Enamel remineralization: controlling the caries disease or treating early caries lesions?§

Abstract: The emphasis currently given to new technologies for enamel remineralization suggests that the changes in the understanding of the dental caries disease, which occurred in the last century, were either not yet adopted or were forgotten. Just like in the past, when the disease was “treated” by restoring cavities, there is presently a misunderstanding on the concept of incipient lesion remineralization. The aim of this paper was to review some concepts about caries, the natural phenomenon of enamel remineralization and the effect of fluoride (F) on it, and also to discuss the clinical relevance of remineralizing products recently launched in the marketplace aiming to “treat early caries lesions”.

Descriptors: Dental caries; Dental enamel; Tooth remineralization; Fluorides.

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Dental caries: the disease and its signals

For many years in the past, the dental profession had difficulty to distinguish dental caries as a disease from its signals: the caries lesions. Therefore when a cavity was filled, not only the professional believed that he was performing the best treatment, but the patient also believed that he/she was being treated for the disease.

A cavity, however, is only the terminal stage of a progressive mineral loss provoked in enamel by caries as a silent disease, dependent on the presence of biofilm and sugar exposure. Biofilm (plaque) and sugar are respectively necessary and determinant factors for the manifestation of the disease, i.e., bacteria need to accumulate on dental surfaces and be frequently exposed to dietary sugars for dental caries to develop. The disease progresses silently until the changes provoked in enamel by biofilm-sugar are clinically visible, i.e., a white spot lesion is present. If the disease is not controlled, dental destruction will not be prevented (Graph 1).

When white spots were recognized as caries lesions that could develop into cavities, a clinical discussion on strategies to repair or revert the changes provoked by the disease in enamel through a non-invasive treatment started in Dentistry. However, if the disease signs are the only factors taken into consideration, the strategies used to treat patients with white spot lesions or early signs of the disease may repeat the same mistakes made in the past when filling was considered the solution for caries.

Furthermore, there is clear evidence showing that the main effect found when a white spot lesion becomes clinically less evident (and does not progress to a cavity) is the result of mechanical abrasion of the enamel surface, and not only of repair (“remineralization”) of the mineral loss suffered by enamel.

Understanding that the “treatment of a white spot” was not only the result of true reprecipitation of minerals (remineralization) was very important because it emphasized the relevance of plaque control not only to arrest the lesion but also to restore the physiological equilibrium between tooth minerals and oral fluids (and allow natural remineralization from saliva) and, as a result, control the disease.

The consequence of disease control would be seen clinically by a change in the appearance of white spots, from chalky and rough (active) to bright and smooth (inactive). Eventually, the white spot could even disappear.

Although arrestment of caries lesions is not merely a “remineralization” of white spots, this should not be used to cast doubt on the natural phenomenon of enamel remineralization which occurs in the oral cavity, and its enhancement by fluoride.

Enamel remineralization

At physiological conditions, the oral fluids (saliva, biofilm fluid) have calcium (Ca) and phosphate (P) in supersaturated concentrations with respect to the mineral composition of enamel and, as a result, these ions are continually deposited on the enamel surface or are redeposited in enamel areas where they were lost. This can be considered a natural defense phenomenon promoted by saliva to preserve the mineral structure of enamel in the mouth. Therefore, remineralization would be best defined as the redeposition of minerals lost by enamel, and this term has been used as a synonymous of enamel repair or rehardening.

Mineral loss (demineralization) or gain (remineralization) by enamel is a dynamic physicochemical process occurring when oral bacteria form a biofilm.
on the enamel surface and this biofilm is exposed to fermentable dietary carbohydrates, sucrose being the most cariogenic of them.\(^7\) Thus, every time sugar penetrates into a cariogenic biofilm and is converted to acids by bacterial metabolism, the biofilm fluid becomes undersaturated with respect to the enamel mineral, and demineralization occurs.\(^8\) A critically low pH for tooth dissolution is maintained for a certain time, but it returns to physiological values when exposure to sugar ceases. Therefore, when the pH is raised and the supersaturating conditions are restored, a certain amount of the mineral lost can be recovered by enamel. This process has been named remineralization. Redeposition of the mineral lost by enamel can occur by Ca and Pi found in the biofilm fluid or by direct action of salivary Ca and Pi soon after the biofilm is removed by toothbrushing. The amount of Ca and Pi gained, however, is lower than that lost, and the net result is a small mineral loss.

