Effect of surface grain boundary density on preosteoblast proliferation on titanium

Terry C. Lowe a, Rebecca A. Reiss b, Patrick E. Illescas b, Casey F. Davis a, Melanie C. Connick b and Johnny A. Sena c

a George S. Ansell Department of Metallurgical and Materials Engineering, Colorado School of Mines, Golden, CO, USA; b Biology Department, New Mexico Institute of Mining and Technology, Socorro, NM, USA; c National Center for Genome Resources, Santa Fe, NM, USA

ABSTRACT
Studies since 2004 have shown that the cytocompatibility of ultrafine grain (UG) commercial purity (CP) titanium exceeds that of coarse grain (CG) CP titanium (Ti) by 30% to 20-fold. To isolate the factors affecting this large reported variability of CP titanium’s cytocompatibility, discs of UG and CG titanium were fabricated with controlled texture and roughness. The discs were seeded with MC3T3-E1 pre-osteoblastic cells and cultured for 72 h. The proliferation of cells on polished UG-Ti exceeded unpolished CG-Ti 3.04-fold. Cell proliferation was found to correlate with a new biophysical parameter, the average grain boundary length per surface-attached cell.

IMPACT STATEMENT
By eliminating the influences of texture and roughness we have isolated the positive correlation between MC3TE-E1 cell proliferation on titanium and grain boundary length per surface-attached cell.

Introduction
Titanium (Ti) alloys are widely used for dental implants [1] and orthopaedic devices [2]. Thus, the design of microstructures and surfaces of Ti medical implants has been the focus of research to enhance the integration of Ti with bone tissue [3]. Surface treatments can create desirable nanoscale structures on surface on Ti, as reviewed by Wang, et al. [4] and Kunčíká, et al. [3], but enhancing titanium’s strength for load-bearing applications also requires fine-scale volumetric features rather than surface-only modifications. The strength [5], corrosion resistance [6], and cytocompatibility [7,8] of commercial purity (CP) Ti can be enhanced by introducing sub-micron and nanometer-scale microstructural features via Severe Plastic Deformation (SPD) methods to produce ultrafine grain (UG) titanium. Continuous Equal Channel Angular Pressing (C-ECAP) [9], a specific form of SPD, can fabricate long, high-strength Ti rods suitable for making medical devices [10]. C-ECAP reduces grain size, alters grain boundary structure, and increases the areal surface density of grain boundaries [11].

The surface of Ti is of special interest since it forms the interface to the physiological environment surrounding medical prosthetics. Studies have demonstrated that the growth of preosteoblasts and other cell types on UG-Ti can be increased from 1.3-fold (30% increase) up to 20-fold (1900%) compared to conventional coarse grain (CG) Ti [7,8]. This enhancement of cell viability on SPD-processed metals has been attributed to the effects of surface energy [12], topography [13], grain size [14], and crystallographic texture [15]. Medvedev et al. [16] measured the effects of all of these variables on the viability of cells grown on CG-Ti and UG-Ti. They found that the viability of SaOS-2 cells and adipose-derived mesenchymal stem cells (adMSC) was...
greatest for substrates polished to nanoscale smoothness (average mean roughness $R_a < 4 \text{ nm}$) for both cell types and they recommended further investigation to quantify non-topological factors on such smooth surfaces. Baek et al. [17] systematically varied the crystallographic texture, grain size, surface energy, and roughness of Grade 2 Ti. They found greater proliferation of MC3T3-E1 cells on UG-Ti surfaces, which had higher surface energy, roughness, and more numerous grain boundaries. However, because they varied surface energy and roughness together, they were not able to differentiate the contributions of surface energy from surface roughness.

Here we follow up on the work of Medvedev, et al. and Baek, et al. by designing our experiments to isolate grain size-related effects on the extent of MC3T3-E1 pre-osteoblast attachment and proliferation on CG-Ti and UG-Ti. We fabricated substrates with nearly identical crystallographic textures and nanoscale smoothness so that we can focus our attention solely on the influence of grain size and the density of grain boundaries on cell response. Our goal is to test the hypothesis that grain boundaries influence cell proliferation, independent of texture and surface roughness.

