Nanotubular Oxide Layer Formed on Helix Surfaces of Dental Screw Implants

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Abstract: Surface modification is used to extend the life of implants. To increase the corrosion resistance and improve the biocompatibility of metal implant materials, oxidation of the Ti-13Nb-13Zr titanium alloy was used. The samples used for the research had the shape of a helix with a metric thread, with their geometry imitating a dental implant. The oxide layer was produced by a standard electrochemical method in an environment of 1M H3PO4 + 0.3% HF for 20 min, at a constant voltage of 30 V. The oxidized samples were analyzed with a scanning electron microscope. Nanotubular oxide layers with internal diameters of 30–80 nm were found. An analysis of the surface topography was performed using an optical microscope, and the Sa parameter was determined for the top of the helix and for the bottom, where a significant difference in value was observed. The presence of the modification layer, visible at the bottom of the helix, was confirmed by analyzing the sample cross-sections using computed tomography. Corrosion tests performed in the artificial saliva solution demonstrated higher corrosion current and less noble corrosion potential due to incomplete surface coverage and pitting. Necessary improved oxidation parameters will be applied in future work.

Keywords: nanotubular oxide; helix surfaces; dental implants; roughness; corrosion properties

1. Introduction

Titanium and its alloys are nowadays among the most popular biomaterials, called the “gold standard” for endosseous dental implants, even if some adverse reactions may be expected. They possess a lot of important properties, such as their low density, suitable fatigue strength, Young’s modulus and specific tensile strength, high resistance to brittle cracking, high corrosion resistance, and the best biocompatibility. Despite that, titanium and its alloys need surface modifications for early osseointegration [1,2]. The type of commercial implant determines surface topography and differences in geometry [3].

Surface modification is nowadays an obligatory treatment of dental implants. Bioactivity of the surface resulting in adhesion of osteoblasts and bone ingrowth can be achieved by the development of surface roughness, creation of bioactive films, and deposition of coatings [2]. Many different methods have been used to change the surface roughness of dental implants, including mechanical techniques such as grinding, polishing, machining, sandblasting and attrition, chemical etching in acids, alkali and fluorides, electrophoretic deposition, and laser treatments [4–11].

The interaction of cells and adsorption of proteins depends on surface structure and is significant in the presence of nanometric pores, which increase the rate of osseointegration and biomechanical fixation [2,7,8,12–17]. A significantly higher bone contact of 27% (p < 0.05) was observed in nanotextured compared to machined implants [18]. However, reproducibility of nanoscale surface profiles of titanium with chemical modifications such as acid-etching is quite difficult to achieve and unreliable, and
knowledge on the ideal surface roughness parameters for rapid osseointegration is still lacking [19,20].

Among various surface treatments, artificial oxidation seems particularly plausible for titanium dental implants resulting in high corrosion resistance and biocompatibility [21]. The oxidation required for dental implants is currently mostly applied by micro-arc oxidation (MAO) [22,23] and electrochemical oxidation [24]. Gaseous oxidation has also been proposed [25]. MAO induced titanium oxide formation in the anatase crystalline phase and also incorporated Ca, P, and Mg in the film [26–33]. An oxide thickness of 600–1000 nm demonstrated significantly stronger bone responses than that of 17 or 200 nm [30]. The coatings comprising nanoTiO2 and nanohydroxyapatite (nanoHAp) demonstrated a torque value of coated screws significantly greater than that of nanoHAp covered screws [34].

The creation of nanotubular oxide layers on titanium and its alloys is well-known. The formation of nanotubular oxide structures on dental implants has not often been investigated and developed. The overly short life of dental implants observed proves the ineffectiveness of the applied surface modifications and provides prompts for further research. The anodization depends to a great extent on the geometry and structure of the surfaces involved. Indeed, the formation of titanium dioxide nanotubes on flat titanium surfaces, provided by well-known suppliers, does not have the same effect on titanium implants, mainly due to the geometry of the implant, which changes the priority, intensity, and interconnection of the electrochemical processes [35]. Nanotubular oxide layers have been reported to increase the bioactivity of titanium implants [36–38], the nucleation and growth of hydroxyapatite coatings [39], and to introduce antibacterial effects after loading the nanotubes with drugs [40,41]. Such a type of surface was already fabricated on the nontoxic Ti-13Zr-13Nb alloy investigated here [42,43]. This research was aimed at an assessment of the creation of nanotubular oxide layers on screw fixed dental implants and the characterization of the layers obtained on the tops and bottoms of helices of implants.

