Fundamentals of Dentifrice: Oral Health Benefits in a Tube

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Continuing Education Units: 2 hours


Disclaimer: Participants must always be aware of the hazards of using limited knowledge in integrating new techniques or procedures into their practice. Only sound evidence-based dentistry should be used in patient therapy.

The course will focus on the most common dentifrice ingredients and the oral health benefits they provide. Upon completion of the course, participants will understand not only the fundamentals of dentifrice ingredients, but also key regulatory aspects of the dentifrice market and the role of professional societies in credentialing consumer dentifrices.

Conflict of Interest Disclosure Statement
• Paula M. Koenigs is a full-time employee of P&G.
• Robert V. Faller is a retired employee of P&G.

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Overview
The majority of patients use dentifrice in their daily hygiene routine. As such, it is a cost-effective and convenient vehicle to deliver ingredients that provide therapeutic benefits, cosmetic benefits, or both. Classification of dentifrice ingredients into these key benefit categories affects how products are regulated.
as well as the types of claims that can be made about a product. Products providing therapeutic benefits are regulated by the US Food and Drug Administration (FDA).

The first therapeutic ingredient to be included in dentifrices was fluoride. Since this first therapeutic advancement in the dentifrice market, many other ingredients have been formulated into dentifrices to provide other benefits, such as plaque and gingivitis reduction, enamel protection, antihypersensitivity benefits, extrinsic whitening, calculus protection and reducing bad breath. These ingredients and their mechanisms of action are described in detail in this course, along with ingredients that provide stability and esthetic benefits to a dentifrice formulation.

Learning Objectives
Upon the completion of this course, the dental professional will be able to:

- Describe the FDA Monograph system.
- Compare the Monograph system with a New Drug Application (NDA).
- Differentiate between claims for therapeutic and cosmetic benefits.
- Describe the American Dental Association (ADA) Seal of Acceptance program.
- Explain fluoride’s mechanism of action and understand key differences in common fluorides.
- List common antiplaque/antigingivitis ingredients in dentifrices and describe their mechanisms of action.
- Compare nerve desensitizing and dentin tubule occluding agents and describe how these agents act to treat dentinal hypersensitivity.
- Name the dentifrice ingredients used to control calculus, stain and bad breath; explain how these agents function.
- Understand the evolution and combination of different benefits in the dentifrice marketplace.
- Describe the role of dentifrice ingredients used to improve esthetics and stability.
- Explain compatibility concerns manufacturers face when formulating a dentifrice.

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Glossary
bioavailability – the degree to which a drug or substance is available to the target tissue following administration.

buffer – chemical system that confers resistance to a change in the pH of a solution (e.g., saliva) when hydrogen ions (H⁺) are added or removed.

carbohydrate – important energy source for the body; a complex molecule made up of one or more simple sugars.

calculus – calcified plaque: a hard yellowish deposit on the teeth, consisting of organic secretions and food particles deposited in various salts, such as calcium carbonate; also called tartar.

cariogenic – contributing to the production of caries.
chelate – action of certain chemical compounds whereby they form several noncovalent bonds to a single metal ion (e.g., Ca²⁺), sequestering it and preventing it from reacting with its surroundings.

chromogen – substance that can be converted to a pigment or dye.

compound – in chemistry, a substance that consists of two or more chemical elements in union.

covalent – in chemistry, a chemical bond formed by the sharing of one or more electrons, especially pairs of electrons, between atoms.

crevicular – a fluid produced by epithelium of the gingival crevice; it contains immunoglobulins and has antimicrobial properties.

enzyme – protein that catalyzes, or facilitates, biochemical reactions.

extrinsic stain – tooth stain on the exterior surface of the tooth that can be removed through routine cleaning procedures. It is generally composed of dietary chromogenic molecules and metal ions which become bound within the salivary pellicle layer that coats exposed tooth surfaces.

gingivitis – inflammation of the gums that often manifests as bleeding during brushing and flossing; mildest form of periodontal disease that is reversible.

heme – a complex red organic pigment containing iron and other atoms to which oxygen binds.

halitosis – the condition of having stale or foul-smelling breath.

hydrophobic – water-resisting; refers to a chemical entity that repels water and prefers oily environments.

ions – atoms or molecules that carry either a positive or a negative electric charge in a solution. For example, sodium chloride (NaCl, common table salt) in water dissociates into Na⁺ and Cl⁻ ions.

intrinsic stain – staining caused by the presence of pigment within the enamel or dentine. Intrinsic stain can often be mediated through bleaching procedures.

lysis – the destruction or dissolution of a cell or molecule, generally through the action of a specific agent.

metabolize – the process through which food is broken down to release energy.

molecule – chemical entity that consists of two or more atoms that have chemically combined to form a single species.

New drug application (NDA) – application requesting FDA approval to market a new drug, drug formulation, or dose.

noncavitated lesion – demineralized, subsurface carious lesion without evidence of discontinuity or break in the enamel surface (sometimes called an early lesion, incipient lesion, or white spot lesion).

organic acids – acid containing at least one carbon atom; also called a carboxylic acid; written chemically as:

\[-\text{COOH} \quad \text{or} \quad \text{CO}_2\text{H} \quad \text{or} \quad \text{OH}\]

Over-the-counter (OTC) – drug products that are generally recognized as safe and effective and are available without a prescription; in oral care, many dentifrices and some rinses are OTC products.

OTC Monograph – a document published by the US FDA that includes lists of ingredients that have proven effectiveness and safety for a particular health concern, as well as information about dosing, drug formulations and labeling.

patency – state or quality of being open, expanded, or unblocked.

pharmacology – study of a drug’s origin, chemistry, effects, and uses.

plaque – an organized community of many different microorganisms that forms itself into a biofilm and is found on the surface of the tongue and all hard surfaces in the oral cavity. Dental plaque is present in all people and can vary from being comprised of totally healthy microorganisms...
(commensals) to being very harmful (pathogenic), predisposing the patient to dental caries or periodontal diseases. Note: Dental plaque is not food debris, nor does it contain food debris. Dental plaque can only be completely removed by mechanical means, such as toothbrushing or prophylaxis.

**subgingival** – located beneath the free margin of gingival tissue.

**supragingival** – located on a portion of the tooth that is not surrounded by gingival tissue.

**surfactant** – compounds such as detergents, emulsifiers, and foaming agents that provide cleaning or help mix substances that prefer to separate (like oil and water). Surfactants typically have a hydrophilic, polar head that interacts with water and a hydrophobic, nonpolar tail that avoids water.

**tartar** – calcified plaque: a hard yellowish deposit on the teeth, consisting of organic secretions and food particles deposited in various salts, such as calcium carbonate; also called calculus.

**toxicology** – study of the unwanted and often adverse effects of substances.

**Dentifrice Market Fundamentals**

The use of dentifrice as part of daily hygiene in the United States is widespread. In fact, there are so many dentifrice options in the oral care aisle today it can be overwhelming. Patients often turn to dental professionals for a product recommendation that will meet their specific oral care needs and desires. Understanding the regulatory environment that guides product claims as well as the process used by credentialing bodies to evaluate products is important as professionals discuss home care options with patients.

In the United States, the Food and Drug Administration (FDA) regulates therapeutic agents to ensure product safety and efficacy. Drugs can enter the market by one of two regulatory pathways. The most common pathway for over-the-counter (OTC) drugs is under the OTC Monograph system. There are several monographs that regulate OTC oral health-related agents.