**Materials and methods**

Rods of mill-annealed 15 mm diameter commercially pure Grade 4 titanium (Carpenter Technology Corporation, Heat H10803) were cut to lengths between 150 and 450 mm for C-ECAP processing. Table 1 lists the alloy composition. The rods were pre-heated to temperatures between 150°C and 200°C and subject to C-ECAP for 4 sequential passes through a 120° die with 90° rotations between each pass to achieve a total shear strain of 4.6. After C-ECAP, all rods were drawn and centerless-ground to a diameter of 13.3 mm.

The rods were sectioned transversely into thirty-two 2 mm thick discs and ground using a progression of SiC abrasive papers (grit 120, 240, 400, 800, 1200, 2500, 4000). Half of the discs were additionally polished to nanometric smoothness using a Buehler Vibromet™ 2 vibratory polisher with a suspension of 20 nm colloidal silica for 12 h to 24 h. Discs were cleaned using organic soap, sonicated for 10 min in high purity isopropanol, rinsed, and surface topographies were documented using a Digital Research Nanoscope IIIa atomic force microscope (AFM).

The discs were sterilized by 12-hour exposure to 100% isopropanol. Mouse preosteoblasts from the cell line MC3T3-E1 sub-clone 4 (American Type Culture Collection (ATCC), Rockville, MD) were cultured at 37°C at 5% CO$_2$ in Alpha Minimum Essential Medium (MEM $\alpha$, GIBCO, Grand Island, NY) supplemented with 10% fetal bovine serum (FBS, GIBCO, Grand Island, NY) and 1% penicillin/streptomycin (Cellgro, Manassas, VA). Discs were seeded in 24-well cell culture plates (USA Scientific, CC672–7524) at a density of $1.5 \times 10^4$ cells per well and incubated for 24 and 72 h. Culture media, along with unattached cells, were removed and fresh media was added at 24 h intervals.

Cell DNA was stained with DAPI (4’,6-diamidino-2-phenylindole) at a concentration of 10 μg/ml and intracellular lipids were stained by Nile Red (9-diethylamino-5H-benzo[alpha]phenoxazine-5-one) at a concentration of 10 μg/ml. Plates were then incubated in darkness at 37°C for 15 min. Cells were counted using a haemocytometer. Each disc was imaged with an Eclipse LV100 fluorescence microscope equipped with a Hamamatsu ORCA-RF camera and Nikon Plan Fluor 10x objective lens using 478–495 nm wavelength filters for DAPI and 532–587 nm wavelength filters for Nile Red. Triplicate colorimetric images were rendered using ImageJ software to quantitate the stained cells [18].

Cells were treated with 2% glutaraldehyde for 15 min and then dehydrated with ethanol rinses. After the final rinse, the cells were dried on the discs overnight. The discs were sputter-coated with platinum and imaged with a Hitachi S-2460N scanning electron microscope (SEM). For three replicates of each Ti substrate condition a JEOL JSM-7000F SEM with an EDAX Hikari Pro Electron Backscatter Diffraction (EBSD) Camera G was used to measure grain size distribution, grain boundary lengths and misorientations, and crystallographic texture. EBSD measurements were repeated at multiple electron microscopy magnifications, between 300x and 35,000x to confirm quantitative microstructural analysis statistics and ensure the absence of scan-size effects. Energy Dispersive Spectroscopy (EDS) was also conducted to measure the possible effects of segregation of elements present in the alloy.

**Results**

Grain orientation maps measured via EBSD show the structures and crystallographic textures for CG-Ti and UG-Ti in Figure 1. Deformation-induced refinement of the grain size is apparent in the inverse pole figure map.
in Figure 1b, though some micron-sized grains remain in the UG-Ti, even after the large strains imposed through C-ECAP.