2. Materials and Methods

2.1. Material

The biphase $\alpha + \beta$ Ti-13Zr-13Nb alloy (SeaBird Metal Materials Co., Baoji, China) with chemical composition presented in Table 1 was investigated in the as-received state.

Table 1. The chemical composition of the Ti-13Zr-13Nb alloy, wt.%. (based on the manufacturer’s certificate).

| Element | Zr | Nb | O  | C  | N  | Ti          |
|---------|----|----|----|----|----|-------------|
| wt.%    | 13.0| 13.0| 0.11| 0.04| 0.019| remainder   |

2.2. Preparation of Specimens

Round specimens of height 9 mm and diameter 8 mm were prepared by precision milling. The metric thread was cut on all specimens. The remaining impurities were cleaned and the surface was prepared by sand blasting with corundum for 15 s. The cleaning was performed at Aesculap Chifa Ltd. in Nowy Tomysl, Poland. Immediately before oxidation, the specimens were washed in an ultrasonic bath (Sonic 3, Polsonic, Warsaw, Poland) in isopropanol (POCH, 99.8%, Gliwice, Poland) for 10 min, in distilled water for 3 min, and methanol (POCH, 99.8%, Gliwice, Poland) for 10 min.

2.3. Electrochemical Oxidation

The oxidation was performed using a direct current power supply (MCM/SPN110-01C, Shanghai MCP Corp., Shanghai, China). The specimen tested was connected to the power supply as an anode and the Pt electrode was used as a cathode. The electrolytic bath contained a solution of 150 mL of distilled water, 20 mL of 1 M H3PO4, and 1.5 mL of
0.3% HF (both from POCH, Gliwice, Poland). The distance between the electrode tested and the Pt electrode was 15 mm. The solution was neither aerated nor deaerated, and non-stirred. The experiments were carried out at ambient temperature. The experiments were performed at a voltage of 20 V for 30 min based on previously conducted experiments [42].

2.4. SEM Surface Examination

Scanning electron microscopy (SEM JEOL JSM-7800 F, JEOL Ltd., Tokyo, Japan) instrument equipped with EDS chemical analyzer (Edax Inc., Mahwah, NJ, USA).

2.5. Light Microscopy Assessment of Roughness

A light microscope (VHX-7000, Keyence, Osaka, Japan) was applied to examine the surface topography. Roughness parameters determined by the 3D Form Measurement software were applied to the Sa area.

2.6. Computer Tomography

Tomographic images were obtained using Phoenix v/Tome/xs computer tomography (General Electric, Lewistown, PA, USA).

2.7. Corrosion Examinations in Simulated Body Fluid

Corrosion tests were performed by a potentiodynamic method in simulated body fluid (SBF) at a temperature of 38 °C. The electrochemical measurements were achieved by using a potentiostat/galvanostat (Atlas 0531, Atlas Sollich, Gdańsk, Poland). An artificial saliva solution (SBF) was prepared according to EN ISO 10993-15 [44] by dissolving reagent grade chemical (NH2)2CO (0.13 g L⁻¹), NaCl (0.7 g L⁻¹), NaHCO3 (1.5 g L⁻¹), Na2HPO4 (0.26 g L⁻¹), K2HPO4 (0.2 g L⁻¹), KSCN (0.33 g L⁻¹), KCl (1.2 g L⁻¹) (POCH, Gliwice, Poland). The potential was measured vs. a saturated calomel electrode (SCE) located in the Haber-Luggin capillary. As a counter electrode, a standard platinum electrode was used. The test specimen was stabilized in a solution of artificial saliva for 30 min at open circuit potential OCP. The potential change rate was 1 mV/s within a scan range of −2000 to 1000 mV. The solutions were agitated with a magnetic stirrer. Using the Tafel extrapolation method, the corrosion potential (E_corr) and corrosion current density (i_corr) values were determined.

