Micro-/nano-topography of selective laser melting titanium enhances adhesion and proliferation and regulates adhesion-related gene expressions of human gingival fibroblasts and human gingival epithelial cells

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Background: Selective laser melting (SLM) titanium is an ideal option to manufacture customized implants with suitable surface modification to improve its bioactivity. The peri-implant soft tissues form a protective tissue barrier for the underlying osseointegration. Therefore, original microrough SLM surfaces should be treated for favorable attachment of surrounding soft tissues.

Material and methods: In this study, anodic oxidation (AO) was applied on the microrough SLM titanium substrate to form TiO2 nanotube arrays. After that, calcium phosphate (CaP) nanoparticles were embedded into the nanotubes or the interval of nanotubes by electrochemical deposition (AOC). These two samples were compared to untreated (SLM) samples and accepted mechanically polished (MP) SLM titanium samples. Scanning electron microscopy, energy dispersive spectrometry, X-ray diffraction, surface roughness, and water contact angle measurements were used for surface characterization. The primary human gingival epithelial cells (HGECS) and human gingival fibroblasts (HGFs) were cultured for cell assays to determine adhesion, proliferation, and adhesion-related gene expressions.

Results: For HGECS, AOC samples showed significantly higher adhesion, proliferation, and adhesion-related gene expressions than AO and SLM samples (P<0.05) and similar exceptional ability in above aspects to MP samples. At the same time, AOC samples showed the highest adhesion, proliferation, and adhesion-related gene expressions for HGFs (P<0.05).

Conclusion: By comparison between each sample, we could confirm that both anodic oxidation and CaP nanoparticles had improved bioactivity, and their combined utilization may likely be superior to mechanical polishing, which is most commonly used and widely accepted. Our results indicated that creating appropriate micro-/nano-topographies can be an effective method to affect cell behavior and increase the stability of the peri-implant mucosal barrier on SLM titanium surfaces, which contributes to its application in dental and other biomedical implants.

Keywords: additive manufacturing, surface modification, soft tissue barrier, dental implants

Introduction
Titanium implants have been widely used in dentistry for the replacement of missing teeth owing to their predictable clinical performance in the clinic.1 The conventional implants are manufactured in cylindrical or tapered shapes with a predetermined length and diameter to match the requirements of most clinical conditions. However, some complicated and irregular anatomies are unsuitable for conventional implants.2
Customized dental implants are therefore in high demand. Furthermore, customized dental implants can also be used for individual treatment and can shorten treatment time. With the development of additive manufacturing (AM), freeform geometries have been produced precisely and individually based on computer-aided design (CAD) data. AM is convenient for fabricating complex and custom-made structures based on a one-step manufacturing process.

As the latest type of AM, selective laser melting (SLM) is widely used in biomedical implant fabrication. Titanium and its alloys fabricated by SLM show great potential in individualized and customized implant manufacturing due to their biocompatibility and biomechanical properties. Reports have shown evidence of good bone regeneration in SLM titanium specimens after surface modification in vivo and in vitro.

However, it is well-known that a major reason for implant loss is peri-implantitis, as a consequence of the penetration of bacterial plaque into the peri-implant sulcus. Therefore, only good bone regeneration is not enough, the barrier function provided by the peri-implant mucosal tissue is equally important. An implant must interface successfully with two types of cells, namely the epithelial cells and the underlying fibroblasts from the soft tissue for a good barrier function. It has been shown that topographic and physicochemical properties of implant surfaces affect both hard and soft peri-implant tissue responses. There is growing evidence that implant surface characteristics may play a role in the attachment of peri-implant soft tissues.

According to current knowledge, nano-structured surfaces have been speculated to influence cell behavior in a different way compared to conventionally sized surfaces. One of the most diffused technologies able to create a nano-structured surface is anodic oxidation. Reports show that a highly crystalline, thin hydroxyapatite (HA) layer coated on the anodized titanium provides a suitable nano-structure surface for clinical oral implants. Among these reports, Takebe et al discovered that anodized-hydrothermally treated titanium affects the adhesion and regulates the gene expression of fibroblasts and epithelial cells. Another report showed that oxidized surfaces enhanced human gingival fibroblast (HGF) adhesion, proliferation, and extracellular matrix deposition, and this could be due to the different structure at micro- and nano-scale levels. All aforementioned reports used commercial pure titanium with a flat surface. However, the surface of raw SLM specimens has a certain roughness due to its layer-by-layer manufacturing process. Few studies have focused on whether nano-structures on the surface of microscale roughness such as SLM specimens can influence cellular behavior of peri-implant soft tissues.

