JOURNAL OF THE AMERICAN DENTAL ASSOCIATION:

Ingested fluoride does not reduce tooth decay, action is topical

Citizens for Safe Drinking Water

1010 University Avenue # 52 San Diego, CA 92103

> Contact: Jeff Green (800) 728-3833 greenjeff@cox.net

he featured Cover Story of the July 2000 *Journal of the American Dental Association* (JADA) is a 13 page article authored by John Featherstone, M.Sc., Ph.D., Professor and Chair, Department of Preventive and Restorative Dental Sciences and Department of Dental Public Health and Hygiene, University of California, San Francisco.

If that sounds like a mouthful, it is. And that is exactly what Dr. Featherstone delivers.

Following in the footsteps of 16 other cited articles that he has either authored or co-authored, Featherstone reminds the reader (in this case the dues-paying members of the American Dental Association) that fluoride's preventive action is topical rather than systemic.

Before you begin jumping up and down in delight or horror, depending on whether you are an avid promoter of using the public water for mass medication or a modern-day Keeper-of-the-Well, hold your horses.

Featherstone continues to give water fluoridation credit for helping reduce tooth decay – not because it is ingested – but because, he states, fluoridated water and other fluoride-containing beverages, foods, and oral care products contribute to the daily topical application of fluoride by bathing the teeth.

In his article, *The Science and Practice of Caries Prevention*, Featherstone concludes that fluoride is a key agent in battling caries, but that it is accomplished by three

Featherstone makes it clear, as he has in other publications, that fluoride incorporated during tooth development is insufficient to play a significant role in caries protection. principal topical mechanisms of action: inhibition of demineralization, enhancement of remineralization, and inhibition of bacterial enzymes. He notes that remineralization of early lesions also requires calcium and phosphate, which are derived primarily from saliva and plaque fluid.

Featherstone makes it clear, as he has in other publications, that fluoride incorporated during tooth development is insufficient to play a significant role in caries protection. He cites, as an example of the weak pre-eruptive effects of fluoride, a study of two groups of Okinawa nursing students which showed that there was no difference in caries status between those

who had received fluoridated water only until about 5 to 8 years of age (and none thereafter) and those who had never received fluoridated drinking water.

"Importantly, this means that fluoride incorporated during tooth mineral development at normal levels of 20 to 100 ppm (even in areas that have fluoridated drinking water or with the use of fluoride supplements) does not measurably alter the solubility of the mineral," writes Featherstone. "Even when the outer enamel has higher fluoride levels, such as 1000 ppm, it does not measurably withstand acid-induced dissolution any better than enamel with lower levels of fluoride."

This is not new information to those who have thoroughly researched the issue, nor is it new for Dr. Featherstone to write or lecture on this and many other findings with the same results. Dr. Featherstone was a featured speaker at the 1997 Canadian conference on controlled-dose fluoride supplements at which he reminded representatives of Canada's dental and medical stakeholders of the reports given at their 1992 conference on the same subject. These reports indicated that ingested fluoride cannot raise the fluoride concentration in the glanular saliva sufficient to meet the bacterial challenge present in the oral cavity.

Ingested fluoride does not reduce tooth decay, cont.

As a consequence of that conference, the Canadian Dental Association no longer recommends fluoride supplements if a child brushes his teeth twice a day with fluoridated toothpaste, and if individual practitioners are determined to increase fluoride exposure for high caries-risk patients in non fluoridated regions, CDA recommends that supplements be used as topical lozenges rather than swallowed.

With a title as all-inclusive-sounding as *The Science and Practice of Caries Prevention,* it is all too easy to erroneously dismiss its importance by pointing to what was not covered, such as irrigation devices, baby bottle tooth decay, and early childhood exposure to oral infections from primary care givers; yet many readers may be surprised at the introduction of other topics that may warrant their further exploration.

Other detractors may be angered by the avoidance of any mention about what constantly bathing the teeth with fluoride will do for total fluoride exposure and the consequences to every other organ and systemic function; but alas the dental industry has no responsibilities in other health arenas, so this publication may never be seen as the appropriate venue for that discussion.

Once again the Journal of the American Dental Association has confirmed that the dental community is not all of one mind... And yes, one topic that is glaring by its omission in a dental discussion is the incidence of dental fluorosis, which has risen to include 66.5% of our children in fluoridated communities displaying the visible signs of fluoride overdose on at least one tooth.

So why is this article so important? Contrary to promotional brochures printed by the dental trade organizations touting the safety and effectiveness of fluoride at virtually any exposure level, previous articles published by JADA

warned of high concentrations of fluoride in chicken and other baby foods, and advised dentists to warn parents to restrict children's consumption of fruit juices because of fluoride pesticide residues. Once again the Journal of the American Dental Association has confirmed that the dental community is not all of one mind, that the portrayal of the oral ingestion of fluoride as magically-effective but never-unsafe can now be corrected by the dental association membership.

It matters not whether Featherstone does or does not have the proof for his contention that 1 ppm in the water flowing past the teeth plays any significant role in caries reduction compared to the 1000 ppm in fluoridated tooth paste. The true question remains:

Will an informed public, a deliberative body such as a water board or city council, or health professional with no other axe to grind than the overall well-being of their patient, continue to support the forced everyday ingestion of a substance that is functional only as a topical application?



THE SCIENCE AND PRACTICE OF CARIES PREVENTION

JOHN D.B. FEATHERSTONE, M.SC, PH.D.

ABSTRACT

Background and Overview. Dental caries is a bacterially based disease. When it progresses, acid produced by bacterial action on dietary fermentable carbohydrates diffuses into the tooth and dissolves the carbonated hydroxyapatite mineral—a process called demineralization. Pathological factors including acidogenic bacteria (mutans streptococci and lactobacilli), salivary dysfunction, and dietary carbohydrates are related to caries progression. Protective factors-which include salivary calcium, phosphate and proteins, salivary flow, fluoride in saliva, and antibacterial components or agentscan balance, prevent or reverse dental caries.

Conclusions. Caries progression or reversal is determined by the balance between protective and pathological factors. Fluoride, the key agent in battling caries, works primarily via topical mechanisms: inhibition of demineralization, enhancement of remineralization and inhibition of bacterial enzymes. Clinical Implications. Fluoride in drinking water and in fluoride-containing products reduces caries via these topical mechanisms. Antibacterial therapy must be used to combat a high bacterial challenge. For practical caries management and prevention or reversal of dental caries, the sum of the preventive factors must outweigh the pathological factors.

