In Vitro Remineralization of Acid-etched Bovine Enamel with Amino Acids/Hydroxyapatite Composite

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Abstract: Remineralization of early enamel caries lesions plays a crucial role in prevention and restoration of dental caries. Based on the crucial amino acids contained in enamel matrix, amino acids /hydroxyapatite (AA/HAP) composite was synthesized in the presence of 10 mmol/L glycine, 10 mmol/L L-serine and 5 mmol/L L-aspartic acid. The physical, chemical and biological properties of the composite were characterized. And its remineralization effect on acid-etched bovine enamel was evaluated. Under the inhibition properties of amino acids, the composite has lower crystallinity and higher biocompatibility compared to the hydroxyapatite (HAP) without amino acids. In vitro remineralization of acid-etched bovine enamel using AA/HAP was conducted in artificial saliva. After remineralization, the surface and cross-section morphology, the components, and the mechanical property of bovine enamel samples were characterized, respectively. The results showed that the AA/HAP could induce the restoration of both surface and subsurface enamel lesions. The amino acids released from the composite could adsorb on the organic matrix residues and induce the formation of parallel arranged HAP crystals, eventually formed significant surface microhardness (SMH) recovering.

Key words: remineralization; enamel; hydroxyapatite; amino acid

Tooth enamel, the hardest tissue in human body, is composed of around 96wt% hydroxyapatite (HAP, Ca₁₀(PO₄)₆(OH)₂), 3wt% of water and less than 1wt% organic matrix[1-2]. Its special properties are ascribed to the structure of parallel arranged HAP prisms (which wrapped in a thin layer of organic matrix)[3-4]. This special structure has an abnormal consistency across all species despite the diversity of other tissues. Such invariability suggests that enamel has a common structural and functional basis between different species[3-5].

Unlike synthetic HAP, the HAP in teeth is often calcium deficient due to cations substitutions, which making enamel vulnerable to acid-etching[6-7]. Enamel is always exposed to acidic food or drink, as well as the acidic by-products produced by cariogenic bacteria. Therefore, enamel caries has been a major oral health problem worldwide. Fluoride is currently the most effective and widely used anti-caries method. It can reduce enamel demineralization and promote enamel remineralization. A relatively high concentration of fluoride could increase abrasion resistance and decrease caries process in enamel[8]. However, excessive exposure to fluoride may induce fluorosis.

In recent years, biomimetic remineralization methods have been paid great attention. For example, amelogenin and its mimetic assembly supramolecular[9-10] was applied in ex vivo remineralization of artificial enamel lesions. Other biomimetic methods, such as amelogenin-derived peptides[11] and repeats of aspartate–serine–serine peptides[12] was also clarified to have biomineralization effect on enamel lesions. However, it is very difficult and too expensive to extract or synthesize these natural proteins or peptides, hence limit the application of these methods. Besides, most of the investigations mentioned above are more concerned about enamel surface remineralization. Regeneration of subsurface and deep enamel lesions was rarely referred to.

As the outermost layer of the tooth, enamel is always exposed to a complex oral environment. Therefore, enamel evolved to be more resistant to demineralization than other mineralized tissues[6]. Furthermore, enamel has a hierarchical graded structure[13]. Its hardness decreases obviously from outer layers to the direction of dentin-enamel junction (DEJ), which is attributed to the less ordered enamel prisms, more crystal defects, lower mineral density and higher organic matrix contents[14].
Once been etched by acid, the subsurface demineralization will form white spot lesions (WSLs) at first. Without effective treatment, WSLs would turn into caries thereby increasing the application of invasive restorative treatments\cite{15}.

As a biocompatible and bioactive material, synthesized HAP has been widely used in remineralization of enamel lesions\cite{16-17}. Nonetheless, these methods mainly utilized the size effect of HAP for the formation of mineralized layers in the cavities of the enamel surface. With a similarity in crystallinity to the apatite in bone and enamel, these HAP materials have poor solubility and only can provide limited Ca\(^{2+}\) and PO\(_4^{3-}\) ions for enamel remineralization. Therefore, the invasive restorative treatments is still badly in need of new strategy.

