Microstructures, mechanical, and biological properties of a novel Ti-6V-4V/zinc surface nanocomposite prepared by friction stir processing

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Background: The interaction between the material and the organism affects the survival rate of the orthopedic or dental implant in vivo. Friction stir processing (FSP) is considered a new solid-state processing technology for surface modification.

Purpose: This study aims to strengthen the surface mechanical properties and promote the osteogenic capacity of the biomaterial by constructing a Ti-6Al-4V (TC4)/zinc (Zn) surface nanocomposites through FSP.

Methods: FSP was used to modify the surface of TC4. The microstructures and mechanical properties were analyzed by scanning electron microscopy, transmission electron microscopy, nanoindentation and Vickers hardness. The biological properties of the modified surface were evaluated by the in vitro and in vivo study.

Results: The results showed that nanocrystalline and numerous β regions, grain boundary α phase, coarser acicular α phase and finer acicular martensite α′ appeared because of the severe plastic deformation caused by FSP, resulting in a decreased elastic modulus and an increased surface hardness. With the addition of Zn particles and the enhancement of hydrophilicity, the biocompatibility was greatly improved in terms of cell adhesion and proliferation. The in vitro osteogenic differentiation of rat bone marrow stromal cells and rapid in vivo osseointegration were enhanced on the novel TC4/Zn metal matrix nanocomposite surface.

Conclusion: These findings suggest that this novel TC4/Zn surface nanocomposite achieved by FSP has significantly improved mechanical properties and biocompatibility, in addition to promoting osseointegration and thus has potential for dental and orthopedic applications.

Keywords: nanocomposite, friction stir processing, nanocrystalline/ultrafine grained, bone marrow stromal cells, cell proliferation, osteogenic differentiation

Introduction

For biomaterials used in human tissue and organ regeneration, the interaction between the material and the organism affects the survival rate of the orthopedic or dental implant in vivo. For example, the implant may cause side effects in the organism, including cytotoxicity, inflammation, and anaphylaxis. Additionally, corrosion and abrasion will gradually occur during the service lifetime of the implant, leading to a decline in its mechanical properties.1–2 In complex biomechanical environments, ideal metal implants should not only possess excellent mechanical properties but also be biocompatible. In recent years, titanium (Ti) alloys have been widely used in various fields, including orthopedics, dental implantation, and tissue engineering, because of their good fatigue resistance, high corrosion resistance, and good biocompatibility. Many commercial products for replacing body parts or functions, such as artificial
joint replacements, bone trauma products, and oral implants, have been fabricated from Ti alloys.5–7

Among all Ti alloys, Ti-6Al-4V (TC4) is the most typical and the most frequently used α+β type Ti due to its high strength-to-weight ratio, low density, high corrosion resistance, and biocompatibility.8,9 Nevertheless, when the metal is implanted in a living body, it inevitably releases ions into the saline solution of bodily fluids.8 When TC4 is used as an implant material, aluminum (Al) and vanadium (V) ions released into surrounding tissues can adversely affect the body.9,10 Moreover, the difference between the elastic modulus of the biomaterial and that of the surrounding hard tissue can result in a stress-shielding effect.11,12 To avoid this disadvantage of TC4 without changing the composition of the whole material, we designed a surface modification process to create a nanocomposite layer on the implant surface, thereby improving its biocompatibility while largely maintaining its mechanical properties.

Friction stir processing (FSP) is considered as a new solid-state processing technology for surface modification.13,14 During FSP, a probe presses into a metallic substrate and rotates and moves along the processing direction. The friction between the probe and base material (BM) provides a heat source to increase the temperature of the metallic substrate. The heat input is accompanied by severe plastic deformation, resulting in recrystallization and the evolution of texture within the stirred zone (SZ) and coarsening within and around the SZ. Based on microstructural characterization, four zones can be distinguished in FSPPed metal, the SZ, the thermo-mechanically affected zone (TMAZ), the heat-affected zone (HAZ), and the BM zone. These various zones show different microstructures, leading to different mechanical properties.15 A new metal matrix nanocomposite (MMNC) structure is achieved by adding reinforcement particles or metal powder into the metallic substrate during plastic flow for participation in grain refinement, metallurgical bond behavior, and plastic deformation. The addition of different elements can promote the evolution of different microstructures, thereby optimizing the surface properties.

Recently, FSP has been increasingly applied to medical Ti. Li et al16 reported that the hardness and wear performance of the TC4 alloy could be significantly improved by FSP. Wang et al17 investigated superelasticity in a novel TiNbTaZr alloy prepared by FSP, providing a new way to obtain desired properties for biomedical applications. In our previous work,17 we prepared a Ti/SiC MMNC by adding SiC particles into the Ti during FSP and reported that the grain-refined surface promoted cell adhesion and osteogenic differentiation of rat bone marrow stromal cells (BMSCs).18 Compared with the coarse-grained surface, the ultrafine-grained Ti surface exhibited improved corrosion resistance.19,20 Estrin et al21,22 have also demonstrated that ultrafine-grained surface can promote cell adhesion and proliferation of stem cells and osteoblasts.

