Effects of bovine milk osteopontin on in vitro enamel remineralization as a topical application prior to immersion in remineralizing solutions with/without fluoride

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The aim of the present study was to investigate the effects of bovine milk osteopontin (OPN) on enamel remineralization as a topical application prior to immersion in remineralizing solutions with/without fluoride. Bovine enamel blocks were demineralized then divided into the following 3 groups: OPN (2.7 and 5.4 µM) solutions and deionized water (control). Each group was divided into 2 groups (remineralizing solution with or without 1 ppm of fluoride (F)). The specimens were analyzed by micro-CT and scanning electron microscope (SEM). The percentage of remineralization was higher in remineralization solution with than without F (p<0.05). The present results suggest that bovine milk OPN inhibits remineralization in solution without F, but 5.4 µM bovine milk OPN does not inhibit remineralization of the demineralized body using solution containing F by interrupting mineral deposition on the enamel surface.

Keywords: Bovine milk osteopontin, Remineralization, Demineralization, Enamel, Fluoride

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INTRODUCTION

Demineralization and remineralization occur on the enamel surface, and the balance between these processes is maintained under an oral environment. A disruption in this balance eventually leads to the development of dental caries. White-spot lesions in the early stage of the dental caries process prior to cavitation are a representative clinical finding that is histologically observed in the mineral-poor region of the enamel subsurface with an intact surface layer. The remineralization of enamel, a natural repair process, occurs at the demineralized region, with plaque/salivary calcium (Ca²⁺) and phosphate (PO₄³⁻) ions being deposited in crystal voids, which results in mineral gain. The presence of free fluoride (F⁻) ions in the oral environment increases the incorporation of Ca²⁺ and PO₄³⁻ ions into the crystal lattice. Therefore, an effective method for dental caries management is to improve the remineralizing process using remineralization products. New improved biochemical materials need to possess properties that achieve enamel remineralization even in saliva with low Ca²⁺ and PO₄³⁻ ions concentrations. These materials include bioactive glass (BAG), casein phosphopeptide-amorphous calcium phosphate (CPP-ACP), and functionalized β-tricalcium phosphate (fTCP). Surface biominerlization is another new approach to improve remineralization with advances in peptide design.

Osteopontin (OPN), a non-collagenous protein, was initially detected in bone, but is also present in teeth, soft tissues, and physiological fluids including the kidneys and mammary and salivary glands. OPN has also been detected in saliva and bovine milk. Milk OPN has multiple functions in early and adult life. A high concentration of OPN is present in human milk, and commercial infant formulas with OPN have recently become available. Previous studies showed that bovine milk OPN bound directly to bacteria and enhanced phagocytosis, and also prevented oral bacterial adhesion to the tooth surface and reduced oral biofilm formation and pH drops. OPN is a phosphorylated glycoprotein, and Kumura et al reported the inhibitory effects of bovine milk OPN and its fragments on the formation of calcium phosphate precipitates. However, the effects of bovine milk OPN on enamel remineralization currently remain unknown.

Topical application of protein essentially inhibits uptake of mineral to enamel. Earlier reports showed that combination of casein and 1 ppm F increase quantity of mineral uptake. We therefore hypothesize the application of OPN to a tooth surface inhibits to intake the mineral, but F in remineralization solution recovers to intake the mineral. In the present study, we investigated the effects of bovine milk OPN, as a potential caries preventive agent in saliva, on the remineralization of enamel lesions using remineralizing solutions with or without fluoride in vitro.

MATERIALS AND METHODS

Ethics approval

The study was performed using food industry animal...
carcass waste, and ethical approval was not required for waste tissues. The bovine teeth used in this study were stored and disposed under the direction of the guidelines and policies of Tokyo Dental College (reference number: 2020-103-02).

**Specimen preparation**

Thirty bovine incisors were purchased from Yokohama Meat Corporation (Yokohama, Japan), and were cryopreserved and defrosted using running tap water immediately before use. After the surrounding periodontal tissues had been mechanically removed, the coronal pulp tissues and roots were also removed. The crowns of teeth were cut into 3×3×2 mm pieces, and enamel-dentin blocks were prepared using a low-speed diamond saw (Isomet, Buehler, Lake Bluff, IL, USA) under water-cooling conditions. These blocks were embedded into epoxy resin (EpoxiCure 2, Buehler), and their enamel surfaces were flatly ground with 600- to 2000-grit silicon carbide (SiC) papers (Fuji Star, Sankyo Rikagaku, Saitama, Japan) under running water. The edge (1 mm) of the surface was coated by applying nail varnish (Revlon Red 680, Revlon, New York, NY, USA) and an observation window (2×2 mm) was created by exposing the polished enamel surface of each specimen. A hole (1 mm in diameter and 0.5 mm in depth) was formed at the side of each specimen using a diamond bur (440SS ISO # 010, Shofu, Kyoto, Japan) as a reference point for micro-computed tomography (micro-CT) scans.

