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The effects of hierarchical micro/nanosurfaces decorated with TiO, nanotubes on the bioactivity of titanium implants in vitro and in vivo

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Abstract: In the present work, a hierarchical hybrid micro/nanostructured titanium surface was obtained by sandblasting with large grit and acid etching (SLA), and nanotubes of different diameters (30 nm, 50 nm, and 80 nm) were superimposed by anodization. The effect of each SLA-treated surface decorated with nanotubes (SLA + 30 nm, SLA + 50 nm, and SLA + 80 nm) on osteogenesis was studied in vitro and in vivo. The human MG63 osteosarcoma cell line was used for cytocompatibility evaluation, which showed that cell adhesion and proliferation were dramatically enhanced on SLA + 30 nm. In comparison with cells grown on the other tested surfaces, those grown on SLA + 80 nm showed an enhanced expression of osteogenesis-related genes. Cell spread was also enhanced on SLA + 80 nm. A canine model was used for in vivo evaluation of bone bonding. Histological examination demonstrated that new bone was formed more rapidly on SLA-treated surfaces with nanotubes (especially SLA + 80 nm) than on those without nanotubes. All of these results indicate that SLA + 80 nm is favorable for promoting the activity of osteoblasts and early bone bonding.

Keywords: nanotopography, osseointegration, dental and orthopedic implant, titanium

Introduction

It is well known that the surface characteristics of biomedical implants influence their interactions with cells and the extracellular matrix (ECM) and are therefore key factors in determining the osseointegration of such implants. In the tissue engineering and biomaterials' field, the interactions between cells and surface microtopographies have been extensively studied, and certain microscale surface topographies and related technologies have already been adopted for commercial implant products. It has been revealed that microtopographies can improve bone-to-implant contact through mechanical interlocking and the promotion of osteoblast functions.² Some microscale surface topographies have already been used on commercial implant products, and good clinical effects have been obtained, including from sandblasting with large grit and acid etching (SLA)-treated titanium implants.³⁻⁵ Nevertheless, though they positively promote osteoblast differentiation, microscale surfaces often inhibit osteoblast proliferation, resulting in a smaller accumulation of bone mass compared with that yielded by a smooth surface.6-11

With recent progress in surface treatment technologies, researchers have the tools to achieve a variety of nanotopographies. The interactions between cells and nanotopographies are of increasing interest, as nanostructures may be more efficient in promoting desirable cell functions. 12 Among the different approaches for producing nanotopographies, titania nanotubes have attracted much attention since they can be fabricated easily

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with precisely controlled diameters and lengths by tailoring the process parameters. ^{13,14} Such surfaces can almost ideally be used as nanoscale spacing models for size-dependent cellular responses. Some studies have reported that titania nanotubes with suitable dimensions can improve bone cell functions, ^{15,16} although some controversy remains. ^{17–21} These nanotubes can also serve as carriers for drugs, such as growth factors, antibacterial agents, and genes, and show good prospects for bone–implant applications. ^{22,23}

Several studies have shown that surface structures composed of micro- and nanoscale components may provide suitable surface topographies for promoting desirable cell functions such as osteogenesis, because these materials can mimic the structure of the natural ECM.^{22,24} Kubo et al showed that a micropit-and-nanonodule hybrid titania topography on a titanium surface enhanced osteoblast attachment, spreading, adhesion, proliferation, and differentiation in vitro,²⁵ while Gao et al fabricated a micro/nanostructured porous surface on titanium and observed increased formation of hydroxyapatite in simulated body fluids and faster adsorption of bovine serum albumin (BSA).¹³

Adding nanotubes to the surfaces of implant materials is expected to alter and likely improve their bioavailability, biocompatibility, and bioreactivity. In particular, the coexistence of micro- and nanotopographies may confer advantageous properties and functionalities to biomaterials. However, the controllable hybrid creation of nano and microstructures has been challenging. Furthermore, the biological synergies of micro and nanostructures, for example, whether further enhancement of cellular behaviors and responses can be obtained by the addition of nanotubes to microstructures without sacrificing their established advantages, and which size range of nanotube microstructures is most effective in increasing bone-to-implant contact in vivo, are largely unknown.

Therefore, in this study, hybrid micro/nanotextured surface topographies were produced on titanium by SLA to produce micropits and subsequent anodization to superimpose nanotubular layers with different nanotube diameters (30 nm, 50 nm, and 80 nm, producing the hybrid surfaces SLA + 30 nm, SLA + 50 nm, and SLA + 80 nm, respectively). We hypothesized that these topographies will enhance bioactivities such as osteogenesis and early bone bonding. First, the bioactivity of the materials was evaluated in vitro using MG63 osteoblasts. The expression of the osteogenesis-related genes runt-related transcription factor 2 (RUNX2), alkaline phosphatase (ALP), and osteocalcin (OCN) was detected using real-time polymerase chain reaction (RT-PCR).

Second, the effect of each surface on osseointegration was determined in vivo using a canine model.

Materials and methods

Specimen preparation

The titanium specimens used for in vitro and in vivo tests were divided into four groups: SLA (control group without nanotubes), SLA + 30 nm, SLA + 50 nm, and SLA + 80 nm.

Before synthesis, titanium discs 15 mm in diameter and 1.5 mm thick were cut from commercially obtained pure titanium plates for cell tests, and titanium cylinder implants (schematic diagram shown in Figure S1) were made for us by a company according to our order (grade 2, Tengxing Metal Materials, Shenzhen, People's Republic of China). All titanium surfaces were ground with silicon carbide sandpapers of 280, 360, 400, 600, 800, and 1,000 grits in series. The titanium specimens were then immersed in 10 mL of acetone (Fuyu, Tianjin, People's Republic of China) at room temperature, ultrasonicated for 20 minutes, ultrasonicated for a further 20 minutes in 10 mL of absolute alcohol (Fuyu), and then rinsed with deionized water. Subsequently, the specimens were dried at room temperature.

The SLA surfaces were prepared according to a previously described method.26 All titanium specimens were sandblasted by 120 µm Al₂O₂ particles from a distance of 50 mm at an angle of 90°. The air pressure used for blasting was set to 0.45 MPa, and the procedure was conducted for 30 seconds. Subsequently, the specimens were etched using a mixture of 18% (v/v) HCl and 49% (v/v) H₂SO₄ at 60°C for 30 minutes and then ultrasonically cleaned in ddH₂O for 15 minutes. Finally, all of the specimens were air dried at room temperature. The nanotubes were superimposed on the surface of the SLA using 0.5% (w/v) hydrofluoric acid (Fuyu) at 10 V, 15 V, and 20 V for 30 minutes to obtain nanotubular layers comprising nanotubes of different diameters to form the surfaces SLA + 30 nm, SLA + 50 nm, and SLA + 80 nm.²⁷ A platinum electrode served as the cathode. After being anodized, the specimens were rinsed with distilled water several times, air dried at 80°C, and heat treated at 450°C for 2 hours to transform the as-anodized amorphous TiO, nanotubes into a crystalline phase. All samples were collected, rinsed with deionized water, and sterilized by autoclaving before use.