The surface properties of a biomaterial directly affect early cell adhesion, which is the primary stage of the interaction between tissue and medical implants, followed by cell proliferation and differentiation. The osteogenic capacity and rapid osseointegration of a biomaterial play vital roles in determining the success rate of an implant.23,24 FSP was used in this study to modify the alloy surface and to add zinc (Zn) particles to optimize its surface properties and to improve its biocompatibility. In this study, Zn was chosen as the reinforcement particle because it is an essential trace element in the human body. Zn participates in the synthesis of various enzymes and plays an indispensable role in bone growth and the maintenance of physiological cellular functions. In addition, Zn oxide has excellent antimicrobial properties and elemental Zn can effectively promote the osteogenic differentiation of BMSCs.25–27

Therefore, we hypothesized that adding Zn to TC4 during FSP could decrease the grain size, lower the elastic modulus, improve the surface hardness, and optimize the biological properties to further satisfy the requirements of in vivo implants. Thus, this study aimed to fabricate a novel TC4/Zn MMNC surface using FSP to strengthen the surface properties and promote the osteogenic capacity of the material. The microstructures and mechanical properties and their relationship were thoroughly explored. In addition, the biological activity of the nanocomposite surface was investigated both in vitro and in vivo.

Methods

Fabrication of TC4/Zn MMNC

Commercially available Zn powder (99.9% purity with an average particle size of 50 nm; Shanghai Shuitian Materials Technology Co., Ltd., Shanghai, People’s Republic of China) and TC4 plates (Gold Borui Titanium Industry Co., Ltd., Baoji, People’s Republic of China) were used as raw materials in this study. The TC4 BM was used as the control group (Con). The TC4 MMNC that underwent FSP without the addition of Zn is denoted as FSP. The TC4/Zn MMNCs that underwent FSP with premade holes of different depths are denoted as Zn-I-FSP (1 mm holes) and Zn-II-FSP (0.5 mm holes). FSP was performed using a professional-grade friction-stir welding machine (CFSW-PL800; Beijing FSW Technology Co., Ltd., Beijing, People’s Republic of China) with a tungsten steel FSP tool in an argon atmosphere to prevent oxidation. The probe was 10 mm in diameter, had a concave shoulder and a 2 mm pin height, and was tilted by 2.5°. When the parameters of the probe were defined, the volumetric ratio of the added Zn nanoparticles could be roughly estimated. The volumetric ratio of Zn in the Zn-I-FSP group was ~5.1%,
whereas the ratio for the Zn-II-FSP group was ~2.48%. A constant traverse speed ($v$) of 50 mm/min at a rotation rate ($\omega$) of 300 rpm was adopted to fabricate TC4/Zn MMNC by FSP. Single-pass FSP was conducted in this study. Samples with dimensions of 1×1 cm were cut from the SZ at the surface of the FSP plate for microstructural examination, mechanical testing, and cell experiments. Before making microstructural observations, samples were cut, ground, and polished according to standard procedures and were then etched with a solution consisting of 2% HF and 6% HNO$_3$, for 5 s. Longitudinal sections of the TC4/Zn MMNC were prepared using a wire cutting machine and were then thinned using an ion thinning instrument from the upper surface to the bottom. Cylindrical implants 2 mm in diameter and 6 mm in length were cut from the SZ using the wire cutting machine for the in vivo study.

**Microstructural change in TC4/Zn MMNC**

The microstructure of FSP-processed samples was analyzed by scanning electron microscopy (SEM) with a Quanta 200 microscope (FEI Company, Eindhoven, the Netherlands) at 20 kV. Transmission electron microscopy (TEM; JEM-200 EX; JEOL, Tokyo, Japan) was used to observe the microstructures of all FSP zones ranging from the upper surface to the bottom of the MMNC (0–1.5 mm). The relative surface wettability was estimated from the contact angle using a contact angle meter (Kruss GmbH, Hamburg, Germany) with water as the medium.

**Surface properties of TC4/Zn MMNC**

Nanoindentation tests were performed using a NANO Indenter G200 Testing System with a diamond Berkovich tip with continuous loading up to a maximum of 10 mN. Before each unloading process, a 30 s dwell time was applied under a fixed load. Load and displacement were continuously monitored and recorded to plot a curve. Using the obtained data, the nanoindentation depth and elastic modulus of the material were calculated. The Vickers hardness was measured using a microhardness tester (Shanghai Taiming Optical Instrument Co., Ltd., Shanghai, People’s Republic of China) under a load of 50 g. Eleven samples were tested.