The specimens were randomly allocated into 6 groups (n=9 each) based on the concentration of OPN (0 µM, 2.7 µM, and 5.4 µM) and F (0 or 1 ppm) in remineralization solution as described below.

**Demineralization**

Each specimen was put in a 2 mL-microtube and immersed in 2 mL of demineralizing solution (17.8 mM CaCl₂, 8.8 mM KH₂PO₄, and 100 mM lactic acid; the pH of the solution was adjusted to 4.3 with 10 mM KOH) at room temperature for 6 to 8 days according to the method by Margolis et al. with modifications. Demineralizing solution was refreshed daily.

We checked all the demineralized specimens by use of micro-CT scan and analysis software Ratoc as described below. The specimens that had extreme deep or shallow demineralization in depth were excluded.

**Application of bovine milk OPN**

OPN derived from bovine milk was purchased from Sigma-Aldrich (St. Louis, MO, USA). OPN was dissolved in deionized water without precipitation. The concentration of OPN was determined in reference to Schlafer et al. After demineralization, 10 µL of OPN (final concentrations of 2.7 and 5.4 µM) was applied to the enamel surface of each specimen at 37°C for 30 min. Specimens were then rinsed off using deionized water for 10 s. As a negative control, deionized water was applied instead of OPN.

**Remineralization**

Remineralizing solution (1.5 mM CaCl₂-2H₂O, 0.9 mM KH₂PO₄, 130 mM KCl, 20 mM HEPES, and NaF 1 mg/L; pH adjusted to 7.0 with 10 mM KOH) was prepared according to the method by ten Cate and Duijsters. Remineralizing solution containing F⁻ (1 ppm) was also used in the present study according to the method by Romero et al.. Each specimen was separately put in the 2 mL-microtube and immersed in 2 mL of remineralizing solution at 37°C for 14 days. The remineralization solution was changed every day. At 7 and 14 days, specimens were scanned by micro-CT and observed under a scanning electron microscope (SEM) at 14 days (Fig. 1).
Micro-CT scanning and image analysis
After demineralization and remineralization, a micro-CT system (InspecXio SMX-100CT, Shimadzu, Kyoto, Japan) was used to evaluate the mineral density (MD) of specimens. The micro-CT system generates polychromatic X-rays with cone-beam geometry. A 0.2-mm-thick brass plate was set in the beam path to reduce the beam-hardening effect. The tube voltage was 100 kV at a current of 100 µA. The specimen was mounted on a turntable to vertically irradiate X-rays on the enamel surface. A series of mineral reference phantoms were also scanned for MD calibration and included three hydroxyapatite (HAP) disks (TRI/3D-BON, Ratoc, Tokyo, Japan); two with different concentrations (0.50 and 0.70 gHAp/cm³) of HAP crystals mixed with epoxy resin (Epoxicure Resin, Buehler) and one pure HAP disk (3.16 gHAp/cm³) (Cellyard, Hoya, Tokyo, Japan). The phantoms were scanned in a plastic tube, which was almost the same as the imaging conditions of the specimens.

Scanned data were reconstructed from 2-dimensional (2D) images with a resolution of 512×512 pixels to a 3-dimensional (3D) image using 3D analysis software (TRI/3D-BON, Ratoc). These images were used for visualization and quantitative volumetric measurements. CT values were converted into MD values (gHAp/cm³) using a linear calibration curve based on the grey values obtained from the mineral reference phantoms. The MD profile of each specimen was obtained by plotting MD (Vol%) and MD using the F (+) group. Maximum MD in each specimen was normalized to 100 vol% against depth. To minimize ring artifacts, air calibration of the detector was carried out prior to each scanning, and 6-frame averaging was also applied in the acquisition phase to improve the signal-to-noise ratio (SNR)³⁴. For correction of beam hardening, a 0.2-mm thick brass (Cu–Zn) filter was installed in the beam path to reduce beam-hardening effect³⁴.

The percentage of remineralization was calculated according to standard procedures³⁴. Lesion depth (LD) was set at the depth from the original enamel surface (covered area with varnish), and mineral loss (ΔZ) (Vol% µm) was integrated mineral loss up to the lesion depth. To evaluate the extent of repair, the relative change in percentage of remineralization (%R) was calculated as follows: %R=(ΔZ in DEM−ΔZ in REM)/ΔZ in DEM×100.