Surface characterization

Following previously described methods, ^{26,28} the morphology of each sample was analyzed using field emission scanning electron microscopy (FE-SEM; FEI-Quanta 400, Hillsboro, OR, USA). Water contact angle measurements were carried

out using the sessile drop technique (OCA 40 Micro; Data-Physics, Filderstadt, Germany) at room temperature. Surface roughness (Ra) was measured by a profilometer (Wyko NT9300; Veeco, NY, USA).

Measurement of protein adsorption

According to a previously described method, 28 the abilities of the SLA + 30 nm, SLA + 50 nm, and SLA + 80 nm surfaces to adsorb a model protein (BSA, fraction V; Sigma-Aldrich Co., St Louis, MO, USA) were compared with that of the SLA-treated surface without nanotubes. First, 300 μ L of protein solution (1 mg of protein/mL of saline) was spread over each specimen using a pipette. After incubation for 1 hour, 2 hours, and 6 hours under sterile humidified conditions at 37°C, all nonadherent protein was removed, and the remaining solution was mixed with microbicinchoninic acid (Sigma-Aldrich Co.) at 37°C for 30 minutes. The protein content of the solution was quantified using a microplate reader (Model 680; Bio-Rad Laboratories Inc., Hercules, CA, USA) at 562 nm.

Cell culture

A human MG63 osteosarcoma cell line was cultured in Dulbecco's Modified Eagle's Medium (Hyclone, Thermo Fisher Scientific, Waltham, MA, USA) supplemented with 10% (v/v) fetal bovine serum (Hyclone) and antibiotics (penicillin [100 U/mL] and streptomycin [100 mg/mL]) at 37°C in a humidified atmosphere containing 5% CO₂. At 80% confluence, the cells were detached using 0.25% (w/v) trypsin in 1 mM ethylenediamine tetraacetic acid 4Na and seeded onto the four different surfaces in 24-well plates at a density of 2.4×10⁴ cells cm⁻². The culture medium was replaced on alternate days. This study was reviewed and approved by the Ethics Committee of Guangdong Provincial Stomatological Hospital, People's Republic of China.

Cell adhesion assay

Cell attachment was initially evaluated by measuring the amount of cells attached to the titanium substrates after 30 minutes, 1 hour, and 4 hours of incubation, as described previously. 26,28 At each prescribed time point, nonadherent cells were removed by gentle rinsing with phosphate-buffered saline (PBS). Adherent cells were fixed with 4% (w/v) paraformaldehyde for 30 minutes and then stained with fluorescent Hoechst 33342 dye for 5 minutes. Cell adhesion was evaluated by counting the number of stained nuclei on each sheet in the fluorescent microscopic images ($100\times$ magnification, counts performed over an area of $1,800\times1,350~\mu\text{m}^2$). Values representing the mean and the standard error of the

number of attached cells were calculated from ten different random fields in each disc using Image-Pro Plus 6.0 software (Media Cybernetics, Rockville, MD, USA; n=3, a total of 30 fields of view for each group).

Immunofluorescent staining

Cytoskeletal changes in the MG63 cells were observed using an immunofluorescent staining technique. After incubation of the cells for 1 hour, 3 hours, and 6 hours, the culture medium was gently removed from the wells. The cells were washed with PBS, fixed with 4% (w/v) paraformaldehyde for 20 minutes, and then washed several times with PBS. The fixed cells were treated with 0.1% (v/v) TritonTM X-100 in PBS and then blocked with 1.0% (w/v) BSA in PBS. The actin cytoskeleton was stained with phalloidin conjugated to Alexa Fluor® 488 dye (Thermo Fisher Scientific), and the nucleus was stained with Hoechst 33342 dye (Thermo Fisher Scientific). The stained cells were observed under a fluorescence microscope (IX51; Olympus Corporation, Tokyo, Japan).

Cell morphology assay

FE-SEM was used to analyze the cell morphology 3 hours and 6 hours after the MG63 cells had been seeded in the four different groups. The culture medium with unattached cells was carefully removed from the wells by a pipette, and the cells grown on the SLA, SLA + 30 nm, SLA + 50 nm, and SLA + 80 nm surfaces were then rinsed two times with PBS. Cells were fixed with 2.5% (v/v) glutaraldehyde in PBS for 1 hour and washed three times with PBS for 5 minutes at room temperature. Each sample then underwent graded dehydration with 30%, 50%, 70%, 90%, and 100% pure ethyl alcohol for 10 minutes at room temperature, after which the samples were dried for analysis by FE-SEM. ^{26,28}

Cell proliferation assay

Cell proliferation was determined by measuring cell density on culture days 1, 3, 5, and 7 using tetrazolium salt-based colorimetry (MTS; Promega Corporation, Fitchburg, WI, USA). 26,28 At each prescribed time point, the specimens were gently rinsed three times with PBS and transferred to a new 24-well plate. Next, $500 \, \mu L$ of Dulbecco's Modified Eagle's Medium was added to each well and incubated with $100 \, \mu L$ of MTS reagent at 37° C for 4 hours. The amount of formazan product was measured using a microplate reader at $490 \, \text{nm}$.

Osteogenesis-related genes' expression

The expression of osteogenesis-related genes was evaluated using RT-PCR as previously described.²² The MG63 cells

were seeded with 2×10⁴ cells/disc and cultured for 7 days and 14 days. The total RNA was isolated using TRIzol® reagent (Thermo Fisher Scientific). One microgram of RNA from each sample was reverse transcribed into complementary DNA using the PrimeScriptTM RT reagent kit (TaKaRa Bio, Otsu, Japan). The expression levels of osteogenesis-related genes including *RUNX2*, *ALP*, and *OCN* were quantified using an iQTM5 Multicolor RT-PCR Detection System (Bio-Rad Laboratories Inc.) with SYBR Premix ExTM Taq II (TaKaRa Bio). Data analysis was carried out using iQTM5 Optical System Software Version 2.0 (Bio-Rad Laboratories Inc.). The relative expression levels for each gene of interest were normalized to that of the housekeeping gene (GAPDH). The primers are listed in Table S1.

Animals and surgery

Animals

Twenty-four healthy male beagle dogs, ~15 months old and weighing 16–17 kg, were used in this study. All of the animals were housed in a temperature-controlled room with a 12 hours alternating light–dark cycle and were given water and food throughout the study. All animals were acclimatized for at least 2 weeks before the study began. The experimental protocol in this study was reviewed and approved by the Animal Care and Use Committees of Guangdong Provincial Stomatological Hospital, People's Republic of China.