If the factors responsible for the disease – biofilm accumulation and frequent sugar exposure – are not controlled, enamel mineral loss cannot be stopped (Graph 1).\(^2\) Repeated events of mineral dissolution will eventually surpass the capacity of oral fluids to repair mineral loss, and the disease will show its first clinical signs: white spot lesions. Differences in the progression rate of caries lesions in different individuals or populations can also be the result of other factors which modulate the caries process, since caries is a “multifactorial” or “complex disease”, and the risk of developing new lesions is never zero.\(^\text{3}\) In some individuals, lesions will progress slowly, and the disease might not be clinically detected throughout their lifetime. In others, they will progress rapidly, and the manifestation of the disease will become clinically evident (white spots) before cavitation (Graph 2). Additionally, the progression from non-cavitated to cavitated lesions could be arrested, and the strategies used should consider if the lesions are “active or not”.\(^9\) This classification is aimed to correctly diagnose caries as a disease, as the basis for treatment decision.\(^9\)

**Fluoride effect on the dynamics of the caries process**

Nowadays there is a consensus that the predominant effect of fluoride (F) is not systemic, by pre-eruptively changing the enamel structure, but mainly locally, by interfering with the caries process.\(^\text{10}\) Hence, F must be present in the right place (biofilm fluid or saliva), and at the right time (when biofilm is exposed to sugar or right after biofilm removal) to interfere with de- and remineralization events. For this effect, even below ppm values of F available are effective.

Thus, as described previously, enamel is dissolved by the lowering of pH in dental plaque due to acid production every time sugar is ingested (Figure 1). However, if F is present in the biofilm fluid, and the pH is not lower than 4.5, hydroxyapatite (HA) is dissolved at the same time that fluorapatite (FA) is formed.\(^6\) The net result is a decrease in enamel dissolution, since a certain amount of Ca and Pi, which was lost as HA, is recovered by enamel as FA. This mineral gain as FA during the pH drop (Figure 1) has not been considered as remineralization but rather as a decrease in demineralization because the mineral redeposited is different from that lost. Furthermore, FA is deposited on the surface layer of enamel while HA is dissolved from the subsurface.

This indirect effect of F reducing enamel demineralization when the pH drops is complemented by its natural effect on remineralization when the pH rises (Figure 2) enhancing the redeposition of Ca and Pi present in the biofilm fluid on demineralized enamel. If the demineralized enamel is cleaned by

![Graph 2 - Enamel mineral loss progression during the lifetime of different individuals or populations (A = slow; B = moderate; C = fast).\(^4\)
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Brushing, saliva is able to remineralize it, but in the presence of F this effect is enhanced. As a result, small amounts of Ca and P lost by enamel during the pH drop can be more efficiently recovered if F is still present in the oral environment (biofilm fluid or saliva) after the cariogenic challenge. This effect should be considered natural, not induced, because it occurs irrespective of patient compliance or dentist intervention if, for example, a F-dentifrice is being used and F is made available to the oral cavity.

By acting on the dynamics of the caries process, F is very effective in slowing down (retarding) the progress of caries lesions. However, since it does not have a direct effect on the etiological factors responsible for the disease (biofilm and sugar), it will not avoid it, and invariably the disease will leave scars on teeth, clinically visible or not (Graph 3).

Nonetheless, the relative contribution of F – reducing demineralization or enhancing remineralization – to the final effect on caries is not known because it is impossible to isolate these effects considering the dynamics of the process. Indeed, some
authors believe that the most important F effect is a reduction of enamel demineralization while others consider that F speeds up the remineralization process, and this is the main mechanism of caries control.

The effect of F on the reduction of enamel demineralization or enhancement of remineralization can be clearly shown in vitro using pH-cycling models which simulate clinical situations of predominance of demineralization (De > Re) or of remineralization (Re > De). Graph 4 shows the effect of fluoride on sound enamel when De > Re (patient under caries activity). The efficacy of F in decreasing enamel surface hardness loss and in reducing lesion depth are shown, as is the gain of F by enamel, which is a consequence of mineral change if F is present during the caries process.

Fluoride effect on caries lesions

Since F enhances enamel remineralization, its clinical use to repair early caries lesions was advocated (“fluoride therapy”). However, the effect of F in the dynamics of the caries process and its success in controlling caries should not be confused with its arrestment or reversal effect on caries lesions. Furthermore, it should be emphasized that shallow demineralized enamel areas remineralize faster than deep ones.