3. Results and Discussion

3.1. Substrate Specimens

The surface of the non-oxidized alloy is shown in Figure 1 at two different magnifications. The relatively smooth surface and screw lines can be seen.

![Figure 1](image1.png)

Figure 1. Surface of reference specimen at different magnification: (a) 130×, (b) 1700×.

3.2. Oxidized Specimens

Figure 2 shows the appearance of the nanotubular oxide layer only in the area at the bottom of the helix. The pores created are spherical and longitudinal. They possess a diameter ranging between 30 and 80 nm. The layer is well adjacent to the substrate and it has a small number of cracks and surface defects. The gradual decrease of the nanoporous
layer and its absence at the top of the helix may be attributed to different current densities, different electrochemical potential, and as a consequence a different course of chemical reactions. The current is screened at the bottom at a given potential and the resultant value is sufficient for electrochemical oxidation to occur. At the tops, the current density is too high and the nanotubes formed undergo fast oxidation, its rate exceeding that of chemical dissolution resulting in nanotubes. The current density is higher at the tops of such surfaces, with the effect attributed to the difficult transport of oxygen to and reaction products from this area, and stepwise depolarization of the area close to the bottom followed by a change in open circuit and corrosion potentials. These processes can shift the current and potential values beyond those necessary to form nanotubular oxide layers.

The EDS examination results presented in Table 2 suggest the obtaining of a layer of titanium oxide on the surface, which is confirmed mainly by the content of titanium and oxygen. High P content results from the absorption of HPO₄²⁻ anion within the layer pores and it is desired for better bioactivity of the surface. Trace amounts of Ca, K, Fe are observed, which most likely were impurities in the distilled water used.

**Table 2.** The EDS examination results of the chemical composition of the oxide layer.

| Element | Wt. Pct. | At. Pct. |
|---------|----------|----------|
| O       | 48.49    | 71.65    |
| Zr      | 5.72     | 2.07     |
| P       | 19.29    | 14.72    |
| Nb      | 6.91     | 1.76     |
| K       | 0.68     | 0.41     |
| Ca      | 0.84     | 0.49     |
| Ti      | 17.77    | 8.77     |
| Fe      | 0.30     | 0.13     |

Results of topography tests are presented in Figure 3. The surface of the top oxide layer is rough and well developed. The roughness profile is 630 nm, the Sa average value
is 1.39 ± 0.79 µm on the tip of the helix, and 5.69 ± 2.98 µm on the bottom of the helix (Table 3). Such values in the nanometric range are also useful. The differentiation of the area comprising small nanotubes and rough pores is important. Surface modification led to smoothing the tip of the helix as a result of dissolving roughness peaks. The influence of roughness on osteointegration has been proven. In the case of long-term implants, a positive osteoblast response is required. With increasing roughness, the possibility of osteoblasts settling increases [45]. High roughness also carries the risk of biofilm formation [46]. The topography results confirmed the obtaining of a surface with a high surface roughness value. The lowest values were obtained for the oxidized sample.

| Specimen                                | Tip of the Helix | Bottom of the Helix |
|-----------------------------------------|------------------|---------------------|
| Reference Ti-13Zr-13Nb (before sand blasting) | 1.81 ± 1.11      | 9.10 ± 4.62         |
| Ti-13Zr-13Nb (after sand blasting)      | 1.63 ± 1.40      | 9.94 ± 5.51         |
| Oxidized Ti-13Zr-13Nb                   | 1.39 ± 0.79      | 5.69 ± 2.98         |

The CT investigations showed the appearance of modifications at the bottoms of the helix and not at the tops (Figure 4). The area of modification can be observed as grey and red areas at the bottoms and base alloy as white metal.