Glycine (Gly), L-serine (Ser) and L-aspartic acid (Asp) are the pivotal amino acids contained in the enamel matrix at different stages of amelogenesis\cite{1,18}. Molecular dynamic simulations showed that the (002) faces of HAP tend to be unstable in Gly solutions\cite{19}. Thus, the ordered assembly of HAP crystals along the c-axis was facilitated for minimizing the total interfacial energy. Meanwhile, Gly has the smallest molecular size in amino acids, and it is easy to diffuse into the deep layer of enamel lesions. Ser with hydroxyl side group and Asp with carboxyl side group both have high affinity for HAP\cite{20}. In addition, HAP is proved to be positively charged when orientated along the c-axis\cite{21}. Therefore, the molecular of Ser and Asp could strongly integrate with the positively charged enamel prisms via electrostatic adsorption, and facilitate the topical supersaturation of Ca\(^{2+}\) and PO\(_4^{3-}\) in the subsurface enamel lesions. In the synthesis of HAP, the adsorption of Gly, Ser and Asp on HAP crystal surface could change the crystalline size and the morphology of HAP, and could significantly increase the solubility of HAP\cite{22}.

The present study verified the remineralization effect of HAP composites which were modified with different combinations of amino acids, and found that the AA/HAP synthesized in the presence of 10 mmol/L Gly, 10 mmol/L Ser and 5 mmol/L Asp showed optimal remineralization effect for both surface and subsurface lesions of acid-etched bovine enamel. Since these three kinds of amino acids all have smaller molecular size and contain different side functional groups, they may act synergistically in the synthesis of AA/HAP and the remineralization of subsurface enamel lesions.

1 Materials and methods

1.1 Materials

Analytical grade NaN\(_3\) was purchased from Fuchen (Tianjin) Chemical Reagent Co, Ltd. Phosphate-Buffered Saline (PBS) was purchased from Corning Mediatech, Inc. The 293T cells were purchased from Kunming Institute of Zoology, Chinese Academy of Sciences. All of the other reagents and solvents if not specified were purchased from Sinopharm Chemical Reagent Co, Ltd and were all analytical grade. Deionized (DI) water was used in all procedures.

1.2 Synthesis of AA/HAP composite

AA/HAP composite was synthesized using a chemical homogeneous precipitation method. Ca(NO\(_3\))\(_2\)-4H\(_2\)O (14.169 g, 60 mmol), Gly (2.580 g, 10 mmol/L), Ser (3.573 g, 10 mmol/L), and Asp (2.287 g, 5 mmol/L) were dissolved in 1600 mL DI water, respectively. The mixture was heated and maintained at (37\(\pm\)1) \(^\circ\)C in the subsequent processes. (NH\(_4\))\(_2\)HPO\(_4\) (4.752 g, 35.98 mmol) dissolved in 800 mL DI water was slowly titrated into the prepared mixture in about 120 min. 2.5wt% ammonia solution was adopted to maintain pH of the mixture at (10\(\pm\)0.5) during the titration process. After continuous stirring for 24 h, the suspension aged at standing state for 24 h and washed with DI water until the pH decreased to about 7.0 to remove the impurities. The AA/HAP powder was obtained following the lyophilized of the precipitate for 12 h. The HAP powder without amino acids was synthesized according to the above procedures.

1.3 Cytotoxicity assay

CCK-8 assay was performed to evaluate the cytotoxicity of the HAP and AA/HAP. The 3rd generation of 293T cells (one typical human renal epithelial cells transfected with adenovirus E1A gene) were cultured in 10% FBS at 37 \(^\circ\)C (under an atmosphere of 5% CO\(_2\) and 95% relative humidity). The cells were seeded in a 96-well culture plate at a density of 5.0\(\times\)10\(^4\) cells/well. After another 48 h incubation, 10 µL CCK-8 solution was slowly added into each well, and the culture plate was incubated for 4 h. Then the culture medium was replaced by 100 µL fresh FBS/well for 24 h. Then the culture medium was replaced by 100 µL FBS/well for 24 h. When OD\(_{570}\) reached (14.169) during the titration process. After continuous stirring for 24 h, the suspension aged at standing state for 24 h and washed with DI water until the pH decreased to about 7.0 to remove the impurities. The AA/HAP powder was obtained following the lyophilized of the precipitate for 12 h. The HAP powder without amino acids was synthesized according to the above procedures.