The second pathway is through a New Drug Application (NDA), which is used for new drug products that fall outside the range of ingredients already included in the OTC Monograph system. NDAs for dentifrices are uncommon, and require the manufacturer to demonstrate that the product is safe and effective through comprehensive clinical testing.

The American Dental Association (ADA), as the leading US dental professional association, takes an interest in educating the public on the safety and efficacy of oral health products. Its primary mechanism for this is by awarding the ADA Seal of Acceptance to qualifying products.

**Description of the US Monograph System**

In 1962, an amendment was passed to the US Federal Food, Drug, and Cosmetics Act (FD&C) requiring that marketed drug products not only had to be safe, but they also had to be effective. At that time, hundreds of thousands of OTC drugs were on the market, and time and resources were too limited to ensure that all these OTC drugs complied with the new regulations.

To ease the approval of OTC therapies considered safe by virtue of their extensive historical use, the monograph drug review mechanism was instituted in 1972. Under this process, the FDA convenes committees to review safety and efficacy data submitted for therapeutic ingredients in the OTC market. The end result of the process is a published document that lists certain therapeutic ingredients (referred to as active ingredients) and the requirements for marketing products that contain those active ingredients. These requirements include a number of parameters, including the intended use, drug dosage or concentration, dosage forms, allowable combinations with other drugs, required labeling, and any special packaging or testing requirements.

There are several classes of monographed OTC drugs for oral use, including anticaries agents, tooth desensitizers, oral antiseptics, anesthetics, and analgesics. Therapeutic dentifrices are regulated by three separate monographs: Anticaries, Antiplaque-Antigingivitis, and Tooth Desensitizer (Table 1).

One factor that differentiates OTC fluoride dentifrices from prescription fluoride dentifrices
is the amount of fluoride they contain as a therapeutic ingredient. Fluoride is a known anticaries ingredient, but it can be toxic if excessive levels are ingested. Although dentifrices are not intended to be ingested, there is enough safety concern to warrant stricter regulations for higher dose products. Most OTC fluoride dentifrices contain 1000–1500 parts per million (ppm) of fluoride and are considered conventional fluoride dentifrices. The maximum allowable fluoride in a monographed OTC dentifrice is 1500 ppm for a sodium monofluorophosphate dentifrice and 1150 ppm for sodium fluoride and stannous fluoride dentifrices. Some prescription-strength fluoride dentifrices contain as much as 5000 ppm of fluoride. Because this level of fluoride is not allowed in an OTC product under the anticaries monograph, these types of products must be prescribed by a dentist.2,3

Table 1. Oral care monographs: Dentifrices are regulated with three separate monographs.

<table>
<thead>
<tr>
<th>Monograph</th>
<th>Current Status</th>
<th>Example Indication</th>
<th>Example Ingredients and Products</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anticaries</td>
<td>FM</td>
<td>Prevents cavities</td>
<td>Sodium fluoride (Aquafresh Extreme Clean)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Sodium monofluorophosphate (Colgate Cavity Protection)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Stannous fluoride (Crest Pro-Health)</td>
</tr>
<tr>
<td>Tooth Desensitizer</td>
<td>TFM</td>
<td>Helps reduce sensitivity</td>
<td>Potassium nitrate (Sensodyne)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Stannous fluoride (Crest Pro-Health)</td>
</tr>
<tr>
<td>Antiplaque-</td>
<td>ANPR</td>
<td>Prevents plaque and</td>
<td>Stannous fluoride (Crest Pro-Health)</td>
</tr>
<tr>
<td>Antigingivitis</td>
<td></td>
<td>gingivitis</td>
<td></td>
</tr>
</tbody>
</table>

* FM—Final Monograph; TFM—Tentative Final Monograph; ANPR—Advanced Notice of Proposed Rulemaking

Table 2. NDA and monograph pathways: Key regulatory differences in these processes are highlighted.

<table>
<thead>
<tr>
<th>NDA</th>
<th>Monograph</th>
</tr>
</thead>
<tbody>
<tr>
<td>Product specific</td>
<td>Active ingredient specific</td>
</tr>
<tr>
<td>Process seeks FDA premarket approval</td>
<td>FDA premarket approval not required if monograph conditions met.</td>
</tr>
<tr>
<td>Product changes may require additional FDA approval</td>
<td>Changes within monograph conditions require no FDA approval.</td>
</tr>
<tr>
<td>Labeling is unique to approved drug product</td>
<td>Labeling is the same for all similar products.</td>
</tr>
<tr>
<td>Possible marketing exclusivity</td>
<td>No marketing exclusivity; monograph is open to everyone.</td>
</tr>
</tbody>
</table>

If a product contains a drug not included in the monograph, it must be approved through the NDA process. Even if the drug is included in the monograph but is being used at a different dose, for a new indication, or in combination with another drug (dual-active product) not specified in the monograph, the product is subject to NDA approval. For example, triclosan is an antibacterial ingredient that is not included in the Antiplaque-Antigingivitis Monograph. Colgate® Total®, a fluoride dentifrice containing triclosan, required approval through an NDA before it could be marketed in the US as an antibacterial dentifrice that treats gingivitis.

Despite their differences, both NDAs and monographs for OTC medicines have very similar standards for safety and efficacy.4

Claims for Therapeutic vs. Cosmetic Benefits
The US Federal Food Drug & Cosmetic Act defines a cosmetic as an article intended to be
applied to the human body to cleanse, beautify, promote attractiveness, or alter the appearance. In contrast, a therapeutic drug is defined as an article intended for use in the diagnosis, cure, mitigation, treatment, or prevention of disease, or article intended to affect the structure or any function of the body. Manufacturer claims for therapeutic vs. cosmetic benefits thus have to follow these definitions.¹

The key therapeutic areas for dentifrices are caries, gingivitis and sensitivity. In addition, several oral care products are marketed with cosmetic claims such as whitens teeth, reduces bad breath, and protects against tartar. If an ingredient is not included in an OTC monograph or is not approved under an NDA, it is not considered a drug, and therapeutic claims cannot be made for it. Many nontherapeutic ingredients are described later in this course.¹,³

CREDENTIALING BODIES
While not a regulatory body, the American Dental Association (ADA) takes a vested interest in informing the public on the safety and effectiveness of oral care products. They do this primarily through their Seal of Acceptance program, which began in 1930. The ADA Seal of Acceptance program is entirely voluntary, whereby manufacturers may submit safety and efficacy data for a product to the ADA Council on Scientific Affairs. Through a review process, the Council decides whether to award its Seal of Acceptance to the product.⁵

The ADA Seal of Acceptance can be a powerful product endorsement in both professionals’ and consumers’ minds, as both of these groups have come to trust the ADA for providing guidance on the safety and efficacy of products. In order to obtain the ADA Seal, manufacturers are required to submit data in accordance with published ADA guidelines for confirmation of each benefit for which the Seal of Acceptance is desired. For some benefits, these requirements include the submission of at least two well controlled clinical studies confirming efficacy, with the clinical trials run according to established ADA protocols. For benefits where clinical studies are not required, the ADA has established a series of specific laboratory tests that must be followed in order to confirm product effectiveness. For a product that wishes to claim multiple benefits, each benefit must be confirmed according to the required guidelines; thus, the overall investment (both clinical and laboratory) to obtain the ADA Seal is considerably higher for a product that is able to claim multiple benefits, compared to a product that claims a single benefit. The ADA Seal is usually awarded for a 5-year period, after which the company must seek renewal. If a dentifrice formulation changes, the manufacturer must submit a new application for the modified product.⁵,⁶

Click here for more information about the “ADA Seal of Acceptance Program & Products” requirements and products that carry the ADA Seal.