The average grain size was 10.9 μm $\pm$ 7.3 μm for CG-Ti samples compared to 0.24 μm $\pm$ 0.38 μm for UG-Ti samples, a 45-fold difference. The difference in grain size distributions for CG-Ti and UG-Ti surfaces is shown in Figure 2. The average grain area for the CG substrates was 125.6 μm² compared to 0.0752 μm² for UG substrates. The inverse pole figures in Figures 1c and 1d show the preference for $\{10\bar{1}0\}$ prism plane normals to align with the rod axis for both CG and UG samples. The similarity in textures was achieved by additional rod drawing reductions as the final deformation processing step after the C-ECAP. The maximum times-random texture intensity was 6.7x for the CG-Ti samples, and slightly higher, 8.3x for UG-Ti samples. For the UG-Ti, the orientation having the maximum intensity was shifted on average approximately 14.6° from the $[10\bar{1}0]$ direction toward the $[0001]$ direction.

Atomic Force Microscopy measurements showed the average arithmetic mean roughness $R_a$ for the as-sectioned unpolished discs was 40.2 nm $\pm$ 24 and 70.2 nm $\pm$ 36 nm for the CG-Ti and UG-Ti, respectively. In contrast, the average $R_a$ for the polished discs was 0.24 nm $\pm$ 0.2 and 0.30 nm $\pm$ 0.2 nm for the CG and UG discs, respectively. The polished discs were nanometrically smooth. For the unpolished discs, the actual surface areas were only marginally greater than the projected areas, 3.02% higher for the CG-Ti samples and 2.59% higher for the UG-Ti samples. For the polished discs the differences in actual and projected surface areas were minuscule, 0.019%. Thus, the effective area available

Figure 1. Rod/extrusion axis inverse pole figure maps of (a) CG-Ti (1000x magnification) and (b) UG-Ti (20,000x magnification); and corresponding rod/extrusion axis inverse pole figures for (c) CG-Ti and (d) UG-Ti.

Figure 2. Grain area distributions for CG-Ti and UG-Ti.
Figure 3. Fluorescent images of mouse preosteoblasts after 72 h for (a) CG-Ti unpolished, (b) CG-Ti polished, (c) UG-Ti unpolished, and (d) UG-Ti polished. Mean cell counts (e) for all four states, with p-values for significant pairwise comparisons shown.
for cells to contact surfaces was not significantly different for the unpolished substrates compared to the polished substrates.

DAPI- and Nile Red-stained preosteoblasts, as shown in Figure 3, were counted on all samples after 24 (not shown) and 72 h. After 24 h, an average of 20 cells was observed on the four surfaces and there were no statistically significant differences.

After 72 h, an average of 72 cells was observed on polished CG-Ti versus 149 on polished UG-Ti, and 49 on unpolished CG-Ti versus 91 on unpolished UG-Ti. The mean number of surface-attached MT3T3-E1 cells was 1.9-fold and 2.1-fold higher compared to CG-Ti after 72 h of cultivation on unpolished and polished substrates, respectively. The differences observed after 72 h are due to increased proliferation, not initial adhesion to the substrate. The combined effects of UG-Ti and polishing increased the average number of attached cells 3.04-fold.

### Discussion

The Ti grain structures imparted by C-ECAP are comparable to those found by others in UG-Ti [19–21]. Likewise, the enhanced cell attachment and proliferation are consistent with prior cell proliferation studies [7,8]. However, in the present study, the crystallographic textures and surface topographies for the CG and UG states were intentionally made similar. The maximum intensity of the $\{10\overline{1}0\}$ texture was 8.3 times-random for the CG samples, comparable to the 6.7 times-random measured for the CG samples. Since the $\{10\overline{1}0\}$ texture fiber is rotated towards the $\{0001\}$ direction in the UG sample, the intensities of the $\{10\overline{1}0\}$ texture fibers for CG and UG are both between 4 and 6 times-random around the $\langle 10\overline{1}0\rangle$ fiber direction. Thus, prospective effects of differences in crystallographic texture on surface energy were minimized. Likewise, the effects of surface topography were eliminated for the nanometrically smooth substrates.

