Evaluation of resin infiltration on demineralized root surface: An in vitro study

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The aim of this study was to evaluate the effect of resin infiltration on root caries induced by Streptococcus mutans biofilms. Human premolar specimens were divided to 5 groups: negative control (NC), Clearfil SE Bond (SEB), Icon-etch120s+Icon-infiltrant (HA120), Icon-etch10s+Icon-infiltrant (HA10) and K-etchant10s+Icon-infiltrant (PA10). The resin penetration was observed by fluorescent microscope. Biofilm-induced demineralization was conducted again and observed by swept-source optical coherence tomography and confocal laser scanning microscope. The maximum resin penetration depth (PDmax), lesion depth increase (∆LD), frequency of cervical enamel loss and dentinoenamel junction separation length were measured and statistically analyzed. HA120 showed 138.00±49.25 µm PDmax that was significantly larger than PA10 and SEB (p<0.05). SEB created 136.58±64.73 µm coating layers. HA120 and SEB showed significantly lower ∆LD than NC (p<0.05). Resin infiltration with 120s-HCl pretreatment has got a good penetration ability and preventive effect on root caries, however, an additional risk factor of cervical enamel loss was identified.

Keywords: Resin infiltration, Root caries, Hydrochloric acid, Phosphoric acid, Biofilm

INTRODUCTION

Root caries usually occurs where root dentine is exposed to oral environment as a result of gingival recession, which may be caused by mechanical toothbrush trauma or chronic periodontitis3). Root caries increases in prevalence with age and will precipitate the breakdown of remaining natural and restored teeth25). Because cementum has half as much mineral as enamel by weight, demineralization on the root surface is approximately twice as rapid as that on enamel and can occur at a higher pH38). Additionally, root dentine surfaces may even present a high risk to caries development49 due to a low salivary flow, which is frequently found in elderly as a side-effect of daily use of medicines69). Various treatment modalities have been proposed for the management of root caries, including the improvement of oral conditions for oral hygiene, topical application of remineralization agents and antimicrobial regimen6), and coating or sealing by resin-based materials7). However, efficient and simple methods, which can protect the exposed root surface from carious attack for a long term and for elderly patients requiring home care, are not yet currently available9).

In recent years, a micro-invasive treatment with a low-viscosity resin-infiltrant has become one of the options for non-cavitated enamel caries. Resin infiltration on enamel caries has been clinically proven to effectively arrest and stabilize lesions5,11). It has also been proven to significantly increase micro-hardness and reduce mineral loss after a demineralization challenge compared with untreated lesions12). The mechanism is that resin-infiltrant exploits capillary forces to transport resins with high penetration coefficients into enamel porosities, and after polymerization the infiltrant occludes diffusion pathways for cariogenic acids and dissolved minerals13). Resin-infiltrant has a significantly deeper penetration in the lesion body than conventional dental adhesives14,15) and the diffusion barrier of caries infiltration is not formed at the lesion surface as caries sealing and coating but created inside the lesion body16). Besides that, caries infiltrants are optimized for rapid capillary penetration and exhibit a very low-viscosity, low contact angle to enamel and high surface tension19). While resin infiltration was efficacious in preventing further demineralization of artificial enamel caries lesions under cariogenic conditions in situ17), its potential effect on root caries has not been studied so far.

A technique called optical coherence tomography (OCT) has been developed for noninvasive cross-sectional imaging of internal biological structures18). In dentistry, OCT was introduced for the evaluation of tooth demineralization and remineralization19,20). A recent study also showed that OCT could quantitatively evaluate the penetration of resin-infiltrant into early dental caries21). The aim of this study was to evaluate the potential effect of resin infiltration with different pretreatment conditions on root caries induced by cariogenic bacterial biofilms in vitro. The null hypothesis was that resin infiltration could not reduce further demineralization of root caries lesions under cariogenic conditions.
MATERIALS AND METHODS