CCK-8 solution was slowly added into each well, and the culture plate was incubated for 4 h. Then the 96-well culture plate was placed on a microplate reader and the optical density (OD) in each well was measured at a wavelength of 450 nm. The relative cell viability (%RCV) of the cells was calculated through the formula:

\[
\text{%RCV} = \frac{OD_t - OD_b}{OD_n - OD_b} \times 100
\]

Where OD\(_t\), OD\(_b\) and OD\(_n\) are the average optical density values of the wells contain experimental materials, the wells without experimental materials (negative control) and the wells without cells (blank control) respectively.
For each concentration of the materials, the OD value of eight wells was measured, and the mean value ± standard deviation was used.

1.4 Preparation of enamel samples and enamel lesions

Sound bovine molars of 2–3 years old cattle were selected in this investigation. After removal of the root and organic contaminants, the molars were cut into 3 mm × 4 mm × 2 mm enamel blocks using sintered diamond saw on a low-speed precision cutting machine. Then the enamel blocks were embedded in PMMA resin with the enamel lip exposed. The enamel surface was polished with water-cooled SiC sandpaper until a mirror-like surface was obtained, and no obvious scratches were observed under a stereomicroscope. The enamel samples were stored in PBS at 4 °C.

The enamel samples were immersed in 3wt% carboxymethyl cellulose gel (pH=4.0) which contains lactic acid (0.1 mol/L) for 24 h to produce acid-etched lesions (the ratio of enamel surface area to solution volume was 2 mm²/mL). After demineralization, the samples were washed with DI water and sonicated for 1 min.

1.5 Remineralization of the enamel in artificial saliva

In order to simulate an oral environment, artificial saliva was prepared by dissolving 1.5 mmol/L CaCl₂, 0.9 mmol/L KH₂PO₄, 130 mmol/L KCl and 20 mmol/L HEPES in DI water (pH was adjusted to 7.0 with 0.1 mol/L KOH), 1.0 mmol/L NaN₃ was added to prevent bacterial growth.

The acid-etched enamel samples were soaked in 60 mL artificial saliva which contained 2.16 g HAP or AA/HAP (the ratio of HAP or AA/HAP to enamel surface is 0.03 g/mm²). Then they were placed on a constant temperature shaker (37 °C) for 12 h. After that, the samples were rinsed with DI water for 20 s and soaked in 60 mL artificial saliva at 37 °C for another 12 h. A total of 21 d were performed in this cycle and the HAP or AA/HAP suspension was replaced every 7 d. Blank group: the enamel samples were soaked in artificial saliva at 37 °C for 21 d (artificial saliva was replaced every 12 h).

1.6 Mechanical property of the enamel samples

The SMH values of the enamel samples were measured by a Vickers microhardness tester (THV-30MD, Lab Testing Technology (Shanghai) Co., Ltd.) under a 9.8 N load for 15 s. The mean values of three points for each sample were recorded at different time (before demineralization, after demineralization, and after remineralization). Before the remineralization, the acid-etched enamel samples were arranged from large to small in order of the SMR values, and the samples were divided into three groups according to the principle of balanced distribution of the SMR values (n=6). The degree of microhardness increase rate (%MHR) after remineralization was calculated through the formula:

\[ \%MHR = \frac{MH_2 - MH_1}{MH_1} \times 100 \]  

where \( MH_1 \) is the mean value of microhardness after demineralization, \( MH_2 \) is the mean value of microhardness after remineralization.

1.7 Characterization

Fourier transform infrared spectroscopy (FT-IR) of the HAP and the AA/HAP was recorded on an FT-IR spectrometer (EQUINOX-55, Germany). Structure and crystallinity analysis were performed on X-ray diffractometer (PANalytical Empyrean, 40 kV, 40 mA, Netherlands). Crystalline size, morphology and interfacial bonding of the HAP and the AA/HAP were characterized by transmission electron microscope (TEM) (JEOL JEM-2100, Japan). High resolution transmission electron microscope (HRTEM) and fast Fourier transform (FTT) results were obtained using the TEM device. The Ca²⁺ concentration in the supernatant of the remineralization solution containing HAP and AA/HAP was measured by inductively coupled plasma-optical emission spectrometry (ICP-OES, USA).