The areal densities of grain boundaries intersecting the surfaces differed for the CG and UG samples. From the EBSD analysis, an average total grain boundary length per unit area of 0.19 $\mu$m/$\mu$m$^2$ was measured on CG-Ti compared to 12.62 $\mu$m/$\mu$m$^2$ measured on UG-Ti. Thus, the average areal density of grain boundaries is 67.5 times higher on the UG-Ti surface compared to the CG-Ti surface. This difference is the greatest contrast between CG-Ti and UG-Ti in this study.

The average distance between grain boundaries with respect to the average dimensions of attached cells differs greatly for CG-Ti and UG-Ti. Recall that the average grain sizes were 10.9 $\mu$m and 0.24 $\mu$m for CG-Ti and UG-Ti, respectively. The typical widths of attached preosteoblast cells, as seen in the images in Figure 3 were between 6 $\mu$m and 20 $\mu$m.

### Table 2. Measured surface-attached cell and grain boundary parameters.

|          | Mean # of Attached Cells | Avg. Substrate Area Covered Per Cell ($\mu$m$^2$) | Avg. Grain Boundary Length per Unit Area ($\mu$m/$\mu$m$^2$) | GBL/Cell, Avg. Grain Boundary Length per Cell ($\mu$m/cell) |
|----------|--------------------------|-----------------------------------------------|-----------------------------------------------------------|-------------------------------------------------------------|
| CG-Ti, Unpolished | 49                      | 518.2                                         | 0.187                                                    | 97.1                                                        |
| CG-Ti, Polished   | 72                      | 1457.5                                        | 0.187                                                    | 273.0                                                       |
| UG-Ti, Unpolished | 91                      | 807.3                                         | 12.62                                                   | 10,189.3                                                   |
| UG-Ti, Polished   | 149                     | 5615.7                                        | 12.62                                                   | 70,881.5                                                   |
underneath the cells. This trend is present regardless of surface roughness, though polished surfaces consistently displayed higher cell counts and larger cell sizes. We expect that the roughness of the unpolished sample surfaces reduces the access of cells to grain boundaries because of geometric effects of the topography. Analysis of SEM micrographs of cells on the substrates revealed that cells occupy concave valleys in the unpolished surfaces, a location preferable to attachment on the convex peaks of surface striations created by machining of the samples.

Figure 4 displays the variation of average cell count with surface roughness measured by AFM. While the polished samples had essentially identical roughness, with $R_a < 1$ nm, significantly higher cell counts were found on the UG-Ti substrates compared to the CG-Ti substrates. Similarly, the unpolished UG-Ti samples also had significantly higher cell counts, though the $R_a$ values were significantly different, 40.2 nm $\pm$ 24 nm for CG-Ti versus 70.2 nm $\pm$ 36 nm for UG-Ti. These results are congruent with Medvedev, et al. for Grade 4 Ti [16] and Baek et al. for Grade 2 Ti [17]. However, in the experiments reported here the effects of surface roughness are absent in the polished samples and cannot contribute to the difference we found in the cell counts for UG and CG samples.

The observed correlation between grain boundary length and cell proliferation shown in Figure 5 reshapes our outlook on hypotheses for the underlying causes and mechanisms. One explanation for this correlation is the prospective influence of the higher energy of grain boundary regions compared to the intra-grain matrix. These energy differences have the potential to alter the work function and distribution of charge within the surface oxide, which may influence the adsorption of proteins or interaction with transmembrane adhesion proteins such as integrins. This possibility has been explored by Ercan, et al. [22]. Also, it is notable that 65.6% of the grain boundaries on UG-Ti had intercept lengths less than 82 nm. This population of closely spaced grain boundary regions on the UG-Ti samples overlaps with the expected average separation distance of cell focal adhesions and their integrin ligands. Le Saux et al. reported this separation to be 44 nm for endothelial cells [23]. This connection supports a hypothesis that grain boundaries have a greater opportunity to influence mechanotransduction and signaling on smooth UG-Ti. This mechanism is also consistent with results from a companion study of the sequencing of the RNA from the cells grown on nanometrically smooth UG-Ti and CG-Ti surfaces. These experiments implicate an interaction of integrins with grain boundaries as the factor that increases cell proliferation through established mechanosensation and transduction pathways (to be reported in a separate manuscript in preparation).