Specimen preparation
Thirty extracted intact human premolars with no visible evidence of caries or cracks were collected after obtaining the informed consent according to a protocol approved by the Institutional Review Board of Tokyo Medical and Dental University (approval no. 725). The teeth were stored in 4°C in water containing 0.1% wt of thymol in order to inhibit bacterial growth for a maximum of 6 months until use. The upper two-third crown and the lower two-third root of the teeth were cut by a low-speed diamond saw (Isomet, Buehler, Lake Bluff, IL, USA), keeping 2 mm crown and 3 mm root away from the enamel-cementum junction (Fig. 1). Then the buccal and lingual cervical parts were vertically cut into half. Sixty square-shaped blocks (5×4×2 mm) were made by a diamond bur (FG 102R, Shofu, Kyoto, Japan) attached to an air turbine headpiece under copious cooling water. Two semi-round holes (depth: 0.5 mm) were marked by a diamond bur (D4010f, GC, Tokyo, Japan) on the upper and lower central top surface of each specimen. The specimens were observed by swept-source optical coherence tomography (SS-OCT, IVS-2000, Santec, Komaki, Japan). Then five cutting surfaces were covered with a thin layer of acid-resistant varnish (Shiseido, Tokyo, Japan) and one edge of the top surface was covered by 1-mm-wide paraffin wax (GC) to keep the baseline from being demineralized.

In vitro root caries formation (First time demineralization)
In vitro biofilms were formed on the tooth surfaces using a laboratory strain of oral cariogenic bacteria Streptococcus mutans (S. mutans) MT8148. A suspension of S. mutans at an optical density of 2.5 at 500 nm (OD$_{500}$=2.5) was prepared from a 16-h fresh culture in Brain Heart Infusion (BHI, Becton Dickinson, Sparks, MD, USA) broth. The suspension was washed three times with phosphate buffered saline (PBS) and stored at 4°C with gentle stirring. For growing biofilms, a solution of Heart Infusion (HI, Becton Dickinson, Sparks, MD, USA) broth with sucrose (at 1–2% final concentration) was used.

The biofilms were formed in an oral biofilm reactor (OBR) according to the previously reported method$^{22,23}$. In brief, OBR is equipped with two chambers, with each chamber containing a warm water jacket to maintain a constant interior temperature, to grow biofilms under anaerobic conditions for 24 h (Fig. 2). A flat-bulb pH electrode is used to monitor the pH beneath the biofilm continuously. The specimens with undisturbed biofilms were further inoculated for three days to produce root caries lesions. All the specimens were kept in separate wells of a 24-well culture plate (Corning, NY, USA) at 37°C and sucrose-HI was supplemented every day.

After a 3-day incubation, specimens were cleaned using 0.25 M NaOH and washed with distilled water to remove biofilms. The paraffin wax was removed gently. Then the specimens were observed by SS-OCT.

Materials application
The composition of materials used in this study was shown in Table 1. The specimens were randomly divided
Fig. 2 Diagram of one of the chambers of the oral biofilm reactor (OBR) within which artificial biofilms were formed on the specimens surfaces. Digital photographs were the specimens before (left) and 24 h after biofilm formation (right).

Table 1 Materials used in this study

<table>
<thead>
<tr>
<th>Material</th>
<th>Code</th>
<th>Composition</th>
<th>Lot No.</th>
<th>Manufacturer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clearfil SE Bond</td>
<td>SEB</td>
<td>Primer: MDP, HEMA, hydrophilic aliphatic dimethacrylate, dl-CQ, N,N-Diethanol-p-toluidine, water</td>
<td>8K0047</td>
<td>Kuraray Noritake Dental, Tokyo, Japan</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Bond: MDP, Bis-GMA, HEMA, hydrophobic aliphatic dimethacrylate, dl-CQ, N,N-Diethanol-p-toluidine, colloidal silica</td>
<td>8D0079</td>
<td></td>
</tr>
<tr>
<td>Icon-Etch</td>
<td>HA</td>
<td>15% HCl, pyrogenic silicic acid, surface-active agents</td>
<td>692332</td>
<td>DMG, Hamburg, Germany</td>
</tr>
<tr>
<td>Icon-Dry</td>
<td></td>
<td>99% ethanol</td>
<td>692328</td>
<td>DMG</td>
</tr>
<tr>
<td>Icon-Infiltrant</td>
<td></td>
<td>methacrylate-based resin matrix (TEGDMA), initiators, additives</td>
<td>692325</td>
<td>DMG</td>
</tr>
<tr>
<td>K-etchant gel</td>
<td>PA</td>
<td>40% H₃PO₄, water, colloidal silica, dye</td>
<td>470022</td>
<td>Kuraray Noritake Dental</td>
</tr>
<tr>
<td>Propidium iodide</td>
<td>PI</td>
<td>red component used for infiltration test of LIVE/DEAD BacLight</td>
<td>1702</td>
<td>Molecular Probes, Invitrogen Detection Technologies, Carlsbad, CA, USA</td>
</tr>
</tbody>
</table>

