N/m² (Pascal, Pa)
- Pound per square inch (psi) 6.895 $\times 10^{3}$ N/m² (MPa)
- Kilogram per square centimeter (kg/cm²) 9.804 $\times 10^{4}$ MN/m² (Pa)
- Pennyweight (dwt) (Troy) = 1.555 g
- 20 dwt = 1 ounce (Troy)
- 1.097 ounce (Avoirdupois)
- Photomicrograph distance: $\mu$m/cm = 10,000 / magnification
- Temperature: $(^\circ C \times 1.8) + 32 = ^\circ F (^\circ F - 32)/1.8 = ^\circ C$

**Prefixes for SI units**
- Multiply by this factor
- Symbol Prefix
- 10⁻¹² T tera
- 10⁻⁹ G giga
- 10⁻⁶ M mega
- 10⁻³ k kilo
- 10⁻² h hecto
- 10⁻¹ da deca
- 10⁻² d deci
- 10⁻³ c centi
- 10⁻⁶ m milli
- 10⁻⁹ µ micro
- 10⁻¹² n nano
- 10⁻¹⁵ p pico
- 10⁻¹⁸ f femto
Appendix E. Answers to Study Questions

Chapter 1 A Comparison of Metals, Ceramics, and Polymers

1. See Table 1-1.

2. The sharper the stress raiser, the greater the concentration of stress around it.

3. Ceramics—Stress concentration at surface scratches and other defects causes ceramics to fail at stresses far below their theoretical strengths.

Metals—The ability of crystal defects called dislocations to move within the crystal structure of metals gives them their ability to bend without fracturing.

Polymers—Strong bonds within the polymer chains and weak bonds between polymer chains are responsible for the low strengths and elastic moduli of polymers.

Chapter 2 Physical Properties and Biocompatibility

1. Knowledge and understanding of physical and mechanical properties help the dentist predict how a material will behave in vivo and how it should be manipulated.

2. Thermal diffusivity is a time-dependent property, and, since the temperature of the oral cavity changes dramatically within a few seconds (drinking hot coffee versus eating ice cream), it predicts the behavior of materials under more realistic conditions. Thermal conductivity, on the other hand, is a steady-state property, and the temperature of the mouth is not constant over time.

3. See Fig 2-4.

![Fig 2-4](image)

Fig 2-4 Measurement of stress and strain on an object being stretched. In the region labeled "elastic deformation," stress is proportional to strain, whereas in the region labeled "plastic deformation," stress and strain are no longer proportional (strain increases faster than stress does). The highest stress at which stress and strain are still proportional is called the proportional limit, and the maximum stress just before the object breaks is called the ultimate tensile strength. The total amount that the object stretches (i.e., the total strain), which is the sum of the elastic deformation and the plastic deformation, is called the percent elongation.
4. Hardness is the resistance of a material to permanent indentation in its surface. It involves complex stresses, so it cannot be directly related to any other physical property.

5. \( a \) is brittle, \( b \) is ductile; \( a \) is stiffer than \( b \); \( a \) is stronger than \( b \), \( b \) is tougher than \( a \).

6. Laboratory testing is necessary to screen out materials that could be obviously harmful before use with patients in a clinical trial. Both might be performed if initial screening tests are favorable.

**Chapter 3 Color and Appearance**

1. Hue, value, and chroma; \( X, Y, \) and \( Z \) tristimulus values; \( L^*, a^*, \) and \( b^* \).

2. A crown that appears gray and nonvital has a low value.

3. The green object absorbs the blue light. Since no light is reflected, the object appears black.

4. The source, the surroundings, and the observer.

5. A high translucency gives a lighter color appearance.

6. A high gloss lightens color appearance.

7. The change in color matching of two objects under different light sources.

8. Metamerism may cause a restoration to have a good color match under one lighting condition but a poor color match under another lighting condition.

9. Dental porcelains are fluorescent in some lighting environments.

**Chapter 4 Gypsum Products**

1. The major constituent is the same in all products—calcium sulfate hemihydrate. Minor differences in composition exist between stones and die stones because of the presence of accelerators, retarders, and coloring matter.

2. Plaster is produced by the dry calcination of ground gypsum (calcium sulfate dihydrate). Partial loss of water of crystallization is not accompanied by a change in shape or size of the individual particles, which are therefore irregular and porous, giving the powder a relatively low apparent density. Stone and die stone are produced by wet calcination, which allows formation of dense regular crystals of hemihydrate, producing powders that have a higher apparent density than plaster. Grinding under controlled conditions increases the apparent density still further.

3. Typical \( W/P \) ratios: plaster, 0.50; stone, 0.30; die stone, 0.20. Lower \( W/P \) ratios can be used with stone and die stone, because the apparent densities of their powders are higher, so less water is needed to make a workable mix. Uncombined water in set
mass: for plaster, 100 g powder + 50 g water = 150 g set mass. Only 18.6 g water combines chemically. Therefore, after the setting reaction is complete, uncombined water is 31.4 g in 150 g, approximately 20%. Similarly, for stone, uncombined water is approximately 9%; for die stone, approximately 1%. (Note: These are theoretical calculations. In actual practice the setting reaction does not reach completion, even in relatively high W/P ratios.)

4. **Chemical change:** Hydration of calcium sulfate hemihydrate to form the dihydrate (gypsum): \(2\text{CaSO}_4 \cdot \frac{1}{2}\text{H}_2\text{O} + 3\text{H}_2\text{O} \rightarrow 2\text{CaSO}_4 \cdot 2\text{H}_2\text{O}\)

**Physical changes:** The mix is first a viscous liquid, then a plastic mass, and then a rigid solid, friable at first but increasing in strength until the reaction has ceased.

5. Set plaster has a higher proportion of inherent microporosity than set stone. The microporosity is caused by (a) residual unreacted water and (b) setting expansion. Both of these factors are greater in the case of plaster. Thus, set plaster is less dense and therefore weaker than set stone.

6. Setting is a continuous process. In practice, any figure given for setting time is arbitrary, unless it refers to the time when the setting reaction ceases. (a) Loss of surface gloss gives an indication of the time when the mix will not pour or flow under vibration. (b) and (c) Both the Gillmore and Vicat initial sets give an indication of the time when the solid material can be handled safely.

7. By the empirical addition of accelerators or retarders to the raw hemihydrate powder.

8. The setting rate is reduced, because increasing the W/P ratio increases the relative amount of aqueous phase present in the mix. Therefore, the physical changes associated with setting occur more slowly because interaction of growing gypsum crystals takes place later.

9. There is a decrease in the volume of the actual solid. However, the gypsum crystals formed are usually long and thin, and crystal growth creates microscopic voids that cause an increase in the total volume of the mass.