The surface and cross-section morphology of the enamel samples were observed by field emission scanning electron microscope (FESEM) (FEI Nova NanoSEM 450, Netherlands). The samples were sputtered with Au for 250 s before the FESEM characterization. The full area Ca/P molar ratio of the enamel surface was obtained from energy dispersive X-ray spectroscopy (EDX, Oxford Instrument Co, UK.). The crystal type and crystallinity of the enamel surface was performed on X-ray diffractometer (PANalytical Empyrean, 40 kV, 40 mA, Netherlands).

1.8 Statistical analysis

The %RCV of the HAP and the AA/HAP composite were expressed as mean±standard deviation. The SMH data of enamel samples were also expressed as mean±standard deviation. One-way ANOVA was applied for overall comparison of the SMH values after demineralization. The comparisons between the SMH value before and after remineralization for each experimental group were analyzed by paired sample t-test. The value of \( p < 0.05 \) is considered significant.

2 Results and discussion

2.1 Characterization of the HAP and AA/HAP composite

Fig. 1 shows the FT-IR spectra of the HAP and
AA/HAP composite. The bands at 1095/1037/962 and 603/565 cm\(^{-1}\) are ascribed to the stretching vibration and the bending vibration of PO\(_4^{3-}\), respectively. The bands at 3450 and 1635 cm\(^{-1}\) are assigned to the stretching vibrations of \(\text{–OH}\) and the bending mode of adsorbed water. These characteristic bands prove that both of the materials are HAP. The vibration bands of N–H and C=O are included in 3450/1635 and 1200–1000 cm\(^{-1}\), respectively. The intensity of the AA/HAP at 1500–1400 and 1200–1000 cm\(^{-1}\) are strengthened by the symmetric/asymmetric stretches of \(\text{–COO}/\text{NH}_2\cdots\text{H}^+\) and the stretching vibrations of C–O, respectively. These results prove the chemical bonding amino acids contained in AA/HAP.

Fig. 2 presents the XRD patterns of the HAP and AA/HAP. Both of the patterns show the characteristic peaks of HAP indexed as (002), (102), (112), (202), (310), (222), (213), (004) and (304) (JCPDS No.09-0432). The diffraction peaks of both materials at 2\(\theta\) of 30°–34° are significantly broadened, indicating the small crystalline size of the synthesized nanoparticles. Meanwhile, AA/HAP diffraction peaks are lower and broader than HAP. It can be concluded that the AA/HAP has lower crystallinity\(^{[23]}\). The passivated and weak peaks of (002) and (004) planes are attributed to the inhibition effect of the complex amino acids on the crystal growth of c-axis.

The \(\alpha\)-carboxyl and hydroxyl group of Ser could tightly adsorb on (002) and (004) lattice planes of HAP, thus the crystal growth perpendicular to c-axis was inhibited\(^{[22-23]}\). The acidic amino acid Asp has a high affinity on the HAP crystal faces parallel to the c-axis\(^{[24-25]}\). With both side terminal carboxyl groups binding to crystal planes, the preferred crystal growth perpendicular to the (310) plane was reduced significantly. The size of a Gly molecule is about 0.3 nm\(^{[26]}\), while the internal channels in the crystal lattice of HAP which are occupied by OH groups are 0.3–0.35 nm in diameter\(^{[27]}\). Therefore, Gly molecules could enter these channels and simultaneously interact with the crystal lattice around these channels, eventually changed the crystallinity of HAP. It can be concluded that the AA/HAP composite with lower crystallinity would help to increase its solubility in the remineralization.