Arrrestment of white spot caries lesions and the changes observed in a demineralized enamel surface and subsurface were well documented 25 years ago. Change in enamel lesions from a whitish to a shiny appearance was explained in terms of wear and polishing of the dull, partly dissolved surface of the active lesions, rather than redeposition of mineral lost. However, the porosity of the deeper parts of the lesions was reduced, suggesting a partial remineralization of the lesion body.

Therefore the presence of a surface layer does not prevent subtle alterations at the crystals level inside the caries lesions after removing the biofilm accumulated on the enamel surface. Re-establishing the physiological conditions between enamel and oral fluids will lead to the redeposition of minerals in the demineralized areas. However, while subsurface lesions remineralize in vitro within weeks, years are required for a complete remineralization in vivo.

The effect of F on enamel remineralization is easily shown in vitro. Graph 5 shows the effect of F on enamel with a “caries lesion” simulating a clinical situation under low caries challenge (Re > De). The enamel surface was rehardened, the lesion depth was reduced and there was an increase in F concentration in the remineralized enamel. However, the lesion was not totally repaired.

Therefore, if the biofilm accumulated on the enamel surface presenting non-cavitated active caries lesions is controlled by brushing with a F-containing dentifrice, it is not surprising that the lesions can not only be arrested, but also partially repaired (Graph 6). Even cavitated lesions can be arrested.

In summary, the effect of F in enhancing enamel remineralization is well known from several in vitro and in situ studies, but considering the best knowledge about dental caries, any “remineralizing therapy” should follow two fundamental principles:

1. Dental biofilm, the necessary factor responsible for caries lesions, should be controlled by toothbrushing.
2. Fluoride should be used either to arrest existing lesions or to reduce the progression of new ones.

It should be emphasized that the source of minerals to rebuild demineralized enamel are salivary Ca and P, since saliva is supersaturated with respect to tooth mineral, favoring mineral precipitation.
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Therefore, the normal endogenous concentrations of Ca and P found in saliva are high enough to induce remineralization, but the latter can be significantly enhanced by exogenous F, supplied by different ways of oral delivery.23

News technologies for enamel remineralization

Enamel remineralization has been studied for about 100 years, and it has been suggested that “the non-invasive treatment of early caries lesions by remineralization has the potential to be the major advance in the clinical management of the disease”.24

This subject was recently reviewed,24 as well as the effect of the three remineralizing products launched in the marketplace after 2000, all based on calcium phosphate remineralization systems. One technology involves casein phosphopeptide stabilized amorphous calcium phosphate (CPP-ACP; Recaldent™), the second is an unstabilized amorphous calcium phosphate (ACP, Enamelon™) and the third is a bioactive glass containing calcium sodium phosphosilicate (NovaMin™). Since all systems rely on calcium and phosphate compounds, their effect is mainly based on an enhancement of the natural capacity of saliva to remineralize mineral loss.

The clinically-based evidence to support these products is currently not abundant. A literature search of the PubMed database conducted in April, 2009 using the key words Recaldent or Enamelon or Novamin alone or combined with remineralization AND caries AND review resulted in the number of papers listed in Table 1.

Recaldent is the most studied system. The available reviews on it state either that it could significantly slow the progression of coronal caries and promote the regression of lesions,24 or that there is insufficient clinical trial evidence, in quantity and also quality, to make any recommendation on...
the clinical use of this product. As for Enamelon, except for a descriptive publication made by the manufacturers of this commercial product, the only other review, published by Reynolds (2008), concluded that there is evidence that Enamelon has an anticariogenic effect on root caries. Except for the paper by Reynolds (2008), no other paper was found on Novamin AND remineralization in PubMed, but a great number of abstracts have been presented in the last IADR meetings (54 in 2008 and 24 in 2009).

As Reynolds (2008) concluded, “calcium phosphate-based remineralization technologies show promise as adjunctive treatments to fluoride therapy in the non-invasive management of early caries lesions”, since they do not have general recommendation, but some people could benefit from them. Otherwise, they would not be a solution to the problem of controlling caries disease.

**Conclusions**

1. Incipient caries lesions regress or clinically dis-

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**Table 1 - Number of publications in PubMed according to product name combined with selected key words.**

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<tr>
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<th>Enamelon™</th>
<th>Novamin™</th>
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**References**

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