10. When setting occurs in water, there is a continuous aqueous phase present and crystal growth is relatively unimpeded. When setting occurs in air, the water content of the mix decreases as the reaction proceeds. Residual liquid in the mix is drawn into voids created between growing gypsum crystals, over which it forms a film. Surface tension forces crowd the growing crystals together and restrain further expansion.

11. It affects the dimensional accuracy of the ultimate dental restoration.

12. Impression plaster, 0.15%; plaster, 0.30%; stone, 0.15%; low-expansion die stone, 0.10%; high-expansion die stone, 0.25%.

13. By the empirical addition of a blend of a suitable accelerator and retarder, both of which reduce setting expansion.
14. Accelerators and retarders also reduce the strength properties of the set material, so when strength is an important consideration the concentration that can be used is relatively low.

15. Setting expansion is reduced, because the relative amount of aqueous phase is increased and interaction of growing gypsum crystals is less effective.

16. Tensile strength is an indication of resistance to fracture; compressive strength is an indication of surface hardness.

17. It reduces both. With a higher W/P ratio there is more residual water remaining after setting is complete, and this causes increased microporosity which reduces strength properties.

18. Drying approximately doubles both tensile and compressive strengths, since the presence of free water in the cast material weakens it.

19. Calcium sulfate hemihydrate acts as a binder. Silica (quartz, cristobalite, or a mixture of both) offsets the contraction of the binder that occurs on heating and often provides a positive thermal expansion. Modifiers may be present in small concentrations to adjust the rate of setting, to give a reducing atmosphere when the mold is heated, or to give an increased thermal expansion.

20. (a) Rate of setting decreases; (b) setting expansion in air increases; (c) setting expansion in water increases; (d) thermal expansion increases; and (e) compressive strength decreases.

21. When the investment powder is mixed with water, the silica is unaffected. The calcium sulfate hemihydrate reacts with water to precipitate gypsum crystals, which bind the silica particles together.

22. The porosity is continuous and provides venting of the mold cavity when the molten alloy is cast.

23. The powder should have a uniform fine particle size (not more than 75 um). A uniform particle size ensures adequate venting of the mold cavity, because the silica particles show least-dense packing. A fine particle size produces a smooth casting; surface roughness of the casting can interfere with its fit.

24. The particles of refractory filler remain unchanged during setting and have a major effect on the smoothness of the mold surface. The particles of binder are dissolved during setting, and gypsum crystals are precipitated that are much smaller than the refractory particles.

25. Expansion, provided by a summation of setting and thermal expansions, is most important. Mold expansion is used to offset the casting shrinkage of the alloy and thus produce an accurate casting.

26. A casting technique in which most of the mold expansion is gained by the increased setting expansion that occurs when the setting investment is exposed to
additional water. The need for thermal expansion of the mold is small.

27. A casting technique in which thermal expansion contributes greatly to mold expansion. In most thermal expansion techniques, setting expansion also plays an important part.

28. At the relatively low burnout temperatures used, oxidation of carbon remaining from the wax pattern is slow. Most of it must be removed to allow adequate mold venting.

29. The refractory can be identified by studying the investment's thermal expansion curve, which should be provided by the manufacturer. Cristobalite investments show a large isothermal expansion at about 250°C. Quartz investments show a smaller isothermal expansion at about 570°C. The type of refractory filler in the investment affects the heating rate to be used and sometimes the burnout temperature.

30. Cristobalite investments undergo a large isothermal expansion at a relatively low temperature (250°C). If a large temperature difference, resulting from rapid heating, exists within the mold at this temperature, nonuniform expansion can cause cracking. This can be prevented by following a slow heating rate below 300°C. The problem does not arise with quartz investments, which have a more gradual thermal expansion. (Note: The need for slow heating during the inversion of cristobalite does not apply to rapid-heat investments.)

31. With quartz investments, the required thermal expansion is available only over restricted temperature ranges (hygroscopic expansion techniques, 350°C to 480°C; thermal expansion techniques, 600°C to 700°C). With cristobalite investments, the thermal expansion is reasonably constant over a wide temperature range (400°C to 700°C).

32. At temperatures above 700°C the calcium sulfate binder decomposes in the presence of carbon, liberating corrosive gases that can adversely affect the casting.

33. It increases it.

34. (a) Setting expansion in air decreases; (b) setting expansion in water decreases; and (c) thermal expansion decreases.

35. By varying the W/P ratio, or by varying the length of time additional water is available to the setting investment. In the latter case this may be achieved by delaying the immersion of the investment, or by adding controlled amounts of excess water to the setting investment instead of immersing it.

36. By varying the W/P ratio.

37. At the casting temperature, the investment mold may be strong enough to restrain the thermal contraction of the solidified casting as it cools. The effect will vary in different directions, depending on the amount of interlocking of casting and mold. This can cause distortion of the casting shape.
An investment with a low hot strength is more likely to allow uniform thermal contraction of the alloy and give an undistorted casting. A safe lower limit for compressive strength at the casting temperature is 1.8 MPa.

38. They should be stored in airtight containers, in a cool dry region of the laboratory. When hemihydrate powders are exposed to the atmosphere, particularly when the water vapor pressure is high, the particles adsorb water from the air. This can react with the hemihydrate to form gypsum crystals. Initially this increases the setting rate of the material, but if deterioration proceeds further, the setting rate is retarded, and eventually the strength properties of the set material are adversely affected.

**Chapter 5 Surface Phenomena and Adhesion to Tooth Structure**

1. The molecules at the surfaces of liquids exert a greater intermolecular force on each other than do interior molecules, due to their greater average separation. This attraction, or pull, results in a contractile surface stress.

2. The degree of capillary penetration decreases as the contact angle increases. Above a contact angle of 90 degrees, a liquid will not penetrate and depression occurs (eg, mercury in a glass tube).

3. Adsorption involves the uptake of one substance at the surface of another. Absorption involves the penetration of one substance into the interior of another.

4. Agar and alginate impression materials, colloidal gold, and detergents.

5. The penetration coefficient includes viscosity, surface tension, and contact angle as follows (PC in cm/sec):

\[
PC = \frac{\gamma \cos \theta}{2 \eta}
\]

6. Major, since previously there was no way to bond materials to dentin.

**Chapter 6 Polymers and Polymerization**

1. Heat- and cold-cured acrylic resins; rubber- and fiber- reinforced acrylic; polycarbonate injection molded; nylon injection molded; and light-activated dimethacrylate.

2. Good appearance, high flexural strength, high impact resistance, high stiffness, long fatigue life, high craze resistance, high creep resistance, high radiopacity, low free-monomer content, good adhesion with teeth and liners, low solubility, low water uptake, dimensional stability, and dimensional accuracy.