In this study, we modified the surface of SLM substrate to form nanotube arrays by an anodizing method, and then deposited nanoscale calcium phosphate (CaP) particles, which are the major elements of HA in the anodized nanotubes or the interval of nanotubes, by an electrochemical technique. We analyzed the surface characteristics and intended to evaluate whether micro-/nano-topography of SLM titanium, especially the anodizing method or the nanoscale CaP particles, could affect the response of HGFs and human gingival epithelial cells (HGEcs) in terms of adhesion, proliferation, and adhesion-related gene expressions.

Material and methods
Specimen preparation and treatment
Specimens were designed by SolidWorks® 12.0 (SolidWorks Corp, Concord, MA, USA) and manufactured by an SLM machine (SLM125HL, SLM solutions GmbH, Lubeck, Germany). Raw materials were commercial grade II titanium powders (Western BaoDe, Xian, PR China) with an average particle size of 30 μm. The SLM processing parameters were adjusted to a laser power of 145 W, laser scanning velocity of 355 mm/s, hatch space of 45 μm, layer thickness of 30 μm, and laser spot size of 50 μm. Specimens were produced as titanium discs (10x10x1 mm³) (Figure 1).

These SLM titanium specimens were degreased by a cleaning solution (0.05 mol/L Na₃SiO₃, 0.1 mol/L Na₃PO₄, and 0.2 mol/L Na₂CO₃) for 15 minutes and then sonicated in deionized water for 15 minutes. These SLM titanium discs were termed SLM samples and set as the control group.

Some of these specimens were polished mechanically by P80, P220, P400, P600, P800, and P1000 sandpaper in sequence and were termed mechanically polished (MP) samples.

Some other SLM samples were first sandblasted with 250 μm ZrO₂ particles to remove residual unmelted titanium spheres, and then chemically polished in a solution of 5% hydrofluoric acid (HF) for 1 minute to remove the native oxide layer before anodization. Thus, the specimens were anodized in an electrolyte consisting of 0.5 wt% HF in deionized water with an applied voltage of 20 V for 45 minutes to create TiO₂ nanotube arrays. These were termed AO samples.

Parts of AO samples were then treated by an electrochemical technique in order to embed nanoscale CaP into TiO₂ nanotubes or the interval of nanotubes. An electrochemical workstation (CHI-660D, CH Instruments, Austin, TX, USA) was used for electrochemical deposition. The solution consisted of 0.20 mmol/L of Ca(NO₃)₂ and 0.12 mmol/L of (NH₄)₂HPO₄. AO samples were anchored onto platinum
cathodes and subjected to a constant voltage of −3 V for 10 minutes, with the counter-electrode also being platinum. The reaction was carried out in a 50°C water bath and the specimens were termed AOC samples.

Afterward, all treated specimens were thoroughly cleaned ultrasonically in deionized water and autoclave sterilized before cell culture studies. There were four groups termed SLM, MP, AO, and AOC.

Surface characterization
The surface topography and chemistry were analyzed by field-emission scanning electron microscopy ([FE-SEM] S-4800, Hitachi, Tokyo, Japan) and X-ray energy dispersive spectrometry (EDS). The crystalline phases of the specimens were investigated by X-ray diffraction ([XRD] X’Pert-PRO, PANalytical, Almelo, Holland). A 3D surface topography analyzer (BMT EXPERT, Lorsch, Germany) was used to scan the specimens’ surface roughness. Water contact angles were measured by the sessile-drop method on an optical contact angle measuring device (OCA40 Micro, Dataphysics, Stuttgart, Germany).