Ithough the prevalence of dental caries in children has declined markedly over the last 20 years in most countries in the Western world,

the disease continues to be a major problem for both adults and children everywhere.

The trends in caries in U.S. children during the last 30 years were recently summarized¹ on the basis of results of four national surveys.²⁻⁶ By the late 1980s, although approximately 75 percent of children aged 5 to 11 years were caries-free, about 70 percent of the 12- to 17-year-olds still had caries. Approximately 25 percent of children and adolescents in the 5- to 17-year age range accounted for 80 percent of the caries in permanent teeth. By age 17 years, however, 40 percent of the population accounted for 80 per cent of the caries.¹⁻⁶ These findings illustrate the need for management of caries by individual risk assessment and for measures more specifically directed to high-risk people and populations.

Although these prevalence rates still leave much to be desired, the overall caries prevalence in children has indeed declined in the United States. Smaller epidemiologic studies in recent years indicate, however, that the decline in caries has not continued during the 1990s and that it may have plateaued.⁶

The reasons for the reductions in caries prevalence during the last 20 years are difficult to pinpoint. Strong evidence exists, however, that the near universal use of fluoride containing products such as dentifrice, mouthrinses and topical gels applied in the dental office have been major contributors.^{7,8} Earlier caries reductions of 40 to 70 percent (before the 1970s) had resulted from the fluoridation of public water supplies in many communities.⁹⁻¹²

Dental caries in adults also continues to be a major problem, as illustrated by a recent U.S. survey.¹³ The survey reported that 94 percent of all dentate adults (aged 18 years or older) had evidence of treat ed or untreated coronal caries.

Caries obviously still is a major problem in adults, as well as children, and we need an improved approach to prevention and therapy. This article reviews and summarizes the scientific basis for and practice of successful intervention in the caries process.

THE CARIES PROCESS

Bacterial plaque and acid production. The caries process is now well-understood; much of it has been described extensively in the dental literature. Some details of the caries process remain to be unraveled, but, in general, we understand the process well enough to initiate better-targeted methods of caries prevention and intervention.

The mechanism of dental caries formation is essentially straightforward.¹ Plaque on the surface of the tooth consists of a bacterial film that produces acids as a byproduct of its metabolism.^{14,15} To be specific, certain bacteria within the plaque are acidogenic—that is, they produce acids when they metabolize fermentable carbohydrates.^{12,14,15} These acids can dissolve the calcium phosphate mineral of the tooth enamel or dentin in a process known as demineralization.¹⁶⁻¹⁸ If this process is not halted or reversed via remineralization the redeposition of mineral via saliva—it eventually becomes a frank cavity.

Dental caries of the enamel typically is first observed clinically as a so-called "white-spot lesion." This is a small area of subsurface demineralization beneath the dental plaque. The

The mutans streptococci and the lactobacilli, either separately or together, are the primary causative agents of dental caries.

body of the subsurface lesion may have lost as much as 50 percent of its original mineral content and often is covered by an "apparently intact surface layer."¹⁹ The surface layer forms by remineralization. The process of demineralization continues each time there is carbohydrate taken into the mouth that is metabolized by the bacteria. The saliva has numerous roles, including buffering (neutralizing) the acid and remineralization by providing minerals that can replace those dissolved from the tooth during demineralization.1,20,21

Any fermentable carbohydrate (such as glucose, sucrose, fructose or cooked starch) can be metabolized by the acidogenic bacteria and create the aforementioned organic acids as byproducts.²² The acids diffuse through the plaque and into the porous subsurface enamel (or dentin, if exposed), dissociating to produce hydrogen ions as they travel.^{17,23} The hydrogen ions readily dissolve the mineral, freeing calcium and phosphate into solution, which can diffuse out of the tooth. Most importantly, lactic acid dissociates more readily than the other acids, producing hydrogen ions that rapidly lower the pH in the plaque.'7 As the pH is lowered, acids diffuse rapidly into the underlying enamel or dentin.

The two most important groups of bacteria that predominantly produce lactic acid are the mutans streptococci and the lactobacilli.'4 Each group contains several species, each of which is cariogenic. Mutans streptococci include Streptococcus mutans and S. sobri nus. The lactobacilli species also are prolific producers of lactic acid and appear in plaque before caries is clinically observed.^{24 25} These two groups of bacteria, either separately or together, are the primary causative agents of dental caries.

HOW FLUORIDE COMBATS THE CARIES PROCESS

The ability of fluoride to prevent and arrest caries has been researched extensively. Fluoride has three principal topical mechanisms of action:

 inhibiting bacterial metabolism after diffusing into the

bacteria as the hydrogen fluoride, or HF, molecule when the plaque is acidified;

- inhibiting demineralization when fluoride is present at the crystal surfaces during an acid challenge;

- enhancing remineralization and thereby forming a lowsolubility veneer similar to the acid-resistant mineral fluorapatite, or FAP, on the remineral ized crystals.

Inhibiting bacterial metabolism. Several investigators have studied the possible effects of fluoride on oral bacteria.²⁶⁻²⁸ The most significant finding reported is that the ion ized form of fluoride, or F⁻, can not cross the cell wall and membrane but can rapidly travel into the cariogenic bacterial cells in the unchanged form as HF. ²⁶⁻²⁸

When the pH in the plaque falls as the bacteria produce acids, a portion of the fluoride present in the plaque fluid then combines with hydrogen ions to form HF and rapidly diffuses into the cell, effectively drawing more HF from the outside.^{1,26-28} Once inside the cell, the HF dissociates, acidifying the cell and releasing fluoride ions that interfere with enzyme activity in the bacterium. For example, fluoride inhibits enolase, an enzyme necessary for the bacteria to metabolize carbohydrates. As fluoride is trapped in the cell, the process becomes cumulative.

In summary, fluoride from topical sources is converted partially to HF by the acid that the bacteria produce and diffuses into the cell, thereby inhibiting essential enzyme activity.

