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Effects of various fluoride solutions on enamel erosion in vitro

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Key words: erosion; fluoride; enamel

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Declaration of interests

The authors declare no conflicts of interest.

Effects of various fluoride solutions on enamel erosion in vitro

Abstract

The objective of this in vitro study was to investigate the effect of different fluoride solutions on enamel erosion. Human enamel specimens were pretreated with 1 of 10 different fluoride solutions (n=20): TiF_4 , NaF, AmF, ZnF_2 , or SnF_2 , each at native pH (pH range: 1.2-7.8) or buffered pH (pH=4). The control group samples received no fluoride pretreatment. All the samples were then eroded by citric acid (pH 2.6) for 6*1 min daily over 5 days. Between erosive cycles, the samples were stored in artificial saliva. Erosion effects were investigated by surface profilometry (n=10), scanning electron microscopy (n=4), and energy-dispersive X-ray spectroscopy (n=6) after fluoride pretreatment and after erosion. To test the effects of pH only, additional experiments were carried out with fluoride-free solutions at similar pH of fluoride solutions. In general, AmF solution was more effective in protecting enamel erosion compared to all other fluoride agents. However, also the application of native TiF_4 , native and buffered SnF_2 , native and buffered AmF solutions resulted in significantly less enamel loss compared to the control group. A Ti-rich coating was formed after application of native TiF_4 , but partially dissolved due to erosive attack. Sample pretreated with SnF_2 showed a significant increase in tin. Surface fluoride concentration was significantly increased by native TiF_4 , native and buffered AmF, buffered ZnF_2 , and buffered NaF application. Under the current experimental setting, the fluoride agents at lower pH had better protective potential. High-concentrated TiF_4 , AmF, and SnF_2 solution was effective in inhibiting erosion in enamel.

Introduction

Dental erosion, which is defined as the loss of tooth substance by chemical processes not involving bacteria, is becoming an increasingly important factor when considering long-term dental health [Lussi et al., 2004]. Considering the presence of dental erosion is growing steadily in the last few decades [Jaeggi and Lussi, 2006], interceptive methods to prevent dental erosion are needed.

Several methods were proposed to prevent the progression of dental erosion, such as fluoride application, modification of acidic beverages, and laser application [Magalhaes et al., 2009]. Although the mechanism of fluoride application in prevention of dental erosion is still controversially discussed, many studies have shown the potential protective effects of fluoride application on dental erosion [Ganss et al., 2004; Lagerweij et al., 2006]. Interestingly, most of the studies were done with the fluoride agents that have shown their anti-caries effects before, such as NaF, AmF, SnF₂ and ZnF₂. It was reported that SnF₂ application led to a much greater reduction of mineral loss compared to NaF and AmF [Ganss et al., 2008]. In an in vitro study investigating TiF₄, SnF₂, and NaF, all the fluoride agents showed protective effects on erosion while TiF₄ protected the enamel surface almost completely [Hove et al., 2006]. With regard to TiF₄, several studies have shown that it had a great inhibitory effect against dental erosion [Hove et al., 2006; Schlueter et al., 2007; Wiegand et al., 2008], while other studies found that TiF₄ reduced erosion only to a small extent compared to the control group [Magalhaes et al., 2008; Vieira et al., 2005]. Furthermore, it has been pointed out that the efficacy of fluoride agents at native and buffered pH in protecting enamel erosion is distinctly different [Arnold et al., 2007; Wiegand et al., 2009b].

In short, in diverse experimental designs the fluoride agents showed a wide range in their efficacy against erosion. Thus, a systematic investigation of pH- and concentration-controlled fluoride preparations on erosion is necessary in order to get better understanding of the effects of different fluoride compounds on erosion. However, relatively little information is available to

date.

Therefore, the objective of this study was to evaluate the effects of different fluoride solutions on human enamel using surface profilometry, scanning electron microscopy (SEM) and energy-dispersive X-ray spectroscopy (EDS). Two hypotheses were proposed: 1) different fluoride agents have a different protective effect on human enamel erosion; 2) fluoride agents at their native and buffered pH differ in their protective ability against human enamel erosion.