**BMSC adhesion and cell proliferation on TC4/Zn MMNC surface**

All animal procedures and experiments were approved by the Animal Research Committee of the Ninth People’s Hospital affiliated with the Shanghai Jiao Tong University School of Medicine, of which the guidelines were followed for the welfare of the animals. BMSCs were isolated from the femurs and tibias of 6-week-old male Sprague Dawley (SD) rats, as previously reported. The cells were cultured in Dulbecco’s Modified Eagle’s Medium (DMEM) with 10% fetal bovine serum in an incubator with a 5% CO$_2$ atmosphere at 37°C. Cell identification was conducted according to our previously published procedures. Cells at passage 2 or 3 were used in subsequent experiments.

Nuclei were stained during the initial seeding period (1, 4, and 24 h) to count the adherent cells on the different samples. At each time point, nonadherent cells were removed using a phosphate-buffered saline (PBS) rinse. The cells were fixed with 4% paraformaldehyde for 30 min at 4°C. Next, nuclei were stained with 4’,6-diamidino-2-phenylindole (DAPI) for 10 min and the samples were then observed using a fluorescence microscope (IX71; Olympus Corporation, Tokyo, Japan). Cell counts were quantitatively analyzed using a Z2 Coulter particle count and size analyzer (Beckman Coulter, Brea, CA, USA).

Integrin β1 immunofluorescence was detected to confirm the expression of this adhesion-related protein. The initial cell seeding density was 2.0×10$^4$ cells/well for 24 h. After fixation with 4% paraformaldehyde for 30 min at 4°C, the samples were sequentially treated with 0.5% Triton X-100 and 3% bovine serum albumin (BSA) at room temperature (RT). A specific primary rabbit-anti-rat antibody targeting integrin β1 (eBioscience Inc., San Diego, CA, USA) was then added and incubated with the cells for 8 h at 4°C. The cells were subsequently incubated with a fluorescein isothiocyanate (FITC)-conjugated antirabbit IgG antibody (Jackson ImmunoResearch Laboratories Inc., West Grove, PA, USA) for 1 h at RT away from light. The cell cytoskeleton was stained with a tetramethylrhodamine (TRITC)-phalloidin-conjugated antibody (KenGEN Biotech, Nanjing, People’s Republic of China). Nuclei were stained with DAPI for 10 min, and the samples were observed using a fluorescence microscope (IX71; Olympus, Tokyo, Japan).

An 3-(4,5)-dimethylthiazihoazo(2-y1)-3,5-diphenytetrazo-
liumromide (MTT) cell metabolic activity assay was used to measure mitochondrial activity to evaluate the proliferation and viability of BMSCs on the samples. First, cells at a density of 2.0×10$^4$ cells/mL were plated onto each specimen in a 24-well plate. After culturing for 1, 4, 7, and 10 days, MTT solution was added to target wells. During 4 h of incubation, formazan was formed and then dissolved in dimethyl sulfoxide. Optical density (OD) absorbance values were measured at 490 nm using an ELX Ultra microplate reader (Bio-Tek, Winooski, VT, USA). The experiment was conducted in triplicate.

**BMSC osteogenic differentiation on the TC4/Zn MMNC**

Osteogenic differentiation was evaluated by alkaline phosphatase (ALP) staining, ALP activity, and real-time PCR.
Cells at a density of 2.0×10^4 cells/mL were plated onto each specimen in a 24-well plate. On the 7th and 14th days of culturing, the cells on the target samples were fixed and stained with BCIP/NBT reagents (Beyotime, Shanghai, People’s Republic of China) according to the manufacturer’s instructions. For the quantitative ALP assay, the cells were lysed and incubated with p-nitrophenyl phosphate (Sigma-Aldrich Co., St Louis, MO, USA) at 37°C for 30 min. ALP activity was estimated by the OD absorbance values at 405 nm and normalized to the total cellular protein. The total protein content was determined using a Bio-Rad protein assay kit (Bio-Rad Laboratories Inc., Hercules, CA, USA) using the OD absorbance values measured at 630 nm. The experiment was conducted in triplicate.

The osteogenesis-related genes examined included Runx2, ALP, OSX, COL-1α, OPN, and OCN. Cells were seeded at a density of 2.0×10^4 cells/mL on the Ti plates (1×1 cm) and were cultured for 7 and 14 days. Each group included five specimens at each time point to ensure that there was sufficient total RNA for PCR. Total RNA was extracted using a TaKaRa MiniBEST Universal RNA Extraction kit (TaKaRa Bio Inc., Otsu, Japan) according to the manufacturer’s instructions, and cDNA was synthesized with a TaKaRa PrimeScript™ 1st Strand cDNA Synthesis kit. The expression levels of the target genes were analyzed using a Bio-Rad Quantitative Real-Time PCR system (Bio-Rad Laboratories Inc.).

The relative expression level was normalized to that of the internal control gene β-actin. The ΔΔCT method was applied to calculate the target gene expression levels, and the data from the Con were used as the control. The experiment was conducted in triplicate.

