Surface Properties and Osteoblastic Cytocompatibility of Two Blasted and Acid-Etched Titanium Implant Systems with Distinct Microtopography

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ABSTRACT

Objectives: The aim of this study is to compare two commercially available screw-type sandblasted and acid-etched (SLA) Ti implant systems from Eckermann Laboratorium S.L., with similar geometry and distinct microtopography, regarding surface properties and osteoblastic cytocompatibility.

Material and Methods: Implant I (referred as a conventional SLA system) and Implant II (a system patented as Eckcyte®) were characterized for macro and microtopography, surface roughness and chemical composition. For the cytocompatibility studies, human bone marrow osteoblastic cells were seeded over the implants’ surface, and the cell response was assessed for cell adhesion and proliferation, alkaline phosphatase (ALP) activity and matrix mineralization.

Results: Implant I presented a rough surface with irregularly shaped and sized cavities among flatter-appearing areas, whereas Implant II exhibited a homogeneous rough microporous surface. Compared to Implant I, Implant II presented higher Ra values (0.8 [SD 0.008] μm and 1.21 [SD 0.15] μm, respectively, P < 0.05) and also increased values of Rz, Rt and Rsm, a more negative value of Rsk, and similar RKu values. XPS showed the expected presence of Ti, O, C and N; Al, Si, F, P and Ca were detected in low concentrations. Implant II exhibited significantly lower Al levels. Both implants supported the adhesion, proliferation and differentiation of osteoblastic cells. Implant II showed a thicker fibrilar cell layer and an earlier onset and more abundant matrix mineralization.

Conclusions: The homogeneous rough and microporous surface of Implant II is most probably a main contributor for its improved cell response.

Keywords: dental implants; surface properties; bone marrow; osteoblasts; differentiation cell; cell culture.

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INTRODUCTION

Titanium (Ti) screw-shaped dental implants are biocompatible, have high corrosion resistance, are lightweight and durable, and can be easily prepared in many different shapes and textures without affecting their biocompatibility [1,2]. Ti surface topography is an essential feature in implant design, but the optimal surface texture remains to be established [3,4], due to the difficulty in correlating surface properties with clinical results [5,6]. In addition to the wide range of possibilities in creating complex topographies, other factors in implant design, namely geometry, i.e. distance between peaks, curvature of the valleys and relative distribution of flat and smooth regions, are also relevant for the short and long-term clinical outcome [1]. Also, other aspects not related with topography, material itself or fabrication process may significantly affect the clinical performance. Each patient exhibits a specific answer to a particular implant depending on bone characteristics, patient’s health status and surgical technique [6,7]. However, studies performed on animals and representative cell culture systems continually provide important information concerning the relevance of surface properties and biological performance of the implants [8-10].

In this context, a wide variety of studies have shown that early fixation and long-term stability is improved by a high roughness profile, compared to a smooth surface, due to the mechanical interlocking between the implant surface and bone ingrowth [11]. It was reported that surface roughness in the range of 1 - 10 μm maximizes the bone/implant interlocking [3,12]. However, a high roughness appears to be associated with the risk of increased peri-implantitis and ionic leakage [13]. Albrektsson and Wennberg have shown that a moderate roughness of 1 - 2 μm may limit this process and improve implant performance [2].

Microtopographic features were also found to play a decisive role on the biological response to dental implants [14-16]. Discontinuities or curvatures associated with microtopographic features seem to induce local changes in the surface free energy, thus modifying the amount and type of adsorbed proteins that can modulate the local cellular response [16]. In vitro studies revealed that topographic properties were found to affect several cellular events, including cell attachment, alignment, direction of proliferation, growth rate and metabolism, and thus influence the osseointegration process [14,15,17]. A theoretical approach suggested that the ideal surface should be covered with hemispherical pits approximately 1.5 μm in depth and 4 μm in diameter [18].

Eckermann Laboratorium S.L. (Alicante, Spain) developed two implant systems with different surfaces. One system is referred by the manufacturer as a conventional SLA surface. The other system, which was launched later and patented as Eckcyte®, is claimed to present improved surface properties. It is important to examine those surface characteristics and osteoblastic cytocompatibility of these two implant systems, in order to relate and compare the two parameters. The aim of this study is to compare two commercially available screw-type sandblasted and acid-etched (SLA) Ti implant systems from Eckermann Laboratorium S.L., with similar geometry and distinct microtopography, regarding surface properties and osteoblastic cytocompatibility.

MATERIAL AND METHODS

Ti Implants

The implants tested in this study were two commercially screw-type sandblasted and acid-etched Ti implant systems manufactured by Eckermann Laboratorium S.L. (Alicante, Spain). The implants were sandblasted with alumina (Al₂O₃) particles, followed by an acid etching treatment involving HCl, H₂SO₄ and HF solutions, and differ on the experimental conditions of the blasting and the etching protocols, however, details of the manufacturing process were not available. The implants will be referred as Implant I (the implant mentioned as a conventional SLA system) and Implant II (the system patented as Eckcyte®). Implants I and II were 3.0 mm in diameter and 8 and 13 mm in length, respectively. The implants were removed from their sterile package just before surface and cytocompatibility studies.

Surface characterization of Ti implants

The surface of the threaded parts of Implants I and II was assessed for macro and microtopography, surface roughness and chemical composition.