DENTIFRICE INGREDIENTS PROVIDING THERAPEUTIC OR COSMETIC BENEFITS
Dentifrices contain ingredients that help reduce caries, plaque, gingivitis, hypersensitivity, calculus, stain, and halitosis. Some ingredients provide a therapeutic benefit, while other ingredients or additives contribute to the cosmetic benefits or physical properties of the dentifrice.

The first dentifrice ingredient clinically proven to provide a health benefit was fluoride, which can be delivered from several different fluoride-based compounds. Over time, dentifrices evolved to provide multiple therapeutic and cosmetic benefits. This section of the course describes the most common dentifrice ingredients used for therapeutic benefits (caries, plaque/gingivitis, and hypersensitivity) as well as cosmetic ones (calculus, whitening, and bad breath), and it provides perspective on how the market evolved to deliver multiple benefits in dentifrice formulations.

Caries Prevention – Fluorides
It is widely accepted that the regular use of fluoride, such as in dentifrice and drinking water, is extremely effective at preventing dental caries. In 1999, the US Center for Disease Control (CDC) issued a statement that water fluoridation is one of the 10 most important public health measures of the 20th century. Fluoride’s presence in low concentration and high frequency is more effective at preventing caries than high levels of fluoride used in low frequency. Because water fluoridation is not available in many countries, dentifrice is considered to be one of
of test markets in 1955, followed by national expansion in January, 1956. In 1960, and again in 1964, the American Dental Association confirmed that Crest effectively prevents tooth decay, reporting that “Crest has been shown to be an effective anticavity dentifrice that can be of significant value when used in a conscientiously applied program of oral hygiene and regular professional care” in granting its Seal of Acceptance (Figure 2).13,14 In 1976, the American Chemical Society recognized Crest with fluoride as one of the 100 greatest discoveries of the previous 100 years.15

The following section will explain the mechanism of action of fluoride, and common fluorides in dentifrices will be described.

Fluoride’s Mechanism of Action

Dental caries is an infectious disease caused by the complex interaction of cariogenic (caries-causing) bacteria with carbohydrates (i.e., sugars) on the tooth surface over time. Cariogenic bacteria metabolize carbohydrates for energy, and produce organic acids as byproducts. The acids lower the pH in the plaque biofilm.16

The hydroxyapatite of tooth enamel is primarily composed of phosphate ions (PO$_4^{3-}$) and calcium ions (Ca$^{2+}$). Under normal conditions, there is a stable equilibrium between the calcium and phosphate ions in saliva and the crystalline hydroxyapatite that comprises 96% of tooth enamel. When the pH drops below a critical level (5.5 for enamel, and 6.2 for dentin), it causes the dissolution of tooth mineral (hydroxyapatite) in a process called demineralization. When the pH is elevated by the natural buffer capacity of saliva, mineral gets reincorporated into the tooth through the process of remineralization.16-18

Point of interest:
When the pH on the tooth surface becomes acidic, phosphate in oral fluids combines with hydrogen ions (H$^+$) to form hydrogen phosphate species (see below). Under these conditions, phosphate is “pulled” from tooth enamel to restore phosphate levels in the saliva, and the hydroxyapatite dissolves. As pH returns to normal, the calcium and phosphate in saliva can recrystallize into the hydroxyapatite, remineralizing the enamel.
Caries is simply the result of a series of demineralization/remineralization cycles where, over time, demineralization conditions prevail. The caries process can be affected in several ways. One of the most effective methods to prevent caries is by promoting remineralization and slowing down demineralization. This can be accomplished with fluoride therapy.2,9,18

When fluoride is present in oral fluids (i.e., saliva), fluorapatite, rather than hydroxyapatite, forms during the remineralization process. Fluoride ions (F⁻) replace hydroxyl groups (OH⁻) in the formation of the apatite crystal lattice (Figure 3). In fact, the presence of fluoride increases the rate of remineralization.

Fluorapatite is inherently less soluble than hydroxyapatite, even under acidic conditions. When hydroxyapatite dissolves under cariogenic (acidic) conditions, if fluoride is present, then fluorapatite will form. Because fluorapatite is less soluble than hydroxyapatite, it is also more resistant to subsequent demineralization when acid challenged (Figure 4).

Caries is a sub-surface phenomenon. With fluoride treatment, a noncavitated lesion can be remineralized with fluorapatite and have greater resistance to subsequent demineralization than hydroxyapatite (Figure 5). Even when available at very low concentrations (as low as 0.02 ppm), fluoride is effective as an anticaries agent.2,19,21

Common Fluorides
Fluoride can be delivered from several different fluoride sources.

- stannous fluoride (SnF₂)
- sodium fluoride (NaF)
- sodium monofluorophosphate (Na₂PO₃F)

The efficacy of fluoride as a caries preventive agent depends largely on its concentration and availability in the oral fluids to affect the demineralization/remineralization balance. Over the years, hundreds of clinical studies have been conducted to test the efficacy of fluoride dentifrices in caries prevention. In general, across all fluoride types, these studies show approximately a 25% reduction in caries over a nonfluoridated control dentifrice.22

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**Figure 3. Fluorapatite Formation.** (A) Fluoride ions (F⁻) replace hydroxyl groups (OH⁻) in hydroxyapatite to form fluorapatite in the tooth enamel. (B) A portion of the apatite crystal lattice is depicted showing the replacement of hydroxide for fluoride.

Adapted from: Posner, 198520

**Figure 4. Fluoride Reactivity.** Under cariogenic conditions, carbohydrates are converted to acids by bacteria in the plaque biofilm. When the pH drops below 5.5, the biofilm fluid becomes undersaturated with phosphate ion and enamel dissolves to restore balance. When fluoride (F⁻) is present, fluorapatite is incorporated into demineralized enamel and subsequent demineralization is inhibited.

Adapted from: Cury, 200919
Figure 5. Demineralization/Remineralization. (A) Plaque acids cause a demineralized, sub-surface lesion. (B) Fluoride treatments remineralize the lesion with a more resistant fluorapatite.