Preparation and identification of HGFs and HGECs
This study was ethically approved by the Ethical Review Committee, Guanghua School of Stomatology, Hospital of Stomatology, Institute of Stomatological Research, Sun Yat-Sen University (no ERC-2016–36). Written informed consent was provided by patients and then sampling was performed. Samples of gingival tissue were harvested from healthy donors who underwent surgery to extract the lower third molar at the Guanghua School of Stomatology, Hospital of Stomatology, Sun Yat-Sen University. Collected tissues were washed three times in PBS (Gibco, Thermo Fisher Scientific, Waltham, MA, USA) supplemented with antibiotics (100 U/mL penicillin and 100 mg/mL streptomycin; Gibco).

For culturing HGFs, the collected tissues were cut into small pieces (1×1×1 mm³). These pieces were digested in 0.1% type I collagenase (Sigma-Aldrich Co., St Louis, MO, USA) at 37°C for 15 minutes and centrifuged at 1,000 rpm for 5 minutes. After centrifugation, the pieces were harvested, supplemented with complete L-DMEM (Gibco) containing antibiotics and 10% FBS (Gibco), and cultured at 37°C in a 5% CO₂ humidified atmosphere. Five days later, the cultured HGFs were collected and fed. The medium was changed every 2 days. Cultures between the fifth and eighth passage were used in the present experiments.

For culturing HGECs, the collected tissues were cut into strips (1×5 mm²) and immersed overnight at 4°C in 0.25% dispase (Sigma-Aldrich Co.). Then, the epithelium was separated with forceps and cut into small pieces (1×1 mm²), which were digested with 0.25% trypsin (Gibco) for 8 minutes at 37°C and centrifuged at 1,000 rpm for 5 minutes. The cells were collected and fed with complete L-DMEM containing antibiotics and 10% FBS. The medium was changed every 2 days. Cultures between the second and fourth passage were used in the present experiments.

An inverted microscope (Axio Observer Z1, Carl Zeiss Meditec AG, Jena, Germany) was used for the morphological identification of cells. An immunofluorescent stain assay was also performed. HGFs and HGECs were respectively seeded onto confocal dishes at a density of 5×10³ cells·mL⁻¹ and cultured for 24 hours. Then, the tissues were fixed in 4% paraformaldehyde (Biosharp, Anhui, PR China) for 10 minutes and 0.2% Triton X-100 (MP Biomedicals, Strasbourg, France) was added for 5 minutes. The cells were
then blocked in 4% BSA (Beyotime, Shanghai, PR China) for 1 hour and stained with pan-cytokeratin (PCK) and Vimentin antibodies (1:100, Boster, Wuhan, PR China) overnight. The secondary antibodies were Alexa Fluor488 Goat anti-Mouse IgG (Invitrogen, Thermo Fisher Scientific). The nucleus was stained with DAPI (Beyotime, Shanghai, PR China). Immunofluorescent staining was performed with a confocal laser scanning microscope (LSM780, Carl Zeiss Meditec AG).

Cell proliferation evaluation by CCK-8
HGFs and HGECs were respectively seeded onto specimens in 48-well plates at a density of $1 \times 10^4$ cells$\cdot$L$^{-1}$ and cultured with L-DMEM containing 10% FBS for 1, 3, 5, and 7 days. At the prescribed time points, the specimens were taken out and washed three times with PBS. The specimens were then transferred to new 48-well plates. The cells were assessed using a cell counting kit-8 assay (CCK-8) (Djindo, Kumamoto, Japan). Twenty-five microliters of 10% CCK-8 fluid with medium was added to each well, and the cells were incubated for 2 hours. Optical absorbance (OD) of the CCK-8 fluid was then measured at 450 nm.

Cell adhesion and morphology evaluation by SEM
HGFs and HGECs were respectively incubated on the specimen surfaces in 48-well plates at a density of $1 \times 10^4$ cells$\cdot$L$^{-1}$ for 24 hours and fixed in 2.5% glutaraldehyde solution for 12 hours. After dehydration in graded ethanol (ie, 50%, 75%, 90%, 95%, and 100%) and subsequent drying, the specimens were observed under field-emission scanning electron microscope.