Fig. 3 reveals the TEM images of the HAP and AA/HAP. The typical TEM image of the HAP (Fig. 3(a1)) shows an enamel prism-like morphology with a length of 50–150 nm and a diameter of 8–12 nm. The interplanar spacing distances of \(d=0.339\) nm (Fig. 3(a2)) are corresponding to the (002) faces of the HAP. This result is consistent with the preferred growth of (002) face shown in the XRD pattern (Fig. 2), and demonstrates that the [001] is the preferred crystallizing direction. The diffraction spots in the FTT image are due to (002) (the brighter spots) and (211) (the darker spots) faces of the HAP, respectively. The TEM image of the AA/HAP (Fig. 3(b1)) shows a prism-like morphology with a length of approximately 100 nm and a diameter of 3–6 nm. This result proves again the inhibition effect of amino acids on the crystal growth along a and b-axis. The diffraction
circles shown in the FTT image of AA/HAP (insert in Fig. 3(b2)) are corresponding to the (002) and (211) faces of AA/HAP, respectively, indicating that the nanoparticles are randomly oriented. The HRTEM image of the AA/HAP proves again that the crystalline of the nanoparticles are significant refinement. The interplanar spacing distances of $d=0.347$ and 0.283 nm are ascribed to the (002) and (211) faces of AA/HAP, respectively. The black spots with a diameter of 2.5–3.5 nm (in the background of Fig. 3(b2)) may be attributed to the carbonization of the amino acids trapped in the crystals of AA/HAP. Both of the XRD and the TEM results illustrate that the AA/HAP has smaller crystalline size and lower crystallinity. These properties could increase the solubility of the AA/HAP in artificial saliva, and may be favorable for the release of Ca$^{2+}$ and PO$_4^{3-}$ ions during the remineralization.

The cytotoxicity of the HAP and AA/HAP composite at various concentrations range from 0.05 g/mL to 0.4 g/mL was evaluated by CCK-8 assay. As shown in Fig. 4, both of the materials present no cytotoxicity within a wide range of concentrations. Nevertheless, %RCV of HAP is far lower than that of AA/HAP. Thus the AA/HAP composite presents better biocompatibility than the HAP.

Fig. 5 shows the Ca$^{2+}$ concentration in the supernatant of remineralization solution containing HAP and AA/HAP.

Once the HAP and AA/HAP were added into artificial saliva, the Ca$^{2+}$ concentrations were both lower than that of artificial saliva (1.5 mmol/L). This result may be attributed to the chelating effect of both materials on Ca$^{2+}$, which reduces the Ca$^{2+}$ concentration of the remineralization solution in a short time. At the same time, the initial value of AA/HAP is lower than that of HAP, which may be due to its higher specific surface area. The Ca$^{2+}$ concentration of the HAP-containing solution gradually decreased to about 0.72 mmol/L in 7 d, while the AA/HAP-containing solution gradually increased to approximately 1.34 mmol/L. This result proves the higher solubility of AA/HAP than HAP.

2.2 In vitro remineralization of acid-etched bovine enamel

After the remineralization, the surface and cross-section morphology of the enamel were measured by FESEM. The EDX, XRD and Vickers microhardness of the enamel surface were characterized, respectively.

2.2.1 Characterization of the surface and cross-section morphology

Fig. 6 presents the surface and cross-section FESEM images of the sound enamel and acid-etched enamel. The sound enamel has a dense surface without obvious gaps (Fig. 6(a1)). Its cross-section is also intact (Fig. 6(a2)). The acid-etched enamel has a scaly-like surface with large gaps between the enamel rods (Fig. 6(b1)), and the enamel prisms can be clearly seen. The cross-section image of the acid-etched enamel (Fig. 6(b2)) presents severe subsurface lesions with a depth up to 30 μm. In the primary hierarchical structure, enamel consists of enamel rod, rod sheath and inter-rod mineral[4]. These rod sheaths are hydrophobic and can protect the enamel rods from acid erosion[5]. The inter-rod mineral has lower crystallinity, and contains a small amount of organic
remnants. Meanwhile, the subsurface enamel prisms contain higher metal ions such as K⁺, Na⁺ and Mg²⁺ than the surface prisms. The potassium, sodium and magnesium phosphates are more vulnerable to acid etching, eventually leading to more serious subsurface lesions.