**In vivo osseointegration of TC4/Zn MMNC implant**

The surgical procedures were performed under sterile conditions. After anesthesia via injection with ketamine (10 mg/kg), the distal femur of 10-week-old SD rats was carefully exposed through a skin incision and periosteum blunt dissection. A hole (⌀=2.0 mm) was drilled into the distal femur along the direction of the medullary cavity. Four types of implants (⌀=2×6 mm) were randomly inserted into each of the holes, and the wounds were conventionally sutured. Each type of implant was tested in triplicate.

Microcomputed tomography (micro-CT), sequential fluorescent labeling observation, and Van Gieson’s staining were used to evaluate new bone formation around the TC4/Zn MMNC implant in vivo. Sequential fluorescent labeling was used to evaluate the rate of new bone formation and mineralization around the TC4/Zn MMNC implant in vivo. Then, at 2, 4, and 6 weeks after the operation, 25 mg/kg of tetracycline (TE; Sigma-Aldrich Co.), 30 mg/kg of alizarin red S (AL; Sigma-Aldrich Co.), and 20 mg/kg of calcein (CC; Sigma-Aldrich Co.) were injected intraperitoneally. At 8 weeks after operation, all SD rats were sacrificed and the femurs with the implants were harvested and fixed in 10% buffered formaldehyde. The micro-CT scanning parameters were set at 65 kV using an Al 1 mm filter with a resolution of 18 mm. The new bone organization was assessed by parameters including the trabecular number (Tb.N), bone volume (BV), and BV/total volume (TV), as analyzed by micro-CT. After micro-CT examination, all samples were dehydrated using graded ethanol, embedded in polymethylmethacrylate (PMMA), and cut into 150 μm thick sections using a Leica SP1600 saw microtome (Leica Microsystems, Wetzlar, Germany). These sections were manually pasted on a slide and carefully polished to a final thickness of ~40 μm. Sequential fluorescent labels were observed using confocal laser scanning microscopy (CLSM; Olympus Corporation). The percentage of the labeling area was calculated using ImageJ 1.48 v (National Institutes of Health, Bethesda, MA, USA).

The bone–implant contact (BIC) was calculated from the histomorphometric measurements by Van Gieson staining.

**Results**

**Microstructures of TC4/Zn MMNC**

Figure 1 shows SEM micrographs of the upper surface of the TC4/Zn MMNC prepared by FSP. As shown in Figure 1A and B, the microstructures of the TC4 alloy are composed of prior α and β phases and the β particles are mainly located on grain boundaries (GBs) or at triple junctions of α grains. This is the typical microstructure of the TC4 alloy, which has been reported in many investigations. After FSP, numerous β regions, GB α phase, coarser acicular α phase, and finer acicular martensite α’ appear (Figure 1C and D). The magnified image clearly shows that the grains are significantly refined after FSP (Figure 1D), with the simultaneous formation of finer lamellar α and β phases. These results indicate that during FSP, the peak temperature of the TC4 alloy can reach the β phase transus temperature due to severe plastic deformation and the alloy completely transforms to the β phase. During the cooling process, the α phase precipitates first from the GB of the β phase and then precipitates and grows within the β phase. Meanwhile, martensite α’ is also formed due to the high cooling rate (Figure 1D). Martensite α’ has been reported to precipitate when the cooling rate is >410°C/s during FSP. The FSP and TC4/Zn MMNC groups exhibit similar microstructural features. However, the β region is significantly larger after adding Zn to the TC4.
The size of the β regions in the FSP, Zn-I-FSP, and Zn-II-FSP groups is ~10, 20, and 15 μm, respectively. Because the volumetric ratio of Zn in the SZ was controlled artificially via controlling the depth of the blind hole, the content of Zn in the Zn-I-FSP group is higher than that in the Zn-II-FSP group. Thus, the β phase region in the TC4/Zn nanocomposite grows with increasing Zn content. In addition, Zn-rich particles are found in the nanocomposite.
some of which are identified as TiZn$_2$, indicating that Zn reacted with Ti and formed a new compound during FSP. Figure 1F and H also clearly show that the quantity of Zn-rich particles increases with the increasing Zn content.

To understand the microstructural evolution in detail, transverse cross-sections of the Zn-I-FSP group alloy were observed using TEM. The results revealed significant differences in the upper surface (within 200 μm) and the middle region of the SZ (~1 mm from upper surface). Figure 2 shows TEM images of the upper surface in the Zn-I-FSP group alloy. Nanocrystals formed during FSP in the upper region (Figure 2A). The lower region, as shown in Figure 2A, has a completely different microstructural morphology and corresponding selected area electron diffraction (SAED) (Figure 2A, inset a2), which indicates that the amorphous phase transformation occurs in this area. Figure 2B shows the TiZn$_2$ phase, and the corresponding SAED also shows discontinuous rings, suggesting the coexistence of both grains and subgrains, as clearly shown in the inset of Figure 2B.