Video 1. Demineralization/Remineralization with fluoride. To view this video, please go to www.dentalcare.com and find this course in the Continuing Education section.
a. Stannous fluoride. Stannous fluoride (SnF₂; also called tin fluoride) was first formulated successfully into a dentifrice to deliver an anticaries benefit in the 1950s.¹¹,¹² Fluoride is highly reactive, and the challenge was finding an abrasive system that had low enough reactivity with fluoride to maintain the bioavailability of the fluoride. The formulation included 0.454% stannous fluoride and the abrasive calcium pyrophosphate; it was marketed as Crest® with Fluoristan™ (see Figure 2). In the past decade, new formulations of stannous fluoride have been introduced with enhanced bioavailability and efficacy.²³

b. Sodium fluoride. Sodium fluoride (NaF) is a fluoride salt commonly used in dentifrices and oral rinses. Sodium fluoride delivers a highly reactive fluoride ion; therefore, formulating it with a compatible abrasive is critically important for achieving the anticaries benefit. In the early 1980s, silica abrasives that were compatible with sodium fluoride became available and allowed dentifrices with stannous fluoride to be reformulated with the more stable sodium fluoride.

c. Sodium monofluorophosphate. Sodium monofluorophosphate (SMFP) was introduced into Colgate’s first fluoridated dentifrice and allowed this brand to obtain the ADA Seal of Acceptance for cavity protection in 1968 (Figure 6).²⁴ Unlike sodium fluoride, SMFP is not an ionic fluoride salt, but rather a covalently bound fluoridated compound that requires enzymatic activation by a salivary enzyme (alkaline phosphatase) to release bioavailable fluoride (Figure 7).²⁵ Because of this lower reactivity, SMFP is compatible with more abrasives than other fluoride sources.²⁶

Enamel Protection
Enamel erosion is characterized by the dissolution and removal of the tooth enamel surface under highly acidic conditions. When the localized pH drops below 4.5, the pellicle cannot protect the enamel surface, and irreversible erosive damage can occur. Enamel erosion has become an important issue with the increased consumption of sports drinks, soft drinks, and citric juices.²⁶ All of these products have a pH below the critical level for dissolving dental enamel. In this context, the enamel erosion benefits of existing dentifrice ingredients has become more relevant and are described below.²⁷-²⁹

Fluoride
The mechanism by which fluoride helps prevent caries is the same mechanism by which it provides some enamel erosion benefit. Fluoride exerts an effect by favoring remineralization and inhibiting demineralization.²,₉,₁₆

The presence of bioavailable fluoride in the oral fluids (i.e., biofilm and saliva) greatly enhances the precipitation of fluorapatite into tooth structure from calcium and phosphate ions present in saliva. Fluorapatite is more resistant to demineralization than hydroxyapatite, and thus provides some enamel erosion benefit; but even fluorapatite will dissolve under highly acidic conditions.

Figure 6. Colgate® with SMFP

Figure 7. Enzymatic activation of SMFP. The covalent bond of SMFP must be broken to release bioavailable fluoride.
**Stannous fluoride**
In addition to the modest level of enamel erosion protection provided by most fluorides, stannous fluoride is unique in its ability to provide significantly more enamel protection than that provided by fluoride alone. This is because stannous fluoride adheres to the surface of tooth enamel and forms a protective layer that is able to shield enamel from the effects of erosive acids.  

**Antiplaque/Antigingivitis**
Adding antibacterial action to dentifrice to reduce plaque and gingivitis was another major therapeutic breakthrough. Several modern dentifrice ingredients have the potential to kill or inhibit bacteria that cause plaque and gingivitis. The mechanism of action varies depending on the specific antibacterial ingredient.

**Stannous fluoride**
Stannous fluoride exerts antibacterial effects by two modes of action. First, stannous fluoride exerts a killing effect on bacteria (bactericidal action). This is probably due to non-specific interaction with the bacterial membrane that causes membrane disruption. The result is leakage of cellular components that leads to cell lysis and death.

The second, and more important, mode of antibacterial activity is through stannous fluoride’s inhibition of metabolic enzymes. The inhibition of metabolic activity affects bacteria in a number of ways, including:

- reduction of bacterial growth
- prevention of bacterial adhesion to oral surfaces (e.g., enamel)
- reduction in bacterial byproducts that boost the inflammatory response leading to gingivitis

Stannous fluoride’s inhibitory effect on bacteria is related to its inhibition of bacterial glycolysis, an energy making process whereby metabolic enzymes break down carbohydrates. In addition, studies have demonstrated that stannous fluoride significantly reduces metabolic toxins produced by bacteria in plaque biofilm.

In early stannous fluoride dentifrice formulations, the stannous fluoride was not fully stable or bioavailable. A key issue in formulating with stannous fluoride is that it is easily inactivated by hydrolysis and oxidation, thus making it difficult to stabilize in a typical dentifrice formula. Stannous fluoride can also have an astringent taste and cause extrinsic staining of teeth and fillings. However, current stannous fluoride-containing dentifrices have been formulated to circumvent instability and bioavailability challenges.

One of the major breakthroughs in stannous fluoride formulation efforts was due to technology innovations that enabled the combination of both whitening and stabilization chemistries to provide highly effective stannous fluoride formulations that are not compromised by common esthetic negatives, such as poor taste or staining, of earlier stannous fluoride products. After more than fifty years of research, P&G successfully formulated a stabilized, consumer-acceptable, stannous fluoride dentifrice with the launch of Crest® Pro-Health® dentifrice in 2005. Crest® Pro-Health® dentifrice has received the ADA Seal of Acceptance for protection against six different conditions: cavities, plaque/gingivitis, sensitivity, calculus, extrinsic stains, and bad breath (Figure 8).

**Triclosan**
Triclosan is a broad-spectrum antibacterial agent that inserts into and disrupts the bacterial membrane. Being a nonpolar molecule, it has an affinity for the hydrophobic environment of the lipid bilayer. This causes leakage of cellular components, ultimately leading to cell death. Since it is an uncharged molecule, alone it has poor retention (substantivity) in the oral cavity (Figure 9).

Triclosan is the antibacterial ingredient in Colgate® Total®, and it provides the plaque and gingivitis benefits of the dentifrice. Colgate® Total® is formulated with a special polymer (Gantrez®), which increases the substantivity of triclosan in the oral cavity.

**Figure 8.** Crest® Pro-Health® (stannous fluoride)**
the oral cavity. Colgate® Total® was introduced outside the U.S. in 1992 and was the first broadly marketed antibacterial dentifrice. Because triclosan is not included in the US Antiplaque-Antigingivitis Monograph, Colgate® Total® had to be approved through an NDA before it could be sold in the US. It received US marketing approval in 1997. Total® also carries the ADA Seal of Acceptance and has been demonstrated effective against multiple indications.47-50 (Figure 10).

Antihypersensitivity

Cervical dentinal hypersensitivity is a condition characterized by sharp pain associated with thermal, evaporative, tactile, osmotic or chemical stimuli. This condition depends on dentin exposure, as well as the patency of the dentinal tubules. It is widely accepted that dentinal hypersensitivity is a result of fluid movement within the dentinal tubules, which stimulates nerve endings in the pulp matrix.41,51-54

Tooth hypersensitivity is a condition patients commonly report to their dental professional; thus, it is a segment of the dentifrice market heavily influenced by professional recommendations. It has been reported that up to 57% of the adult population suffers from this condition.41,55
The nerve cell is stimulated, these ions cross the nerve cell membrane through channels and move from an area of high concentration to an area of lower concentration (referred to as the concentration gradient). Thus, potassium ions flow from the inside to the outside of the cell and the sensation of pain is transmitted.

Potassium ion is a desensitization agent because it diffuses through dentin tubules and increases the extracellular potassium concentration at the nerve ending, eliminating the potassium ion concentration gradient across the nerve cell membrane. Without this concentration gradient, the nerve cell will not depolarize and will not respond to stimuli; thus the sensation of pain will not be transmitted. Potassium ion can be delivered in a variety of salt forms (e.g., potassium nitrate, potassium citrate). The most common

A segment of the fluoride dentifrice market has emerged to specifically address the needs of patients suffering from sensitive teeth. One of the first dentifrice products to enter this segment of the market was Sensodyne®, which was introduced in 1961. More recently, tooth sensitivity has become a very dynamic area, as several new products have entered the market with proprietary ingredients to treat dentinal hypersensitivity.