![Figure 3. The topography of reference Ti-13Zr-13Nb before (a) and after (b) sand blasting and oxidized Ti-13Zr-13Nb surfaces (c) obtained by light microscopy; the bottom of the helix (left) and top (right).](image-url)
The corrosion results are presented in Figure 5 and Table 4. The creation of the oxide layer became difficult because of much higher current values and a presumed shift of electrochemical potential into more anodic values resulting in the dissolution of metal rather than the oxidation of the surface. The local appearance of the nanotubular and highly rough surface is evidence that some microcells are formed due to change in potential. The local changes in pH value influence the anodization rate, the thickness of the oxide layer and its structure, or even its formation. In case of too low or too high acidity, the oxide layer is unable to achieve the nanotubular structure [47]. Here the anodization was made at the proper HF concentration enhancing the stabilization of the appropriate low pH value and resulting in a short oxidation time, thin nanotubes, a short distance between them, and scarce surface cracks. The roughness of the oxidized surface was close to that observed in similar experiments [48].

Figure 5. Potentiodynamic polarization curve of reference and oxidized Ti-13Zr-13Nb specimens.
Table 4. Corrosion properties of reference and oxidized Ti-13Zr-13Nb specimens.

| Specimen          | Current Density (nA/cm²) | Corrosion Potential (mV) |
|-------------------|--------------------------|--------------------------|
| Reference Ti-13Zr-13Nb | 503.25                  | -392.46                  |
| Oxidized Ti-13Zr-13Nb | 1451.00                 | -1174.13                 |

The open-circuit potential (OCP) of the non-oxidized specimen was about −199 mV/SCE. The anodic polarization exhibits a narrow plateau between 300 and 1150 mV, which can be attributed to the presence of natural titanium oxide on a specimen surface. The passive current value in this area ranged between 200 and 300 μA. For the previously oxidized specimen, the OCP was about −616 mV. The anodic curve shows a very stable passive region between −200 and 2000 mV. The passive current was about 200 μA in the entire region. However, despite high passive regions, the increase in corrosion current density after oxidation shows that the surface has not been uniformly covered with oxide layers and many microcells could appear in these oxidation conditions. The titanium dioxide layer formed on the surface of the titanium can provide increased corrosion resistance only if it is continuous over the entire surface of the alloy. The layer presented in the paper is characterized by cracks and a lack of continuity. This results in the formation of so-called “corrosion channels”, which accelerate the degradation of the material. A similar effect was obtained in research [49]. The occurrence of this phenomenon may explain the deterioration of the corrosive properties compared to the reference sample.

The microscopic investigation of reference specimens after corrosion tests showed effects of pitting, some discontinuity of material, and a heterogeneous structure at the bottoms of the helix and at the tops (Figure 6). The microscopic investigation specimens oxidized in an electrochemical way and showed a network of cracks in the surface of the helix top and corrosion pitting in the bottom of the helix (Figure 7), it is probably related to the grater thickness of the obtained modification, which was confirmed by CT tests—Figure 4b.

![Figure 6](image-url)  
**Figure 6.** Surface of reference specimen after corrosion test at different magnifications: (a) surface of helix top ×200, (b) surface of helix top ×1000, (c) bottom of the helix ×200, (d) bottom of the helix ×1000.
**Figure 7.** Surfaces of specimens oxidized in an electrochemical way after corrosion tests: (a) surface of helix top ×200, (b) surface of helix top ×1000, (c) bottom of the helix ×200, (d) bottom of the helix ×1000.

**4. Conclusions**

In summary, nanotubular oxidation on the helix lines of titanium dental implants is possible, but it depends heavily on the geometric shape of the implant, anodization parameters, and environment composition. The parameters proposed here make it possible to obtain the nanotubes on the bottom of the helix and distinctly roughen almost all remaining surfaces. However, the applied conditions applied indicate that future investigations be oriented towards oxidation of the whole surface by introducing slightly higher HF contents and lower current values and mixing the electrolyte bath.

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