Surgical procedure

The animals were randomly divided into four groups (n=6 in each group). In each beagle dog, eight implants of one type (SLA, SLA + 30 nm, SLA + 50 nm, or SLA + 80 nm) were inserted into two tibias, and each tibia received four implants. Therefore, a total of 192 implants were inserted into the tibias of 24 beagle dogs (Figure S2). All surgical procedures were performed under general anesthesia. Su Mian Xin 2 was used as a preanesthetic (composition: 2,4-dimethylaniline thiazole, ethylenediamine tetraacetic acid, dihydroetorphine hydrochloride, and haloperidol; 0.1 mL/kg; Animal Husbandry Research Institute, Jilin, People's Republic of China) through intramuscular administration. General anesthesia was then obtained after an intramuscular injection of 3% (w/v) pentobarbital sodium (1 mL/kg, Sigma-Aldrich Co.). Surgery was performed under sterile conditions. After hair shaving, skin exposure, and antiseptic cleaning with iodine solution at the surgical and surrounding areas, a 5 cm incision at the skin level was made. Then, a flap was retracted to expose the tibia diaphysis. Using a series of implant drills (a 2.3 mm round bur, a 2.5 mm pilot drill, a 2.8-3.2 mm reamer drill, and a 3.3 mm profile drill), four identical holes were produced from

proximal to distal along the central region of the bone for implantation and were drilled at a low speed (800 rpm) under profuse cooling normal saline (0.9% [w/v] NaCl) irrigation. Four implants were inserted by press-fit. The periosteum and skin of the wound were sutured in two layers. After surgery, the animals were allowed to move freely and received antibiotics for 3 days to prevent postsurgical infection. After a healing period of 2 weeks or 4 weeks, the animals were sacrificed by overdose of 3% (w/v) sodium pentobarbital.

Histological preparation

The tibias containing implants were harvested at weeks 2 and 4 of the healing process and fixed in 10% (w/v) buffered formalin at 4° C for 2 weeks. All of the specimens were dehydrated in an ascending series of alcohol rinses and embedded in light-curing epoxy resin (Technovit 7200VLC; Heraeus Kulzer, Wehrheim, Germany) without decalcification. The specimens were cut along the long axis, and thus every section had four chambers that were used for tests. The specimens were ground to a thickness of 30 μ m with a grinding system (Exakt Apparatebau, Norderstedt, Germany). The sections were stained with toluidine blue and observed by light microscopy ($40\times$ magnification).

Histomorphometry

To evaluate the response of the bone surrounding the implants, histological and histomorphometrical analyses were performed using a light microscope (BX41; Olympus Corporation), and image analysis software (Image-Pro Plus 6.0) was used for the histomorphometrical evaluation, following previously described methods.^{29–31} Figure S3 shows a schematic representation of the analyses, which included evaluation of the following variables:

- 1. Bone–implant contact (BIC; %) = (sum of the length of BIC)/(circumference of the implant chamber region) $\times 100$, where the BIC was defined as the interface where the bone tissue was located within 20 μ m of the implant surface without any intervention of soft tissue;
- 2. Bone area (BA; %) = (sum of the BA in the implant chamber region)/(area of the implant chamber region) $\times 100$.

Statistical analysis

The data were analyzed using SPSS 13.0 software (SPSS Inc., Chicago, IL, USA). Q–Q plots were used to test the distribution of data. A one-way analysis of variance followed by a Student–Newman–Keuls post hoc test was used to determine the level of significance. Factorial analysis was used to evaluate the effects of group and time. Values of P < 0.05 were considered to be significant, and values of P < 0.01 were considered to be highly significant.

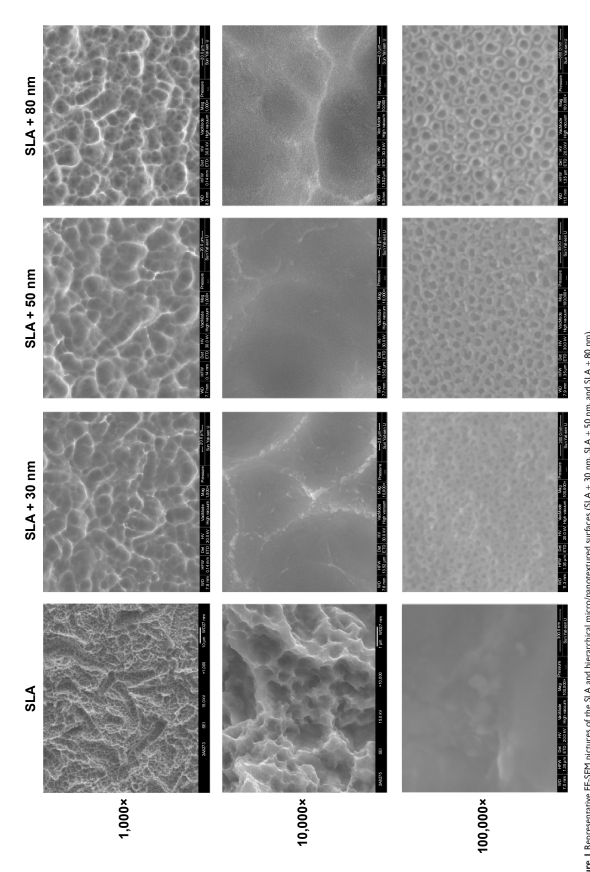


Figure I Representative FE-SEM pictures of the SLA and hierarchical micro/nanotextured surfaces (SLA + 30 nm, SLA + 50 nm, and SLA + 80 nm).

Notes: The nanotubes were found to have 30 nm, 50 nm, and 80 nm outer diameters. The sharp edges can be smoothed. Nanotubes of different sizes were found to have 30 nm, 50 nm, and 80 nm outer diameters. The sharp edges can be smoothed. Nanotubes of different sizes were found to have 30 nm, 50 nm, and 80 nm outer diameters. uniformly.

Abbreviations: FE-SEM, field emission scanning electron microscopy; SLA, sandblasting with large grit and acid etching.

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Results

Surface characterization of different groups

Multilevel pores characterized by 10–30 μm pits and 1–3 μm micropits were initially formed on the specimen surfaces by SLA. The surface structure consisted of relatively sharp edges (Figure 1). Next, the nanotubular layers were fabricated on the microstructured surface. After anodization at 10 V, 15 V, or 20 V and annealing at 450°C, the morphology of the resulting TiO₂ nanotube array thin films was characterized using FE-SEM to ascertain the nanotube dimensions (Figure 1). The nanotubes produced at 10 V, 15 V, and 20 V were found to have 30 nm, 50 nm, and 80 nm outer diameters, respectively. The sharp edges were smoothed, and the micropits became blunt after adding the nanotubular surfaces. Nanotubes of different sizes formed on the microstructured surface and were distributed relatively uniformly.

The Ra of the different groups was assayed, and the results are displayed in Figure 2. The Ra values of the SLA, SLA + 30 nm, SLA + 50 nm, and SLA + 80 nm groups were 2.06 \pm 0.13 μ m, 2.87 \pm 0.21 μ m, 2.74 \pm 0.20 μ m, and 2.44 \pm 0.31 μ m, respectively. There were no statistically significant differences in Ra between the SLA + 50 nm and SLA + 80 nm groups, but these groups had significantly higher Ra values than those of the other two groups, and the SLA + 80 nm surface was significantly rougher than that of the SLA-treated surface without nanotubes.

The contact angle measurements are shown in Figure 3. The average contact angle of the SLA surface was ~11°. The

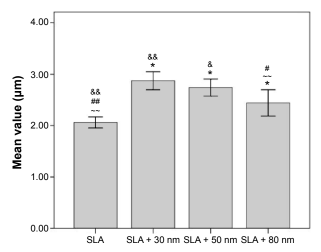


Figure 2 Comparisons of the roughness parameter (Ra) of different surfaces. **Notes:** There were no statistically significant differences in the roughness parameters between the SLA + 50 nm and SLA + 30 nm groups, but they were significantly higher than the other two groups, and the SLA + 80 nm surface was significantly more rougher than the SLA group. Ra: description of height variation; $^*P<0.05$ compared with SLA; $^*P<0.01$; $^*P<0.05$ compared with SLA + 50 nm; $^*P<0.01$; $^*P<0.05$ compared with SLA + 80 nm; and $^{88}P<0.01$.

other three groups showed superhydrophilicity at a water contact angle of 0° .