As noted above, exposure of dentinal tubules to external stimuli is a common cause of tooth sensitivity. Dentinal hypersensitivity is generally treated in one of two ways.

1. chemical desensitization of the tooth nerve endings
2. tubule occluding agents or barriers to reduce dentin permeability

Antihypersensitivity treatments with these mechanisms are described below.

**Nerve Depolarization Agents**

To understand how a chemical desensitization agent works, one must first understand how a nerve cell transmits pain stimuli. Potassium ion (K+), sodium ion (Na+), and chloride ion (Cl−) are important ions involved in the electrical activity of nerve cells. When the nerve cell is at rest, the potassium ion concentration is higher on the inside of the cell than on the outside, while the sodium ion concentration is higher on the outside of the cell than on the inside (Figure 11).
potassium salt used in sensitivity dentifrices is potassium nitrate (KNO₃).⁵⁶

**Tubule Blocking or Occluding Agents**
Another strategy to treat/prevent dentinal hypersensitivity is to reduce the permeability of the dentin by occluding or blocking the exposed dentin tubules. This prevents stimuli from causing fluid flow in the tubules, thereby preventing the nerve endings inside the tooth from being stimulated.⁵²,⁵³,⁵⁷

a. **General mechanism of action.** Several dentifrice ingredients can be used to occlude or block the dentinal tubules. All of these agents have similar mechanisms of action, forming salt precipitates on the surface of the exposed dentin and inside the dentinal tubules. These precipitates effectively reduce or block the fluid flow in the tubules and exert a desensitization effect. Strontium chloride was the desensitizing ingredient used in the original Sensodyne® dentifrice, and it acted via this mechanism by forming strontium salt precipitates; however, it is rarely used anymore because of its strong metallic taste and incompatibility with fluoride.

b. **Newer tubule-occluding agents.** Other tubule-occluding agents new to the market include arginine with calcium carbonate (Pro-Argin™), strontium acetate, and calcium sodium phosphosilicate (Novamin®).

Arginine and 8% calcium carbonate (Pro-Argin™; Colgate® Sensitive Pro-Relief™). Pro-Argin™ is claimed to block tubules upon its application by depositing calcium- and phosphate-containing minerals within the dentinal tubules. Both arginine and calcium carbonate (CaCO₃) are required for this action (Figure 12). Arginine, found naturally in saliva, may help usher calcium to open tubules for incorporation of calcium phosphate into dentin. Calcium carbonate creates a basic environment, and calcium phosphate salts are less soluble at higher pH (more basic). The combination of high local calcium concentration at the dentin tubule at basic pH is designed to promote precipitation of calcium phosphate salts.⁵⁶

Calcium sodium phosphosilicate (Novamin®, Sensodyne® Repair & Protect). In saliva, Novamin® releases calcium and phosphate ions and raises the pH (Figure 13). Under these conditions, calcium phosphate salts precipitate from solution to not only block dentin tubules but also to form an insoluble calcium phosphate layer on the surface of enamel.⁵²,⁵⁸,⁵⁹

c. **Stannous fluoride.** Stannous fluoride is also a tubule-occluding agent that can treat dentinal hypersensitivity. Stannous fluoride, through hydrolysis and oxidation reactions, forms many insoluble metal salts that can precipitate in dentinal tubules and on the dentin surface (see video) to provide effective relief against hypersensitivity.⁵⁵,⁵⁹,⁶¹ Stannous fluoride is the only fluoride delivering protection from caries and plaque/gingivitis as well as hypersensitivity.
subgingival calculus forms either from saliva or crevicular fluid. Dental calculus that forms from crevicular fluid can contain heme and some breakdown products which make it pigmented. It is called serumnal calculus. Calculus forms most readily in areas which are adjacent to the openings of the salivary ducts, where the calcium phosphate in saliva is least stable. In populations with poor oral hygiene supragingival calculus can be extensive and result in gingival recession. Calculus formation can be controlled by adding mineralization inhibitors to dentifrices and mouthrinse. The chemical agents used most often for calculus control in dentifrice are described briefly below.  

a. **Pyrophosphate.** Phosphate is a ubiquitous chemical group found in biological systems. As shown in (Figure 14), two phosphate groups combine chemically to form a molecule called pyrophosphate (P$_2$O$_7^{4-}$). Pyrophosphate occurs naturally in saliva and plays a role in inhibiting calculus formation. These molecules chelate calcium (Figure 15),

**Cosmetic Benefits**

While delivering fluoride for cavity protection was a major therapeutic advance in the dentifrice market, researchers saw an opportunity, over time, to expand the benefits offered by dentifrice. By the 1980s, additional innovations were having an impact. Agents were discovered that could provide protection against calculus and stain, and this opened an era where improved cosmetic benefits spurred the dentifrice market.

**Calculus Control**

Dental plaque calcifies when calcium phosphate begins depositing in it. Under normal conditions, the oral fluids are saturated with calcium and phosphate, which is important for maintaining sound enamel. However, this abundance of mineral ions also contributes to calculus formation on the tooth surface (i.e., calcification of plaque biofilm). The amount and type of calcium phosphate salts present vary greatly but include brushite, octacalcium phosphate (OCP), tricalcium phosphate (TCP) and apatite. While supragingival calculus forms from saliva, subgingival calculus forms either from saliva or crevicular fluid. Dental calculus that forms from crevicular fluid can contain heme and some breakdown products which make it pigmented. It is called serumnal calculus. Calculus forms most readily in areas which are adjacent to the openings of the salivary ducts, where the calcium phosphate in saliva is least stable. In populations with poor oral hygiene supragingival calculus can be extensive and result in gingival recession. Calculus formation can be controlled by adding mineralization inhibitors to dentifrices and mouthrinse. The chemical agents used most often for calculus control in dentifrice are described briefly below.

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![Figure 14. Pro-Argin™. Arginine is a naturally occurring amino acid, and calcium carbonate creates a basic (high pH) environment.](image-url)
slowing the rate of nucleation (crystal formation) and calcification of plaque. The pyrophosphate binds to calcium in a growing crystal, essentially slowing further crystal growth at that site and effectively decreasing calculus build-up (Figure 16). Original Crest® Tartar Control dentifrice contained 3.3% pyrophosphate. It was the first tartar control dentifrice introduced to the market, and the first tartar control dentifrice to receive the ADA Seal of Acceptance.

b. Sodium hexametaphosphate (SHMP). SHMP is a large polyphosphate molecule and has multiple calcium binding sites in one molecule. It is a very effective calculus inhibitor. Because it works only on the surface, it is sometimes called a calcium surface active builder. SHMP is susceptible to hydrolysis, and must be formulated in a low water dentifrice to be stable (Figure 17). When sodium hexametaphosphate (SHMP) is included in a low water dentifrice, the SHMP particles may be perceived as “gritty.” These particles will begin to dissolve immediately upon brushing and are not abrasive.

c. Zinc. Zinc salts (e.g., zinc citrate, zinc chloride, zinc lactate) are used in some tartar control dentifrices and oral rinses, and have been shown to be moderately effective at controlling calculus. Positively charged zinc ion (Zn2+) inhibits crystal growth by substituting for calcium in the crystal lattice of calcium phosphate. This interferes with the crystal formation and slows crystal growth. As a result, calculus formation is reduced. However, zinc containing dentifrices have

**Figure 15.** Pyrophosphate. Negatively charged pyrophosphate molecules bind (chelate) positively charged calcium ions.

**Figure 16.** Anticalculus Action. Pyrophosphate inhibits calculus formation by inhibiting calcium phosphate deposition in plaque.