MDP: 10-methacryloyloxydecyl dihydrogen phosphate; HEMA: 2-hydroxyethyl methacrylate; CQ: camphorquinone; Bis-GMA: bisphenol-A-diglycidyl methacrylate; TEGDMA: triethyleneglycol dimethacrylate

into 5 groups with 12 specimens in each and were treated as follows:
(1) NC: served as a negative control group without using any materials.
(2) SEB: treated by Clearfil SE Bond (Kuraray Noritake Dental, Tokyo, Japan). The surface was applied by the primer for 20 s and mild air was blown. The bond was applied and air was gently blown. Then they were irradiated with a halogen light-curing unit with 600 mW/cm² output (Optilux 501, Kerr, Orange, CA, USA) for 10 s.
(3) HA120: treated by 120 s 15% HCl (Icon-etch, DMG, Hamburg, Germany), 99% ethanol (Icon-
Dry, DMG) and a low-viscosity resin (Icon-infiltrant, DMG). The surface was etched with Icon-etch for 120 s, rinsed for 30 s and dehydrated with Icon-dry for 30 s. Icon-infiltrant was applied on the specimens with a sponge. After 3 min, gentle air was blown. The resin was light-cured (Optilux 501, Kerr) for 40 s at 600 mW/cm². Icon-infiltrant was then applied a second time for an additional 1 min and light-cured for 40 s.

(4) HA10: treated by 10 s 15% HCl (Icon-etch), 99% ethanol (Icon-dry) and a low-viscosity resin (Icon-infiltrant). Except the etching time, other steps were the same as the application of group (3).

(5) PA10: treated by 10 s 40% H₃PO₄ (K-etchant gel, Kuraray Noritake Dental), 99% ethanol (Icon-dry) and a low-viscosity resin (Icon-infiltrant). Except the etching material and etching time, other steps were the same as the application of group (3).

**Fluorescent microscope (FM) observation**

In order to observe the resin thickness and the resin penetration depth, 4 specimens in group (2) were applied by the mixture of the bond of SE Bond and propidium iodide (PI, LIVE/DEAD BacLight Bacterial Viability Kit, Molecular Probes, Invitrogen Detection Technologies, Carlsbad, CA, USA), and 4 specimens in group (3) to (5) were applied by the mixture of Icon-infiltrant and PI as pretreatments. The left 0.5 mm cutting surface of the stained specimens were gently trimmed off using wet 2000-grit silicon carbide papers to expose a fresh cross section. Then they were observed by fluorescent microscope (FM). A custom code in the image analysis software (Image J, version 1.48, National Institutes of Health, Bethesda, MD, USA) was used to measure the maximum resin thickness and the maximum resin penetration depth (PDmax) (Fig. 3). Then the specimens were observed by SS-OCT.

**Demineralization challenge of materials (Second time demineralization)**

One edge of the top surface of root dentin was covered again by 1-mm-wide paraffin wax to keep the baseline from being demineralized. All specimens were demineralized again the same way as mentioned above. After a three-day incubation, specimens were cleaned using 0.25 M NaOH and washed with distilled water to remove biofilms. The paraffin wax was removed gently. Then the specimens were observed by SS-OCT.

**Swept-source optical coherence tomography (SS-OCT) observation**

SS-OCT (IVS-2000, Santec) was used to examine every step of the experiment process. The stages were labeled as DeM0 (before demineralization), DeM1 (after the 1st time demineralization), Mat (after immediate materials application) and DeM2 (after the 2nd time demineralization). This system is a frequency-domain OCT technique that interprets the magnitude and coherence of the light reflected from the subject into the depth-profile of the subject. The system incorporates a high-speed frequency, swept external-cavity laser. The wavelength ranged from 1,260 to 1,360 nm (centered at 1,310 nm) at a 20-kHz sweep rate. The optical resolution was 20 µm transversally and 12 µm axially in the air (7–8 µm in tissues with a refractive index around 1.5). Every time, SS-OCT examination was performed along the same plane between the marked two holes and then central cross-section images were obtained. The demineralized specimens were cross-sectionally scanned with SS-OCT after gentle blot-drying of the surface leaving it moist without any visible water-droplets. To ensure the reproducibility of the scan, the specimens were placed at the same orientation as accurately as possible.
A custom code in the image analysis software (Image J) was used to read the raw data of SS-OCT. The obtained SS-OCT image was rotated to compensate for the tilting during the scan to reach a horizontal surface. A binarization process (Fig. 4) was applied to the raw data (a', b'). The lesion depth was measured vertically from the surface to the bottom of the lesion. The lesion depths of 20 positions from each specimen were measured with a width of 2,000 µm and an interval of 100 µm. The average was calculated as the value of lesion depth of one specimen. LD1 was the lesion depth after DeM1. LD2 was the lesion depth after DeM2. ∆LD was the increase in lesion depth from LD1 to LD2.