ELISA of hEGF protein secreted by HGECs
HGECs were incubated on the specimen surfaces in 48-well plates at a density of $5 \times 10^4$ cells$\cdot$L$^{-1}$ for 7 days. Then, the cells were washed in PBS three times. RIPA Lysis Buffer (Cwbio, Beijing, PR China) was used to collect the hEGF protein as per the manufacturer’s instructions. An ELISA kit (Cloud-clone, Wuhan, PR China) was used for hEGF detection.

ELISA of type I collagen synthesis by HGFs
HGFs were incubated on the specimen surfaces in 48-well plates at a density of $5 \times 10^4$ cells$\cdot$L$^{-1}$ for 7 days. The medium was collected and an ELISA kit (Chondrex, Redmond, WA, USA) was used to detect type I collagen synthesis.

RT-PCR evaluation of gene expression by HGECs and HGFs
HGECs and HGFs were respectively incubated on the specimen surfaces in 48-well plates at a density of $5 \times 10^4$ cells$\cdot$L$^{-1}$ for 7 days. Then, the total RNA was isolated from the cells using TRIZol reagent (Invitrogen, Thermo Fisher Scientific) based on the single-step method. First-strand cDNA was synthesized from the total RNA using the PrimeScriptRT MasterMix (TaKaRa, Kusatsu, Japan), followed by qRT-PCR carried out on a Light Cycler 480 (Hoffman-La Roche Ltd., Basel, Switzerland) with SYBR Premix Ex Taq II (TaKaRa) and specific oligonucleotide primers. The primers for HGECs were used as the template for PCR with primers specific to integrin-$\alpha_3$, integrin-$\beta_3$, laminin-$\alpha_2$, laminin-$\beta_3$, and laminin-$\gamma_2$; the primers for HGFs were used as the template for PCR with primers specific to integrin-$\alpha_5$, integrin-$\beta_3$, fibronectin, and vinculin. The mRNA levels for cells were normalized for GAPDH mRNA. The primers used are shown in Table 1.

Table 1 Primers used for RT-PCR

<table>
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<tr>
<th>Gene</th>
<th>Forward primer sequence (5′-3′)</th>
<th>Reverse primer sequence (3′-5′)</th>
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<tr>
<td>integrin-$\alpha_3$</td>
<td>ATCCCAAGCGGAATAGCC</td>
<td>TCTCATGCGAGCCCTCAT</td>
</tr>
<tr>
<td>integrin-$\beta_3$</td>
<td>ACTTGTAGGCCACGTCTCG</td>
<td>GGCCTCCGCTTACCTTG</td>
</tr>
<tr>
<td>Laminin-$\alpha_2$</td>
<td>CCACCCAGAGGAAAGAC</td>
<td>CTCCTCATACAGGAACCA</td>
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<tr>
<td>Laminin-$\beta_3$</td>
<td>AAAGCCGCGGATGAGT</td>
<td>TGTGGACAGAGTCGTAC</td>
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<tr>
<td>Laminin-$\gamma_2$</td>
<td>GCCTGGGTATTGAGCAG</td>
<td>AGACAACATGATGGGATT</td>
</tr>
<tr>
<td>integrin-$\alpha_5$</td>
<td>CAGGGTACAGAAGCTGGAGT</td>
<td>AAGTCAAGAATGGCAGAGT</td>
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<td>integrin-$\beta_3$</td>
<td>ATTAGCAAATCTCCCTC</td>
<td>CACTGGTCAGACATCAT</td>
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<tr>
<td>Fibronectin</td>
<td>ACGGCCATATGGGAAACT</td>
<td>CACAGCGAGCATTAAGT</td>
</tr>
<tr>
<td>Vinculin</td>
<td>TGATGTCATTGCCCTGC</td>
<td>AGACCTTTGCAAAACCTCCGACTA</td>
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<tr>
<td>GAPDH</td>
<td>GAGGCTGAAGGGCCTCTAG</td>
<td>GTACCCCAAGATGCGCTTGA</td>
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</table>

Abbreviations: integrin-$\alpha_3$, ITGA6; integrin-$\beta_3$, ITGB4; laminin-$\alpha_2$, LAMA3; laminin-$\beta_3$, LAMB3; laminin-$\gamma_2$, LAMC2; integrin-$\alpha_5$, ITGA3; integrin-$\beta_3$, ITGB1; fibronectin, FN1; vinculin, VCL

Statistical analysis
The OD values from CCK-8, concentration of hEGF and type I collagen from ELISA, and relative quantity of...
Results

Surface characterization

The surface topographies of the four groups are shown in Figure 2. SLM samples showed flat surfaces. MP samples showed fine surface features. AO samples showed arrangement of ordered nanotubular structures. AOC samples showed amorphous nanoparticles embedded in the nanotubes or the interval of nanotubes.