3. *Advantages:* Good appearance, high glass-transition temperature, ease of fabrication, low capital costs, good surface finish
**Disadvantages:** Free-monomer content or formaldehyde can cause sensitization, low impact strength, flexural strength low enough to penalize poor denture design, fatigue life too short, radiolucency

4. **Advantages:** Easy to deflask, dimensionally accurate, can have higher flexural strength than heat-cured materials

**Disadvantages:** No cheaper over the long term, increased creep, increased free-monomer content, color instability, reduced stiffness, tooth adhesion failure

5. **Advantage:** Improved impact strength

**Disadvantage:** Reduced stiffness

6. **Advantages:** High stiffness, very good impact strength, good fatigue life, polypropylene fibers: good translucency and good surface finish

**Disadvantage:** Carbon and Kevlar fibers: poor color and poor surface

7. **Advantages:** Dimensional accuracy, low free-monomer content, good impact strength

**Disadvantages:** High capital costs, difficult mold design problems, less craze resistance, less creep resistance

8. **Advantages:** No methyl methacrylate monomer, reduced polymerization shrinkage, time savings, possible better fit than conventional denture base materials, processing procedure requiring little equipment

**Disadvantage:** Somewhat increased elastic deformation during mastication

9. Acrylic, silicone (RTV), and heat-cured silicone.

10. **Advantage:** Resilient

**Disadvantages:** Low tear strength, low bond strength to dentures, attacked by cleansers, buckle in water, poor abrasion resistance

11. **Advantages:** Resilience, adequate bond strength to acrylic, more resistant to aqueous environment and cleansers than RTV

**Disadvantages:** Low tear strength, poor abrasion resistance

12. **Advantages:** High peel strength to acrylic denture base, high rupture strength, some can be polished if cooled, reasonable resistance to damage by denture cleansers

**Disadvantages:** Poor resilience, loses plasticizer in time, some buckle in water

13. Flow under constant force, resilience at high rates of deformation, remain viscous for several days, have high tack to aid retention to denture base.
Chapter 7 Impression Materials

1. Natural resins and waxes provide thermoplastic properties. Stearic acid acts as a lubricant and plasticizer. Inorganic fillers and pigments give control of flow and color, respectively.

2. Thermoplastic.

3. Impression compound has a higher flow. Tray material, with a lower flow, does not record fine detail.

4. Avoid burning in a direct flame or heating in a water bath for long periods.

5. Low thermal conductivity requires time for thorough cooling to prevent distortion during removal.

6. The amounts of filler and water incorporated during kneading control the flow properties.

7. Zinc oxide and eugenol.

8. ZnO + eugenol \rightarrow Zn eugenolate, which forms a solid matrix holding unreacted ZnO.

9. Increases in temperature and/or humidity shorten the setting time.

10. Agar acts as a gelling agent. Borax improves strength. Potassium sulfate provides good surfaces on models or dies. Alkylbenzoates are preservatives.

11. The liquefaction and gelation temperatures are different.

12. Storage in air results in shrinkage; water, in expansion; 100% relative humidity, in shrinkage (syneresis); and potassium sulfate solution, in shrinkage or expansion, depending on the ionic strength of the solution.

13. Sodium alginate and calcium sulfate are reactants to give calcium alginate. Sodium phosphate is a retarder; filler (e.g., diatomaceous earth) controls stiffness; alkali zinc fluorides provide good surfaces on dies and models; and coloring and flavoring are for esthetics.

14. A low water/powder ratio and/or high temperature shorten the setting time.

15. CaSO_4 + Na_3PO_4 \rightarrow Na_2SO_4 + Ca_3(PO_4)_2 provides working time. Na alginate + CaSO_4 \rightarrow Ca alginate + Na_2SO_4, in the presence of water, provides setting time.

16. Alginates do not revert to a sol on heating or by chemical means.

17. A decreased W/P ratio increases strength, tear resistance, and consistency, but decreases working time, setting time, and flexibility.
18. Insufficient spatulation gives a grainy mix and poor recording of detail. Adequate spatulation yields a smooth, creamy mix with a minimum of voids.

19. Decreased water temperature increases the working and setting times of alginites.

20. Alginites and agars have similar properties.

21. Polysulfide cures by condensation of terminal mercaptan groups catalyzed by lead peroxide or other catalysts. Condensation silicones cure by condensation of terminal hydroxyl groups by orthoalkylsilicates to form polymer and alcohol. Addition silicones cure by free-radical polymerization with a platinum catalyst. Polyethers cure by ring opening of the ethylene-imine group.

22. 

<table>
<thead>
<tr>
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<th>Proportioning</th>
<th>Increased Temperature</th>
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<tbody>
<tr>
<td>Polysulfide</td>
<td>Decreases with increased amount of catalyst</td>
<td>Decreases</td>
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<tr>
<td>Silicone</td>
<td>Minimal</td>
<td>Decreases</td>
</tr>
<tr>
<td>Polyether</td>
<td>Decreases with increased amount of accelerator</td>
<td>Decreases</td>
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23. Addition silicones.

24. Addition silicone ≥ condensation silicone ≥ polyether > polysulfide. A large elastic recovery value indicates smaller distortion of the impression on removal.

25. Polysulfide > condensation silicone ≥ addition silicone ≥ polyether.

26. Follow the instructions given by the manufacturer for the impression material used, because the recommended procedure depends on the material.

27. (a) Hydrophilic materials will wet the tooth surface more readily during impression taking and are more readily wet by dental stone. Therefore, they are less likely to entrap air. Also, their wetting characteristics make them easier to electroplate. (b) A single material can be used as both the low- and high-viscosity material for the syringe-tray impression technique. (c) This method of mixing is quick and produces bubble-free mixes.

Chapter 8 Polymeric Restorative Materials

1. Resin composites contain two major components: a polymer matrix and a ceramic filler. The polymer consists of a dimethacrylate oligomer (bis-GMA or UDMA). Larger filler particles may be quartz or any of a number of glasses. The colloidal-size particles of microfine composites are silica. In addition, a silane coupling agent is attached to the filler surface to create a particle-matrix bond during polymerization.

2. Resin composites are commonly used for the restoration of Class 3 and Class 5 cavity preparations, for replacement of fractured incisal edges (Class 4), and for
veneering of facial surfaces of natural teeth. Continued improvement has led to their use in selected, conservative posterior restorations. They are also used in the repair of porcelain and for the cosmetic recontour of anterior teeth.

3. Longevity of conventional resin composites varies. In general, composite restorations in anterior teeth provide adequate serviceability for up to 10 years. Class 3 restorations tend to last longer than those placed in cervical regions because they are exposed to less mechanical abrasion and stress due to tooth flexure. Posterior resin composites have a shorter service life, generally requiring replacement within 7 years. Reasons for replacement of any composite include deterioration of esthetics, wear, microleakage, and recurrent decay.