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\Delta LD = LD_{2\text{OCT}} - LD_{1\text{OCT}}
\]

The length of dentino-enamel junction (DEJ) separation was measured and the frequency of specimens with cervical enamel loss was counted.

Confocal laser scanning microscope (CLSM) observation

After the SS-OCT observation of the 2nd demineralization stage, all specimens were fixed in epoxy resin (EpoxiCure, Buehler). After 8 h, a low-speed diamond saw (Isomet) was used to cut each specimen in the following way.
half along the two central holes and obtain discs with thickness of approximately 2 mm. Then, the slices were trimmed off using wet 2000-grit silicon carbide papers and further polished with diamond paste down to 0.25 µm under running water. The cross-sectional slice examined using the nondestructive SS-OCT imaging was physically separated and examined under CLSM (1LM21H/W, Lasertec, Yokohama, Japan). The lesion depth from CLSM (LD2CLSM) was measured in the same manner as that from SS-OCT by Image J (Fig. 5).

Dentin surface removal by pretreatments
Additional specimens were prepared for observing the amount of normal dentin surface removal by different pretreatments. Four intact human root surfaces were polished by wet 2000-grit silicon carbide papers to expose flat and fresh root dentin surfaces (4×4×3 mm). After cleaning by tap water, half surfaces were covered by stick tapes to keep the baselines and another half surfaces were applied by 15% HCl (Icon-Etch) 120 s, 15% HCl (Icon-Etch) 10 s, 40% H₃PO₄ (K-etchant gel) 10 s and SE Bond primer 20 s, respectively. Then the tapes were removed and air was blown according to manufacturer instructions. After SS-OCT observation, the specimens were fixed in epoxy resin and cut in half by a low-speed diamond saw. The cross-sectioned surfaces were polished and observed by CLSM. The dentin surface removal was measured vertically from the baseline to the eroded surface by Image J.

Statistical analysis
The maximum resin penetration depth (PDmax) and the increase in lesion depth (∆LD) were analyzed by one-way ANOVA with Tukey’s post hoc tests. The correlation between LD2OCT and LD2CLSM was analyzed by Pearson’s correlation test. The DEJ separation length of DeM and that of Mat was compared by Wilcoxon signed ranks test. The frequency of specimens with cervical enamel loss of DeM1 and that of Mat was analyzed by Fisher’s exact test. All statistical analyses were performed with 95% level of confidence using Statistical Package for Medical Science (Version 16.0 for Windows, SPSS, Chicago, IL, USA).

RESULTS
The penetrated resin in the lesion body by the treatment of Icon-infiltrant and SE Bond could be detected from FM (Fig. 3). HA120 showed 138.00±49.25 µm PDmax that was significantly larger than PA10 and SEB (p<0.05). A maximum thickness of 136.58±64.73 µm coating layer in SEB could be detected on the lesion surface, while resin-infiltrant could not be detected on the lesion surface. Moreover, resin-infiltrant formed inhomogeneous penetration layers in demineralized dentin, while SE Bond formed homogeneous coating layers, which were stable even after the 2nd demineralization (Figs. 3 and 6).