Cell proliferation evaluation by CCK-8

Figure 7A shows results of the HGECs proliferation of each group. On the first, third, fifth, and seventh days, the MP and AOC groups showed more proliferation than the SLM and AO groups (P < 0.05). However, there were no significantly different proliferation rates among the MP, AOC, and AO groups.

Abbreviations: SEM, scanning electron microscopy; SLM, selective laser melting; MP, mechanically polished; AO, anodic oxidation; AOC, anodic oxidation composited with electrochemical deposition.

Figure 2 SEM images showing the surface topography of samples (scale bar of A, C, E = 2 µm, scale bar of D, F = 400 nm).

Notes: (A) SLM sample with unmelted titanium particles and spheres on the rough waving surface; (B) MP sample with flat surface; (C, D) different magnifications of AO sample showing the ordered nanotubular structures on the surfaces; (E, F) different magnifications of AOC sample showing the amorphous nanoparticles embedded in the nanotubes or the interval of nanotubes.

Identification of HGFs and HGECs

The results of the morphological observation of HGECs showed a typical "paving stone like" keratinization (Figure 6). The immunofluorescent staining assay of HGECs showed positive expression of PCK, while no expression of vimentin was observed. On the other hand, the results of HGFs showed positive expression of vimentin, while no expression of PCK was observed.

Water contact angles of SLM, MP, and AO samples were 73.9°, 76.3°, and 40.7°, respectively, while AOC samples showed favorable hydrophilicity with a relatively lower water contact angle of approximately 18.3° (Figure 5).

Abbreviations: SLM, selective laser melting; MP, mechanically polished; AO, anodic oxidation; AOC, anodic oxidation composited with electrochemical deposition.
higher rates of cell proliferation between SLM and AO groups ($P>0.05$), nor between MP and AOC groups.

Figure 7B shows the results of the HGFs’ proliferation of each group. The surface-treated MP, AO, and AOC groups showed significantly higher rates of cell proliferation than SLM on the first, third, fifth, and seventh days ($P<0.05$). Moreover, AO and AOC groups showed more proliferation than MP at each time point ($P<0.05$), and the AOC group showed the most proliferation ($P<0.05$).

### Cell adhesion and morphology evaluation by SEM

The HGECs converged and showed typical “paving stone like” keratinization in each group (Figure 8). On the surface of MP, the HGECs were flatter and extended in the direction of the polishing lines. Although the HGECs were less stretched on the surface of AO and AOC, we could see multiple contact points with tiny protrusions extending from the lamellipodia and wrapping around the nanotubes.

The HGFs converged as platy on each group (Figure 9). The HGFs were not only more stretched but also showed extended lamellipodia anchored to the surface of AO and AOC.

### ELISA of hEGF protein secreted by HGECs

We used ELISA to determine the influence of different surface treatments on the expression of hEGF. Our data showed that the SLM and AO groups had significantly lower expression than the MP and AOC groups after 7 days’ inoculation of HGECs ($P<0.05$; Figure 10). There was no evidence of differences between SLM and AO groups. The MP and AOC groups showed the same results ($P>0.05$).

### ELISA of type I collagen synthesis by HGFs

Type I collagen synthesized by HGFs was measured by ELISA and the data (Figure 11) showed increasing amounts of collagen synthesized by HGFs in the SLM, MP, AO, and AOC groups, in that order. The data of each group showed a statistically significant difference ($P<0.05$).