Inhibiting demineralization. The mineral of our teeth (enamel, cementum, dentin)



Figure 1. High-resolution electron microscope images (magnification approximately x2,000,000) of individual enamel crystals. The black lines are rows ot calcium atoms, which are visualized by this tech nique. A. Normal enamel crystal showing white patches (arrows), which are calcium-deticient and carbonate-rich detect regions. B. Demineralized crystal trom the body ot a natural caries lesion showing "large" hexagonal holes coinciding with the "small" detect regions seen in normal enamel. (Adapted trom Featherstone and colleagues.30,31)

and bones is a carbonated hydroxyapatite²⁹ that can be approximately represented by this simplified formula:

 $\begin{array}{c} Ca_{10\text{-}x}(Na)_x(PO_4)_{6\text{-}y}(CO_3)_z\\ (OH)_{2\text{-}u}(F)_u\end{array}$

The substitutions in the hydroxyapatite crystal lattice (the arrangement of atoms and ions in the crystal) occur as the mineral is first laid down during tooth development, with the carbonate (CO₃) ion in particular causing major disturbances in the regular array of ions in the crystal lattice.^{30,31} During demineralization, the carbonate is lost, and during remineralization it is excluded from the newly formed mineral. The calcium-deficient, carbonate-rich regions of the crystal are espe-



Figure 2. Typical pH curves for normal subJocts with normal salivary flow and for subjects with xerostomia (mean for each group) after ingestion of sucrose. A curve for ingestion of a sugar-free sweetened product is shown for comparison. (Reproduced from Featherstone' with permission of the publisher. Copyright © 1999, Munksgeard.)

cially susceptible to attack by the acid hydrogen ions during demineralization, as has been shown by several investiga tors.^{21,29-33} High-resolution lattice imaging, which images crystals almost to atomic resolution (viewed at about x2,000,000 magnification), was used to illustrate the appearance of hexagonal holes in the early stages of enamel crystal dissolution in dental caries (Figure 1), which coincided with the calcium-deficient, carbonate-substituted regions of the crystal.30-33

The carbonated hydroxyapatite, or CAP, of our teeth is much more soluble in acid than hydroxyapatite, or HAP (HAP = $Ca_{10}(PO_4)_6(OH)_2$), and that in turn is much more soluble than fluorapatite, or FAP (FAP = $Ca^{10}(PO^4)_6F_2)_{,21}$ in which the OH⁻ ion in pure hydroxyapatite is completely replaced by an F⁻ ion. The resulting mineral FAP is highly resistant to dissolution by acid.

Fluoride inhibits demineralization. Sound enamel, except in its outer few micrometers, generally contains fluoride at levels of about 20 to 100 parts per million, or ppm, depending on the fluoride ingestion during tooth development.34 Teeth in children who lived in areas with fluoridated drinking water during tooth development have fluoride content toward the higher end of this range. The outer few micrometers of enamel can have fluoride levels of 1,000 to 2,000 ppm.³⁴

Fluoride in the solution surrounding CAP crystals has been shown to be much more effective in inhibiting demineraliza-

tion than fluoride incorporated into the crystals at the levels found in enamel.^{21,35} Ten Cate,²¹ Nelson and colleagues³⁵ and Featherstone and colleagues^{35,37} found no measurable reduction in the acid solubility of synthetic CAP (3 percent CO₃ by weight, comparable to that of dental enamel mineral) with about 1,000 ppm fluoride incorporated. Importantly, this means that fluoride incorporated during tooth mineral development at normal levels of 20 to 100 ppm (even in areas that have fluoridated drinking water or with the use of fluoride supplements) does not measurably alter the acid solubility of the mineral. Even when the outer enamel has higher fluoride levels, such as 1,000 ppm, it does not measurably withstand acidinduced dissolution any better than enamel with lower levels of fluoride. Only when fluoride is concentrated into a new crystal surface during remineralization is it sufficient to beneficially alter enamel solubility. The fluoride incorporated developmentally—that is, systemically into the normal tooth mineralis insufficient to have a measurable effect on acid solubility.^{21,38}

In contrast to the lack of effect of fluoride incorporated into the CAP crystals of tooth mineral developmentally, as little as 1 ppm in the acid solution reduced the dissolution rate of CAP to a rate equivalent to that of HAP.³⁶ Further increases in fluoride in the acid solution in contact with the CAP mineral surface decreased the solubility rate logarithmically. These results indicate that if fluoride is present in the aqueous solution surrounding the crystals, it is adsorbed strongly to the surface of CAP carbonat-

ed apatite (enamel mineral) crystals and thus acts as a potent protection mechanism against acid dissolution of the crystal surface in the tooth's subsurface region. If fluoride is in the plaque fluid at the time that the bacteria generate acid, it will travel with the acid into the subsurface of the tooth and, therefore, adsorb to the crystal surface and protect it against being dissolved.

In summary, fluoride present in the water phase at low levels among the enamel or dentin crystals adsorbs to these crystal surfaces and can markedly inhibit dissolution of tooth mineral by acid.^{21,36} Fluoride that acts in this way comes from the plaque fluid via topical sources such as drinking water and fluoride products. Fluoride incorporated during tooth development is insufficient to play a significant role in caries protection. Fluoride is needed regularly throughout life to protect teeth against caries.

Enhancing remineralization. As the saliva flows over the plaque and its components neutralize the acid, raising the pH (Figure 2), demineralization is stopped and reversed. The saliva is supersaturated with calcium and phosphate, which can drive mineral back into the tooth.^{21,39} The partially demineralized crystal surfaces within the lesion act as "nucleators," and new surfaces grow on the crystals (Figure 3). These processes constitute remineralization-the replacement of mineral in the partially demineralized regions of the carious lesion of enamel or dentin (including the tooth root).^{20,21} Fluoride enhances remineralization by adsorbing to the crystal surface and attracting calci-



Figure 3. High-resolution electron microscope images (magnification approximately x2,000,000) of individual enamel crystals that visualize remineralization at the atomic level. The black lines are rows of calcium atoms, which are visualized by this technique. A. Normal enamel crystal dissected from the inner region of enamel, showing "small" white patches of calcium-deficient, carbonate-rich regions. B. Crystal on which a "remineralized" surface veneer has been grown after treatment with fluoride, calcium and phosphate. (Adapted from Featherstone and colleagues, 1981.³⁰)

um ions, followed by phosphate ions, leading to new mineral formation. The newly formed "veneer" excludes carbonate and has a composition somewhere between HAP and FAP as described above (Figure 4). FAP contains approximately 30,000 ppm F and has a very low solubility in acid. The new







Figure 5. The caries balance: a schematic diagram of the balance between pathological and protective factors in the caries process. (Reproduced from Featherstone' with permission of the publisher. Copyright ©1999, Munksgaard.)

remineralized crystal now will behave like low-solubility FAP rather than the highly soluble CAP of the original crystal surface.³⁶

In summary, fluoride in solution from topical sources enhances remineralization by speeding up the growth of a new surface on the partially demineralized subsurface crystals in the caries lesion. The new crystal surface veneer is FAP-like, with much lower solubility than the original CAP tooth mineral. Subsequent acid challenges must be quite strong and prolonged to dissolve the remineralized enamel.