The SS-OCT images were taken in wet condition, and the lesion surface appeared flat with no remarkable shrinkage or loss of surface as presented in Fig. 4. In all the SS-OCT images after demineralization, a boundary was seen below the lesion surface that suggested the lesion front as bright zones with increased signal intensity. LD2OCT was remarkably higher than LD1OCT in all groups (Figs. 4 and 7). HA120 and SEB showed significantly lower ∆LD than NC (p<0.05), while HA10 and PA10 did not show significantly lower ∆LD than NC (p>0.05) (Fig. 7). LD2OCT and LD2CLSM showed a significant correlation (r=0.865, p<0.05) (Fig. 8).
DEJ separation showed on SS-OCT images as a white line with intensified brightness and changed in length during different experimental stages. The length of DEJ separation reduced in HA120, HA10 and PA10, while that increased in SEB after immediate materials application. In HA120, DEJ separation length of Mat

**Fig. 8** Significant correlation between LD2\_OCT and LD2\_CLSM (r=0.865, p<0.05, Pearson’s correlation test).

**Fig. 9** DEJ separation changed in length during different experimental stages. The blank arrows showed the DEJ separation. After the 1st demineralization (DeM1), DEJ separation formed and showed on SS-OCT images as a white line with increased signal intensity. After immediate application of resin-infiltrant with 120 s HCl pretreatment (HA120), DEJ separation disappeared. However, after the 2nd demineralization (DeM2), DEJ separation formed again and increased in length.

**Fig. 10** Length of DEJ separation from SS-OCT analysis. The length of DEJ separation was compared between DeM1 and Mat within each group of SEB, HA120, HA10 and PA10. Bars showed significant differences between DeM1 and Mat within each group (p<0.05, Wilcoxon signed ranks test). DeM1: after the 1st time demineralization; Mat: after immediate materials application; DeM2: after the 2nd time demineralization.

**Fig. 11** The frequency of specimens with cervical enamel loss from SS-OCT analysis. The frequency was compared between DeM1 and Mat within each group of SEB, HA120, HA10 and PA10. No significant difference was detected between DeM1 and Mat within each group (p>0.05, Fisher’s exact test). DeM1: after the 1st time demineralization; Mat: after immediate materials application; DeM2: after the 2nd time demineralization.
Fig. 12 CLSM images of surface removal of normal human root dentin with different pretreatments.
The dot line was the baseline. After etching with 15% HCl (Icon-etch) for 120 s, approximately 25–30 µm dentin surface was removed. After etching with 15% HCl (Icon-etch) for 10 s, approximately 5 µm dentin surface was removed. After etching with 40% H₃PO₄ (K-etchant) for 10 s, dentin surface removal could not be detected. After applying by SE Bond primer for 20 s, dentin surface removal could not be detected.

was significantly lower than that of DeM1 (p<0.05). However, in groups SEB, HA10 and PA10, DEJ separation length did not show significant difference between DeM1 and Mat within each group (p>0.05) (Figs. 9 and 10).

From SS-OCT images, although no significate difference in the frequency of specimens with cervical enamel loss was detected between DeM1 and Mat within each group (p>0.05), the frequency increased in HA120 and HA10 after immediate materials application. Finally, approximately half specimens of HA120 and HA10 and only one specimen of SEB had cervical enamel loss (Fig. 11).

The thicknesses of normal root dentin surface removal by different pretreatments were ordered by follows: 15% HCl 120 s (25–30 µm)>15% HCl 10 s (5 µm)>40% H₃PO₄ 10 s (0 µm)=SE Bond primer 10 s (0 µm) (Fig. 12).

DISCUSSION
Resin infiltration has been introduced as an alternative treatment option for non-cavitated enamel lesions that are not expected to remineralize or arrest by non-invasive measures alone. It could effectively prevent the further demineralization of artificial enamel caries lesions under cariogenic conditions in situ. On the other hand, with the increase of a dentate elderly population, the occurrence of root caries is also increasing. Preventing and arresting root caries is one of the essential issues to maintain the quality of life of elderly people. Therefore, the potential effect of resin infiltration on root caries was evaluated in this in vivo study. As a comparison, a conventional resin adhesive without composite resin filling was used to seal or to coat on root caries.