### Table 2 X-ray energy dispersive spectrometry (EDS) analysis of the elemental composition (wt%)

<table>
<thead>
<tr>
<th>Samples</th>
<th>Titanium</th>
<th>Oxygen</th>
<th>Calcium</th>
<th>Phosphorus</th>
<th>Fluorine</th>
</tr>
</thead>
<tbody>
<tr>
<td>SLM</td>
<td>100</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>MP</td>
<td>100</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>AO</td>
<td>67.74</td>
<td>29.64</td>
<td>–</td>
<td>2.62</td>
<td>–</td>
</tr>
<tr>
<td>AOC</td>
<td>49.20</td>
<td>48.00</td>
<td>0.22</td>
<td>0.14</td>
<td>2.44</td>
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</tbody>
</table>

**Abbreviations:** SLM, selective laser melting; AO, anodic oxidation; AOC, anodic oxidation composited with electrochemical deposition.

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**Figure 3 EDS spectra of the samples.**

**Notes:** (A) SLM and (B) MP samples were composited with 100% titanium. (C) Some oxygen and a little fluorine besides titanium on the surface of AO samples. (D) Titanium, oxygen, calcium, phosphorus, and fluorine were present on the surface of AOC samples.

**Abbreviations:** EDS, X-ray energy dispersive spectrometry; SLM, selective laser melting; MP, mechanically polished; AO, anodic oxidation; AOC, anodic oxidation composited with electrochemical deposition.
RT-PCR evaluation of gene expression by HGECs and HGFs

The relative gene expression levels of adhesion molecules (integrin-α, integrin-β, laminin-α, laminin-β, and laminin-γ) in HGECs after 7 days of incubation on different samples showed the following results: AOC group > MP group > AO group > SLM group (Figure 12A).

After culturing for 7 days, the relative expression levels of fibronectin and vinculin in HGFs on different samples followed the order of AOC group > AO group > SLM group > MP group. For integrin-α and integrin-β, the expression levels followed the order of AOC group > AO group > MP group > SLM group. The results are shown in Figure 12B.

Discussion

There is an increasing demand for customized implants given their convenient and individual application. This has led to a rapid development of SLM technology. Titanium is one of the most commonly used materials for SLM implants. The interface between implant and soft tissue has received increasing attention owing to its influence on successful implantation rate. Surface modifications are used on titanium to enhance this soft tissue barrier. In this study, we tested in vitro whether micro-/nano-topography of SLM titanium surface might positively affect the interaction between the dental implant and the surrounding soft tissue, by evaluating the adhesion, proliferation, and adhesion-related gene expressions of HGECs and HGFs. We tried to establish a new composition treatment technology for improved attachment to soft tissues. We chose the SLM surface as a control group to evaluate the function of anodization and electrochemical deposition. Furthermore, the MP group was selected, because the polished surface represents the first and more extensively tested implant surface, and long-term success rate of rehabilitation realized with these surfaces continues to be the reference point for all implant treatments.

Surface roughness usually plays an important role in the adhesion and proliferation of cells. In most research studies, the HGECs seem to prefer smooth surfaces. However, a microrough surface is recommended, which may be the best for HGFs. In our study, we produced the original SLM specimen with a 7.57 µm in Ra value. Except for the layer-by-layer SLM manufacturing process, the roughness is also influenced by the partially melted particles. Therefore, we used sandblasting technology and chemical polishing first, to decrease the roughness during the process of surface modification. Lastly, the AO and AOC specimens measured 2.15 µm in terms of Ra value and likely match the requirement for micro-roughness.

When we talk of surface features in the nanometer scale, we refer to topographical characteristics with dimensions less than 100 µm. In our study, we observed nanotubes with dimensions of approximately 70–90 µm. Nanotopography can regulate cell behavior. For example, nanotopography can promote neuronal differentiation of human induced pluripotent stem cells. Carbon nanotubes can induce fibrogenesis on nanostructured substrates. Kato et al reported that collagen fibers can integrate into the nanotubes resulting in enhanced cellular adhesion and collagen synthesis. Yang et al delineated the relationships between focal adhesions, nucleus and cell function and highlighted that nanotopography could regulate cell phenotype and function by modulating nuclear deformation. Furthermore, Takebe et al considered that a thin HA layer and anodic titanium oxide film with nanotopographic structure can regulate the gene expression of fibroblasts and epithelial cells. In this study, we adopted an electrochemical approach to deposit the nano-CaP into TiO2 nanotubes or the interval of nano-tubes, expecting a better function than the HA layer. Our SEM results showed lamellipodia anchoring on the AO and AOC surfaces of HGECs and HGFs. This may enhance the