Fig. 7 shows the FESEM images of the enamel surface and cross-section after remineralization. The blank group also formed a mineral layer on its surface (Fig. 7(a1)). This may attribute to the self-repair ability of the acid-etched enamel in artificial saliva. Despite the relatively intact surface, the blank group still presents obvious defects beneath the surface area. Both of the limited Ca²⁺ ions in artificial saliva and the restricted Ca²⁺ bioavailability of the acid-etched enamel restrained the subsurface remineralization. The surface image of the HAP treated group (Fig. 7(b1)) shows many inter-rod gaps. And only a small amount of minerals adsorb on the surface. This result means that the HAP is mainly utilizing the adsorption properties to fill the cavities of the enamel surface. The cross-section image of the HAP group (Fig. 7(b2)) shows a similar defect to the blank group, but presents deeper unrepaired subsurface lesions. As mentioned above, the crystallinity of the HAP is relatively high, resulting in lower solubility and limited Ca²⁺ and PO₄³⁻ ions concentration in the remineralization. As shown in Fig. 5, the HAP particles may adsorb Ca²⁺ and PO₄³⁻ ions, and re-grow in artificial saliva, therefore, leading to a worse repair effect than the blank group.

The AA/HAP treated group obtained the densest mineral layer on the enamel surface (Fig. 7(c1)). The insert image proves the existence of newly formed nanoparticles. The AA/HAP synthesized in the presence of Gly, Ser and Asp shows poor crystallinity and higher solubility. Therefore, more Ca²⁺ and PO₄³⁻ ions could be provided for surface remineralization of the enamel samples, just as shown in Fig. 5. Meanwhile, the molecules of amino acids could adsorb on the enamel surface and attract Ca²⁺ and PO₄³⁻ ions to sustain the topical supersaturation, ultimately promote the HAP crystal growth. The cross-section image of the AA/HAP group (Fig. 7(c2)) shows an intact morphology with no defect exists. Furthermore, parallel arranged crystals formed in the inter-rod area (red arrows in Fig. 7(c2)). The molecular size of amino acids is very small compared with macromolecular components (such as polypeptides and amelogenin). This feature facilitates the rapid diffusion of amino acids molecules from remineralization solution to deep lesions. Then the amino acids could adsorb on the organic matrix residues and promote the orderly growth of HAP crystals. In summary, if dissolved in solution, amino acids would inhibit the crystal growth of HAP by chelating Ca²⁺ and PO₄³⁻ ions or binding to the nuclei of calcium phosphate. When adsorbed on organic residues, the side functional groups (such as carboxyl and hydroxyl) of amino acids would become nucleation sites. These nucleation sites would regulate the crystal growth of HAP along parallel orientation.

2.2.2 Components analysis

Table 1 reveals the elemental composition results of the enamel surface. The Ca/P molar ratio of sound enamel surface is 1.62, which is lower than that of empirically derived HAP (Ca/P=1.67). This result indicates that enamel is composed of calcium-deficient HAP. Calcium ions in the phosphorus-oxygen tetrahedron in apatite are more susceptible to acid corrosion than phosphate ions. Therefore, the Ca/P molar ratio of the acid-etched enamel decreased significantly to 1.48. After the remineralization, the Ca/P molar ratio of the blank group, HAP group and AA/HAP group increased to 1.58, 1.63 and 1.65, respectively. The Ca/P molar ratio of the AA/HAP group is higher than that of sound enamel, and is very close to 1.67, indicating that the surface of the acid-etched enamel is well repaired. Meanwhile, the higher component of O

| Sample            | Elemental composition/ at% | Ca/P ratio |
|-------------------|---------------------------|------------|
| Polished enamel   | 19.86 17.97 29.04         | 1.62       |
| Acid-etched enamel| 25.95 19.34 28.59         | 1.48       |
| Blank group       | 14.07 17.59 27.85         | 1.58       |
| HAP group         | 19.63 16.05 26.20         | 1.63       |
| AA/HAP group      | 40.61 17.87 29.49         | 1.65       |
in the AA/HAP treated group may attribute to the carboxyl and side functional groups of amino acids.