4. Fine-particle composites generally have superior physical and mechanical properties compared to microfine composites, which have a lower filler content.

5. Microfilled composites contain colloidal silica (0.01 to 0.12 um) as filler. Because the diameter of the particles is less than 0.05 um, they cannot be detected visually, giving the composite a translucent quality and allowing polishing to a high luster.

6. A hybrid composite contains colloidal silica in addition to larger filler particles. The colloidal silica is commonly added to improve certain handling characteristics, such as flow and packability. Hybrid composites can be used for both anterior and posterior restorations, because they combine the physical properties of fine-particle composites and the esthetics of microfills.

7. Polymerization of the polymer matrix after placement results in shrinkage of the matrix and places stresses on the bond at the tooth-composite interface. At present, the enamel-composite bond is more likely to withstand these stresses than is the dentin-composite bond. Loss of marginal integrity results in an influx of oral fluids and greatly increases the possibility of microleakage, postoperative sensitivity, and recurrent decay.

8. Glass ionomers, resin-modified glass ionomers, compomers, and fluoride-releasing composites. Additionally, the materials with higher fluoride release have higher fluoride recharge capacity.

9. The success of a cervical restoration may increase with increased flexibility. Current research suggests that restorative materials with higher flexibility (lower elastic modulus) may be desirable. Flowable composites and resin-modified glass ionomers are more flexible than most all-purpose composites or compomers, and are therefore the materials of choice.

Chapter 9 Dental Cements

1. About 55 MPa (8,000 psi).


3. The solubility in organic acid solutions, such as citric or lactic acid, is 20 to 30
times higher than in water.

4. These cements contain monomers with polar groups, such as phosphate or carboxyl, that improve wetting and are potentially adhesive to the tooth or restorative material surface. Adhesive cements may also utilize a dentin bonding primer system before application of the cement.

5. They are combinations of glass ionomers with resin additions, providing easier handling, dual curing, and higher flexural strength.

6. Water, zinc salts such as acetate and sulfate, and other acidic materials.

7. The high solubility of zinc oxide-eugenol cements is due to hydrolytic breakdown of zinc eugenolate and the extraction of eugenol from the set cement.

8. Mineral fillers, such as silica or alumina; natural resins, such as pine rosin; and synthetic polymers, such as poly(methyl methacrylate), polystyrene, or polycarbonate.

9. Zinc oxide-eugenol cements inhibit the polymerization of resin restorative materials, resulting in softening and discoloration.

10. The powder of EBA cements contains more mineral filler, such as alumina, than does the powder of zinc oxide-eugenol cements, and in the liquid only about one third is eugenol; the remainder is ethoxybenzoic acid.

11. The powder/liquid ratio, the reactivity and particle size of the zinc oxide, the presence of additives, and the molecular weight and concentration of the polyacrylic acid.

12. The mild effect of polycarboxylate cements on pulp may be due to the relatively high pH of the setting cement; localization of the polyacrylic acid molecules; and/or the minimal osmotic effects on fluid in the dentinal tubules.

13. The components of the cement should be carefully proportioned and mixed on a cooled slab. All the powder may be added to the liquid at one time so the total mix time is no more than 30 to 40 seconds. The mix should be used while it is still glossy, before the onset of cobwebbing.

14. They set by both acid-base reaction and polymerization of monomer groups, giving higher early strength, improved physical properties, and water resistance.

15. High strength and stiffness, adhesion, translucency, leachable fluoride and potential cariostatic effect, and good resistance to dissolution in the mouth.

16. The two types of resin cement are acrylic resin cements, based on poly(methyl methacrylate) and methyl methacrylate monomer, and resin composite materials, based on a ceramic filler and a bis-GMA dimethacrylate monomer.

17. Short working time, viscous behavior of the mix, microleakage and pulpal irritation, and difficulty of removal of excess cement.
18. Long-term clinical studies were not performed on hybrid ionomers before marketing. Furthermore, the physical properties studies, including dimensional changes on setting, were performed dry and indicated a shrinkage. Under moist conditions, these cements probably expand by water absorption. Although useful, physical property tests cannot always identify phenomena that clinical studies can locate.

19. Long-term, large-scale clinical studies were not done at different clinics. Short-term studies with small samples performed at one clinic would not detect this problem, since the problem involves overdrying of the tooth as well.

Chapter 10 Abrasion, Polishing, and Bleaching

1. A material that causes wear of another material through mechanical means.

2. Hardness, particle size and shape, speed and pressure, lubrication.

3. Chalk, rouge, pumice, cuttle, sand, aluminum oxide, silicon carbide, diamond.

4. Little effect; it is too soft to be a good polishing agent for dental porcelain.

5. For esthetic and functional reasons (to retard plaque accumulation).

6. Coarse particles would leave scratches on the polished surface. Avoid contamination of the instruments used and have the patient rinse between the grinding and polishing steps.

7. The tongue can distinguish scratch depths between 20 and 2 um.

8. (a) Pumice, sand, zirconium silicate, and chalk. (b) Chalk, dibasic calcium phosphate dihydrate, anhydrous dibasic calcium phosphate, tricalcium phosphate, calcium pyrophosphate, and hydrated alumina. (c) Aluminum oxide, silicon carbide, and sand.

9. Degree of staining, toothbrushing habits, soft restorative materials present in the oral cavity, and amount of exposed cementum or dentin.

10. An amalgam restoration should be polished at least 24 hours after placement. Use flour of pumice, extra-fine silex, or tin oxide on a rotating cup, brush, or felt.

11. They are composed of two phases with greatly different hardnerness. A rough surface would be desirable if more composite material or surface glaze had to be added to a previously set composite restorative material.

12. Usually the evidence is based on one observer comparing the affected teeth before and after treatment with an uncalibrated shade guide. Many results are not published.

Chapter 11 Structure and Properties of Metals and Alloys
1. Pure metals have a melting temperature, whereas alloys usually exhibit a melting temperature range. Alloys of eutectic composition are an exception—they melt and solidify at a single temperature.

2. If a casting alloy is incompletely melted or not heated above its melting temperature range, it will not flow into all areas—especially not thin areas—of the investment mold of a wax pattern. An incomplete casting will result. In the case of soldering, the solder will not flow adequately, and voids may be left in the solder joint.

3. The density of a molten metal is less than that of its crystalline solid, because the atoms pack more closely together in a crystalline solid's crystal lattice than in the disordered liquid state.

4. If more molten liquid cannot flow from the casting button to take up the volume shrinkage in the casting as it solidifies, porosity will result.