As shown in Figure 2C, several nanotwins formed during...
FSP in the upper surface of the sample. Several high-density dislocation areas were also observed (not shown). In addition, some Zn-rich particles appeared, as shown in Figure 2D, and the energy-dispersive X-ray spectroscopy (EDS) composition analysis of the matrix revealed that the matrix alloy also contained Zn (Figure 2E and F), suggesting that during FSP, the added Zn can transform into a second phase or dissolve into the matrix crystal structure to form a solid solution.

Figure 3 shows the TEM images of middle region of the SZ (~1 mm from upper surface) in the Zn-I-FSP group alloy. Figure 3A clearly exhibits deformed and rod-like α grains, suggesting that severe plastic deformation occurs in this area during FSP and that the grains are elongated and refined. Figure 3B shows that new recrystallization is also formed during FSP, which indicates that dynamic recrystallization occurred during FSP, thus forming new grains. This recrystallization occurred because the alloy was simultaneously subjected to severe plastic deformation and the high temperature produced by the stirring effect of the FSP tool. Moreover, grains of the TC4 alloy are fractured or elongated. Meanwhile, grains may have nucleated from areas with high amounts of stored energy, which then grow to form new grains.33

The contact angles of water droplets on the surfaces were 61.23° (Con), 42.43° (FSP), 30.5° (Zn-I-FSP), and 39.13° (Zn-II-FSP) (Figure 4). Thus, the surface of Zn-I-FSP showed the highest hydrophilicity and surface wettability with the addition of Zn particles. As mentioned earlier, the volumetric ratio of Zn affected the evolution of microstructural characteristics, which explains the different contact angles of Zn-I-FSP and Zn-II-FSP. Grain refinement seems to be another contributing factor, because the contact angle was smaller for the FSP group than the Con. The difference in contact angle is attributable to the sub-micron/nanoscale grain size and different surface microstructures induced by the different volumetric ratios of added Zn.22,24

Surface properties of TC4/Zn MMNC
Figure 5 illustrates the force–displacement plots from the nanoindentation tests in each group. The elastic modulus of the Con is 126.23±19.48 GPa. After FSP, the elastic modulus of the FSP group (116.86±16.61 GPa) decreased and that of the TC4/Zn MMNC surface further decreased with the addition of Zn particles. The elastic moduli of the Zn-I-FSP and Zn-II-FSP groups are 106.93±18.99 and 107.21±22.43 GPa, respectively (Table 1).

The FSP group (395.5±29.44 HV) showed the highest microhardness value among the four groups (Table 1). Compared with the Con (344.6±15.52 HV), the microhardness of the Zn-I-FSP group (375.7±13.00 HV) was 9.02% higher and that of the Zn-II-FSP group (389.3±26.81 HV) was 12.97% higher.
higher. The addition of Zn particles lowered the effect of FSP; however, the increased addition of Zn decreased the microhardness.

**TC4/Zn MMNC promotes cell adhesion and proliferation**

Figure 6A and B shows the adhesion ability of rat BMSCs on the surface of each sample at 1, 4, and 24 h. More cells adhered to the Zn-I-FSP and Zn-II-FSP samples than to the Con samples after 4 and 24 h (Figure S1). The quantitative data further demonstrate that cell adhesion was significantly enhanced on the surface of Zn-I-FSP and Zn-II-FSP groups (Figure 6A). Cell adhesion on surfaces after 24 h was ranked in the following decreasing order: Zn-I-FSP > Zn-II-FSP > FSP > Con.

To further confirm the expression of an adhesion-related protein and to observe initial cell spreading, integrin β1 and the cytoskeleton of BMSCs in each group were examined by immunofluorescence after staining with a FITC-tagged antibody and TRITC-phalloidin to visualize integrin β1 and cytoskeletal actin expression, respectively (Figure 6B). The nuclei of BMSCs were stained with DAPI. The BMSCs on each sample exhibited a multipolar spindle-like morphology and extended pseudopodia on all surfaces. The cells seeded on the Zn-I-FSP surface expressed the highest level of integrin β1, and those on the FSP and Zn-II-FSP group expressed higher levels of integrin β1 than those on the Con.

The MTT results are shown in Figure 6C. No statistically significant differences were observed among the four groups on day 1. On day 4, there was no significant difference in the

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**Table 1** Mechanical properties of each group

<table>
<thead>
<tr>
<th>Group</th>
<th>Elastic modulus (GPa)</th>
<th>Microhardness (HV)</th>
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</thead>
<tbody>
<tr>
<td>Con</td>
<td>126.23±19.48</td>
<td>344.6±15.52</td>
</tr>
<tr>
<td>FSP</td>
<td>116.86±16.61</td>
<td>395.5±29.44</td>
</tr>
<tr>
<td>Zn-I-FSP</td>
<td>106.93±18.99</td>
<td>375.7±13.00*</td>
</tr>
<tr>
<td>Zn-II-FSP</td>
<td>107.21±22.43</td>
<td>389.3±26.81*</td>
</tr>
</tbody>
</table>

**Notes:** Data are presented as mean ± SD. No statistical differences in elastic modulus were found among the groups. *The microhardness showed significant differences among the groups (P<0.05 compared with the Con).