Saliva and caries. Saliva has a critical role in the prevention or reversal of the caries process; it provides calcium, phosphate, proteins that maintain supersaturation of calcium in the plaque fluid, proteins and lipids that form a protective pellicle on the surface of the tooth, antibacterial substances and buffers.⁴⁰ The saliva components neutralize the acids produced by bacterial metabolism in the plaque, raise the pH and reverse the diffusion gradient for calcium and phosphate. Thereby, they return calcium and phosphate to the subsurface lesion, where these ions can regrow new surfaces on the crystal remnants that were produced by demineralization. These so-called "remineralized" crystals have a veneer of much less soluble mineral. Saliva also clears carbohydrates and acids from the plaque.

In the case of salivary dysfunction,⁴¹ all of the above benefits of saliva are reduced or eliminated (as is illustrated partially in Figure 2 by the pH profile of the subjects with xerostomia).

THE CARIES BALANCE

Fluoride's three extensively studied and documented principal mechanisms of action rely on the presence of fluoride in saliva, in the plaque at the tooth surface and in the fluid among the crystals in the subsurface of the enamel or dentin. The clinical effects of fluoride, therefore, can be optimized by using delivery methods that bring fluoride to the surface of the tooth and into the plaque rather than incorporating fluoride into the tooth mineral crystals during tooth development. These topical delivery methods are equally applicable to adults and children and include fluoride in beverages and foods, dental products and drinking water. The benefits of continually providing low levels of fluoride in the saliva and plaque from the aforementioned topical sources are described more fully

in a recent review article.1

Pathological and protective factors in the caries balance. Caries progression, as opposed to reversal, consists of a delicate balance between the aforementioned factors-namely, a bacterially generated acid challenge and a combination of demineralization inhibition and reversal by remineralization.^{1,42} The balance between pathological factors (such as bacteria and carbohydrates) and protective factors (such as saliva, calcium, phosphate and fluoride) is a delicate one that swings either way several times daily in most people (Figure 5).

Protective factors. Saliva is essential for the protection of the tooth against dental caries and provides many natural protective factors summarized earlier,^{40,41} including calcium, phosphate, antibacterial components and other proteins with various functions. Extrinsic antibacterial agents such as chlorhexidine also can be considered as protective factors in this balance, as can fluoride from external sources. The mechanisms of action of fluoride described in this article apply primarily to fluoride from topical sources; systemically incorporated fluoride has only a minor role in protecting against dental caries. This conclusion is supported not only by laboratory data as described previously, but also by epidemiologic studies. For example, a four-year study in England found a 27 percent lower caries incidence among children who were 12 years old when water fluoridation began in their communities, relative to the incidence in control subjects of the same age in nonfluoridated areas.43 This was a wellconducted study, and it clearly

showed the posteruptive (topical) effects of fluoride in the drinking water. Other studies have illustrated the weak preeruptive effects of fluoride. For example, in two groups of Okinawa nursing students aged 18 to 22 years, there was no difference in caries status between those who had received fluoridated water only until about 5 to 8 years of age (and none thereafter) and those who had never received fluoridated drinking water.44 The cariostatic effects of fluoride are, in part, related to the sustained presence of low concentrations of ionic fluoride in the oral environment,^{1,21,38}

> There is a mistaken belief that drilling out a caries lesion and placing a restoration eliminates the bacteria and thereby stops caries progression.

derived from foods and beverages, drinking water and fluoride-containing dental products such as toothpaste. Prolonged and slightly elevated low concentrations of fluoride in the saliva and plaque fluid decrease the rate of enamel demineralization and enhance the rate of remineralization.^{21,26,38,45-48} For example, fluoride at 0.04 ppm in saliva can enhance remineralization. Remineralization of early lesions also requires calcium and phosphate, which are derived primarily from saliva and plaque fluid.

Pathological factors. Pathological factors obviously include cariogenic bacteria and the frequency of ingestion of fermentable carbohydrates that sustain these bacteria. The importance of mutans streptococci (which includes S. mutans and S. sobrinus) in the development of dental caries has been reviewed extensively.^{12,14,15,49,50} Numerous cross-sectional studies in humans have shown that greater numbers of mutans streptococci and lactobacilli in saliva or plaque are associated with high caries rates.15,25,49,51-54 Longitudinal studies have shown that an increase over time in numbers of both of these bacterial groups is associated with caries onset and progression.24,55,56

CARIES INTERVENTION

The methods of caries intervention can be summarized by joining the principal components of the caries process with the interventional possibilities (Table).

Cariogenic bacteria and high bacterial challenge. Dental caries is a transmissible, bacterially generated disease. There is the mistaken belief that drilling out a caries lesion and placing a restoration eliminates the bacteria and thereby stops caries progression. Although traditional restorative work may eliminate the bacteria at the site of the restoration, the remainder of the mouth is left untouched, caries continues unchecked in the remainder of the mouth and recolonization commences rapidly at the margins.57

It is logical, therefore, to use

TABLE

SUMMARY: THE CARIES PROCESS AND METHODS OF CARIES INTERVENTION.

CARIES COMPONENT	INTERVENTION METHOD
Bacteria	Antibacterial therapy such as treatment with chlorhexidine gluconate (see text.)
Carbonated Hydroxyapatite	Make the mineral less soluble by transforming it to other crystalline forms such as hydroxy- apatite without carbonate (future caries- preventative treatments by specific laser irradia- tion will enable this to be done ^{69,70})
Fermentable Carbohydrates	Reduce the frequency of ingestion; substitute with noncariogenic sweeteners (this method is well-accepted and used in patient education.
	Recommend use of sugar-free chewing gum, which reduces frequency of fermentable carbo- hydrate ingestion and also enhances remineral- ization
Organic Acids Produced by Oral Bacteria	Neutralize the acid by providing extra buffer- ing or enhancing saliva; sugar-free gum assists in this as well
Saliva	Enhance the saliva flow and function
Fluoride	Exploit its known effects on bacteria, inhibi- tion of demineralization and enhancement of remineralization by using "topical" fluoride delivery by means of dental products, drinking water, beverages and foods

antibacterial therapy—such as treatment with chlorhexidine gluconate rinse—as a cariespreventive measure. Although this has been proposed for many years⁵⁸⁻⁶⁰ and used in several European countries, an antibacterial approach almost never is used in the United States for the prevention of the progression of dental caries.