5. Each corner atom is shared with eight other unit cells, so one eighth of each corner atom belongs to a unit cell. There are eight corner atoms, so together they contribute one atom to the unit cell. Each face-centered atom is shared between two unit cells, so one half of each face-centered atom belongs to each unit cell. The six face-centered atoms together contribute three atoms to the unit cell. The face-centered cubic unit cell thus contains four atoms.

6. The density is equal to the mass divided by the volume of the unit cell.

\[ \text{Density} = \frac{\text{Number of atoms} \times \text{Atomic weight}}{(\text{Lattice parameter})^3} \]

7. Copper, iron, and gold.

8. Work hardening, hardening heat treatment, and a decrease in grain size.

9. Increase the number of grain boundaries (ie, smaller grains); increase the number of dislocations (ie, cold working); or treat with heat to create phase changes to lattices that are more resistant to dislocation motion.

10. Small-grain metals have a higher yield stress and more uniform plastic deformation, which result in a higher ultimate strength.

11. a. The smallest grain visible would equal the resolution of the microscope, ideally 0.5 um.

   b. The atoms at the grain boundaries are not as chemically stable because they do not pack together as well as interior atoms of the grain. Different grains etch at different rates because their orientations present planes of atoms of different atomic densities to the polished surface. If the metal is multiphased, the different compositions of the grains will respond differently to the acid etch.

12. The eutectic temp. is 779°C, and the composition is 71.9% Ag and 28.1% Cu.
13. It begins to solidify (liquidus temperature) at 833°C and is completely solidified (solidus temperature) at 779°C.

14. The atomic percentage of silver in Ag₃Sn is 75%: \[ \frac{3 \text{ Ag atoms}}{3 \text{ Ag} + 1 \text{ Sn}} = 0.75 \]. The weight percentage of silver in Ag₃Sn is 73.2%: \[ \frac{3(107.9) \text{ g}}{3(107.9) + 1(118.7) \text{ g}} = 0.732 \].

15. A crystal lattice indicates the location and periodic spacing of atoms in a crystalline solid. A space lattice indicates the way in which mathematical points can be located in space so that every point has a similar grouping of points surrounding it. This requirement develops a repeat pattern of points that correlates with the periodic table of elements.

16. During the recovery stage, the yield strength decreases only slightly, and the percentage elongation begins to increase. During recrystallization, the yield strength drops rapidly, and the percentage elongation increases rapidly. There is a further small increase in percentage elongation and decrease in yield strength with grain growth.

17. By definition, it takes 1 hour for a metal to recrystallize at its recrystallization temperature; at a higher temperature it would take less time and at a lower temperature it would take more. The melting temperature of iron in degrees Kelvin is 1,535°C + 273°C (1,808 K). The recrystallization temperature in degrees Kelvin is 450°C + 273°C (723 K). The recrystallization temperature divided by the absolute melting temperature is 0.42.

**Chapter 12 Dental Amalgams**

1. It is best not to alter the trituration conditions because other properties may be degraded. Choose an amalgam that has the working time characteristics desired.

2. Zinc in the presence of moisture decreases the performance of amalgams. Proper moisture control will give a zinc-containing amalgam better performance than a zinc-free system.

3. Yes, fast-setting alloys. Read the instructions for use.

4. The small spherical-shaped particles tend to slip past one another. Less mercury, however, is required for plasticity. These amalgams also require less packing force to condense.

5. Creep is a bad predictor for clinical marginal fracture in high-copper amalgams, but it has some value as a predictor for traditional systems. The best predictor, however, is proof from clinical trials.

6. The primary reason for the sealing/bonding of amalgam restorations is the attempt to reduce postoperative sensitivity seen in teeth restored with some high-copper spherical amalgams.

7. Except for allergic reactions affecting a small segment of the population, there is
no credible scientific evidence that dental amalgam restorations cause disease in humans. Environmental concerns regarding mercury in amalgam can be addressed by proper office procedures for handling, dispensing, and waste management.

Chapter 13 Precious Metal Casting Alloys

1. Gold and the platinum group metals (platinum, palladium, iridium, rhodium, ruthenium, and osmium).

2. Gold, copper, and silver. The platinum metals also contribute some hardening.

3. Type I alloys are used for one-surface restorations that will be subjected to slight stress; Type II are for two- and three-surface inlays; Type III are for crowns and fixed partial dentures; and Type IV are for fixed and removable partial dentures.

4. Iron, tin, and indium.

5. Palladium has a strong whitening effect on the color of gold alloys.

6. Heat for 10 minutes at 700°C followed by water quenching.

7. Heat for 10 minutes at 350°C followed by water quenching or rapid cooling in air. Cooling a casting in a mold to room temperature (bench cooling) will produce hardening, because the alloy remains in the 350°C to 400°C temperature range long enough.

8. Iridium. Smaller grains result in a stronger, more ductile and homogeneous casting.

9. The gold alloys containing iron, indium, and tin are heated from about 700°C to 950°C in air, and then air cooled. The purpose is to burn off organic contamination and produce an adherent oxide for bonding.

10. Palladium, silver, gold, tin, and indium.

11. Ordering is a crystal structure organization in which the atoms of an element (eg, copper) are regularly arranged in a repeating pattern as opposed to a random distribution (ie, disordered).

12. The present theory holds that hardening in gold-copper-silver alloys involves a separation of silver-rich and copper-rich gold phases within the grain structure. Ordering of a gold-copper phase also occurs, but is not as important in these ternary alloys. Silver, therefore, is a hardener when used in proper amounts along with copper.

13. The frequency of tarnish increases as the noble metal content decreases. Laboratory tests indicate that below about 50% palladium or gold, tarnish is very likely.

14. The ductility or percentage elongation measures the degree to which the alloy can
be burnished (spread). Other properties involved are hardness and yield strength, which indicate a resistance to burnishing.

15. Silver produces a green discoloration of the porcelain.

**Chapter 14 Alloys for Porcelain-Fused-to-Metal Restorations**

1. The term *precious* refers to higher-cost alloys. The term *semiprecious* has been applied to alloys that are mixtures of precious and nonprecious ingredients.

2. Nonprecious alloys are composed of nonprecious ingredients, except for small amounts of beryllium. Most are nickel-chromium; some are cobalt-chromium or iron based.

3. Rational selection of a specific alloy should be based on a balanced consideration of cost and intended use. For single crowns, strength and sag resistance are less important than they are for fixed partial dentures. Castability, biocompatibility, tarnish and corrosion resistance, porcelain color, and hardness are usually equally important for both alloy uses. For fixed partial dentures, solder and joining behavior, sag resistance, strength, elastic modulus, and the porcelain's thermal expansion compatibility become increasingly important as the span increases.