**Abbreviations:** Con, control group; FSP, friction stir processing; Zn, zinc.
total cell metabolic activity between the FSP and Zn-I-FSP groups; however, that of both groups was higher than that of the Con. On day 7, the total cell metabolic activity in the Zn-I-FSP group was significantly greater than that in the other groups. On day 10, all three modified groups showed more stimulated total cell metabolic activity than the Con. The MTT results indicate that the novel surface nanocomposite positively affects the viability and proliferation of the BMSCs.

**TC4/Zn MMNC enhances in vitro osteogenic differentiation**

As shown in Figure 7A, all groups showed intense ALP staining on days 7 and 14 of culture, except for the Con, which showed light staining. Quantitative ALP activity assay of the Zn-I-FSP group on day 7 showed significantly higher activity than that in the Con, whereas BMSCs in the Zn-II-FSP group showed higher ALP activity on day
14 than those in the Con. The ALP activity results indicate enhanced osteogenic differentiation of BMSCs seeded on the TC4/Zn MMNC surface with the addition of Zn particles (Figure 7B).

The expression levels of osteogenic differentiation marker genes, including Runx2, ALP, OSX, COL-1α, OPN, and OCN, were examined by PCR after 7- and 14-day cultures. As shown in Figure 8, after a 7-day culture, the expression of Runx2, ALP, COL-1α, and OPN was higher in the Zn-I-FSP and Zn-II-FSP groups than that in the Con, whereas Runx2 and COL-1α showed higher expression in the FSP group than in the Con. The highest expression of COL-1α was observed in the FSP group. The expression of COL-1α was ranked in the order of FSP > Zn-II-FSP > Zn-I-FSP > Con. After a 14-day culture, the expression of Runx2, OPN, and OCN in the Zn-II-FSP group was significantly upregulated compared with expression in the other three groups. The expression of ALP in the Con was lower than that in the other three groups. The expression of COL-1α was downregulated in the experimental groups compared with expression in the Con after culturing for 14 days. Compared with the Con, the Zn-I-FSP group showed upregulated OCN expression.

**TC4/Zn MMNC promotes rapid osseointegration**

An in vivo study was performed to further detect the osseointegration of the TC4/Zn MMNC implant. Figure 9A
shows images of the intersecting surface of the implant captured by micro-CT. The new bone around the implant surface appeared to have a high-density structure, and this structure was larger round the implant surfaces in the marrow cavity of the FSP, Zn-I-FSP, and Zn-II-FSP samples than in that of the Con samples. The BV, BV/TV, and Tb.N around the implant (within 2 mm from the implant) were determined (Figure 9B–D). The Zn-II-FSP group demonstrated a higher BV than that of the other groups. In terms of the Tb.N, the Zn-I-FSP group exhibited obvious advantages among the four groups. Both the FSP and Zn-II-FSP groups exhibited a higher BV/TV than the other two groups.

The sequential fluorescent labeling analysis results are shown in Figures 10 and 11. After 2 weeks, the percentage of

Figure 8 Expression levels of osteogenic differentiation marker genes.
Notes: Gene expression of Runx2, ALP, OSX, COL-1α, OPN, and OCN genes on days 7 and 14 assayed by real-time PCR; *P<0.05, **P<0.01.
Abbreviations: ALP, alkaline phosphatase; Con, control group; FSP, friction stir processing; Zn, zinc.
TE labeling area in the Zn-I-FSP group was found to be the largest among the four groups and the labeling area of the Con was significantly lower than that of the other groups. After 4 weeks, the percentage of AL labeling in the Zn-II-FSP group was obviously larger than that in the other groups. At the last time point, the percentage of CC labeling area in the Zn-I-FSP group and Zn-II-FSP group was larger than that in the other two groups. The sequential fluorescent labeling results indicate that the TC4/Zn MMNC stimulates more new bone formation at every time point, especially in the form of Zn-II-FSP.

Figure 12A shows the position of the implant in the distal femur of the SD rats. Histological analyses are shown in Figure 12B. The partial magnifications of histological stain reveal the formation of fibrous connective tissue around the Con and FSP implants, which separated the new bone tissue from the implant surface (marked by red arrows). The BIC was calculated as the quotient of the length of the direct BIC and the length of the implant. The BIC index of Zn-I-FSP and Zn-II-FSP was 56.0% ± 7.26% and 58.5% ± 5.45% (Figure 12C), respectively, both of which are significantly larger than that of Con (35.25% ± 6.24%) and FSP (42.75% ± 4.65%), suggesting that the two TC4/Zn MMNC groups induced better BIC among the four groups.
Figure 10 Sequential fluorescent labeling observations of TE, AL, and CC at 8 weeks after implantation.