Protein adsorption

The protein adsorptive rates of the SLA, SLA + 30 nm, SLA + 50 nm, and SLA + 80 nm surfaces over different time periods were analyzed using a bicinchoninic acid assay, and the results are shown in Figure 4. The SLA + 30 nm group had the highest protein adsorption rate compared with those of the other groups, regardless of incubation time. After 1 hour of incubation, the protein adsorptive rate of the SLA + 30 nm group reached 18%, and after 6 hours of incubation, it reached ~38%. Among the other three groups, more protein was adsorbed on the SLA + 50 nm surface than on the SLA or SLA + 80 nm surfaces. The SLA group had the lowest protein adsorption rate at different incubation times, only reaching 2% after incubation for 1 hour and 15% after incubation for 6 hours.

Cell adhesion

Initial cell adhesion on the SLA, SLA + 30 nm, SLA + 50 nm, and SLA + 80 nm surfaces was estimated by counting the number of MG63 cell nuclei stained with Hoechst dye as shown in Figure 5. The results were calculated using Image-Pro Plus 6.0 software and are displayed as histograms in Figure S4. At each time interval adopted in this study, the numbers of adherent cells on the three micro/nanotextured surfaces were dramatically larger than that on the SLA surface (P<0.05). The cell number on the surface of the SLA + 30 nm group was significantly higher than those on the other two groups (P<0.01). For each incubation time, the surfaces of the SLA group had the lowest number of cells among groups.

Immunofluorescent staining

MG63 cell adhesion and spreading on the SLA, SLA + 30 nm, SLA + 50 nm, and SLA + 80 nm surfaces were further investigated by immunofluorescent staining. Fluorescence images of the MG63 cells adhered to the four groups of substrates are shown in Figure 6. After 1 hour of incubation, most of the cells maintained a round shape. Actin around the blue nuclei was stained green and was not yet arranged on the different surfaces. At 3 hours after seeding, the cells had begun to spread and formed an irregular shape. Particularly on the SLA + 30 nm surface, the actin of the cells was organized along the spreading direction and had formed a number of filopodia. Interestingly, most of the cells on the surface of the SLA + 80 nm group maintained a round or oval shape, but the cells also stretched out many filopodia. In accordance with the FE-SEM results, after 6 hours of incubation, cells

Abbreviation: SLA, sandblasting with large grit and acid etching.

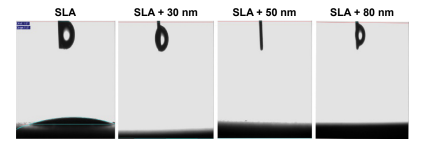


Figure 3 Water contact angles on the SLA, SLA + 30 nm, SLA + 50 nm, and SLA + 80 nm surfaces.

Notes: The average contact angle of the SLA surface was ~11°. The other three groups showed superhydrophilicity at a water contact angle of 0°.

Abbreviation: SLA, sandblasting with large grit and acid etching.

on the three micro/nanotextured surfaces had stretched out many filopodia and some lamellipodia. In particular, the cellular cytoskeleton of cells on the SLA + 80 nm surface achieved a more extensive and homogeneous arrangement compared with those of the other three groups. The shapes of cells grown on the SLA + 30 nm and SLA + 80 nm surfaces were clearly different. We observed that those grown on the SLA + 80 nm surface were the most irregularly shaped, while those grown on the SLA + 30 nm surfaces had relatively regular shapes.

Cellular morphology on different surfaces

As shown in Figures 7 and 8, the morphology of the MG63 cells obviously changed over time after seeding on different

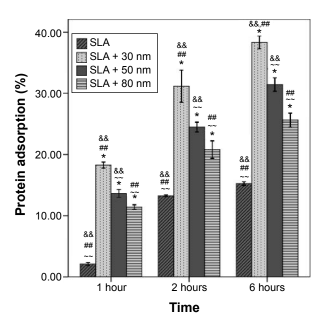


Figure 4 Protein adsorptive rates of the SLA, SLA + 30 nm, SLA + 50 nm, and SLA + 80 nm surfaces assessed using a model protein.

Notes: The SLA + 30 nm group had the highest protein adsorption rate compared with the other groups regardless of incubation time. Among the other three groups, more protein was adsorbed on the surface of the SLA + 50 nm group relative to the other two groups. The SLA group had the lowest protein adsorption rate at different incubation times. *P<0.01 compared with SLA + 30 nm; *P<0.01 compared with SLA + 30 nm; Abbreviation: SLA, sandblasting with large grit and acid etching.

surfaces. After 3 hours of incubation, MG63 cells attached to the 10– $30~\mu m$ pits generated by sandblasting, especially on the surfaces of the SLA group. On the other three micro/nanotextured surfaces, the cells had more extended filopodia from their leading edges. Six hours after seeding, the cells on the micro/nanotextured surfaces still showed more extended filopodia and some lamellipodia, especially in the SLA + 80~nm group, in which the cells formed extraordinarily irregular shapes. In contrast, the MG63 cells on the SLA surface showed few extended filopodia grasping the edges of the pits. Thus, we concluded that the shape of the cells on the SLA surface was decided by the shape of the pits.

Cell proliferation

Cell proliferation was evaluated by MTS assay as shown in Figure 9. Cell proliferation increased over time in all four groups. Cells grown on the SLA + 30 nm surface had significantly higher proliferative activity than did those on the other three surfaces, regardless of incubation time (P<0.05). No statistically significant differences were found among the other three groups.

Gene expression analysis

Gene expression on the titania surfaces was quantified using RT-PCR as shown in Figures 10–12. Generally speaking, ALP expression was detected at a high level at week 1 and was subsequently greatly decreased at week 2 (P<0.01). The expression level of ALP in cells grown on the SLA + 80 nm surface was higher than that in cells grown on other surfaces at all time points (P<0.01). The lowest level of ALP expression was observed in the SLA + 30 nm group, but ALP expression was not significantly different between cells grown on the SLA + 30 nm and SLA + 50 nm surfaces at week 1. RUNX2 expression was upregulated at week 1 and dropped to a low level at week 2 (P<0.01). The group showing the highest level of RUNX2 expression was SLA + 50 nm (P<0.01), followed by the SLA + 80 nm group at two time points, while SLA + 30 nm and SLA were

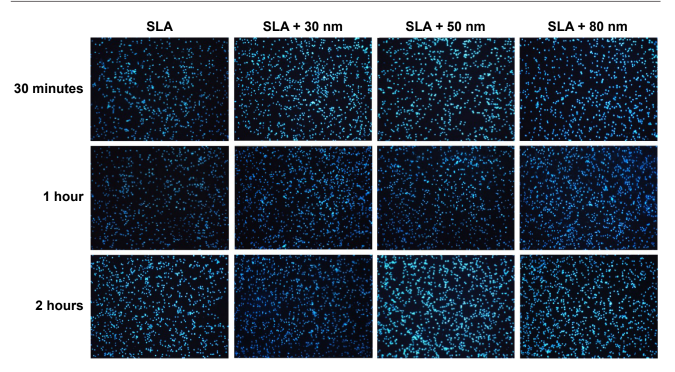


Figure 5 The number of MG63 cell nuclei stained with Hoechst dye on the SLA, SLA + 30 nm, SLA + 50 nm, and SLA + 80 nm surfaces at 30 minutes, 1 hour, and 2 hours (100×).