4. Excellent clinical working characteristics.

5. Advantages are good clinical working characteristics, excellent strength, and no porcelain color problems. Disadvantages include slightly more difficult melting and casting than with palladium-silver, and some soldering difficulties.


7. Nonprecious alloys offer lower cost, higher hardness (more wear resistance), higher tensile strength, and higher elastic modulus. Elongation is about the same as for precious metals, but is negated by the high yield strength, which makes it difficult or impossible to work the metal. Some nickel-chromium alloys, especially those containing beryllium, have mold-filling abilities that are superior to all other alloy groups, permitting easier casting of thin sections and producing sharp margins on castings.

8. Silver-free alloys containing about 50% gold and 40% palladium. Highly successful commercially, they have favorable yield strength and hardness and higher elastic modulus than high-gold alloys. Cost is comparable to the gold-palladium-silver group. The only disadvantage is thermal expansion incompatibility with some of the higher-expansion porcelains.

9. Strict control of grinding dust (with suction, masks, etc) and screening of patients for possible nickel allergy (eg, pierced ear posts and other jewelry).
Chapter 15 Dental Porcelain

1. Crystalline ceramics have an orderly, repetitive arrangement of atoms. Vitreous ceramics have an amorphous structure without the orderly pattern of a crystal.

2. Alkali ions (ie, Na\(^+\), K\(^+\), Li\(^+\)) disrupt the silicate structure to form glasses.

3. Metallic oxides are added for color in dental porcelains.

4. A frit is powdered glass made by fusing the constituents together in a furnace and then quenching and grinding.

5. Sintering involves increasing the density of a powdered mass by bonding at points of contact rather than by melting particles.

6. The driving force for sintering is the reduction of surface area by the force of surface tension. Therefore, a fine particle size and high glass surface tension promote rapid sintering.

7. To reduce porosity created by entrapped air.

8. The porcelain is not heated enough to produce glazing until the final bake in order to limit firing shrinkage. Excessive glazing also produces a rounding of edges.

9. About 30% to 40%.

10. The temperature below which glass becomes very rigid or behaves like a solid.

11. Glasses and other brittle solids fail by crack propagation. Tensile stresses cause cracks to spread, whereas compressive stresses do not.

12. Residual compressive stresses at the surface of a ceramic inhibit surface-crack propagation and increase strength. Tensile stresses at the surface lower strength. Therefore, the location and direction of residual stresses determine their effect on properties.

13. According to the cohesive plateau theory, the maximum measurable bond strength is equal to the cohesive strength of the porcelain (ie, 5,000 psi [35 MPa] in tension).

14. The nature and thickness of the oxide layer formed on alloys are critical to the bond strength. Bonding to pure gold produces only a relatively weak bond. Tin, indium, and iron oxides adhere strongly to a gold-alloy surface, reduce the contact angle, and produce cohesive porcelain fractures.

15. Deduction from scientific theory: Engineering mechanics would predict less failure for stronger materials; however, a full clinical study is still needed. Other factors, including the strength of the outer porcelain layer, will also affect longevity.
Chapter 16 Base Metal Casting Alloys


2. Most chromium-type alloys are harder and stronger than conventional gold fixed partial denture alloys. Strength and hardness of some materials are comparable to those of chromium-type removable partial denture alloys.

3. High modulus of elasticity (stiffness) and yield strength (resistance to permanent deformation) suggest the usefulness of chromium-type alloys for the fabrication of long-span fixed appliances.

4. Excessive oxidation of substrate castings inhibits porcelain-to-metal bonding.

5. Proportional limit, offset yield strength, ultimate strength, rupture strength, and elastic modulus of cast titanium alloys are lower than those of chromium-type alloys.

**Removable partial denture alloys**

1. True: Principal strengthening elements are molybdenum, tungsten, and carbon.

2. False: Nitride inclusions and excessive metallic carbides can cause alloy embrittlement.

3. True: Low density makes chromium-type alloys especially useful for the fabrication of large maxillary appliances. Lighter devices are more resistant to displacement by gravitational forces and are, therefore, less likely to subject abutment teeth to unnecessary stresses.

4. False: Chromium-type removable partial denture alloys are about 30% harder than Type IV golds. Strengths of the chromium-type alloys and Type IV gold alloys, however, are comparable.

5. False: The modulus of elasticity (stiffness) of chromium-type removable partial denture alloys is about twice that of the Type IV golds. Thus, sufficient stiffness can be obtained with the use of relatively thin chromium-type castings.

6. False: Strong oxidizing agents should not be used for cleaning appliances fabricated from chromium-containing alloys.

7. True: Ethyl silicate- or phosphate-bonded investments are required for the casting of alloys with fusion temperatures higher than $1,315°C (2,400°F)$.

8. True: Casting temperature affects microstructure and mechanical properties.

9. False: The alloys are very hard. Conventional equipment and procedures consume excessive amounts of time and are relatively ineffective.

10. False: Improper design and fit of chromium-type appliances are the major causes of adverse tissue reactions.
**Fixed partial denture alloys**

1. False: Chromium-type fixed partial denture alloys employ the cobalt-chromium as well as the nickel-chromium system.

2. False: Chromium-type alloys are harder and stronger than conventional gold fixed partial denture alloys. Strength and hardness of some materials are comparable to those of chromium-type removable partial denture alloys.

3. True: High modulus of elasticity and relatively high yield strength (resistance to permanent deformation) suggest the usefulness of chromium-type alloys for the fabrication of long-span fixed appliances.

4. True: The tensile strength and yield strength of some titanium alloys are similar to those of chromium-type fixed partial denture alloys. When compared to chromium-type alloys, modulus of elasticity values for titanium alloys are relatively low.

5. False: Excessive oxidation of a substrate casting can inhibit porcelain-to-metal bonding.

6. False: The biocompatibility of titanium and some titanium alloys is well documented, but the long-term biocompatibility of chromium-type fixed partial denture alloys remains to be determined.

**Surgical casting alloys**

1. False: The nickel-chromium-cobalt base alloy (Surgical Ticonium) is much softer and much more ductile. The cobalt-chromium-based material (Vitallium) is exceptionally strong and hard.

2. False: Physical and mechanical factors also influence biologic tolerance to alloy implants.

3. True: The lungs, liver, and spleen are principal target organs for metallic ions.

4. True: Large implants are more prone to failure than smaller ones. Large surface areas create spaces between the implant and the tissue that may be transformed into undesirable bursae.

5. True: Complete rejection is characterized by development of fistulae or frank exposure of the implant.

**Chapter 17 Casting**

1. A wax pattern is used to form a refractory mold for the casting of a molten metal.

2. Wax shrinkage + gold shrinkage = setting expansion + hygroscopic expansion + thermal expansion of the investment + wax expansion.
3. A wax pattern can be made using the direct technique, in which the wax pattern is formed directly on the prepared tooth, or the indirect technique, in which the wax pattern is fabricated on a gypsum replica of the prepared tooth.