Materials and Methods

Study design

Enamel samples were pretreated with distilled water (negative control) or 1 of 10 different fluoride solutions (TiF_4 , SnF_2 , AmF , ZnF_2 , NaF , each at native pH or pH 4, each $n=20$) and subjected to a five-day cyclic de- and remineralization procedure. Demineralization was performed with citric acid at pH 2.6 for 6*1 min daily and the specimens were immersed in artificial saliva during the remaining time. The effects of fluoride treatment and erosion were analyzed using surface profilometry ($n=10$), SEM ($n=4$), and EDS ($n=6$). Additionally, to test the effects of pH alone on erosion, enamel specimens were pretreated with 4 fluoride-free buffer solutions at pH 1.2, 2.7, 4, 7.8, respectively, analogous to the pH of the fluoride solutions used in this study. Subsequently, the samples were subjected to the same cyclic de- and remineralization procedure and analyses as described above.

Specimen preparation

300 enamel samples (3 mm in diameter) were obtained from the labial and palatal surfaces of 150 previously extracted, caries-free human molar teeth. The samples were embedded in ring-shaped ceramic moulds (3 mm diameter, 3 mm thickness) with acrylic resin (Paladur, Heraeus Kulzer, Wehrheim, Germany). The ceramic moulds were cut from a ceramic tube (Degussit,

Friatec/Degussa, Düsseldorf, Germany) using a water-cooled low speed saw (Isomet, Buehler, Lake bluff, IL, USA). The embedded specimens were ground flat and polished with water-cooled carborundum discs (1200, 2400 and 4000 grit (FEPA-P), Water proof silicon carbide paper, Stuers, Erkrat, Germany). This procedure resulted in the removal of about 200 μm depth of enamel, which was controlled with a digital micrometer (Horex, Nuremberg, Germany).

The polished specimens were cleaned in distilled water in an ultrasonic cleaner (M.Scherrer, Wil, Switzerland) for 1 min to remove any debris. Before use, all the specimens were stored in 100% humidity.

Fluoride solution preparation and pretreatment

Native TiF_4 solution (0.48 M F, pH 1.2) was prepared by mixing 1.5 g titanium tetrafluoride powder (Stream Chemicals, Newburyport, MA, USA) with 100 ml distilled water. The TiF_4 solution was adjusted to pH 4 by adding 2.45 g sodium citrate/100 ml. Native SnF_2 solution (0.48 M F, pH 2.7) was obtained by adding 3.75 g stannous fluoride powder (Riedel-de Haen, Seelze, Germany) in 100 ml distilled water. The SnF_2 solution was adjusted to pH 4 by adding 2.5 g sodium citrate/100 ml. Native AmF solution (0.48 M F, pH 4.6) was prepared with mixing 65.1 g Olaflur solution (GABA International, Münchenstein, Switzerland) and 34.9 g distilled water. The AmF solution was adjusted to pH 4 by adding 4.25 ml 5 M H_3PO_4 /100 ml. Native ZnF_2 solution (0.20 M F, pH 5.1) was obtained by mixing 1.6 g $\text{ZnF}_2 \cdot 4\text{H}_2\text{O}$ powder (Aldrich Chem, Milwaukee, WI, USA) in 100 ml distilled water. The ZnF_2 solution was adjusted to pH 4 by adding 0.3 ml 5 M H_3PO_4 /100 ml. Native NaF solution (0.48 M F, pH 7.8) was prepared by mixing 2.02 g NaF powder (Merck, Darmstadt, Germany) with 100 ml distilled water. The NaF solution was adjusted to pH 4 by adding 7.5 ml 5 M H_3PO_4 /100 ml. All the solutions were prepared freshly prior to applications to the specimens.

For fluoride pretreatment, the specimens were separately fixed in plastic chambers. The chambers were then filled with 5 ml respective fluoride solution

and left undisturbed for 3 min at room temperature. After treatment, all the samples were rinsed with distilled water for 30 s. Specimens of the control group were treated with distilled water only.

For the additional experiment regarding the effect of pH of pretreatment solution on enamel erosion alone, specimens were pretreated with fluoride-free solutions at different pH (pH 1.2, 2.7, 4 and 7.8, n=20) for 3 min. Then the samples were rinsed in the same manner as mentioned above.