Notes: The color in yellow (TE), red (AL), and green (CC) indicates the rate of new bone formation at 2, 4, and 6 weeks after the operation, respectively. Scale bar: 200 μm.

Abbreviations: AL, alizarin red S; CC, calcein; Con, control group; FSP, friction stir processing; TE, tetracycline; Zn, zinc.

Figure 11 The area percent of fluorochrome stained bone in each group.

Notes: The TE, AL, and CC indicated the rate of new bone formation at the 2, 4, and 6 weeks after the operation, respectively; *P<0.05, **P<0.01).

Abbreviations: AL, alizarin red S; CC, calcein; Con, control group; FSP, friction stir processing; TE, tetracycline; Zn, zinc.

Discussion

Microstructural evolution and mechanical properties of TC4/Zn MMNC

Microstructural observations indicate that the TC4 alloy undergoes intense plastic deformation at the elevated temperature during FSP and that the peak temperature exceeds the phase transformation point. Thus, the microstructures transform completely to the β phase. During cooling, the α phase preferentially nucleates at triple junctions and boundaries of β grains with a low energy orientation relationship, leading to the formation of a GB α phase (Figure 1D, F, and H). Thereafter, the acicular α and martensite α’ phases form in the interior of β grains. Martensite α’ can form when the TC4 alloy is cooled from above the β transition temperature at a cooling rate of >20°C/s. Therefore, we believe that the finer microstructures formed during FSP are
the main reason for the higher microhardness of the FSPed TC4 alloy. This result is consistent with those of Li et al’s \(^7\) investigation, in which the greater surface microhardness of the TC4 alloy after FSP was attributed to finer \(\alpha/\beta\) lamella. These researchers also concluded that the martensite \(\alpha'\) phase could enhance the hardness of the FSPed TC4 alloy. \(^7\) In addition, especially on the upper surface of the FSPed samples, nanocrystalline, amorphous, nanotwins, high-density dislocations, among other features, form during FSP due to more severe plastic deformation (Figure 2A, C, and E). These microstructural features can significantly increase the microhardness of the TC4 alloy, which has been reported in many investigations. \(^36-40\) For example, Liu et al\(^{16,37}\) considered that the nanocrystallinity and high-density dislocations of the TC4 alloy formed via shot peening could result in remarkable grain refinement strengthening and work-hardening of the alloy, which would significantly improve their surface microhardness. The amorphous metal has no dislocations or GBs, resulting in higher microhardness, showing extraordinary yield strength and microhardness. \(^38-40\)

After adding Zn into the TC4 alloy, Zn can transform into a second phase or dissolve into the matrix crystal structure to form a solid solution (Figure 2B, D, and F). However, the addition of Zn into the TC4 alloy will also result in an increase in the size of the \(\beta\) phase region (Figure 1E and G), which may explain why the FSP-Zn group has a lower microhardness than the FSP group and why the microhardness decreases with the increase in the Zn content (Table 1). However, the reason that elemental Zn can accelerate the growth of \(\beta\) grains is not clear and requires further investigation.

The nanoindentation test results showed that the FSP group and FSP/Zn groups have lower elastic moduli than the Con, which can be attributed to the relatively lower \(\beta\) phase fraction in the TC4 alloy due to the higher content of \(\alpha\) phase-stabilizing element Al. After FSP, more \(\beta\) phase forms due to the faster cooling rate (Figure 1D, F, and H), while the Ti alloy with the higher \(\beta\) phase content has a lower elastic modulus. \(^16,17,41\) On the other hand, the MMNC produced by FSP has very small grains and, in our previous study, the material with more refined grains exhibited...
a lower elastic modulus. Therefore, the FSP and MMNC groups have lower elastic moduli than the Con. After adding Zn into the TC4 alloy, although the size of the β region increased, more β phase may simultaneously form, which may be why FSP/Zn MMNC has a lower elastic modulus than the FSP group.

In medical implants, metal ion release from the Ti alloy inevitably occurs due to wear or corrosion and may cause an immune response, such as inflammation around or an allergy to the implant. The wear properties of the material are positively correlated with hardness according to the Archard principle. Through grain refinement and the change in the α/β ratio caused by FSP, the microhardness of the Zn-I-FSP group was significantly increased by 9.02% and that of the Zn-II-FSP group was increased by 12.97%. These changes indirectly indicate that fewer metal ions would be released. In terms of corrosion, human body fluids are complex electrolyte media that contain a variety of organic acids, oxides, and ions. Corrosion reactions of medical implants will occur inevitably over the long term. Multiple studies have reported that nanocomposites modified by FSP have enhanced corrosion resistance. In the corrosive environment of a sodium chloride solution, the electronegativity of Zn is stronger than that of Al, meaning that Zn is more likely to be released. Nevertheless, further investigations are required to evaluate the corrosion resistance of this novel material.