One of the difficulties in persuading clinicians to use the antibacterial approach is that there have not been rapid and accurate methods of determining the levels of cariogenic bacteria in the mouth. Furthermore, although numerous studies have indicated that mutans streptococci and lactobacilli definitely are risk factors for dental caries, there is no one-to-one direct correlation between levels of these bacteria and caries progression.24,49 However, it now is well-established that high levels of mutans streptococci, high levels of lactobacilli or both constitute a "high bacterial challenge."24 This bacterial challenge can be balanced by the protective factors described earlier, which include salivary componentsespecially calcium, phosphate and fluoride-and the amount of saliva present.42 Figure 5 illustrates the balance between pathological factors (including cariogenic bacteria, reduced salivary function and frequency of use of fermentable carbohydrates) and protective factors. If these pathological and protective factors are in balance, caries does not progress. If they are out of balance, caries either progresses or reverses.

Antibacterial therapy for caries control. Currently, the most successful antibacterial therapy against cariogenic bacteria is treatment by chlorhexidine gluconate rinse or gel.47,61 Chlorhexidine is available by prescription in the United States. Studies have shown that a daily dose of chlorhexidine rinse for two weeks can markedly reduce the cariogenic bacteria in the mouth and that, as a result, recolonization takes place in three to six months rather than immediately.58 In patients with high levels of bacteria, therefore, chlorhexidine treatments at three-month

intervals are indicated.

The problem faced by clinicians is how to determine, in a timely fashion, whether the bacterial challenge is high, medium or low. For many years, commercial "dip slides" have been available in Europe, and they recently became available in the United States.⁵⁸ A saliva sample is taken from the patient and incubated on the dip slide; two days later, a result is provided of the levels of S. mutans and lactobacilli bacteria in the mouth.58 Although these slides are a major advance in convenience and are the best tools available at the time of this writing, it has been shown that this technology is not well-correlated with traditional bacterial plating. It is anticipated that methods of rapid chairside assessment of bacterial challenge, based on molecular biology, will be available in the future.

Several investigators have explored the possibility of using modern molecular biology for better and more rapid methods of bacterial assessment,⁶² but they were unable to overcome a number of complications. An exciting development is work by Shi and colleagues,63 who recently published a method using speciesspecific monoclonal antibodies that recognize the surface of cariogenic bacteria. With this technology, it is not necessary to split open the bacterial cells to assess the internal DNA or RNA. These probes can be tagged either with a fluorescent molecule or with a marker that can be measured quantitatively in a simple spectrophotometer.

It is anticipated that these

probes will be available commercially in the near future, and that clinicians will be able to use them chairside and obtain results within a few minutes. This will enable clinicians to determine the quantitative levels of bacteria in a patient's mouth while he or she is in the operatory and to factor these numbers into an overall risk assessment of caries for that patient. It is envisaged that computer programs will be available that will include the assay numbers, as well as other data. The practitioner will receive guidance as to the level of

> Methods of rapid chairside assessment of bacterial challenge, based on molecular biology, will be available in the future.

caries risk and what regimen or regimens to use to prevent further caries and to reduce the bacterial challenge. With the new monoclonal antibody probes, the levels of bacteria and success of the intervention could readily be followed over time. This is an exciting, innovative tool that may become widely used and accepted within a few years.

CARIES RISK ASSESSMENT

Several studies have attempted to determine risk factors that can be reliably used to assess the level of risk of caries progression in individual patients. Studies still are under way, and there is no definitive formula yet available. The status of risk assessment was summarized, however, by the authors of a special supplement to The Journal of the American Dental Association in 1995; this publication can be used as a guide until more definitive information is available.⁶⁴ Figure 5 represents a basis for determining caries risk with the information currently available.

It has been established that high-risk patients include those who have a high bacterial challenge, which may consist of a combination of high numbers of mutans streptococci, lactobacilli or both. Although fluoride has excellent properties in terms of balancing caries challenge, if the challenge is too high, then fluoride-even at increased concentrations, with increased use or both-cannot balance that challenge. Therefore, in the case of high bacterial challenge, the bacterial infection must be dealt with, typically with a chlorhexidine rinse, as well as the enhancement of salivary action by topical delivery of fluoride. These principles apply equally well to adults and children. Accurate detection of early caries can increase the reliability of caries risk assessment, particularly if those measurements are made at three- or six-month intervals and caries progression can be measured. In the case of caries progression, obviously, intervention is needed either antibacterially, with fluoride or





with other techniques, some of which are described in the following material.

Caries management by risk assessment. As the caries risk assessment methodologies are refined, we will have more definitive biological and chemical risk assessment measures to guide clinical decision making. These measures form the basis for assessing the direction in which the caries balance is likely to move for a particular patient. Early caries detection, especially in occlusal surfaces, is an essential part of caries management by risk assessment.

Caries management by risk assessment now is receiving considerable attention, and software programs are being developed that will aid practitioners in assessing risk and lead them to the use of current and new technologies by specifying treatments recommended for the various risk categories.^{59,60} As we move into the future, tooth restorations will become less and less desirable as a treatment and will be used only as a final resort when new intervention measures have failed or when people have not participated in caries intervention programs such as those indicated previously.

CARIES MANAGEMENT TOOLS FOR THE FUTURE

Several technological advancements are currently close to clinical reality and will be embraced if they are proven successful.

Assessment of bacterial challenge by chairside molecular probes. The use of chairside bacterial probes for assessing a patient's cariogenic bacterial challenge will be an essential component of caries management by risk assessment.

Caries immunization. In a program of caries management by risk assessment, it is logical that all available tools should be used. One such tool that has been investigated for many years is an immunization against caries. There are many obstacles to the success of immunization, as caries is not a systemic infection that can be dealt with simply by administering a specific antibiotic. The infection must be dealt with in the mouth, where the internal body fluids do not pass and, therefore, the normal immune response is not relevant. However, IgA that is produced by the saliva naturally can interfere with the colonization of the surface of the tooth by specific bacteria.