Cyclic erosive treatment

The cycling de- and remineralization regimen was performed 6 times daily. The samples were first eroded by immersion in 5 ml citric acid (pH 2.6) for 1 min. After erosion, the samples were rinsed with distilled water for 30 s and stored for 1 h in 5 ml artificial saliva until the next erosion challenge. The artificial saliva was mixed according to the formulation given by Kilmek et al. [1982]. After 6 daily cycles, treated specimens were stored in artificial saliva overnight. Specimens of the control group were maintained in artificial saliva for the entire experimental period. The artificial saliva was renewed every day.

Profilometric analysis

Enamel loss was measured by a stylus profilometer (Perthometer S2/GD 25, Mahr, Göttingen, Germany) after fluoride pretreatment and after erosion. The procedure has been described in detail previously [Yu et al., 2009].

Three profiles were performed on each specimen via scanning from the reference (ceramic mould) surface to the treated surface. An average of these three readings (μm) was obtained and used for data analysis.

SEM and EDS

After fluoride pretreatment, 4 samples from each group were randomly selected for SEM observation. The specimens were mounted on aluminum stubs and sputter coated with platinum, and then examined using a Supra 50

VP Scanning Electron Microscope (Carl Zeiss NTS, Oberkochen, Germany) with an acceleration voltage of 2 kV. Likewise, after erosion, 4 samples from each group were selected and observed under SEM.

Furthermore, quantitative changes of the surface composition of the fluoride-treated and eroded samples were evaluated using EDS. The EDS analysis was done with the same SEM equipped with an EDAX PV7715/89 ME energy-dispersive X-ray spectrometer. Six specimens from each group were sputter coated with carbon and the respective EDS spectra were obtained in a 200*200 μm area with an acceleration voltage of 15 kV. The spectra were later analyzed using the EDAX Genesis Spectrum software package (EDAX, Mahwah, NJ, USA).

Statistical analysis

The data were analyzed using the SPSS statistical software package (SPSS 13.0 for Windows, SPSS, Chicago, IL, USA). The Kolmogorov-Smirnov test was used for checking the normal distribution of data. The results of profilometric analysis and EDS analysis were analyzed by one-way analysis of variance (ANOVA) followed by Tukey multiple comparison tests. Since data for F content (after pretreatment and after erosion) and P content after erosion were not normally distributed, Mann-Whitney test was performed for comparison of respective contents in different groups. All statistical analyses were carried out at a significance level of 0.05.

Results

Profilometric analysis

Table 1 shows the enamel loss in different groups after fluoride application and after erosive attack. After fluoride treatment, the native TiF_4 - treated samples showed the most pronounced enamel loss (0.25 μm), although no significant differences were found among all groups including the control group. In most cases, all the fluoride applications reduced erosive substance loss.

The application of AmF at pH 4 resulted in almost no erosive loss (97% reduction compared to controls), while the other fluoride group led to reduction of enamel loss from 15% to 82% compared to the control. However, only the application of native TiF_4 , native and buffered SnF_2 , and native and buffered AmF was able to significantly decrease erosive enamel loss compared to the control group. With regard to the total loss due to fluoride treatment and erosive attack, similar results were found. Total erosive substance loss was significantly smaller for TiF_4 application at native pH than at the buffered pH. All other fluoride solutions at lower pH reduced the erosive loss, but not significantly, compared with the respective solutions at higher pH.

For the samples pretreated with fluoride-free solutions, the application of the solution at pH 1.2 and pH 2.6 led to significantly greater erosive enamel loss compared with the ones at pH 4 and pH 7.8, as well as the control specimens treated with distilled water. However, no significant differences were found in substance loss due to erosion among these four groups and the control group (table 2).

SEM and EDS

Representative SEM images and EDS results are shown in Figs 1-6. The application of native TiF_4 , native and buffered AmF, and buffered NaF resulted in distinct alteration in the surface morphology of human enamel. After application of native TiF_4 , a surface layer with some globular materials was formed on the enamel surface. This layer was found to have some microcracking and spacing (fig. 1a). The underlying enamel was extremely porous and granular in appearance (fig. 1b). After erosion, the coating was partially disappeared and the rest of the surface demonstrated severe demineralization (fig. 1e). The morphologic appearance of the enamel surfaces revealed the presence of precipitates in both native and buffered AmF treated samples. The precipitates varied in appearance. The globules on the native AmF treated specimens were larger than the ones on the buffered