The improved surface wettability of the material contributed to the different cell adhesion and osteogenic responses observed. The surface properties of a biomaterial directly affect early cell adhesion. In this study, the improved surface wettability is thought to contribute to the high numbers of cells adhering to the surface of Zn-I-FSP and Zn-II-FSP. Cellular adhesion is the primary stage of the interaction between the tissue and the medical implants, followed by cell proliferation and differentiation. A series of specific proteins are thought to be involved in signaling transduction during cell adhesion. Integrin, as an extracellular matrix receptor, participates in the signal transduction involved in responding to external stimuli, regulating gene expression, and influencing cell migration, proliferation, and differentiation. The detection of the expression of integrin β1 confirmed the crucial role of this adhesion-related protein in the process of cell adhesion. Surface wettability has been reported to be an important factor for controlling the enhanced attachment and spreading of stem cells. However, further investigations are required to elucidate the exact mechanism.

The surface nanocomposite modified by FSP promoted the proliferation of BMSCs, as previously reported. ALP is an early marker of osteogenesis, and its upregulation indicates enhanced phosphate metabolism. In this study, the ALP upregulation in Zn-I-FSP and Zn-II-FSP groups indicated positive regulation via Zn release. Zn is an important cofactor for ALP; moreover, it can activate Zn-containing metalloenzymes, including collagen enzyme, DNA polymerase, and RNA polymerase, which are directly involved in the expression of nucleic acids and protein that are closely associated with bone metabolism. Multiple studies have shown that Zn can upregulate the osteogenic differentiation of BMSCs. Huo et al added Zn to the coating of a Ti surface using anodization and hydrothermal treatments and achieved enhanced osteogenic activity and antibacterial properties. ERK1/2 signaling was found to be involved in these processes.

The PCR results confirmed the enhanced ALP activity at the gene level. Early enhanced expression of Runx2, a key gene in osteogenesis, can further activate and upregulate the expression of other related genes. OSX accelerates the process of osteogenic differentiation and new bone formation. In this study, the role of OSX was not obvious. It has been reported that Col-1α can promote adhesion behavior mediated by integrin β1. Col-1α exerts a considerable influence on the early deposition of inorganic molecules. OPN is associated with adhesion and matrix synthesis before mineralization and is a relatively late marker. OCN is a later marker of osteogenesis. The upregulation of OCN contributes to mineral deposition in the extracellular matrix and new bone mineralization. The enhanced expression of these osteogenic differentiation marker genes prompted in vitro BMSCs osteogenic differentiation at different stages.

TE, AL, and CC fluorescent markers can integrate with calcium ions in bone tissue and form complexes deposited in situ. The speed of calcium deposition around implants can be recorded using sequential fluorescent labeling to evaluate new bone formation. The enhanced rapid osseointegration of Zn-I-FSP and Zn-II-FSP was closely related to the positive effects of proliferation and osteogenic differentiation of BMSCs, which imbed the local environment with more osteoblasts around the implant.

Hydroxyapatite in natural bone takes the form of acicular nanocrystals (<100 nm). The nanocrystalline structure of TC4/Zn MMNC was similar to the crystal structure of hydroxyapatite in natural bone; thus, it could biomimetically promote the growth of hydroxyapatite. In addition, the release
of Zn in the body can promote extracellular matrix-related protein secretion and new bone mineralization.

According to the guidelines of classical implant dentistry, before the prosthetic procedure and loading, at least 3 months are required for adequate osseointegration after implant surgery. In this study, rapid osseointegration in vivo of the novel TC4/Zn MMNC surface was enhanced compared with the smooth TC4 surface. However, further research comparing this new implant material with classical modified implants should be conducted. Furthermore, how the nanocrystalline structure of TC4/Zn MMNC elicits responses in rat BMSCs requires further elucidation.

Conclusion
To strengthen the surface mechanical properties and promote the osteogenic capacity of this biomaterial, we modified the surface of TC4 through FSP with the addition of Zn particles and successfully constructed a TC4/Zn MMNC surface with grain refinement. Based on the characterization of the microstructural evolution, the mechanical properties were improved. Nanocrystalline and numerous β regions, GB α phase, coarser acicular α′ phase, and finer acicular martensite α′ appeared because of the severe plastic deformation caused by FSP, resulting in a decrease in the elastic modulus and an increase in the surface hardness. With the addition of Zn particles and the enhancement of hydrophilicity, biocompatibility was greatly promoted in terms of cell adhesion and proliferation. The in vitro osteogenic differentiation of rat BMSCs on the novel TC4/Zn MMNC surface and the in vivo osseointegration of the TC4/Zn MMNC implant were enhanced. The TC4/Zn MMNC surface modified using FSP not only exhibited superior mechanical properties but also provided improved surface and physicochemical properties for cell attachment and osseointegration. This novel TC4/Zn MMNC shows great potential for dental and orthopedic applications.

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

References
Supplementary material

Figure S1 Nuclear staining of each group at 1, 4, and 24 h.

Abbreviations: Con, control group; FSP, friction stir processing; Zn, zinc.