Notes: At each time interval adopted in this study, the adherent cell numbers on the three micro/nanotextured surfaces are dramatically larger than that on the SLA surface (P<0.05). The cell number on the surface of the SLA + 30 nm group is significantly higher than those on the other two groups (P<0.01). The surfaces of the SLA group have the lowest number of cells.

Abbreviation: SLA, sandblasting with large grit and acid etching.

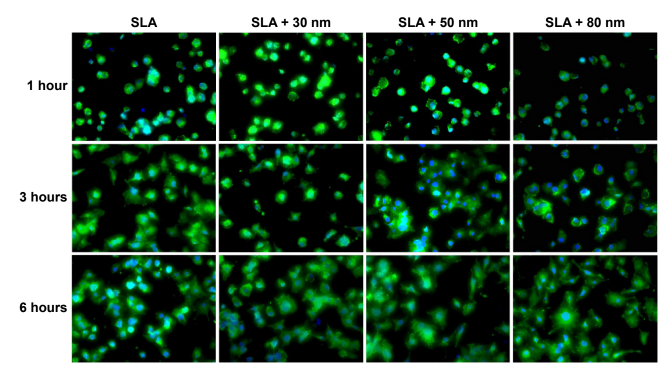
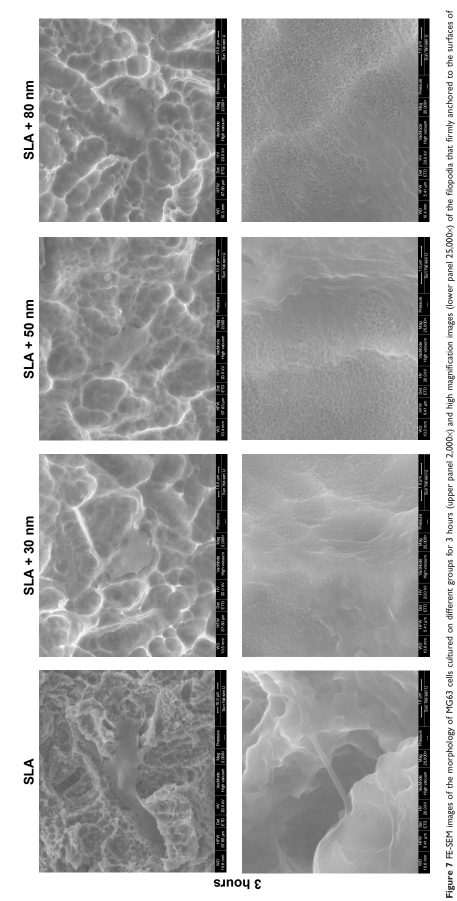


Figure 6 Representative confocal microscopic images of cells stained with rhodamine phalloidin for actin filaments (green) and Hoechst (blue) at different times (400×).

Notes: From small (30 nm) to larger (80 nm) diameter nanotubes, the shape of the cells had changed. After 6 hours of incubation, we observed that the MG63 cells on the SLA + 80 nm surface were most irregular, while the cells on the SLA + 30 nm group surfaces looked relatively regular.

Abbreviation: SLA, sandblasting with large grit and acid etching.

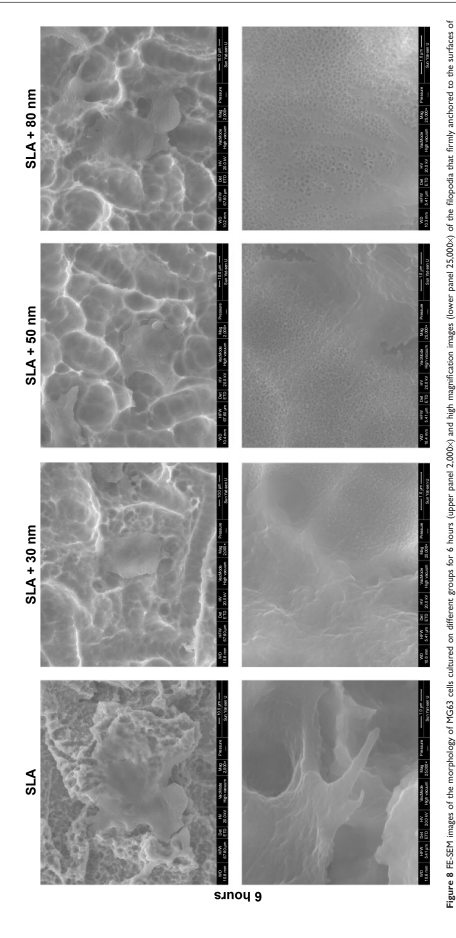


different groups.

Notes: MG63 cells attached to the 10–30 µm pits generated by sandblasting. On the three micro/nanotextured surfaces, the cells show more filopodia extending from the leading edges with increasing nanotube diameters compared

with the SLA group. **Abbreviations:** FE-SEM, field emission scanning electron microscopy; SLA, sandblasting with large grit and acid etching.

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Notes: The cells on the micro/nanotextured surfaces still had more filopodia and some lamellipodia stretching out, especially in the SLA + 80 nm group in which the cells formed extraordinarily irregular shapes. In contrast, the MG63 cells on the SLA surface had few filopodia stretched out and grasping the edges of the pits.

Abbreviations: FE-SEM, field emission scanning electron microscopy; SLA, sandblasting with large grit and acid etching. different groups.

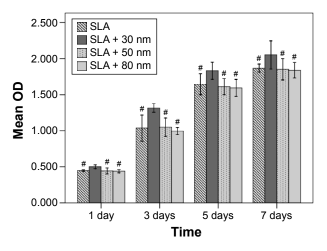


Figure 9 The proliferation of MG63 cells seeded onto SLA, SLA \pm 30 nm, SLA \pm 50 nm, and SLA \pm 80 nm surfaces as measured by the MTS assay.

Notes: The SLA + 30 nm group had significantly higher proliferative activity than those on the other three surfaces regardless of incubation time (P<0.05). There were no statistically significant differences among the other three groups at the time points used in this study. $^{\#}P$ <0.05 compared with the SLA + 30 nm.

Abbreviations: SLA, sandblasting with large grit and acid etching; OD, optical density.

the groups showing the lowest levels of RUNX2 expression at weeks 1 and 2, respectively. For OCN, a low level of expression was observed in all groups at week 1, followed by upregulation during week 2 (P<0.01). The groups with the highest and lowest expression levels of OCN, respectively, were the SLA + 80 nm and SLA + 30 nm groups at two time points (P<0.01).