AmF treated specimens. However, similar surface morphology with distinctly larger globules was found in both AmF-treated samples after 30 min erosion (fig. 2). The globular precipitation was also found on the buffered NaF-treated samples, but was completely dissolved after erosion (fig. 3). SnF₂ treatment did not produce globular deposits on the enamel surface. The native SnF₂-treated surfaces revealed small pits, while larger pores were found after erosion on the surface (fig. 4a, b). Although no visible difference was found after treatment with buffered SnF₂, moderate demineralized surface with some pores were found after erosion (fig. 4c, d). Apart from the phenomenon mentioned above, the sample surfaces appeared smooth and structureless after distilled water or fluoride application, while the severe etched surfaces were found after erosive challenges (figs. 5 and 6).

Table 3 presents the concentration (wt%) of the respective elements. All the eroded samples showed a decrease in surface Ca and P concentration, although some of the differences were not statistically significant. Treatment with native TiF₄ led to a significant increase in Ti compared to treatment with buffered TiF₄. Likewise, a larger amount of Sn was found after applying native SnF₂. The concentration of these metal ions (Ti and Sn) was reduced after erosion. The application of native TiF₄, native and buffered AmF, buffered ZnF₂, and buffered NaF resulted in a significant increase in F, which was significantly higher for native AmF than for the other solutions. After erosion, a decrease was found in F concentration accompanying with an increase in C concentration.

All the samples pretreated with fluoride-free solutions at different pH showed a similar surface tomography and composition as distilled water after 30 min erosive attack (Table 4).

Based on the above results, the hypotheses that different fluoride agents have a different effect on human enamel erosion and that the pH of fluoride agents affects enamel erosion under the testing conditions, were therefore accepted.

Discussion

In literature, many techniques have been used to investigate the effects of erosive attacks on dental hard tissues. Micro-indentation, surface profilometry, microradiography, chemical analysis and SEM were considered the most established laboratory assessment in evaluating enamel erosion [Attin, 2006; Barbour and Rees, 2004]. In the present study, surface profilometry and SEM were selected to analyze the mechanical effect of the fluoride solutions on enamel erosion in order to measure substance loss accurately and provide visual information on surface precipitates and change of the surface morphology. On the other hand, EDS provides a specific method to determine the concentration of chemical elements on substratum surfaces, being largely used in engineering and chemistry, but not in many studies in dentistry [Paradella et al., 2008]. Since the detection limits of EDS is about 0.1 wt% [Kuisma-Kursula, 2000], EDS seems to be a sufficient tool to evaluate the chemical changes of fluoridation and erosion on the enamel surface.

The application of native TiF_4 resulted in minor enamel loss ($0.25 \mu\text{m}$), which was much lower than the application of fluoride-free buffer solution at the same pH ($3.17 \mu\text{m}$). Moreover, this application led to a build-up of a layer on the enamel surface containing Ti which was in accordance with previous studies [Magalhaes et al., 2008; Wiegand et al., 2009b]. The mechanism of the layer formation following application of TiF_4 is still not clear. It is likely that within a dissolution-precipitation process a new compound (hydrated hydrogen titanium phosphate) is formed [Ribeiro et al., 2006]. During the application calcium in the hydroxyapatite (HAP) lattice may be replaced by titanium [Leadley et al., 1997]. Further, the SEM images (fig. 1b) showed severely demineralized enamel under the surface coating, suggesting that enamel might suffer from a dissolution-precipitation process after the application of native TiF_4 solution. It was reported that the application of TiF_4 at pH 1.2 led to a dense surface layer [Wiegand et al., 2009b]. However, in the present study, the layer was found to have microcracks most likely due to specimen

preparation prior to EDS. The development of cracks during desiccation may be an indication of a gel-like composition of this layer. Apart from Ti, the surface F concentration increased after native TiF_4 treatment. Therefore, we may assume that some fluoride-containing compounds are also formed apart from titanium phosphate. The composition of the globular deposits, having higher F and Ti concentration and lower Ca concentration compared with the surface layer (fig. 1c, d), can add some support to this hypothesis. Contrasting to the results of the previous study [Wiegand et al., 2009b], the formed layer was partially dissolved after erosion. This might be due to the more intensive erosive attacks adopted in the present study (30 min vs. 10 min acid challenge). This layer did provide a protective barrier to the enamel surface capable of providing protection against erosive attack for a certain period of time as shown by profilometry. Furthermore, it is noteworthy that the application of TiF_4 at pH 4 failed to form this Ti rich coating on the enamel surface, thus leading to a similar high enamel loss as the control group after erosion.