Table 3 Surface roughness of the samples

<table>
<thead>
<tr>
<th>Samples</th>
<th>SLM</th>
<th>MP</th>
<th>AO</th>
<th>AOC</th>
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<tr>
<td>Ra (µm)</td>
<td>7.57±0.32</td>
<td>0.39±0.01</td>
<td>2.15±0.04</td>
<td>2.15±0.06</td>
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<td>Rq (µm)</td>
<td>11.20±0.66</td>
<td>0.50±0.01</td>
<td>2.72±0.03</td>
<td>2.71±0.05</td>
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</table>

Abbreviations: SLM, selective laser melting; MP, mechanically polished; AO, anodic oxidation; AOC, anodic oxidation composited with electrochemical deposition.
adhesion and affect the proliferation and adhesion-related gene expressions as we expected.

In order to evaluate the biological features of these surfaces, we chose primary cultures of HGECs and HGFs as the cellular model, which are the main resident cell populations in the peri-implant connective attachment.

The data showed that after 7 days of incubation, HGECs on the AOC surface upregulated gene expressions of integrin-α6β4 and laminin-α3β3γ2 (laminin-5), the major components of the hemidesmosome and basement membrane, respectively, as compared to the SLM and AO surfaces. The MP surface is always considered best for HGECs but showed similar

Figure 5 Water contact angles of the samples.

Note: The (A) AO samples and (B) AOC samples showed better hydrophilia than (C) SLM samples and (D) MP samples.

Abbreviations: AO, anodic oxidation; AOC, anodic oxidation composited with electrochemical deposition; SLM, selective laser melting; MP, mechanically polished.

Figure 6 (A) Morphological observation of HGECs (scale bar = 200 μm); (B) Immunofluorescent staining showing PCK-positive HGECs (scale bar = 50 μm); (C) Immunofluorescent staining showing vimentin-negative HGECs (scale bar = 50 μm); (D) Morphological observation of HGFs (scale bar = 200 μm); (E) Immunofluorescent staining showing PKC-negative HGFs (scale bar = 50 μm); (F) Immunofluorescent staining showing vimentin-positive HGFs (scale bar = 50 μm).

Abbreviations: HGECs, human gingival epithelial cells; PCK, pan-cytokeratin; HGFs, human gingival fibroblasts.
Topography of SLM titanium affects gingival fibroblasts and epithelial cells

Figure 7 Cell proliferation of HGECs (A) and HGFs (B) on the surfaces of each group after 1, 3, 5 and 7 days.
Notes: *P<0.05, **P<0.01, ***P<0.001.
Abbreviations: HGECs, human gingival epithelial cells; HGFs, human gingival fibroblasts; SLM, selective laser melting; MP, mechanically polished; AO, anodic oxidation; AOC, anodic oxidation composited with electrochemical deposition.

expressions as the AOC surface. Because hemidesmosome and basement membrane ensure efficient attachment of epithelial cells to the implant surface, these results predict that HGECs on the AOC surface could achieve better attachment to the implant than the SLM and AO surfaces, and as good as the MP surface.40,41 In addition, similar results were seen in proliferation and hEGF protein secretion activity of HGECs. Thus, implants with AOC surfaces may achieve...

Figure 8 Representative SEM images of HGECs’ adhesion to the surfaces of each group after 24 hours (scale bar of A–C, E=5 µm, scale bar of D, F=600 nm).
Notes: (A) SLM; (B) MP; (C, D) AO; (E, F) AOC.
Abbreviations: SEM, scanning electron microscopy; HGECs, human gingival epithelial cells; SLM, selective laser melting; MP, mechanically polished; AO, anodic oxidation; AOC, anodic oxidation composited with electrochemical deposition.
faster epithelial sealing around the transmucosal component than that with SLM or AO surfaces. The CaP nanoparticles could be predicted to play a key role in enhancing adhesion, proliferation, and regulation of adhesion-related gene expressions in that the AOC samples showed stronger ability than AO samples and SLM samples. At the same time, the AOC surface was equally favorable for HGECs compared to the standard established MP surface.