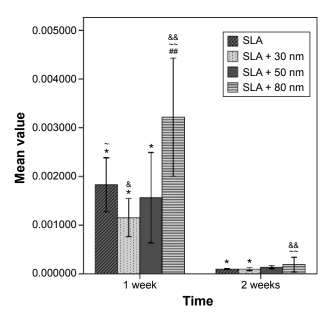


Figure 10 Comparison of ALP mRNA expression of different types of cells (n=3, $\overline{X} \pm S$).

Notes: ^{-}P <0.05 and ^{-}P <0.01 compared with the SLA + 30 nm group; $^{\#}P$ <0.01 compared with the SLA + 50 nm group; ^{8}P <0.05 and ^{88}P <0.01 compared with the SLA group; and $^{*}P$ <0.01 compared with the SLA + 80 nm group.

Abbreviations: ALP, alkaline phosphatase; SLA, sandblasting with large grit and acid etching.

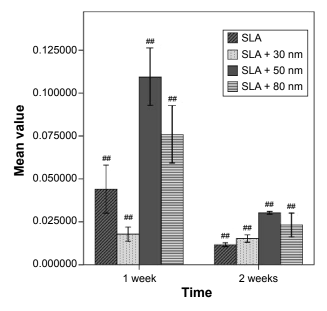


Figure 11 Comparison of RUNX2 mRNA expression levels of different types of cells (n=3, $\overline{X} \pm S$).

Note: ##A significant difference in comparison with all other groups (P<0.01). **Abbreviations:** RUNX2, runt-related transcription factor 2; SLA, sandblasting with large grit and acid etching.

Histological observations/ histomorphometrical analysis

Postoperative healing was uneventful in all dogs. No complications, such as allergic reactions, abscesses, and infections, were observed throughout the whole study period.

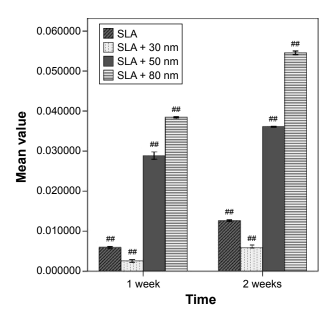


Figure 12 Comparison of the OCN mRNA expression level of different types of cells (n=3, $\overline{X} \pm 5$).

Note: ##A significant difference in comparison with all other groups (P<0.01). **Abbreviations:** OCN, osteocalcin; SLA, sandblasting with large grit and acid etching.

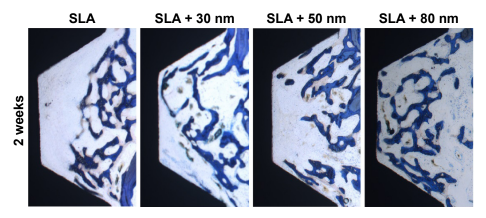


Figure 13 Optical micrographs obtained at 40× original magnification of the four different groups after 2 weeks of implantation time.

Notes: The healing chambers were partially filled with woven bone, especially in the SLA + 80 nm group, which had more new woven bone that was observed to be in close contact with the implant surface and was filling the healing chambers. In the SLA group, a small amount of new bone was observed to be in contact with the implant surface and was filling the healing chambers; toluidine blue stain.

Abbreviation: SLA, sandblasting with large grit and acid etching.

Two weeks and 4 weeks after the implantation, all implants in the control and test groups were histologically in direct contact with the surrounding bone, with no signs of inflammation at the bone-implant interface (Figures 13 and 14). In general, the healing chambers were partially filled with woven bone at 2 weeks in all groups. The SLA + 80 nm group had more new woven bone than the other groups, and this new bone was observed to be filling the healing chambers and in close contact with the implant surface. In the SLA group, a small amount of new bone was observed to be in contact with the implant surface and was filling the healing chambers. At 4 weeks after implantation, we observed more newly formed bone extending from the surface of the implant into the roof of the chamber, and some was in direct contact with the implant surface. Distinct demarcation lines were clearly observed between darker stained new bone and lighter stained old bone, particularly in the SLA + 80 nm group. This finding indicated that the new bone in the chambers had undergone remodeling and had become mature lamel-

At 2 weeks, the proportion of direct BIC with the SLA implants was 12.4%±0.008%. In the implants with 30 nm, 50 nm, and 80 nm TiO, nanotube surfaces, the bone was more continuous along the implant surfaces, and the BIC values were 14.4%±0.008%, 18.3%±0.012%, and 23.3%±0.014%, respectively. The SLA + 80 nm group had the highest BIC value, and this was significantly higher than those of the other three groups (P<0.01). Group SLA + 50 nm had a higher BIC value than that of SLA + 30 nm, while the SLA group had the lowest level. The BIC values of the SLA, SLA + 30 nm, SLA + 50 nm, and SLA + 80 nm groups were 24.1%±0.007%, 32.1%±0.010%, 39.4%±0.010%, and 48.8%±0.012%, respectively, at 4 weeks after implantation. The SLA + 80 nm group still had the highest BIC level (P < 0.01), followed by the SLA + 50 nm group, and the SLA remained the lowest group at 4 weeks after implantation.

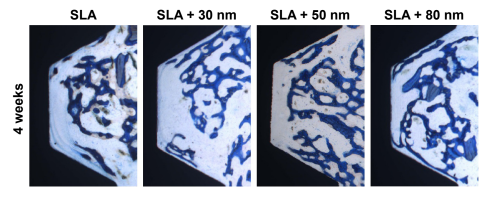


Figure 14 The osseointegration of the different groups at 4 weeks (40×).

Notes: More newly formed bone was extending from the surface of the implant into the roof of the chamber, and some was in direct contact with the implant surface. Distinct demarcation lines were clearly observed between the darker stained new bone and the lighter stained old bone, particularly in the SLA + 80 nm group. **Abbreviation:** SLA, sandblasting with large grit and acid etching.

The mean BAs of the SLA, SLA+30 nm, SLA+50 nm, and SLA+80 nm groups were 17.3% \pm 0.023%, 22.5% \pm 0.023%, 25.2% \pm 0.020%, and 31.7% \pm 0.042%, respectively, at 2 weeks. At 4 weeks after implantation, mean BAs had increased in all groups compared with baseline values (P<0.01). Specifically, the BAs of the SLA, SLA+30 nm, SLA+50 nm, and SLA+80 nm groups were 37.4% \pm 0.03%, 42.9% \pm 0.028%, 45.1% \pm 0.036%, and 53.1% \pm 0.023%, respectively. At each time interval adopted in this study, the mean BAs of the three micro/nanotextured surfaces were dramatically larger than that on the SLA surface (P<0.01), and the SLA+80 nm group always had the highest mean BA (P<0.05). The SLA group always had the lowest mean BA at each time point.

Thus, at weeks 2 and 4, significantly higher new bone growth (as measured by BIC and BA) was observed with the TiO₂ nanotube-coated implants, especially those coated with 80 nm TiO₂ nanotubes (Figures S5 and S6), while the SLA group always had the lowest BIC and BA.