SEM observations
Specimens were prepared in the same manner as described for micro-CT observations. The enamel surface at the treatment window was observed under a field emission SEM (SU6600, Hitachi High Technologies, Tokyo, Japan) at x3,500 magnification, following a specimen processing protocol for SEM, which included desiccation and carbon sputter coating. After observation of enamel surface, the specimen was cut with diamond pointed burs and nippers, and cross-sectional surface was observed under SEM. Cross-sections were also observed at the same magnification.

Statistical analysis
The results of percentage of remineralization (%R) were expressed as the mean and standard deviation. Differences within and between the mean values of groups were statistically analyzed using a two-way repeated-measures analysis of variance (ANOVA) with application and period as factors and Tukey’s test. The significance level of all tests was set at α=0.05.

RESULTS
Micro-CT analysis
An X-ray transmission image of the specimen immersed in demineralizing solution for 6 to 8 days showed demineralization on the surface layer of enamel (Fig. 2, left column). In all groups, radiopacity was observed in the demineralized lesions after 7 and 14 days. Radiopacity became noticeable at 14 days in specimens in solution with fluoride (the F (+) group) compared to that without F (the F (−) group).

Percentage of remineralization
Table 1 shows the mean percentage of remineralization of all groups.

In all groups, the percentage of remineralization was higher after 14 days than after 7 days. The percentage of remineralization was higher in the F (+) group than in the F (−) group (p>0.05) (Table 1).

In the F (−) group after 7 days, no significant difference was observed in the percentage of remineralization among the three groups (Table 1). After 14 days, the percentage of remineralization was significantly higher in the control group than in the OPN 2.7 µM and OPN 5.4 µM groups (p<0.05) (Table 1). No significant differences were observed between the OPN 2.7 µM and OPN 5.4 µM groups.

In the F (+) group after 7 days, the percentage of remineralization was significantly lower in the OPN 2.7 µM group than in the control group after immersion in remineralizing solution (p<0.05), and no significant differences were observed between the control and OPN 5.4 µM groups. After 14 days, the percentage of remineralization was significantly lower in the OPN 2.7 µM group than in the control and OPN 5.4 µM groups (p<0.05), and no significant differences were observed between the control and OPN 5.4 µM groups.

SEM observations
After 14 days, compared to the groups of demineralization (Fig. 3a) and control without F (Fig. 3b), a crystal-like deposit of approximately 1 µm in size was observed in OPN (+) of the F (−) groups (Figs. 3c and d). At OPN 5.4 µM, more deposits were detected on the enamel surface than at OPN 2.7 µM (Figs. 3c and d). Rod-like deposits were observed in the F (+) group (Figs. 3e, f, and g).

In cross-sectional images, a sparse structure was confirmed at the subsurface region after demineralization (Fig. 4a). In the F (−) group after 14 days, the sparse structure disappeared in the control group (Fig. 4b), but remained in the OPN (+) groups (Figs. 4c and d). In the
Fig. 2 2D micro-CT images of single specimens from each group. 2D images of the control, OPN 2.7 µM, and OPN 5.4 µM groups are shown after demineralization and after 7 and 14 days of remineralization. After demineralization, an X-ray transmission image was obtained showing lesions on the surface layer of enamel. After 7 and 14 days of remineralization, the radiopacity of the areas was increased in all groups. 7d: after 7 days of remineralization; 14d: after 14 days of remineralization; OPN: osteopontin; F: remineralizing solution containing F (1 ppm).

Table 1 Mean percentage of remineralization (%R) for each experimental group

<table>
<thead>
<tr>
<th>Group</th>
<th>7 d</th>
<th>14 d</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>24.27±3.84^Aa</td>
<td>41.90±2.84^Bd</td>
</tr>
<tr>
<td>F, Control</td>
<td>42.29±2.81^Ab</td>
<td>51.40±2.97^Bf</td>
</tr>
<tr>
<td>OPN 2.7 µM</td>
<td>20.70±4.89^Aa</td>
<td>36.91±2.84^Be</td>
</tr>
<tr>
<td>F, OPN 2.7 µM</td>
<td>36.22±4.10^Ac</td>
<td>42.53±3.59^Bd</td>
</tr>
<tr>
<td>OPN 5.4 µM</td>
<td>20.17±3.68^Aa</td>
<td>33.87±2.78^Be</td>
</tr>
<tr>
<td>F, OPN 5.4 µM</td>
<td>37.38±2.08^Ac</td>
<td>47.39±4.04^Bf</td>
</tr>
</tbody>
</table>

Data were expressed as mean (%)±standard deviation (SD). The percentage of remineralization was higher in the F (+) group than in the F (−) group in the control, OPN 2.7 µM, and OPN 5.4 µM groups at 7 and 14 days. The same capital letters in 7d and 14d are not significantly different (p>0.05). The same small letters in each group are not significantly different (p>0.05). 7d: after 7 days of remineralization; 14d: after 14 days of remineralization; OPN: osteopontin; n=9 per group; F: remineralizing solution containing F (1 ppm).