Recent studies by Ma and colleagues65,66 have illustrated the effectiveness of specific IgA in the inhibition of recolonization of mutans streptococci. The next logical step is to use this technology as one of the tools for caries intervention. It is possible to use genetically engineered plants, such as tobacco or alfalfa, to produce immunoglobulins.66,67 A study is in progress at the University of California, San Francisco, to test IgA that has been produced using genetically engineered tobacco plants. At press time, the results were not known, but if the trial is successful, this IgA can be applied to the teeth after chlorhexidine treatment has removed the cariogenic bacteria, with the aim of inhibiting future recolonization by mutans streptococci.

Early caries detection and intervention. Successful use of the innovative methods described here for caries intervention will require accurate methods for the early detection of dental caries in enamel and dentin. Early-detection methods such as fluorescence, optical coherence tomography, electrical impedance and ultrasonography are likely to become available for use by clinicians in the near future.⁶⁸ It will be possible to detect lesions in the occlusal surface and to determine whether they have progressed into the dentin and, if so, how far. This is not possible with current radiographic technology.

Once new methods are introduced for the early detection of caries, they can be used in two opposing fashions. Clinicians with traditional training are likely to use these methods to intervene physically at an earlier stage with carious lesions—drilling, filling and placing restorations. This outcome is of concern, as many more restorations would be placed than may be necessary, which weakens the tooth structure. Early detection and intervention by placing a restoration also does not take advantage of the body's natural protective mechanisms of inhibition of demineralization and enhancement of remineralization via saliva.

Alternatively, early detection of caries can be used as an opportunity to promote remineralization via salivary enhancement, use of topical fluoride and chlorhexidine and meticulous oral hygiene. In addition, as innovative methods for early caries intervention are introduced, the need for restorations may be eliminated for many patients, thereby preserving the tooth structure and halting or reversing progression of dental caries.

Caries prevention by laser treatment. In May 1997, the U.S. Food and Drug Administration approved the use of an erbium:yttriumaluminum-garnet, or Er:YAG, laser for use on teeth. This was the first approval for laser use on dental hard tissues. This approval by the FDA was for this particular laser to be used for the removal of dental caries and the cutting of sound tissue before the placement of restorations. This event has ushered in a new era for lasers in dentistry. Since then, other lasers have been approved for the same purpose, and additional hard-tissue uses are likely to be approved in the future, including the use of lasers for the inhibition of progression of dental caries by altering the composition of surface enamel

> As innovative methods for early caries intervention are introduced, the need for restorations may be eliminated for many patients.

or dentin mineral.

Kantorowitz and colleagues69 and Featherstone and colleagues70 have studied the effects of lasers on hard tissues for almost 20 years. The overall objective of these studies is to establish the scientific basis for the choice of laser parameters that can be used clinically for the prevention, removal or treatment of caries lesions. Their studies have demonstrated that specific pulsed carbon dioxide, or CO₂, laser treatment of dental enamel can inhibit subsequent carieslike progression in a severe demineralization-remineralization model in the laboratory by up to 85 percent. They have demonstrated that carbonate is lost from the CAP mineral of the tooth during specific laser irradiation, making the mineral highly resistant to dissolution by acid. Although they have demonstrated in the laboratory, using pH cycling models, that as little as 20 pulses of 100 microseconds each can produce a preventive effect similar to daily use of fluoride dentifrice, these promising and exciting results have not yet been tested in human mouths.70

For practical purposes, it would be desirable to develop a laser that can remove carious tissue and subsequently be used to treat the walls of the area from which carious tissue is removed to make them resistant to subsequent caries challenge⁷¹ (Figure 6). Fried and colleagues⁷² recently published a report on a new CO2 laser that efficiently removes carious tissue. After caries and a minimal amount of surrounding tissue are removed, it will be possible to change the laser parameters to perform cariespreventive treatment on the same area. This would be followed by placement of a resinbased composite restoration, thereby inhibiting subsequent caries around that restoration. For example, if an early occlusal lesion was detected (by the new methods described previously) that was deemed to be beyond hope of remineralization, this lesion could be conservatively removed with an appropriate laser. Then the surrounding cavity preparation walls could be treated for caries prevention by the laser and a small conservative restoration

placed. The cavity walls will be highly resistant to acid attack and therefore resistant to secondary caries. Providing bacterial intervention via chlorhexidine rinse was also part of the treatment in the same patient, future caries would be unlikely.

SUMMARY AND CONCLUSIONS

The mechanism of dental caries is well-established to the point where new approaches are being made for caries prevention based on a scientific understanding of the processes involved. Several existing methodologies are available to enable successful management of dental caries by risk assessment. Understanding the balance between pathological factors and protective factors is the key. Beyond the wellestablished and currently used methods, some innovative and exciting techniques have shown early research successes that most likely will be used for early caries intervention in the future. These methods include fluoride therapy for inhibition of demineralization and enhancement of remineralization, molecular probes for the quantitative detection of cariogenic bacteria at chairside, computerized caries risk assessment programs, genetically engineered IgA for inhibition of recolonization of cariogenic bacteria, specific lasers for conservative removal of carious tissue and specific lasers for the prevention of caries progression.

The use of these technologies will require extensive retraining of clinical dentists. But it will dramatically alter the way in which dentists diagnose, intervene, treat and manage caries, with major benefits to the oral

health of their patients.

Dr. Featherstone is a professor and the chair, Department of Preventive and Restorative Dental Sciences and Department of Dental Public Health and Hygiene, University of California, San Francisco, 707 Parnassus Ave., Box 0758, San Francisco, Calif. 94143, e-mail "jdbf@itsa.ucsf.edu". Address reprint requests to Dr. Featherstone.

The author sincerely acknowledges contributions from numerous colleagues over many years to much of the work reviewed here.

1. Featherstone JD. Prevention and reversal of dental caries: role of low level fluoride. Community Dent Oral Epidemiol 1999; $27(1) \cdot 31 - 40$

2. Kaste LM, Selwitz RH, Oldakowski RJ, Brunelle JA, Winn DM, Brown LJ. Coronal caries in the primary and permanent dentition of children and adolescents 1-17 years of age: United States, 1988-1991. J Dent Res 1996:75:631-41

3. National Institute of Dental Research Epidemiology and Oral Disease Prevention Program. Oral health of United States children: The National Survey of Dental Caries in U.S. School Children, 1986-1987-national and regional findings. Bethesda, Md.: U.S. Department of Health and Human Services Public Health Service, National Institutes of Health; 1989. NIH publication 89-2247

4. The prevalence of dental caries in United States children 1979-1980: The National Dental Caries Prevalence Survey. Bethesda Md.; U.S. Department of Health and Human Services, Public Health Service, National Institutes of Health; 1981. NIH publication 82-2245

5. National Center for Health Statistics. Decayed, missing and filled teeth among youths, 12-17 years: United States, 1974. Vital and Health Statistics, Series 11-No. 144. Washington: Government Printing Office; 1974. DHEW publication (HRA) 75-1626.