Another alternative mechanism by which grain boundaries may influence cell proliferation is the possibility of microsegregation of alloying elements or impurities. Locally high concentrations of these elements could present chemical differences in grain boundary regions. However, in our study Energy Dispersive Spectroscopy (EDS) measurements at 20,000x to 35,000x magnifications did not show evidence of microsegregation of Fe, O, N, or C on any substrate (data not shown).

Following the work of Ercan et al [22], it is plausible that the underlying mechanisms and effects of varying surface roughness differ for surfaces with roughness $R_a$ in the 1–10 nm range compared to the 10–100 nm range. However, Figure 5 shows that the number of cells relative
to the grain boundary length correlates with cell proliferation for the entire range of surface roughness in the present study. For the polished samples with $R_a < 1\text{ nm}$, the grain boundaries presented no significant topographical features to proteins or cells on the surface, indicating that the grain boundaries themselves have a role in cell attachment and proliferation. Additional experiments are needed with more levels of surface roughness and grain boundary densities to verify this conclusion. However, the present experiments are the first to isolate the positive correlation of grain boundary length with cell attachment and proliferation through the parameter GBL/Cell which represents the average total grain boundary length per surface-attached cell.

**Conclusion**

Preosteoblasts from the MT3T3-E1 line were cultured for 72 h on four states of CP Ti substrates: CG and UG, and with or without polishing to nanometric smoothness. The mean numbers of surface-attached MT3T3-E1 cells on UG-Ti were 1.9-fold and 2.1-fold higher compared to CG-Ti after 72 h of cultivation on unpolished and polished substrates, respectively. The combination of having ultrafine grains and nanosmooth surfaces produced the greatest increase in the number of surface-attached cells, by a factor of 3.04 compared to unpolished CG-Ti. A new biophysical parameter, the average grain boundary length per surface-attached cell (GBL/Cell), was proposed to explain the results.

Grain boundary length per cell was computed by combining data from image analysis of surface-attached cells with the EBSD data on grain boundary length. The average grain boundary length per unit surface area was 13.8 times greater for UG-Ti compared to CG-Ti. The average number of attached MT3T3-E1 cells increased monotonically with GBL/Cell. A linear correlation was found between the average number of surface-attached cells and the logarithm of GBL/Cell (linear least square correlation coefficient $R^2 = 0.88$), showing that cell attachment and proliferation correlate with the density of grain boundaries on the surface of Ti.

**Acknowledgements**

Partial funding was provided by the National Center for Research Resources: [Grant Number 5P20RR016480-12]; National Institute of General Medical Sciences: [Grant Number 8 P20 GM103451-12].

**Disclosure statement**

No potential conflict of interest was reported by the author(s).

**Funding**

This work was supported by National Center for Research Resources: [Grant Number 5P20RR016480-12]; National Institute of General Medical Sciences: [Grant Number 8 P20 GM103451-12].