4. Geometry of the casting, or more specifically the surface/volume ratio, is the primary factor.

5. A high-expansion form of silica.

6. To increase the setting, hygroscopic, and thermal expansions.

7. High heat, water immersion (low heat-hygroscopic), controlled water added, and phosphate bonded.


9. By varying the amount of the silica sol component of the liquid mixed with the phosphate-bonded investment powder.

10. The sprue is used as a mount for wax pattern, a channel for wax escape during burnout, a channel for filling the mold with molten gold, and a compensation for shrinkage during solidification

11. To remove oxides, sulfides, and particles of investment that were formed or added during the first melt.

12. Centrifugal, pressure, and pressure/vacuum.

13. These solutions comprise acids or a combination of acid with a HCl base.


15. Gold alloys possess a relatively low modulus of elasticity and proportional limit. They are relatively soft, and their density is about twice that of the cobalt-chromium alloys.


**Chapter 18 Soldering, Welding, and Electroplating**

1. Copper will lower the fusion temperature, increase the strength, and make it susceptible to age hardening. Silver is added primarily to improve the free-flowing qualities of the solder.

2. The appliance would probably come apart at the joint during firing of the porcelain because of the high temperatures. In addition, the porcelain near the joint would have a greenish tinge because of the copper in the solder.

3. The fluoride flux is required to remove the chromium oxide coating, which gives the alloy its corrosion resistance. Regular borax flux will not remove chromium oxide.
4. Solder will not wet a surface covered with antiflux. Antiflux can be used to confine the solder to a given region.

5. Graphite from a lead pencil; iron rouge suspended in alcohol.

6. The fineness of the solder should be less than the fineness of the parts.

7. 580-fine.

8. Silver solder has a sufficiently low fusion temperature so that carbide precipitation can be minimized or avoided with proper soldering procedures.

9. Solder will not wet a surface covered with thick casting oxides and organic films. These films must be removed prior to soldering, since the action of the flux is not sufficient to remove them.

10. Investment soldering is used whenever exact positioning of parts is required.

11. Fracture of the sticky wax indicates that the parts have been moved in relationship to each other and that the finished appliance probably will not fit.

12. Preheating eliminates moisture and provides thermal expansion to compensate for the expansion of the metal. Underheating the investment will lead to poor fit of the appliance. Overheating the investment could fracture it, leading to a poorly fitting appliance, and it could release sulfur, which would contaminate the surface and prevent the flow of solder over the parts.

13. All regions of the appliance adjacent to the joint will be yellowish red.

14. If one part is significantly hotter than the other, the solder will flow to and wet the hot surface and not the other surface.

15. In free-hand soldering, because of the small size of wires, greater care must be exercised to avoid overheating. In addition, wires should be in contact, compared with the 0.1-mm gap distance recommended for investment soldering of larger parts.

16. Diffusion between solder and wire, recrystallization, and grain growth are all possible if a wire is overheated. Overheated stainless steel wires can lead to excessive carbide precipitation.

17. If allowance is not made for the fact that an object at a given temperature will appear much brighter in a dark room than in subdued light, the parts are likely to be too cool for the solder to flow. The lack of flow could mistakenly be attributed to contamination. If the torch is concentrated on the solder, it could become overheated before the parts come to the proper temperature.

18. Surface porosity was probably due to overheating. Excessive use of flux probably resulted in the incorporation of the flux into the solder. The large pore at the center of the joint probably indicates that the parts were in contact in this area.
19. The solder will not wet the oxidized surfaces. If soldering operations are continued, overheating of the work is likely.

20. Oxidation is due to one or more of the following: insufficient flux; improperly adjusted torch; using the oxidizing portion of the flame; and/or removing the flame before soldering operations are completed.

21. A poorly fitting appliance could be caused by one or more of the following: fracturing of the sticky wax, fracture of the investment by improper heating, failure to preheat the investment, and use of a high water/powder ratio for the investment.

22. A flux is required to remove surface oxides and helps to protect the parts and solder from oxidation at soldering temperatures.

23. The large reduction of elongation upon age hardening indicates that the solder has become considerably more brittle. The work is quenched after 5 minutes to prevent it from staying at the age hardening temperature too long. Waiting 5 minutes allows some hardening, but the increased proportional limit is more important than the slight increase in brittleness.

24. Because the copper parts to be welded and the copper electrodes have similar resistances, the electrodes are likely to be welded to the work.

25. A greater amount of distortion would be required for copper because of its surface oxides. In addition, the stress and energy required for this distortion would be much higher because copper has a higher proportional limit and lower malleability than pure gold.

26. With a laser, the region of heat input can be localized to a very small area, and the time of application can be reduced so the total heat required for melting the metal at the junction is insufficient, when dissipated throughout the parts, to produce a temperature high enough to cause distortion or destruction of the cast.

27. It is, in principle, possible to weld other metals or alloys besides pure gold. In practice, the stresses are considerably higher than those required for pure gold, and in many cases heat is also required to obtain good welds.

28. It would increase the resistance of the contact areas and reduce the current, which varies inversely with the resistance. Since the heat generated is proportional to the square of the current and only directly proportional to the resistance, a large reduction in current could make the available heat insufficient to melt the metal.

29. Electroplated dies for fixed partial dentures.

30. Electroplated dies are more abrasion resistant. However, electroplating takes 10 or more hours for completion, requires special equipment and solutions, and may introduce distortions of the impression's surface. Some plating solutions are very toxic and should only be used under a hood.
31. A solution with a high throwing power will produce more uniform plating of an impression and therefore preserve accuracy.

Chapter 19 High-Temperature Investments

1. The reaction time can be altered by changing the temperature of the reacting components (higher temperature results in faster reactions) or by changing the acidic concentration or the amount of water available (lower concentrations of these ingredients result in slower reaction rates).

2. (a) Yes. (b) The temperature may be lowered until the liquid begins to freeze, at which point precipitation of gelled silica hydrate renders the liquid incapable of properly binding investment particles via a gelation process. The temperature of 4.4°C has practical significance, as this is a temperature achieved in the main storage compartment of a refrigerator.

3. To avoid higher costs because of the federal tax on spirits, and to eliminate the chance it will be imbibed.

4. Advantages: (1) Rapid setting rate; (2) useful for lower burnout temperatures because much of the expansion is achieved as a result of the setting reaction rather than temperature increase; (3) high green strength; (4) high fired strength, which results in less mold cracking and fewer fins on castings; (5) liquid formed by mixing colloidal silica with water may be used immediately.