6. Speechley M, Johnston DW. Some evidence from Ontario, Canada, of a reversal in the dental caries decline. Caries Res 1996; 30(6):423-7

7. Jenkins GN. Recent changes in dental caries. Br Met J 1985;291(6505):1297-8.

8. Hargreaves JA, Thompson GW, Wagg BJ. Changes in caries prevalence in Isle of Lewis children between 1971 and 1981. Caries Res 1983;17(6):554-9.

9. Burt BA, Fejerskov O. Water fluoridation. In: Fejerskov O, Ekstrand J, Burt BA, eds. Fluoride in dentistry. Copenhagen,

Denmark Munksgaard, 1996:275-90. 10. Murray JJ. Fluorides in caries prevention. 3rd et. Oxford, England; Boston: Butterworth-Heinemann, 1991.

11. Newbrun E. Effectiveness of water fluoridation J Public Health Dent 1989,49: 279-89.

12. Newbrun E. Cariology. 3rd ed. Chicago: Quintessence; 1989:63-87, 331-49.

13. Winn DM, Brunelle JA, Selwitz RH, et al. Coronal and root caries in the dentition of adults in the United States, 1988-1991. J Dent Res 1996,75:642-51.

14. Loesche WJ. Role of Streptococcus mutans in human dental decay. Microbiol Rev 1986;50(4):353-80.

15. Loesche WJ, Hockett RN, Syed SA. The predominant cultivable flora of tooth surface plaque removed from institutionalized subjects. Arch Oral Biol 1972;17(9):1311-25. 16. Featherstone JD. An updated understanding of the mechanism of dental decay and its prevention. Nutr Q 1990,14:5-11.

17. Featherstone JD, Rodgers BE, Effect of acetic, lactic and other organic acids on the formation of artificial carious lesions. Caries Res 1981;16(5):377-85

18. Featherstone JD, Mellberg JR. Relative rates of progress of artificial carious lesions in bovine, ovine and human enamel. Caries Res 1981;15(1):109-14

19. Silverstone LM. Structure of carious enamel, including the early lesion. Oral Sci Rev 1973.3:100-60.

20. ten Cate JM, Duijsters PP. Influence of fluoride in solution on tooth demineralization. II. Microradiographic data. Caries Res 1983; 17(6):513-9.

21. ten Cate JM, Featherstone JD. Mechanistic aspects of the interactions between fluoride and dental enamel. Crit Rev Oral Biol Med 1991;2(3):283-96

22. Geddes DA. Acids produced by human dental plaque metabolism in situ. Caries Res 1975;9(2):98-109.

23. Featherstone JD. Diffusion phenomena and enamel caries development. In: Guggenheim B, ed. Proceedings of the Cariology Today International Congress, September 1983, Zurich, Switzerland, Basel Switzerland: Karger; 1984:269-68.

24. Leverett DH, Proskin HM, Featherstone JD, et al. Caries risk assessment in a longitudinal discrimination study. J Dent Res 1993.72(2).538-43

25. Leverett DH, Featherstone JD, Proskin HM et al. Caries risk assessment by a cross sectional discrimination model. J Dent Res 1993:72(2):529-37.

26. Hamilton IR, Bowden GH. Fluoride effects on oral bacteria. In: Fejerskov O, Ekstrand J, Burt BA, eds. Fluoride in dentistry. Copenhagen, Denmark: Munksgaard 1996:230-51.

27. Whitford GM, Schuster GS, Pashley DH, Venkateswarlu P. Fluoride uptake by Streptococcus mutans 6715. Infect Immunol 1977;18(3):680-7.

28. Van Louveren C. The antimicrobial action of fluoride and its role in caries inhibition. J Dent Res 1990;69:676-81.

29. LeGeros RZ. Calcium phosphates in oral biology and medicine. In: Myers HM, ed. Monographs in oral science. Basel

Switzerland: Karger, 1991:1-201 30. Featherstone JD, Nelson DG, McLean JD. An electron microscope study of modifications to defect regions in dental enamel and synthetic apatites. Caries Res 1981,15(4):

278-88.

31. Featherstone JD, Goodman P, McLean JD. Electron microscope study of defect zones in dental enamel. J Ultrastruc Res 1979;67: 117-23

32. Nelson DG, McLean JD. High-resolution electron microscopy of octacalcium phosphate and its hydrolysis products. Calcif Tissue Int 1984;36(2):219-32

33 Nelson DG, McLean JD. Direct observation of near-atomic details in synthetic and biological apatite crystallites. In: Fearnhead RW, Suga S, ed. Tooth enamel IV: Proceedings of the Fourth International Symposium on the Composition, Properties, and Fundamental Structure of Tooth Enamel.

Amsterdam, Netherlands: Elsevier Science Publishers; 1984:47-51.

34. Robinson C, Kirkham J, Weatherell JA. Fluoride in teeth and bone. In: Fejerskov O, Ekstrand J, Burt BA, eds. Fluoride in dentistry. Copenhagen, Denmark: Munksgaard; 1996:69 87

35. Nelson DG, Featherstone JD, Duncan JF, Cutress TW. Effect of carbonate and fluoride on the dissolution behaviour of synthetic apatites. Caries Res 1983;17(3):200-11.

36. Featherstone JD, Glena R, Shariati M, Shields CP. Dependence of in vitro demineralization of apatite and remineralization of dental enamel on fluoride concentration. J Dent Res 1990;69:620-5.

37. Featherstone JD, Shields CP, Khademazad B, Oldershaw MD. Acid reactivity of carbonated apatites with strontium and fluoride substitutions. J Dent Res 1983; 62(10):1049-53.

38. Fejerskov O, Thylstrup A, Larsen MJ. Rational use of fluorides in caries prevention. A concept based on possible cariostatic mechanisms. Acta Odont Scand 1981;39(4):241-9.

39. Moreno EC, Kresak M, Zahradnick RT. Physicochemical aspects of fluoride-apatite systems relevant to the study of dental caries. Caries Res 1977;11:142-71.