**Figure 17.** SHMP Hydrolysis. (A) SHMP is a polyphosphate created from a chain of repeating phosphate units. (B) The hydrolysis or breakdown of SHMP proceeds to single phosphate molecules, although many intermediate products are also produced.
some limitations, such as poor bioavailability and an astringent taste.\textsuperscript{68}

d. **Gantrez**. Gantrez is a copolymer of methylvinyl ether (PVM) and maleic acid (MA) and is an ingredient in Colgate\textsuperscript{®} Total\textsuperscript{®} dentifrice. The mechanism of action for this co-polymer is to bind (chelate) calcium ions, thus inhibiting plaque mineralization (Figure 18).

**Stain Control/Whitening Agents**

Stain control and whitening are key benefits of modern dentifrices. These are accomplished via ingredients that target specific types of tooth stain. Stains can be classified as extrinsic (surface stains) or intrinsic (below the enamel surface), and their management is based primarily on that classification (Table 3). Dentifrices primarily work against extrinsic stains. Bleaching products that contain hydrogen peroxide (i.e., whitening strips) or carbamide peroxide (i.e., dental office bleaching trays) address intrinsic stains as well as extrinsic stains.\textsuperscript{36,69}

**Figure 18.** Gantrez\textsuperscript{®} calcium chelation.

**Video 9.** Formation of extrinsic stain. To view this video, please go to www.dentalcare.com and find this course in the Continuing Education section.

**Table 3. Tooth stain: Sources and treatments of extrinsic and intrinsic stains.**\textsuperscript{36}

<table>
<thead>
<tr>
<th>Stain Type</th>
<th>Sources of Stain</th>
<th>Agents to Remove/Prevent Stain</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Extrinsic tooth stain</strong></td>
<td>Daily diet (coffee, cola, tea, red wine) Tobacco Oral therapeutics Poor oral hygiene</td>
<td>Abrasives • Aluminum oxide • Calcium carbonate • Dicalcium phosphate dihydrate (DCPD) • Silica Chemical agents • Sodium pyrophosphate • Sodium tripolyphosphate • Sodium hexametaphosphate Peroxide (hydrogen, carbamide)</td>
</tr>
<tr>
<td><strong>Intrinsic tooth stain</strong></td>
<td>Natural aging of dentition Tetracycline use in children Prolonged tobacco use</td>
<td>Peroxide (hydrogen, carbamide)</td>
</tr>
</tbody>
</table>
The abrasivity of dentifrice is measured in terms of Relative Dentin Abrasivity, or RDA. This rating was introduced in the early 1970s and is used by professional dental societies and boards of health to rate the abrasivity of commercial dentifrices.\textsuperscript{70,71}

RDA values are obtained in the laboratory by comparing the amount of tooth structure worn away by a standardized tooth brushing protocol using any given dentifrice with that of a standard dentifrice. The standard protocol integrates factors such as pressure, time, temperature, and humidity. Because dentifrices with lower RDA values are less abrasive, they also tend to have less potential to remove surface stain. The International Standards Organization (ISO) specification states that a dentifrice should not exceed an RDA of 250, which is considered safe for hard tissues. Although there is a wide range of RDA values for various dentifrices, there are no relative degrees of safety between 0 and 250. In other words, a dentifrice with an RDA of 200 is as safe as one with an RDA of 50. Having an effective abrasive system in a dentifrice is important for cleaning the teeth and removing extrinsic stain.

Fluoride ions are very reactive and can interact with common dentifrice abrasives, rendering the fluoride inactive for caries control. Also, many dentifrice abrasives have a very porous, negatively charged surface that can bind many dentifrice ingredients (e.g., stannous), lowering their bioavailability. For this reason, formulating dentifrices with the right abrasive is critical to achieving the desired benefits of other ingredients (Table 4).

**Chemical action.** Polyphosphates, which are dentifrice ingredients used to control calculus, also target extrinsic stain. One of the most effective ingredients with this dual action is SHMP, which is used in several marketed dentifrices. SHMP controls stain with a chemical action. Stain molecules, or chromogens, are usually negatively charged molecules; they have an affinity for positively charged ions like calcium (Ca\textsuperscript{2+}) that reside in the tooth enamel and cross-link pellicle proteins. SHMP is also negatively charged, with a strong affinity for calcium. SHMP can displace stain molecules from calcium binding sites. It binds to the tooth surface and integrates into the pellicle to prevent additional stain molecules from binding (Figure 19).\textsuperscript{36,37,72}

### b. Intrinsic stain

Intrinsic stains are stains and discolorations that are located below the enamel surface. Due to their diverse etiology, intrinsic stain treatment varies with the cause. Bleaching is usually used to remove or minimize intrinsic discolorations. The most popular bleaching agent is hydrogen peroxide (H\textsubscript{2}O\textsubscript{2}), in concentrations that range from 3% – 35%. Typically the higher concentration products are used for in-office bleaching, whereas products with 15% or lower concentrations are often used in home-applied whitening protocols (e.g., whitening strips).\textsuperscript{69,73,74} Regular dentifrices are generally not used to deliver agents to remove intrinsic tooth stain because longer contact time is typically needed.

**Halitosis (Bad Breath)**

Halitosis, otherwise known as fetor oris, oral malodor or simply bad breath, is universally considered to be a socially unacceptable condition.\textsuperscript{75} Although prevalence estimates vary, it is clear that the condition affects a significant portion of the population, with an estimated 20-30% of adults reported to suffer from chronic breath malodor.\textsuperscript{75,81} Although a limited percentage of halitosis cases result from extraoral factors,
such as diabetes, liver, kidney and other metabolic diseases, the highest percentage of cases are the result of intraoral causes, and are characterized by the production of gaseous volatile sulfur compounds (VSCs) associated with unpleasant bad breath. Hydrogen sulfide, methyl mercaptan and dimethyl sulfide are frequently cited as exhaled VSCs most commonly associated with unpleasant breath.

Certain foods, especially ones like garlic and onions that contain pungent oils, can contribute to bad breath because the oils from these foods are eventually carried to your lungs and out through your mouth. Another source of bad breath can be individuals that experience ongoing sinus conditions, as sinus conditions can lead to a dry mouth. People with sinus conditions often have stuffed up noses and therefore need to breathe through their mouth. The drying effect of mouth breathing can create an environment that promotes bad breath. Additionally, sinus sufferers are likely to be taking antihistamines, a type of medicine that is known to create mouth dryness.

Even people who don’t have an ongoing problem with bad breath easily notice that their breath is least pleasant in the morning when they first wake up. During the night, a person’s salivary flow is reduced when a person sleeps. Saliva flow helps maintain mouth moisture, and it helps cleanse debris, bacteria and bacterial by-products that cause bad breath every time we swallow. As that effect is reduced overnight when we sleep, the result can be stale breath in the morning; a condition that is similarly noticed by people whose
mouth becomes dry after speaking for long periods of time. Smoking is also considered to be a major cause of bad breath.