Discussion

In modern dental and orthopedic implant materials, microroughness, microirregularity, and microtopography have become standard surface features required for eminent mechanical fixation and clinical performance. ^{32,33} SLA treatment is considered to be a reliable procedure for achieving a modified titanium surface with suitable topography and roughness for use in implants. It is believed that these microscale surface topographies are beneficial for the mechanical fixation of the implant by increasing the contact area with bone tissue and the in-growth of new bone. However, the early osseointegration of the implants and cell behaviors, such as proliferation, intracellular total protein synthesis, and ALP activity, as well as ECM deposition and mineralization, are obviously reduced on these rough surfaces. ⁴⁻⁸

Nanoscale surface morphological modification has been applied to titanium-based materials. 34–37 This surface possesses unique properties that influence molecular and cellular activities and alter the process of osseointegration by direct and indirect mechanisms. 1,13–15 Several previous studies have demonstrated the beneficial effects of TiO₂ nanotubes on osteoblast differentiation and bone formation around implants in vivo and in vitro. 16–18,21 In this study, we first obtained a classical microscale surface by SLA treatment, and then nanotubes with different diameters (30 nm, 50 nm, and 80 nm) were superimposed on the micron-scale surface via simple anodization. We aimed to achieve a

hybrid surface possessing the advantages of both micro- and nanostructured surfaces.

First, the protein adsorptive rates for each of the tested surfaces (SLA, SLA + 30 nm, SLA + 50 nm, and SLA + 80 nm) were analyzed because proteins are suggested to play vital roles in cell-biomaterial interactions. Our results showed that the SLA + 30 nm group had a higher protein adsorption rate than those of the SLA + 50 nm, SLA + 80 nm, and SLA groups, regardless of the duration of incubation. It is speculated that this higher protein adsorption is due largely to a higher specific surface area. Furthermore, among the various sizes of TiO, nanotubes, the smaller dimension nanotubes of 30 nm diameter and to some extent the 50 nm diameter showed higher protein adsorption rates when compared with nanotubes of 80 nm diameter. The reason for the greatly reduced protein aggregate adsorption observed on the larger diameter nanotubes of 80 nm diameter is most likely because 30 nm-sized protein aggregates are too small to anchor onto 80 nm diameter nanotubes, which have much larger open pore spaces, because the protein aggregates initially attach only to the available surfaces of the top portion of the nanotube walls.19

The phenomenon of initial protein adsorption is very important in understanding why MG63 cells developed more irregular shapes when grown on the SLA + 80 nm nanotube surface than when grown on SLA-treated surfaces with smaller sized or no nanotubes. ECM proteins are required for cells to adhere to surfaces and spread out. During incubation with a micro- or nanostructured surface, proteins in the culture media (derived from the serum) adsorb to the surface and act as a preliminary ECM, which can help cell adhesion considerably at the earliest stages of culture before cells begin secreting ECM. MG63 cells cultured on 30 nm TiO₂ nanotubes can more easily attach to the surface because of the high population of ECM proteins deposited across the specimen surface.¹⁹ However, MG63 cells cultured on SLA + 80 nm would probably have to search more widely for areas containing protein aggregates to establish initial contact due to larger spacing among protein aggregates resulting from the large pore sizes of the nanostructure. This larger spacing among aggregates also means that MG63 cells have to extend their filopodia and lamellipodia excessively and move to find proteins, thus becoming more mobile and forming more irregular shapes.

The cell adhesion and proliferation aspects were further studied to elucidate the optimum nanotube size for positively influencing the behavior of MG63 cells grown on the micro-nano-hybrid titanium surface. It is notable that MG63

cells adhered to the smallest tested nanotubes (SLA + 30 nm surface) more efficiently in the first 2 hours than after that time, probably because of the high protein population initially adsorbing to the surface. Of course, this finding also suggests that the numbers of adhered MG63 cells initially adhering onto the experimental substrata decreased almost inversely with the pore size of the TiO, nanotubes. 19 Surfaces with constructed microtopographies are known to promote osteoblast differentiation but reduce osteoblast proliferation.^{8,38} Park et al demonstrated that the adhesion, proliferation, migration, and differentiation of mesenchymal stem cells (MSCs) were maximally induced on 15 nm nanotubes but were prevented on 100 nm nanotubes, which induced cell death. 16-18 However, some uncertainty about the broad applicability of these results persists. 19-21 Our study revealed that the addition of small diameter nanotubes to a microtopographic surface (SLA + 30 nm) led to enhanced osteoblast proliferation, and these results are in agreement with the findings of Oh et al.¹⁹ In contrast, on larger diameter nanotubes (SLA + 80 nm), the proliferation of osteoblasts was found to be unaffected. This may be due to the synergistic effects of the micro and nanotopographies. Kubo et al have suggested that micro- and nanotopographies may play a synergistic role in promoting osteoblast proliferation.25

Although the interaction between cells and biomaterials is observed to be regulated by nanoscale features, the optimal size range of nanotubes on microtopographic surfaces is still poorly understood. In this study, the expression of osteogenesis-related genes (OCN, RUNX2, and ALP) by osteoblasts was studied by quantitative RT-PCR analysis after 7 days and 14 days of incubation. Our results show that the SLA + 80 nm surface generally elicited the highest osteogenic gene expression compared with the other sizes of nanotubes. It is well known that various types of physical stresses from the substrate morphology and topography can improve osteoblast differentiation.^{39,40} Immunofluorescent and FE-SEM images of MG63 cells on larger sizes of TiO, nanotubes (SLA + 80 nm) showed the formation of a higher number of filopodia, lamellipodia, and cellular extensions compared with those of cells grown on surfaces coated with smaller diameter or no nanotubes. It is most probable that this elongated morphology caused higher levels of cytoskeletal tension and stress for the MG63 cells cultured on the SLA + 80 nm surface compared with those of cells grown on a surface coated with smaller diameter nanotubes. It has been reported that an extracellular stimulus can pass through the cytoskeletal components to the nucleoskeleton, resulting in alterations to the structures of the interphase chromosomes positioned inside the nucleus and changes in gene expression.^{41,42} Our data show a scenario similar to those described by other studies relating to the modulation of the extrusion of cellular extensions and cell differentiation by material surface properties.

To further support the RT-PCR results described above, a canine model was used to perform an in vivo evaluation to determine the optimum diameter of TiO, nanotubes that should be superimposed on the microtopography for early osseointegration. Not surprisingly, the in vivo results further demonstrated the enhanced early osseointegration behavior of the implants with SLA + 80 nm surfaces. In native bone tissues, stem cells and osteoblasts are exposed to the surface of the implant. As the cells encounter the SLA + 80 nm surface, they are forced to elongate and stretch to search for protein aggregates, activating intracellular signaling cascades that promote their differentiation into mature osteoblasts. The in vivo results clearly demonstrated the synergistic effects of the larger diameter nanotubes (80 nm) and the microscale rough surface. Similar results were previously demonstrated by Yun et al.²⁹ New bone tissue had apparently formed on the SLA + 80 nm surface, while only a little new bone tissue could be found on the SLA + 30 nm and SLA surfaces. However, the mean BA and BIC of the SLA + 30 nm group were higher than those of the SLA group, and overall, the effectiveness of the hybrid surfaces in promoting new bone growth exceeded our expectations. The increased new bone growth may be due to more proteins and cells being attached to the surface of the SLA + 30 nm than to the SLA surface. However, the level of osteogenic gene expression of cells grown on the SLA + 30 nm surface was the lowest of the four groups.