Both native SnF_2 and buffered SnF_2 were able to reduce the erosive enamel wear. The application of SnF_2 resulted in an increase of Sn, indicating the possible reaction between Sn and the HAP lattice [Hove et al., 2008]. Since there was only a minor increase in the F concentration on the SnF_2 -treated surface, the protective effect of the SnF_2 solution might be largely due to the Sn-containing reaction products (possibly Sn_2OHPO_4 , $\text{Sn}_3\text{F}_3\text{PO}_4$, and $\text{Ca}(\text{SnF}_3)_2$) [Babcock et al., 1978]. However, in contrast with previous studies [Ganss et al., 2008; Wei, 1974], the so called Sn-rich coating was not found after application of SnF_2 . Possible explanation could be the different tin concentration and application time of SnF_2 solution used in this study.

In accordance with a previous study [Rosin-Grget et al., 2000], globular CaF_2 -like globular precipitates were found on the enamel surface after application of AmF at pH 4.7 and 4. The appearance of the precipitates was found to be pH-dependent. The treatment of AmF at 4.7 resulted in larger

globular precipitates and higher F concentration on the surface than at pH 4. Less amount of phosphate was found in the larger precipitates after topical application of AmF at pH 4.7 than at pH 4 (10.43 wt% vs. 14.54 wt%, fig2. b, f). Similarly, for both native AmF and buffered AmF-treated samples, the P concentration showed a decrease after erosion. The differences in the appearance of the globular precipitates seem to be related to phosphate contamination [Rolla and Saxegaard, 1990]. After erosion, a distinct surface alteration with larger globular precipitates of CaF₂-like deposits was found in both AmF-treated surfaces. It is well known that the solubility of CaF₂-like deposit decreases with an increase in size of the globules [Nelson et al., 1983]. Thus, it can be speculated that the smaller CaF₂-like materials dissolved during demineralization process and the mineral and fluoride tended to fuse into bigger CaF₂-like materials. Interestingly, some polishing traces on the native and buffered AmF-treated samples were still visible after erosion, correlating well with the profilometric data and corroborating the protective effect of the AmF. However, it is noteworthy that the thickness of the analyzed layer is a few microns under the usual conditions of EDS analysis [Verita M et al., 1994]. As seen from the Figs 1-3, the globular precipitates varied in size on the enamel surfaces. The EDS spectra might contain the signal not only from the precipitates, but also from the underlying enamel.

After erosion, all samples showed a reduced surface Ca and P concentration, indicating that demineralization happened on the enamel surface due to erosion. In addition, an interesting change in surface fluoride concentration should be noted. After application of native TiF₄, native and buffered AmF, buffered ZnF₂, and buffered NaF, the F concentration of the enamel surface increased. This phenomenon was probably related to the formation of CaF₂-like materials on the surface, although some of the changes were not detectable under SEM. After erosion, the surface F concentration was reduced, possibly indicating the dissolution of the CaF₂-like precipitation. Interestingly, for the samples treated with buffered AmF, the surface fluoride

concentration remained stable after erosion, although a distinct difference was found under SEM observation. Nevertheless further studies are needed to clarify this issue.

In accordance with a previous study [Schlueter et al., 2009], NaF and ZnF₂ were not able to reduce erosive enamel loss, although the application of buffered NaF and ZnF₂ led to the distinct formation of CaF₂-like materials on the surfaces. Possibly, the CaF₂-like precipitates formed by the application of buffered NaF and ZnF₂ dissolved quickly in the beginning of the acid attack. This hypothesis could be confirmed by the fact that the substance loss of buffered NaF and ZnF₂-treated samples were only a bit lower than in the control samples.

A further point to consider is that the pH of fluoride solution plays an important role in the efficacy of fluoride agents against erosion. Based on the findings of this study, the fluoride solutions at lower pH increased the surface F concentration (except for AmF) and provided better protection against erosive enamel loss. This phenomenon could be partially explained by that the increased formation of CaF₂-like deposit [ten Cate, 1997] and better incorporation of metal ions (Ti and Sn) into enamel under lower pH condition.