In the majority of cases, bad breath is caused by the presence of oral bacteria and oral debris. The bacteria and oral debris associated with breath malodor are largely found on the tongue as well as in subgingival and interproximal niches that are difficult to clean. In the absence of regular, thorough brushing and flossing, bacteria can accumulate on the bits of food left between the teeth, in the mouth and on the tongue. Sulfur compounds released by these bacteria give breath an unpleasant smell; with halitosis occurring when the unpleasant odor is then expelled from the mouth when exhaling. In addition to halitosis being an undesirable condition to have, it clearly has the potential to make social situations particularly unbearable.

Oral inflammatory diseases such as caries, gingivitis and periodontitis are also associated with halitosis; a result of bacteria hiding in diseased tissues, producing foul gases. While meticulous oral hygiene in conjunction with scrupulous tongue brushing could theoretically help prevent persistent malodor, studies and surveys have shown that few adults regularly remove enough dental plaque through mechanical oral hygiene alone to alleviate the problem. In most cases, good professional oral care combined with a daily regimen of oral hygiene, including interdental cleaning, deep tongue cleaning and optional use of efficacious oral care products specially formulated to combat the germs that can cause bad breath, will lead to improvement.

**a. Flavors to freshen breath.** Bad breath sufferers often seek out help in the form of commercial products marketed to freshen objectionable breath. While some of these products are able to deliver a brief masking of the halitosis, most do not have the potential to provide long-term benefits. They are designed simply to temporarily mask odors. Unfortunately, many are quickly washed away by the natural flow of saliva. Methods used to help reduce bad breath, such as mints, mouth sprays, mouthwash or gum, may only temporarily mask the odors created by the bacteria on the tongue. These methods, however, cannot cure bad breath because they do not remove the source of the bad breath.

**b. Antibacterial agents to reduce malodor.** Antibacterial approaches can provide substantially longer-term breath efficacy than that provided by odor masking agents. As opposed to flavor agents that simply mask odor, antibacterial agents actually treat the source of the problem by targeting the malodor producing bacteria.

The use of zinc and triclosan are two dentifrice ingredients that have been identified for their ability to target bacteria and help reduce volatile compounds responsible for bad breath. Stannous fluoride, a well-studied antimicrobial agent with concurrent anticaries, desensitizing...
and anti-plaque and gingivitis benefits\textsuperscript{97} has been found to be a particularly effective as a dentifrice ingredient compared to other approaches for its ability to provide both germ kill\textsuperscript{98} and breath protection.\textsuperscript{99}

**Dentifrice Ingredients Providing Stability or Esthetic Benefits**

This section will review key dentifrice ingredients that provide stability and esthetic benefits. These nontherapeutic dentifrice components are called inactive ingredients, additives, or excipients and include binders, surfactants, buffering agents, humectants, preservatives, sweeteners, flavorings, and dyes (Figure 20). These components are essential to keep the dentifrice properly mixed with a smooth consistency, and they make the product palatable to the consumer. Three dentifrice ingredients (abrasive, humectants, and solvent) typically represent about 95% of the dentifrice ingredients.

**Humectants**

Humectants retain moisture so that the dentifrice does not dry out. Humectants function by binding and holding the solvent in the dentifrice. Water is the solvent used in most dentifrices. Humectants, such as glycerin and sorbitol, also inhibit bacterial growth and provide flowability to the dentifrice. Humectants and solvent combined represent approximately 75% of a typical dentifrice formulation.

**Binders**

Binders, also referred to as thickeners, provide texture and determine how “thick” or “runny” the dentifrice is. Binders are used for cohesiveness, to provide body, and to prevent ingredients from separating. Xanthan gum, carboxymethyl cellulose (CMC) carbomers, carrageenan, and synthetic cellulose are all commonly used dentifrice binders. Binders are usually large polymeric polar molecules that form strong interactions with water. These interactions change the consistency and flowability of the dentifrice. Without the binder, the toothpaste would separate into different phases, a liquid portion and a solid-like portion, and would have to be stirred before each use.

![Dentifrice Ingredients](image)

**Figure 20.** Dentifrice Ingredients. Dentifrices contain a number of ingredients that stabilize the product and/or provide esthetic benefits, in addition to the ingredients that provide therapeutic or cosmetic benefits.

**Buffers**

Crest\textsuperscript{®} Oral-B\textsuperscript{®} at dentalcare.com Continuing Education Course, January 16, 2013
Colors/Visuals
Finally, coloring agents are added to provide dentifrice with pleasing colors. The opacity of a paste dentifrice comes from the addition of titanium dioxide. Dentifrices formulated without titanium dioxide result in the formation of a gel dentifrice, rather than an opaque paste. Mica is used to provide a sparkly appearance in some dentifrices, such as those marketed to children. A summary of dentifrice additives can be found in Table 5.

Compatibility Concerns
As noted throughout this course, a dentifrice is a very complex aggregate of chemicals with very specific functions. Not only do these ingredients have to be effective individually, they also have to be compatible with one another. All of these requirements demand very careful formulation and processing in order to be able to manufacture a high quality dentifrice.

A primary concern when formulating a dentifrice is the need to prevent the inactivation of therapeutic agents by other ingredients. This is commonly the case with abrasive components (e.g., silica) that can bind to and limit the bioactivity of therapeutic ingredients. Other concerns involve the degradation or inactivation of ingredients by water. For example, SHMP can be hydrolyzed by water into individual phosphates or other intermediate breakdown products.

Conclusion
The FDA uses two mechanisms to regulate OTC drugs: drug monographs and NDAs. A drug monograph identifies active ingredients that are deemed to be safe and effective for a specific therapeutic need. Most OTC fluoride-containing dentifrices are regulated through the Anticaries, Antiplaque-Antigingivitis, and Tooth Desensitizer monographs. If a dentifrice contains a drug that is not included in a monograph, it must be approved through an NDA. Therapeutic dentifrices brought to market under one of these two regulatory pathways can make claims related to treating or preventing disease.

The ADA is a professional society that takes great interest in informing the public on the safety and efficacy of oral care products. This is done primarily by awarding its Seal of Acceptance. The ADA Seal of Acceptance program is a rigorous,
agents used in newer antihypersensitivity
dentifrices.

Fluoride was the first therapeutic ingredient
used in dentifrice. Fluoride helps prevent caries
by increasing remineralization and inhibiting
demineralization. The three fluoride salts
approved by the FDA for use in dentifrices are
stannous fluoride (SnF₂), sodium fluoride (NaF),
and sodium monofluorophosphate (Na₂PO₃F).

Since the introduction of early fluoride dentifrices,
many other ingredients have been discovered
and added to dentifrice to provide multiple
additional benefits, including the following:

- **Plaque/gingivitis/malodor reduction:**
  Plaque, gingivitis, and halitosis are caused by
  bacteria. Antibacterial-containing dentifrices
can help prevent these conditions. The exact
  mechanisms by which different agents exert
  antibacterial actions may differ.

- **Antihypersensitivity:** Dentinal
  hypersensitivity can be treated by chemically
depolarizing nerve endings in the tooth or by
blocking dentinal tubules. Potassium nitrate
is the most common nerve desensitizing
agent. Stannous fluoride, arginine + calcium
carbonate, strontium acetate, and calcium
sodium phosphosilicate are tubule occluding

Table 5. Nontherapeutic dentifrice ingredients:
Summarized below are common dentifrice ingredients and their functions.