Developing approaches to promote the faster and firmer fixation of implants into bone have been a persistent challenge for developing implants to expand therapeutic indications, reduce patients' morbidities, and improve the success rate of the treatments. 43,44 The enhancement of multiple functions related to new bone growth of cells grown on the hierarchical micro/nanotextured surfaces compared with that of cells grown on the microstructured surface led to faster bone maturation around the implant without compromising the amount of new bone mass. Moreover, the mechanical interlocking feature of the microtopography was maintained in this micro/nanostructure. It was observed in our experiments that the microtopography itself was not excessively altered after anodization, except that sharp edges became blunter. All of the three micro/nanotextured surfaces had similar hydrophilicities, and there was no obvious correlation

between Ra and their biological effects. Therefore, the hierarchical SLA + 80 nm surface described in this study may be a useful surface for implants due to yielding of improved growth of surrounding bone.

Conclusion

In this report, hierarchical hybrid micro/nanotextured titanium surface topographies were produced by simple acid etching followed by anodization with titania nanotubes of different diameters. The SLA + 30 nm surface induced enhanced protein adsorption and initial cell adhesion and proliferation of osteoblasts. On the SLA + 80 nm surface, the shape of the cells appeared more irregular and osteogenesis-related gene expression was dramatically enhanced. In the experimental in vivo model, the SLA + 80 nm surface yielded improved new bone growth and implant integration compared with those induced by the other tested surfaces, which featured smaller diameter or no nanotubes. In light of these findings, the SLA + 80 nm surface presented in this study represents a pivotal advancement and appears to offer advantages over the titanium surfaces currently used for orthopedic and dental implants.

Acknowledgment

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Disclosure

The authors report no conflicts of interest in this work.

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Supplementary materials

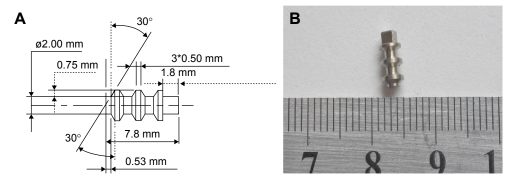


Figure S1 The blueprint of our experimental implant (A) and turned surface implant (B).

Table SI PCR primer sequences and product sizes (base pairs)

Gene	Primer sequence	Amplicon size (bp)
ALP	S: 5'-CATGCTGAGTGACACAGACAAGAA-3'	141
	A: 5'-ACAGCAGACTGCGCCTGGTA-3'	
OCN	S: 5'-GACGAGTTGGCTGACCACA-3'	138
	A: 5'-CAAGGGGAAGAGGAAAGAAGG-3'	
RUNX2	S: 5'-TCCACACCATTAGGGACCATC-3'	136
	A: 5'-TGCTAATGCTTCGTGTTTCCA-3'	
GAPDH	S: 5'-TGGCACCCAGCACAATGAA-3'	186
	A: 5'-CTAAGTCATAGTCCGCCTAGAAGCA-3'	

Abbreviations: PCR, polymerase chain reaction; ALP, alkaline phosphatase; RUNX2, runt-related transcription factor 2; OCN, osteocalcin.

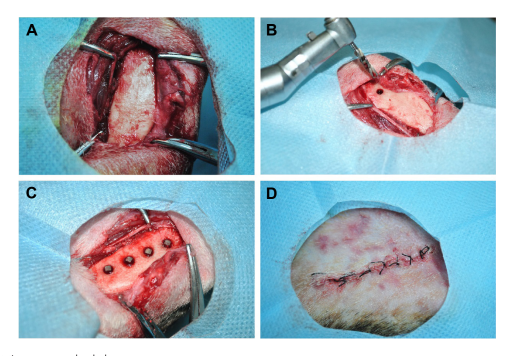


Figure S2 Implantation surgery on a beagle dog.

Notes: (A) The incision and exposure of the surgical fields; (B) preparation of holes for implantation; (C) placement overview of the four implants; (D) suture of the wound.

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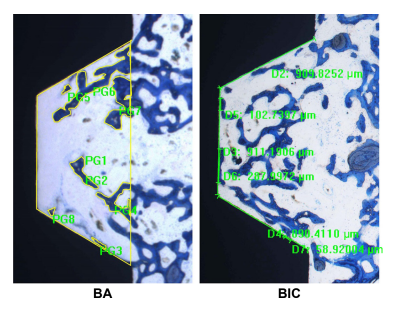


Figure S3 The schematic representation of BA and BIC (40×). Abbreviations: BA, bone area; BIC, bone—implant contact.

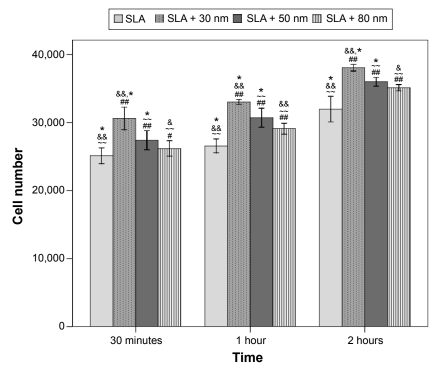


Figure \$4 Cell numbers (mean ± standard deviation) on different surface at different times (n=3, ten different random fields of each disc). Notes: ##P<0.01 and #P<0.05 compared with the SLA group; P<0.01 compared with the SLA + 30 nm group; 88P<0.01 and 8P<0.05 compared with the SLA + 50 nm group; and *P<0.05 compared with the SLA + 80 nm group. Abbreviation: SLA, sandblasting with large grit and acid etching.

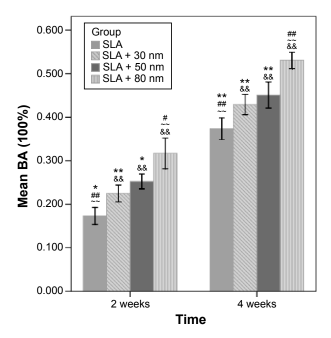


Figure S5 A comparison of the BA of the different groups at 2 weeks and 4 weeks.

Notes: ^{88}P <0.01 compared with the SLA group; ^{-}P <0.01 compared with the SLA + 30 nm group; $^{##}P$ <0.01 and $^{#}P$ <0.05 compared with the SLA + 50 nm group; and $^{**}P$ <0.01 and $^{*}P$ <0.05 compared with the SLA + 80 nm group.

Abbreviations: BA, bone area; SLA, sandblasting with large grit and acid etching.

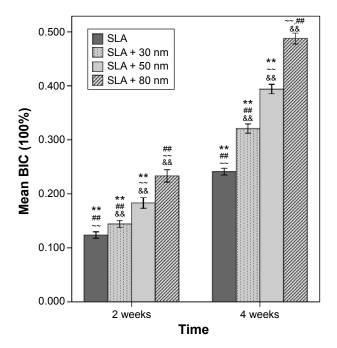


Figure S6 A comparison of the BIC of the different groups at 2 weeks and 4 weeks.

Notes: ^{88}P <0.01 compared with the SLA group; ^{-}P <0.01 compared with the SLA + 30 nm group; $^{##}P$ <0.01 compared with the SLA + 50 nm group; and $^{**}P$ <0.01 compared with the SLA + 80 nm group.

Abbreviations: BIC, bone–implant contact; SLA, sandblasting with large grit and acid etching.

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