DISCUSSION

The present results demonstrated that remineralization of demineralized bovine enamel occurred even in the presence of bovine milk OPN. Salivary phosphoproteins, such as acidic proline-rich proteins (PRPs) and statherins, stabilize high concentration of Ca^{2+} and PO_{4}^{3−} to facilitate ion transport and the formation of calcium-phosphate crystals.
bioavailability for mineralization. They inhibit phase change in solution, but upon interaction with the tooth surface bound ions can be made bioavailable. Advantages of introducing phosphoprotein to simulate the mineral-stabilizing properties of saliva have been proposed. Saliva and bovine milk contain phosphorylated OPN, and thus, the application of OPN to the enamel surface may be made bioavailable. However, the
effect has been unknown. Our micro-CT images showed radiopacity became noticeable at the demineralized areas despite OPN application at 7 and 14 days, and the percentage of remineralization (%R) at 14 days was higher than that at 7 days in corresponding group. The results suggest percentage of remineralization in demineralized enamel gains despite the application of bovine milk OPN. This is the first study to examine the effects of bovine milk OPN on enamel remineralization. However, the degree of mineralization was affected by fluoride added to artificial saliva; the percentage of remineralization was higher in F (+) solution than in F (-) solution.

In F (-) solution, the percentage of remineralization was significantly higher in the control group than in the OPN groups at 14 days, suggesting that OPN inhibited remineralization. The SEM analysis showed the accumulation of crystal-like deposits on the surface of enamel in the 2.7 µM and 5.4 µM groups. The enamel surface is covered by salivary proteins and molecules, such as statherin and PRPs, which are important constituents of acquired pellicles 4). OPN has Ca 2+-binding sites 40) and binds to HAp on the enamel surface. Therefore, bovine milk OPN binds not only to HAp on the enamel surface, but also to Ca 2+ in remineralization solution. As a result, the entrance of Ca 2+ into the subsurface area was interrupted at the enamel surface in F (-) solution.

The percentage of remineralization was higher in F (+) solution than in F (-) solution. At 14 days, SEM images showed a firm structure at the subsurface region and rod-like structural precipitations, which may be images showed a firm structure at the subsurface region and rod-like structural precipitations, which may be images showed a firm structure at the subsurface region and rod-like structural precipitations, which may be explained by F (+) solution. As a result, the enamel surface was covered by salivary proteins and molecules, such as statherin and PRPs, which are important constituents of acquired pellicles 4). OPN has Ca 2+-binding sites 40) and binds to HAp on the enamel surface. Therefore, bovine milk OPN binds not only to HAp on the enamel surface, but also to Ca 2+ in remineralization solution. As a result, the entrance of Ca 2+ into the subsurface area was interrupted at the enamel surface in F (-) solution.

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The application of protein to the enamel surface at a high concentration has been shown to inhibit enamel remineralization 40). We measured the concentration of OPN based on previous findings showing that 1.5 and 2.7 µM of bovine milk OPN effectively inhibited initial oral bacteria adhesion 27) and also reduced biofilm formation and pH drops in dental biofilms 28,29,42,43). We used 1 ppm of F, which was previously reported to be effective for dental caries prevention 30,41). Our results showed 2.7 µM OPN inhibited remineralization whereas 5.4 µM OPN recovered remineralization in solution with F. The present results indicate that 5.4 µM of bovine milk OPN with 1 ppm of F solution would contribute to facilitate ion bioavailability for remineralization.

CONCLUSION
In conclusion, the remineralization of demineralized enamel was achieved despite the application of bovine milk OPN prior to immersion in solution with F. Therefore, the present results suggest that bovine milk OPN inhibits remineralization in solution without F, but 5.4 µM bovine milk OPN does not inhibit remineralization of the demineralized body using solution containing F by interrupting mineral deposition on the enamel surface.

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CONFLICTS OF INTEREST
The authors have explicitly stated that there are no conflicts of interest.

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