40. Lamkin MS, Oppenheim FG. Structural features of salivary function. Crit Rev Oral Biol Med 1993;4:251-9.

41. Mandel ID. Relation of saliva and plaque to caries. J Dent Res 1974;53(2): 246-66.

42. Featherstone JD. Clinical implications: New strategies for caries prevention. In: Stookey GK, ed. Indiana Conference 1996: Early Detection of Dental Caries. Indianapolis: Indiana University School of Dentistry, 1996:287-95.

43. Harwick JL, Teasdale J, Bloodworth G. Caries increments over 4 years in children aged 12 at the start of water fluroridation. Br Dent J 1982;153:217-22.

44. Kobayashi S, Kawasaki K, Takagi O, et al. Caries experience in subjects 18-22 years of age after 13 years' discontinued water fluoridation in Okinawa. Community Dent Oral Epidemiol 1992;20:81-3.

45. ten Cate JM, Mundorff-Shrestha SA. Working group report: laboratory models for caries (in vitro and animal models). Adv Dent Res 1995,9:332-4.

46. Zero DT. In situ caries models. Adv Dent Res 1995;9:214-30.

47. Featherstone JD, Zero DT. Laboratory and human studies to elucidate the mechanism of action of fluoride-containing dentifrices. In: Embery G, Rolla R, eds. (Clinical and biological aspects of dentitrices. Oxford, England: Oxford University Press; 1992: 41-50.

48. Arends J, Nelson DG, Dijkman AG, Jongebloed WL. Effect of various fluorides on enamel structure and chemistry. In: Guggenheim B, ed. Proceedings of the Cariology Today International Congress, September 1983, Zurich, Switzerland. Basel, Switzerland: Karger, 1984:245-58.

49. Krasse B. Biological factors as indicators of future caries. Int Dent J 1988;38(4): 219-25.

50. Ellen RP. Microbiological assays for dental caries and periodontal disease susceptibility. Oral Sci Rev 1976;(8):3-23.

51. Alalnusua S, Kleemola-Kujala E, Nystrom M, Evalahti M, Gronroos L. Caries in the primary teeth and salivary *Streptococcus mutans* and lactobacillus levels as indicators of caries in permanent teeth. Pediatr Dent 1987;9(2):126-30.

52. Klock B, Krasse B. Microbial and salivary conditions in 9- to 12-year-old children. Scand J Dent Res 1977,85(1):56-63.

53. Klock B, Emilson CG, Lind SO, Gustavsdotter M, Olhede-Westerlund AM. Prediction of caries activity in children with today's low caries incidence. Community Dent Oral Epidemiol 1989;17(6):285-8.

54. Seppa L, Hausen H. Frequency of initial caries lesions as predictor of future caries increment in children. Scand J Dent Res 1988;96(1):9-13.

55. Loesche WJ, Eklund S, Earnest R, Burt B. Longitudinal investigation of bacteriology of human fissure decay: epidemiological studies in molars shortly after eruption. Infect Immunol 1984;46(3):765-72.

56. Kingman A, Little W, Gomez 1, et al. Salivary levels of Streptococcus mutans and lactobacilli and dental caries experiences in a U.S. adolescent population. Community Dent Oral Epidemiol 1988,16(2):98-103.

57. Wright JT, Cutter GR, Dasanayake AP, Stiles HM, Caufield PW. Effect of conventional dental restorative treatment on bacteria in saliva. Community Dent Oral Epidemiol 1992;20(3): 138-43.

58. Anderson MH, Bales DJ, Omnell KA. Modern management of dental caries: the cutting edge is not the dental bur. JADA 1993;124(6):37-44.

59. Anusavice KJ. Treatment regimens in preventive and restorative dentistry. JADA 1995:126(6):727-43.

60. Anusavice KJ. Effcacy of nonsurgical management of the initial caries lesion. J

Dent Educ 1997;61(11):895-905.

61. Lagerlof F Oliveby A. Clinical implications: new strategies for caries treatment. In: Stookey GH, Beiswanger B, eds. Indiana Conference 1996: Early Detection of Dental Caries. Indianapolis: Indiana University School of Dentistry; 1996.

62. Cangelosi GA, Iversen JM, Zuo Y Oswald TK, Lamont RJ. Oligonucleotide probes for mutans streptococci. Mol Cell Probes 1994;8(1):73-80.

63. Shi W, Jewett A, Hume WR. Rapid and quantitative detection of *Streptococcus mutans* with species-specific monoclonal antibodies. Hybridoma 1998;17(4):365-71.

64. Caries diagnosis and risk assessment: a review of preventive strategies and management. JADA 1995,126(suppl):1-24S.

65. Ma JK, Hunjan M, Smith R, Lehner T. Specificity of monoclonal antibodies in local passive immunization against *Streptococcus mutans*. Clin Exp Immunol 1989;77(3):331-7.

66. Ma JK, Hikmat BY, Wycott K, et al. Characterization of a recombinant plant monoclonal secretory antibody and preventive immunotherapy in humans. Nat Med 1998;4(5):601-6.

67. Ma JK, Lehner T, Stabila P, Fux Cl Hiatt A. Assembly of monoclonal antibodies with IgG1 and IgA heavy chain domains in transgenic tobacco plants. Eur J Immunol 1994;24(1):131-8.

68. Stookey GK. Practical applications of early caries detection methods. In: Stookey GK, ed. Early detection of dental caries II: 1999. Indianapolis: Indiana University School of Dentistry (in press).

69. Kantorowitz Z, Featherstone JD, Fried D. Caries prevention by CO₂ laser treatment: dependency on the number of pulses used. JADA 1998;129:585-91.

70. Featherstone JD, Barrett-Vespone NA Fried D, Kantorowitz Z, Seka W. CO2 laser inhibitor of artificial caries-like lesion progression in dental enamel. J Dent Res 1998; 77(6):1397-403.

71. Featherstone JD. Innovative methods for early caries intervention. In: Stookey GK ed. Early detection of dental caries II: 1999 Indianapolis: Indiana University School of Dentistry (in press).

72. Fried D, Murray MW, Featherstone JD et al. Dental hard tissue modification and removal using sealed TEA lasers operating at $\lambda = 9.6$ and 10.6 µm. Proceedings of Lasers in Dentistry V: Jan. 24-25, 1999, San Jose, Calif. Bellingham, Wash.: SPIE Press; 1999:196-203.