The HGFs spread well and grew rapidly on all surfaces. AOC showed the best ability for cell proliferation, while SLM surface fared the worst at all time points. The type I collagen synthesis of HGFs showed the same results.

**Figure 9** Representative SEM images of HGFs' adhesion on the surfaces of each group after 24 hours (scale bar of A–C, E = 5 µm, scale bar of D, F = 600 nm).

**Notes:** (A) SLM; (B) MP; (C, D) AO; (E, F) AOC.

**Abbreviations:** SEM, scanning electron microscopy; HGFs, human gingival fibroblasts; SLM, selective laser melting; MP, mechanically polished; AO, anodic oxidation; AOC, anodic oxidation composited with electrochemical deposition.

**Figure 10** Concentration of hEGF in HGECs after 7 days.

**Note:** *P < 0.05, **P < 0.01, ***P < 0.001.

**Abbreviations:** HGECs, human gingival epithelial cells; SLM, selective laser melting; MP, mechanically polished; AO, anodic oxidation; AOC, anodic oxidation composited with electrochemical deposition.

**Figure 11** Concentration of type I collagen in HGFs after 7 days.

**Note:** *P < 0.05, **P < 0.01, ***P < 0.001.

**Abbreviations:** HGFs, human gingival fibroblasts; SLM, selective laser melting; MP, mechanically polished; AO, anodic oxidation; AOC, anodic oxidation composited with electrochemical deposition.
Type I collagen is the major component of the extracellular matrix of connective tissues and plays a central role in the architecture of the peri-implant tissue, which is formed by both epithelial and connective structures. In particular, the connective attachment underlying the junctional epithelium comprises scar-tissue-like tissue with a high density of high turn-over fibroblasts (approximately one third in volume) and collagen fibers (approximately two thirds in volume) disposed in close contact with the titanium surface of the implant neck. The expression of integrin-α3 and integrin-β1 was consistent with these results. However, these cells expressed higher amounts of fibronectin and vinculin on the SLM than MP surface. It is known that fibroblasts adhere to implant surfaces by focal adhesion. Vinculin is a focal adhesion linker protein, and increased vinculin expression is associated with enhanced adhesion strength. A possible explanation is that the rougher SLM surface was easier to adhere to than the smooth MP surface, although the AOC surface showed the best adhesion.

Taken together, with respect to proliferation and adhesion, AOC was the best for HGFs and HGECs. Not only anodic oxidation but also CaP nanoparticles come into play when comparing with the original SLM surface. When compared to the classic MP surface, AOC was advantageous for HGFs and not bad for HGECs. Thus, implants with AOC surface may result in an early and effective formation of soft tissue barrier, which would be beneficial for the long-term maintenance of osseointegration. However, our results cannot be extrapolated to a clinical scenario yet because of the lack of animal experimentation, which forms the basis of future research.

**Conclusion**

In this study, anodic oxidation technology was successfully used to form nanotubes on the rough SLM surface to create a favorable micro-/nano-topography. Furthermore, CaP nanoparticles were deposited into the nanotubes or the interval of nanotubes with an electrochemical technique. This surface modification highly enhanced the adhesion, proliferation, and regulation of adhesion-related gene expressions of HGECs and HGFs, as compared to the original SLM surface. This treated surface can compare favorably with accepted MP surface for HGECs and is even better than MP surface for HGFs. These results indicate that forming appropriate micro-/nano-topographies can be an effective method to affect cell behavior and increase the stability of the peri-implant environment.
mucosal barrier which contributes to its application in dental and other biomedical implants. Further studies, however, are needed to investigate the in vivo and clinical implications of such encouraging preliminary results.

Acknowledgments
The authors would like to thank research staff members of the Department of Oral Implantology, Guanghua School of Stomatology and the Institute of Advanced Technology, Chinese Academy of Science for their support. This work is funded by National Natural Science Foundation of China (no 81501600) and Science & Technology Plan Project of Guangzhou (201710010193).

Disclosure
The authors report no conflicts of interest in this work.

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