Due to the low solubility of zinc fluoride (1.6 g/100 g water) [International Union of Pure and Applied Chemistry, 2009], the fluoride concentration of ZnF₂ solution was 0.20 M. Since there is some evidence that the protective effects of fluoride agents were not strongly fluoride concentration dependent [Wegehaupt et al., 2009; Wiegand et al., 2009a], the relatively low fluoride concentration of ZnF₂ may not be a critical problem in this study.

In this study, various high-concentrated fluoride agents were tested and the distinct protective effect of TiF₄, AmF and SnF₂ application on the development of erosive lesions were shown. However, further *in situ* or clinical studies are needed to see whether it is valid to recommend high-concentrated fluoride application for prevention of dental erosion.

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Legends

Table 1. Means and standard deviations (SD) of substance loss (μm) for the fluoride-treated groups and control group.

Table 2. Means and standard deviations (SD) of substance loss (μm) for the buffer-treated groups and control group.

Table 3. Means and standard deviations (SD) of concentration (wt%) of the respective elements in fluoride-treated groups and control group by EDS analysis ($200 \times 200 \mu\text{m}$).

Table 4. Means and standard deviations (SD) of concentration (wt%) of the respective elements in buffer-treated groups and control group by EDS analysis ($200 \times 200 \mu\text{m}$).

Fig 1. Representative images of enamel treated with native TiF_4 (pH 1.2): a. SEM images of enamel surface after native TiF_4 application (10,000x); b. SEM images of underlying enamel at 80,000x; c. EDS spectra of the surface layer; d. EDS spectra of the globular materials; e. SEM images of native TiF_4 -treated enamel surface after erosion (10,000x); f. EDS spectra of the surface layer after erosion; g. EDS spectra of the surface globules after erosion; h. SEM images of enamel surface after buffered TiF_4 (pH 4) application (10,000x); i. SEM images of buffered TiF_4 -treated enamel surface after erosion (10,000x). Arrows are marking the corresponding surface area.

Fig. 2. Representative images of enamel treated with AmF: a. SEM image of enamel surface after native AmF (pH 4.7) application (40,000x); b. EDS spectra of surface globules after native AmF application; c. SEM images of native AmF-treated enamel surface after erosion (40,000x); d. EDS spectra of surface globules after erosion; e. SEM image of enamel surface after buffered AmF (pH 4) application (40,000x); f. EDS spectra of surface globules after buffered AmF application; g. SEM images of buffered TiF_4 -treated enamel surface after erosion (40,000x); h. EDS spectra of surface globules after erosion. Arrows are marking the corresponding surface area.

Fig. 3. Representative images of enamel treated with NaF: a. SEM image of enamel surface after native NaF (pH 7.8) application (40,000x); b. SEM image of native NaF-treated enamel surface after erosion (40,000x); c. SEM image of enamel surface after buffered NaF (pH 4) application (40,000x); d. EDS spectra of surface globules after buffered NaF application (arrows are marking the corresponding surface area); e. SEM image of buffered NaF-treated enamel surface after erosion (40,000x).

Fig. 4. Representative images of enamel treated with SnF₂: a. SEM image of enamel surface after native SnF₂ (pH 2.7) application (40,000x); b. SEM image of native SnF₂-treated enamel surface after erosion (40,000x); c. SEM image of enamel surface after buffered SnF₂ (pH 4) application (40,000x); d. SEM image of buffered SnF₂-treated enamel surface after erosion (40,000x).

Fig. 5. Representative images of enamel treated with ZnF₂: a. SEM image of enamel surface after native ZnF₂ (pH 5.1) application (40,000x); b. SEM image of native ZnF₂-treated enamel surface after erosion (40,000x); c. SEM image of enamel surface after buffered ZnF₂ (pH 4) application (40,000x); d. SEM image of buffered ZnF₂-treated enamel surface after erosion (40,000x).

Fig. 6. Representative images of enamel treated with distilled water (control): a. SEM image of enamel surface after distilled water application (40,000x); b. SEM image of control enamel surface after erosion (40,000x).