Disadvantages: (1) The investment powder will react with moisture, imposing limitations on the shelf life of opened containers. (2) High setting expansion is an impediment to using the investment for producing refractory models that will be used for articulation against "stone" models (the high expansion results in a model that will not properly match the opposing model of stone). (3) High tendency for reaction with nonprecious alloys produces oxides that are difficult to remove from castings. (4) Lower permeability yields a tendency to produce short castings in a gas entrapment. (5) The colloidal silica liquid cannot be shipped under conditions that would result in freezing of the liquid and agglomeration of colloidal silica. (6) Higher-temperature castings (investment temperature higher than 980°C [1,800°F]) have poorer surfaces owing to refractory loss.

5. Advantages: (1) High permeability yields sharply defined dental castings. (2) Low setting expansion (contraction) renders refractory partial denture models that may be articulated against stone models. (3) A nearly flat expansion vs temperature curve at high temperature (approx. 1,150°C [2,100°F]) and a low expansion vs time effect at that temperature make possible precise control of total expansion. (4) The investment is more refractory, which results in smoother castings. (5) Lower burnout strength results in easier removal of castings and cleaning of oxides from the casting.

Disadvantages: (1) Limited shelf life of liquid. (2) Substantial waiting period prior to using freshly mixed liquid. (3) Potential of cracking during burnout due to high thermal expansion.

6. The hygroscopic technique to generate the maximum expansion of the investment,
thereby helping to compensate for the greater shrinkage of nonprecious alloys as opposed to gold.

7. (a) Phosphate. (b) Ethyl silicate or mixed ethyl silicate, colloidal silica, phosphate-bonded. (c) Phosphate.

8. (1) Increased colloidal silica for phosphate-bonded systems, increased ethyl silicate for ethyl silicate system—both increase expansion. (2) Hygroscopic technique (phosphate). (3) Higher burnout temperatures. (4) Increased soak time. (5) Modification of the grain size of the investment. (6) Alteration of the basic mineral composition of the investment, ie, more cristobalite or increased glass content.

9. One advantage might be a lower setting shrinkage (possible expansion) than a straight ethyl silicate system. Also, smoother casting surfaces eliminate the need for pattern precoats, because the fine colloidal silica yields a loss of refractory capability, causing sintering of particles on the mold cavity surface.

10. A properly burned out carbon-filled investment is the best choice for each alloy. However, if the investment is made by a laboratory, this control is difficult. A carbon mold for the gold alloy reduces oxides on the casting.

For the nonprecious alloy, a noncarbon investment would be advised if control of burnout is difficult, since carbon can interact rapidly with the nonprecious alloys. This may reduce the strength of the PFM bond and could also increase hardness of a nickel-based alloy.

**Chapter 20 Waxes**

1. Paraffin, beeswax, carnauba wax, spermaceti, cerasin.

2. The properties of natural waxes vary with the conditions under which they are produced. For this reason, these waxes are not consistent in their properties. The properties of synthetic waxes are much more uniform, as the manufacturer may impose quality control procedures during their production.


4. Solidification shrinkage and contraction on cooling to room temperature.

5. Increase temperature and apply force.

6. Heat wax uniformly; invest the pattern without delay; store the uninvested pattern at a low temperature.

7. It must have high flow above oral temperature to reproduce detail of cavity preparation, and it must have low flow at oral temperature to reduce distortion when the pattern is removed.
8. Memory is the return of waxes to their original shapes over time. This produces distortion.

9. Incomplete burnout leaves wax residue, which leads to poor castings from either inclusions or incomplete margins.

**Chapter 21 Orthodontic Wires**


2. Force delivery characteristics, elastic working range, ease of manipulation by permanent deformation to desired shapes, capability of joining individual segments to fabricate more complex appliances, corrosion resistance and biocompatibility in the oral environment, and cost. The beta-titanium and nickel-titanium archwires are much more expensive than the traditional stainless steel alloys, but they offer unique properties that should be carefully considered when selecting wires.

3. The composition and structure of the wire alloy, which determine the elastic modulus, and the wire segment geometry—cross-section shape and size (moment of inertia) and the length.

4. Excellent clinical corrosion resistance in the oral environment, excellent formability, and low cost.

5. A cobalt-chromium-nickel alloy very similar in appearance, physical properties, and joining characteristics to stainless steel wires, but with a much different composition and considerably greater heat treatment response. Available in four tempers: soft, ductile, semiresilient, and resilient.

6. The two commercially available products have very similar compositions: titanium, 78% to 79%; molybdenum, 11%; zirconium, 6% to 7%; and tin, 4%. The addition of alloying elements to pure titanium causes the body-centered cubic beta polymorphic phase to be retained at room temperature, rather than the hexagonal close-packed alpha phase, which results in excellent formability or the capability for permanent deformation.

7. An intermediate force delivery between stainless steel or cobalt-chromium-nickel and nickel-titanium wires, excellent formability, and true weldability. Because of their lower elastic modulus, beta-titanium archwires more nearly fill the bracket slots, as compared to stainless steel or cobalt-chromium-nickel archwires. The ductility allows arches or segments with complicated loop configurations, which are not possible with nickel-titanium wires.

8. Special techniques are required for permanent bending, and the wires cannot be bent over a sharp edge or into a complete loop. The wires cannot be soldered or welded but must be joined by a mechanical crimping procedure.

9. The complex proprietary strategies involve the amount of cold work and the heat treatment temperatures used during wire processing, along with varying the alloy composition. The latter may involve slight variations in the relative atomic
percentages of nickel and titanium, in addition to the incorporation of slight amounts of other alloying elements such as copper and chromium. For the shape-memory orthodontic wires, the austenite-finish temperature, where the transformation from martensitic NiTi to austenitic NiTi is completed, must be at the temperature of the oral environment.

Chapter 22 Endodontic Materials

1. (c) K-file.

2. (a) Access, biomechanical instrumentation, and obturation.

3. (a) They work best by rotation.

4. (d) EDTA.

5. (b) Spreader.

Chapter 23 Implant and Bone Augmentation Materials

1. A working definition of osseointegration is a direct structural and functional connection between ordered, living bone and the surface of a load-carrying implant.

2. Commercially pure titanium, titanium alloy (Ti-6Al-4V), bioactive glasses and glass ceramics, and calcium phosphate ceramics.

3. Surgical technique, bone quality, minimization of interfacial motion, and surface chemistry, among other factors.

4. Materials and material processing; mechanisms of implant-tissue attachment; mechanical properties; implant design; loading type; tissue properties; stress analysis; initial stability and mechanisms of enhancing osseointegration; biocompatibility of the implant materials; surface chemistry, mechanics, and bone-bonding ability of the implant.