**ORCID**

Terry C. Lowe http://orcid.org/0000-0001-9288-1076

**References**

[1] Buser D, Sennerby L, De Bruyn H. Modern implant dentistry based on osseointegration: 50 years of progress, current trends and open questions. Periodontol 2000. 2017;73(1):7–21.
[2] Rack H, Quzi J. Titanium alloys for biomedical applications. Mater Sci Eng C. 2006;26:1269–1277.
[3] Kunčická L, Kocich R, Lowe TC. Advances in metals and alloys for joint replacement. Prog Mater Sci. 2017;88:232–280.
[4] Wang M. Surface modification of metallic biomaterials for orthopaedic applications. Mater Sci. 2009;618–619:285–290.
[5] Valiev RZ, Estrin Y, Horita Z, et al. Fundamentals of Superior Properties in Bulk NanoSPD Materials. Mater Res Lett. 2016;4(1):1–21.
[6] Balyanov A, Kutnyakova J, Amirkhanova NA, et al. Corrosion resistance of ultra fine-grained Ti. Scr Mater. 2004;51(3):225–229.
[7] Lowe TC, Reiss RA. Understanding the biological responses of nanostructured metals and surfaces. IOP Conf Ser Mater Sci Eng. 2014;63(1):1–17.
[8] Bagherifard S, Ghelichi R, Khademhosseini A, et al. Cell response to nanocrystallized metallic substrates obtained through severe plastic deformation. ACS Appl Mater Interfaces. 2014;6(11):7963–7985.
[9] Raab GI, Valiev RZ, Lowe TC, et al. Continuous processing of ultrafine grained Al by ECAP-Conform. Mater Sci Eng a-Structural Mater Prop Microstruct Process. 2004;382(1–2):30–34.
[10] Raab GI, Valiev RZ, Gunjurov DV, et al. Long-length ultrafine-grained titanium rods produced by ECAP-Conform. In: Estrin Y, Maier HJ, editors. Nanomaterials by severe plastic deformation IV, Pts 1 and 2. Vol 584–586. Weinheim, Germany: Wiley-VCH; 2008. p. 80–85.
[11] Valiev R, Islamgaliyev R, Alexandrov I. Bulk nanostructured materials from severe plastic deformation. Prog Mater Sci. 2000;45(2):103–189.
[12] Misra RDK, Thein-Han WW, Pesacreta TC, et al. Cellular response of preosteoblasts to nanograined/ultrafine-grained structures. Acta Biomater. 2009;5(5):1455–1467.
[13] Puckett SD, Taylor E, Raimondo T, et al. The relationship between the nanostructure of titanium surfaces and bacterial attachment. Biomaterials. 2010;31(4):706–713.
[14] Faghihi S, Vali H, Tabrizian M. Effects of crystal size and orientation of substrates on cell adhesion: Implication for medical implants. Int J Mod Phys B. 2008;22(18–19):3069–3081.
[15] Hoseini M, Bocher P, Azari F, et al. Effects of grain size and texture on the biocompatibility of commercially pure titanium. In: Tewari A, Suwas S, Srivastava D, editors. Textures of Materials, Pts 1 and 2. Vol 702–703. Stafa-Zurich, Switzerland: Trans Tech Publications; 2012. p. 822–825.

[16] Medvedev AE, Neumann A, Ng HP, et al. Combined effect of grain refinement and surface modification of pure titanium on the attachment of mesenchymal stem cells and osteoblast-like SaOS-2 cells. Mater Sci Eng C. 2017;71:483–497.

[17] Baek SM, Shin MH, Moon J, et al. Superior pre-osteoblast cell response of etched ultrafine-grained titanium with a controlled crystallographic orientation. Sci Rep. 2017;7(1):44213.

[18] Schneider CA, Rasband WS, Eliceiri KW. NIH image to ImageJ: 25 years of image analysis. Nat Methods. 2012;9(7):671–675.

[19] Stolyarov VV, Zhu YT, Lowe TC, et al. Microstructure and properties of pure Ti processed by ECAP and cold extrusion. Mater Sci Eng A. 2001;303(1–2): 82–89.

[20] Gunderov DV, Polyakov AV, Semenova IP, et al. Evolution of microstructure, macrotexture and mechanical properties of commercially pure Ti during ECAP-conform processing and drawing. Mater Sci Eng A-Structural Mater Prop Microstruct Process. 2013;562:128–136.

[21] Dyakonov GS, Zemtsova E, Mironov S, et al. An EBSD investigation of ultrafine-grain titanium for biomedical applications. Mater Sci Eng A. 2015;648:305–310.

[22] Ercan B, Khang D, Carpenter J, et al. Using mathematical models to understand the effect of nanoscale roughness on protein adsorption for improving medical devices. Int J Nanomedicine. 2013;8(Suppl 1):75–81.

[23] Le Saux G, Magenau A, Gunaratnam K, et al. Spacing of integrin ligands influences signal transduction in endothelial cells. Biophys J. 2011;101(4):764–773.