<table>
<thead>
<tr>
<th>Ingredients</th>
<th>Examples</th>
<th>Functions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Humectants</td>
<td>Sorbitol, glycerin</td>
<td>Provide flowability, prevent dehydration, prevent microbial growth</td>
</tr>
<tr>
<td>Solvents</td>
<td>Water</td>
<td>Provide flowability, solvate polar ingredients, hydrate binders</td>
</tr>
<tr>
<td>Abrasives</td>
<td>Hydrated silica, alumina, calcium carbonate</td>
<td>Remove stains, increase viscosity</td>
</tr>
<tr>
<td>Surfactants</td>
<td>Sodium laurel sulfate, cocamidopropyl betaine, poloxamer</td>
<td>Create foam, emulsify flavors, clean</td>
</tr>
<tr>
<td>Buffers</td>
<td>Trisodium phosphate, sodium hydroxide, sodium citrate</td>
<td>Maintain pH to maximize stability and maintain efficacy.</td>
</tr>
<tr>
<td>Binders</td>
<td>Xanthan gum, carboxymethyl cellulose (CMC) carbers, carrageenan</td>
<td>Provide structure and thickening.</td>
</tr>
<tr>
<td>Flavors and sweeteners</td>
<td>Menthol, peppermint, spearmint, green tea, sodium saccharin</td>
<td>Mask taste of unpleasant ingredients. Provide consumer acceptability.</td>
</tr>
<tr>
<td>Colors and visuals</td>
<td>Titanium dioxide, dyes, pigments, mica</td>
<td>Enhance esthetics</td>
</tr>
</tbody>
</table>

voluntary process in which manufacturers can
choose to participate for specific products.

Additional dentifrice ingredients include
humectants, binders, buffers, flavors, sweeteners,
and surfactants. These ingredients stabilize
the product and create esthetic benefits for the
consumer. They are needed to keep the dentifrice
properly mixed with a palatable consistency. Not
all dentifrice ingredients are compatible, however,
so manufacturers must formulate products in a way
that does not interfere with the bioavailability of
the therapeutic ingredients. Creating a dentifrice
that delivers important therapeutic and cosmetic
benefits, while at the same time being acceptable
to the consumer, requires the manufacturer to
delicately balance the overall formulation.
Course Test Preview
To receive Continuing Education credit for this course, you must complete the online test. Please go to: www.dentalcare.com/en-us/dental-education/continuing-education/ce410/ce410-test.aspx

1. **The maximum allowable fluoride in a NaF-containing OTC dentifrice is ________ ppm.**
   a. 850
   b. 1150
   c. 1300
   d. 1700

2. **NDAs are typically used for which of the following?**
   a. All oral medications
   b. Food supplements
   c. New drug indications
   d. New flavor formulations

3. **The ADA Seal of Acceptance program is a required process to market a fluoride dentifrice in the US.**
   a. True
   b. False, the ADA Seal of Acceptance program is a voluntary process to provide assurance to consumers of the safety and efficacy of a product.
   c. False, the ADA Seal of Acceptance program is a voluntary program to monitor the efficacy of dentifrices.

4. **Fluoride salts approved by the FDA for use in US dentifrices include which of the following?**
   a. stannous fluoride
   b. sodium fluoride
   c. sodium monofluorophosphate
   d. Only B and C
   e. A, B and C

5. **Dental calculus control is considered to be what type of benefit?**
   a. therapeutic
   b. cosmetic
   c. It can be either cosmetic or therapeutic, depending on the product.

6. **The ISO standard specifies that dentifrices with an RDA of less than ________ are safe on hard tissues.**
   a. 100
   b. 150
   c. 250
   d. 300

7. **Which of the following is less soluble at pH 5?**
   a. fluorapatite
   b. hydroxyapatite

8. **Potassium nitrate is the most commonly used tubule-occluding agent used to treat dentinal hypersensitivity.**
   a. True
   b. False
9. Dentifrice ingredients with antibacterial activity can help reduce all of the following EXCEPT ___________.
   a. gingivitis
   b. intrinsic stain
   c. oral malodor
   d. plaque biofilm

10. Triclosan is a broad-spectrum antibacterial agent.
    a. True
    b. False, triclosan is a chemical whitening agent.
    c. False, triclosan is a tubule-occluding agent.

11. What is the most common nerve depolarizing agent used in sensitivity dentifrice?
    a. amine fluoride
    b. potassium nitrate
    c. sodium hexametaphosphate
    d. strontium chloride

12. Humectants allow dentifrices to retain moisture.
    a. True
    b. False

13. What is the most common surfactant used in dentifrices?
    a. polyoxyethylene glycol
    b. sodium dodecyl sulfate
    c. sodium hydrogenate
    d. sodium lauryl sulfate

14. Which of the following ingredients add opacity to a dentifrice?
    a. mica
    b. titanium dioxide
    c. xanthan gum
    d. xylitol

15. The active ingredient in the first dentifrice to receive the ADA Seal of Acceptance was ____________.
    a. sodium fluoride
    b. sodium monofluorophosphate
    c. stannous fluoride

16. The primary mechanism of action for fluoride includes _____________.
    a. promotion of remineralization
    b. inhibition of demineralization
    c. Both A and B
    d. None of the above.

17. Three dentifrice ingredients (abrasive, humectants, and solvent) typically represent about ___________ of the dentifrice ingredients.
    a. 25%
    b. 50%
    c. 75%
    d. 95%
18. Stannous fluoride is the only fluoride that delivers anticaries, antiplaque/gingivitis, enamel protection and anti-hypersensitivity benefits from one ingredient.
   a. True
   b. False

19. Effective anticalculus agents work by chelating calcium and inhibiting plaque calcification.
   a. True
   b. False

20. Halitosis, or bad breath, can be caused by which of the following?
   a. dry mouth
   b. bacteria
   c. oral debris
   d. oral diseases
   e. All of the above.

21. With fluoride treatment, a noncavitated lesion can be remineralized with fluorapatite and have greater resistance to subsequent demineralization than hydroxyapatite.
   a. True
   b. False

22. Sodium fluoride can be formulated with almost any of the available dentifrice abrasives.
   a. True, sodium fluoride is easily formulated with almost any abrasive.
   b. False, only compatible abrasive systems can be used, due to the high reactivity of sodium fluoride.

23. The key therapeutic areas for dentifrices are:
   a. caries
   b. gingivitis
   c. sensitivity
   d. whitening
   e. A, B and C

24. Without the binder, the toothpaste would separate into different phases, a liquid portion and a solid-like portion, and would have to be stirred before each use.
   a. True
   b. False

25. Which of the following reduces plaque and gingivitis?
   a. stannous fluoride
   b. potassium nitrate
   c. triclosan
   d. A and C
   e. A and B
References
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Dr. Koenigs is a Principal Scientist at The Procter & Gamble Company. Dr. Koenigs attended the University of Kansas as a Watkins-Berger Scholar where she received a Bachelor of Science in Chemistry and was awarded the Clark E. Bricker Scholarship for Chemistry. She attended Duke University to receive her doctorate in Physical Organic Chemistry where she researched the inhibition and photo-reactivation of serine protease enzymes in the blood coagulation cascade. While at Duke University she was the recipient of an NIH Pharmacology Training Grant. Dr. Koenigs has worked for P&G for 20 years. Her first 10 years at P&G were spent in drug discovery research with P&G Pharmaceuticals in the area of bacterial infectious disease. Since that time she has been involved with technical communication and training in the areas of osteoporosis, respiratory sciences and oral care. Dr. Koenigs currently holds the title of Scientific Information Manager in P&G’s Global Oral Health Professional & Scientific Relations Organization.

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