In this study, the null hypothesis that resin infiltration could not reduce further demineralization of root caries lesions under cariogenic conditions was rejected, because resin infiltration pretreated by 120 s HCl was shown to have a significantly less increase in lesion depth than the control group according to SS-OCT analysis. Whereas, resin infiltration pretreated by 10 s HCl and 10 s H₃PO₄ showed similar lesion depth increase as the control group. Also, resin-infiltrant with HCl pretreatments could penetrate deeper than that with H₃PO₄ pretreatment and SE Bond. Resin infiltration results in considerably deeper resin penetration whilst the pre-treatment with hydrochloric acid seems to be more suitable compared with the use of a phosphoric acid. However, the application of 15% HCl for 120 s apparently removed more dentin structure from the root surface (approximately 25–30 µm) than other pretreatments. Similar result was reported previously that 120 s 15% hydrochloric acid gel treatment could erode the surface layer more effectively (around 50
lesion body than conventional dental adhesives\(^{14,15}\). The resin infiltrant has a significantly deeper penetration in the lesion surface, but only 18.00±12.36 µm maximum coating layers on the lesion surface. In reverse, the later resin infiltration groups, while they increased in length in the demineralization of root caries model formed on human teeth using the major cariogenic bacteria by simulating an oral environment and before the second biofilm attack. Moreover, the substructure for the application of resin infiltration on root caries, which may provide some possibilities and information to explore a wider application of the material.

Although resin infiltration had risks of cervical enamel loss and dentin over removal because of the strong acid (15% HCl) pretreatment. Although resin infiltration (pretreated by 120 s HCl) had less lesion depth increase than the control group and deeper resin penetration into the lesion body and DEJ separation than SE Bond, more specimens in caries-infiltrant groups (pretreated by HCl for 120 s and for 10 s) had cervical enamel loss than the specimens in SE Bond group. The stronger acid with a longer time application caused more damage to the normal dental hard tissues. In case of soft tissues, although short-term contact of this strong acid with mucosa has been shown to be harmless\(^{30}\), it may cause severe burns on contact with soft tissues even with a rubber dam in clinical practices. Also, a strong acid can cause sensitivity on exposed dentin surface after a gingival recession.

Propidium iodide (PI, LIVE/DEAD BacLight Bacterial Viability Kit), the red-fluorescent nucleic acid stain is usually used to detect dead bacteria and also dead animal cells. In the present study, PI was used as a fluorescent dye mixed with Icon-Infiltrant or with SE Bond. We successfully detected them by a FM to get clear images inside the demineralized dentin body (as PI has an affinity to bind with resin and has been used in ion-exchange biotechnology at a molecular level). There are some discrepancies in resin infiltration that caused incomplete and inhomogeneous filling into lesions. The caries model formed on human teeth using the major cariogenic bacteria by simulating an oral environment in this study expected to be very similar to the natural caries. Therefore, incomplete surface layer erosion (even for those etched with HCl) in natural lesions and organic materials, such as proteins and carbohydrates, which might contaminate the pores of natural caries, could be responsible for those discrepancies\(^{20}\). Moreover, inorganic components of dentin (e.g. collagens) might have resisted proper penetration of the resin-infiltrant and interfered its solidification as it happens in case of non-cavited enamel chemically combining with enameleprisms. Other factors, including residual biofilms, bacteria, culture media, other solutions, residual chemical compounds, reduced wettability and entrapped air might hamper resin penetration as well.

Four specimens in each group were stained instead of all specimens because the main purpose was to evaluate the lesion depth using SS-OCT considering that mixing with the stain may cause some disturbances to the infiltrant, that did not occur actually though. Another technical difficulty of this staining was that the FM images needed to be taken just after the material application and before the second biofilm attack. Moreover, the substructure for the application of resin infiltration was caries-infected dentin rather than non-cavited enamel caries. The current version of Icon-infiltrant is actually indicated for early enamel caries treatment only, not for root caries treatment; rather contra-indicated under the same application protocol of pretreatment with 120 s 15% HCl. As obvious, risks of damaging delicate structures at the cervical parts of vital or even non-vital teeth have been considered. To minimize those risks, the application of strong acids for the shortest possible time (10 s) was also tested in this study. Unfortunately, results were not expectedly effective both in terms of the resin penetration and the prevention of demineralization progression. However, this is the first study till now to evaluate the effect of resin infiltration on root caries, which may provide some possibilities and information to explore a wider application of the material.

Consequently, modified protocols and studies on
the pretreatments for cervical hard and soft tissues protection and for root caries against the further demineralization seem to be necessary for the application of resin infiltration in future.

**CONCLUSION**

Within the limitations of this *in vitro* study, it can be concluded that resin infiltration pretreated by 120 s HCl has a better penetration ability than that pretreated by H3PO4 and SE Bond. Also it has a preventive effect on root caries as similar as SE Bond. However, the strong acid (15% HCl) pretreatment has risks of cervical enamel loss and over removal of dentin structure as well.

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**REFERENCES**