The XRD patterns of the enamel surface are shown in Fig. 8. All of the patterns show the characteristic peaks of HAP indexed as (002), (211), (112), (202), (222), (213), (004) and (304). The diffraction peak of (310) lattice plane at $2\theta=39.82^\circ$ almost disappear, indicating that the crystals locating in the enamel surface have tiny crystalline size along the crystallographic $a$-axis. The sharp and intense (002) and (004) lattice planes of the sound enamel (Fig. 8(a)) illustrate that the enamel surface is comprised of parallel arranged crystals in the orientation of $c$-axis direction$^{[31]}$. The enamel prisms in the surface and subsurface area are severely corroded, thus the peak intensities of the acid-etched enamel (Fig. 8(b)) decrease greatly. Meanwhile, its XRD pattern shows a broadened and weakened peak at $2\theta=32–35^\circ$. These results demonstrate that the surface prisms of the acid-etched enamel have lost crystal lattice$^{[32]}$. After remineralization, the peak intensities of (002) and (004) faces strengthen in all the three experimental groups. This result proves the restoration of the enamel prisms along $c$-axis direction. For the characteristic peaks at $2\theta=31^\circ–33^\circ$, the AA/HAP group (Fig. 8(e)) shows lower intensity and broadened pattern. This result is consistent with the FESEM image shown in Fig. 7(c1). It proves the abundant regenerated nanoparticles formed on the enamel surface.

2.2.3 Mechanical property

The mechanical property of the enamel represents changes in microstructure. Fig. 9 shows that the SMH values of all the enamel samples decreased significantly after acid-etch. One-way ANOVA shows that there is no significant difference in SMH among the experimental groups after demineralization ($p>0.05$). The SMH of the blank group before and after remineralization are almost identical. Paired sample $t$-test analysis shows that there are significant difference between the SMH before and after remineralization both for the HAP group and the AA/HAP group ($p<0.05$). Nonetheless, the SMH of the HAP treated group is still lower than that of the AA/HAP group, which could also be verified by the poor subsurface remineralization in the cross-section morphology (Fig. 7(b2)). In summary, the %MHR of the blank group, HAP group and AA/HAP group are 5.52%, 35.07% and 148.42%, respectively. However, compared with the SMH of sound enamel, the SMH of AA/HAP group is still very low. Enamel has a complex hierarchical structure reinforced by the precise arrangement of rods and the organic remnants$^{[4]}$. Once eroded into the inner area, this delicate structure would be destroyed and the complete recovery of the SMH could not be realized$^{[33]}$.

3 Conclusions

In this study, amino acids/hydroxyapatite (AA/HAP) composite was synthesized to evaluate its remineralization effect on acid-etched bovine enamel. Compared with the HAP without amino acids, the AA/HAP composite with refined crystalline size and lower crystallinity was obtained under the inhibition effect of Gly, Ser and Asp. Cytotoxicity assay proved better biocompatibility of the AA/HAP composite than the HAP. In artificial saliva, AA/HAP could sustain a calcium supersaturation circumstance and induce restoration of both surface and subsurface lesions in acid-etched bovine enamel. The molecular size of Gly, Ser and Asp are all very small, thus facilitate the rapid diffusion of these molecules into deep lesions and the adsorption on organic matrix residues. The parallel arranged HAP crystals which are beneficial for the recovery of SMH may be promoted by these adsorbed amino acids. The current approach may provide a safe, economical and effective strategy for the treatment of early enamel lesions.

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氨基酸/羟基磷灰石复合材料应用于酸蚀牛牙釉质体外再矿化

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摘 要：早期牙釉质龋病的再矿化在龋齿的预防和修复中起着至关重要的作用。基于牙釉质基质中所含的主要氨基酸，在 10 mmol/L 甘氨酸、10 mmol/L L-丝氨酸和 5 mmol/L L-天冬氨酸的存在下制备了氨基酸/羟基磷灰石(AA/HAP)复合材料，表征了其物理、化学和生物学性质，并评估了其对酸蚀牛牙釉质的再矿化作用。与不含氨基酸的羟基磷灰石(HAP)相比，在氨基酸的抑制作用下，AA/HAP 复合材料具有更低的结晶度和更高的生物相容性。在人工唾液中用 AA/HAP 对酸蚀牛牙釉质进行体外再矿化。再矿化后，分别表征了牛牙釉质样品的表面和横截面形态、成分和力学性能。结果表明，AA/HAP 可以诱导表面和深层牙釉质病变的修复。复合材料中释放的氨基酸可以吸附在有机基质残基上并诱导平行排列的 HAP 晶体的形成，从而使牙釉质表面显微硬度(SMH)得到显著恢复。最后，讨论了 AA/HAP 复合材料对酸蚀牛牙釉质的再矿化机制。

关 键 词：再矿化; 牙釉质; 羟基磷灰石; 氨基酸

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