Macro and microtopography

Morphology and texture were assessed by scanning electron microscopy (SEM) with a JEOL JSM 6301F scanning electron microscope equipped with an X-ray energy dispersive spectroscopy (EDS) microanalysis capability (Voyager XRMA System, Noran Instruments, Madison, WI, USA). Implants were fixed to an aluminium sample holder on a conducting carbon surface. Observation was performed at 15 kV, and magnification varied from x30 to x2000.
Surface roughness and optical profilometry

Roughness measurements were performed with a T8000 Hommel profiler (Hommel AG, Hamburg, D). The measured length was 1.00 mm, cut-off was 0.08 mm, the radius tip was 5 mm, and the filter used was M1 (according to ISO 11562). The measurements were performed in randomly selected regions; in each region, three measurements were done. The implants were assessed for Ra, Rt, Rz, Rsm, Rku and Rsk. The three-dimensional-surface roughness was assessed with an optical profiler coupled to a confocal laser scanning microscope (CLSM, Leica SP2 AOBSE, Leica Microsystems, Germany). The images were taken on the peak and on the valley of the thread, and were worked out with a specific software (Leica Microsystems Heidelberg GmbH, 2.61 Build 1538 LCS Lite, Leica Microsystems, Germany).

Chemical composition

The chemical composition was analysed by X-ray photoelectron spectroscopy (XPS, ESCALAB 200A, VG Scientific, UK) by using an Al source operated at 15 KV and 300W and a pressure lower than 10^-6 Pa. The area of the surface analysed was approximately 10 mm². This analysis allows for the identification of residual contamination and for the semi-quantification of the composition of the external layers of the surface. The following peaks were considered: C 1s, O 1s, Ti 2p, Al 2p, N 1s, Si 2p, F 1s, P 2p, Ca 2p and Pb 4f.

Osteoblastic cytocompatibility of Ti implants

Human bone marrow cell cultures

Cytocompatibility studies were performed with human bone marrow osteoblastic cell cultures. Bone marrow was obtained from orthopaedic surgery procedures, after Local Ethical Committee approval and patient informed consent. Bone marrow was cultured in α-Minimal Essential Medium (α-MEM) containing 10% foetal bovine serum, 50 μg/ml ascorbic acid, 100 IU/ml penicillin, 2.5 μg/ml streptomycin and 2.5 μg/ml fungizone, at 37 °C in a humidified atmosphere of 5% CO₃ in air.

Primary cultures were maintained until near confluence (10 - 15 days) and, at this stage, adherent cells were enzymatically released (trypsin-EDTA solution). Second-passage cells were seeded (5 x 10⁴ cell/cm²) over the surface of the implants. Seeded implants were cultured for 30 days in the presence of 50 μg/ml ascorbic acid, 10 mM β-glycerophosphate and 10 nM dexamethasone.

Colonized implants were evaluated for cell adhesion, pattern of cell growth, alkaline phosphatase (ALP) activity and matrix mineralization.

Alkaline phosphatase activity (ALP)

The colonized materials were treated with 0.1% triton in water (to lyse the cell layer) and the cell lysates were evaluated for ALP activity, at days 7, 14, 21, 24 and 28. ALP was assayed by the hydrolysis of p-nitrophenyl phosphate in alkaline buffer solution, pH 10.3, 30 min at 37 °C, and colorimetric determination of the product (p-nitrophenol) at λ = 405 nm, in an ELISA plate reader (Synergy HT, Biotek, Vermont, USA). Enzyme activity was normalized to total protein content, at the same time-points. The total amount of protein was assayed by the Lowry’s method with bovine serum albumin used as a standard. ALP activity is expressed as nanomoles of p-nitrophenol produced per min per μg of protein (nmol.min⁻¹/μg protein).

Scanning electron microscopy (SEM) observation

SEM was performed at 3 and 24 h and 28 days. Seeded implants were fixed (1.5% glutaraldehyde in 0.14 M sodium cacodylate buffer, pH 7.3, 10 min), dehydrated in graded alcohols, critical-point dried, sputter-coated with gold and analysed in a JEOL JSM 6301F scanning electron microscope.

Concentration of ionized calcium in the culture medium

Culture medium from the cultured implants was collected every 2 - 3 days (and cultures were refed with fresh medium). Analysis of ionized calcium (Ca_i) content in the culture medium was conducted using Sigma Diagnostics Kit (Sigma-Aldrich Corporation, St. Louis, MO, USA) procedure number 587. Results were expressed as millimoles of Ca_i per litre of medium (mmol/l).

Statistical analysis

Results regarding surface characterization and cytocompatibility studies were obtained from three independent experiments. In each experiment, and
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for each time-point, 6 implants were characterized. In the cytocompatibility studies, the three experiments were performed with cell cultures established from different donors. Quantitative data are presented as mean (SD). Groups of data were evaluated using one-way analysis of variance (ANOVA) and Bonferroni procedure for post-hoc comparison using SPSS (Apache software foundation, version 15.0). Values of $P \leq 0.05$ were considered significant.

RESULTS

Surface characterization

Macro and microtopography

Implant I and Implant II contained 6 and 11 threads, respectively, and a similar spacing and peak-valley distance (Figure 1A, B). At low magnification SEM images, both implants showed a relatively homogeneous appearance (Figure 1A - D). At high magnification (Figure 1E - H), Implant I presented a rough surface with irregularly shaped and sized cavities among flatter-appearing areas of various sizes. Implant II showed a regular rough surface with a relatively homogeneous topography, displaying abundant and randomly distributed rounded pits with an average diameter of 1 - 4 μm.

Surface roughness

The roughness profiles of the implants (Figure 2A) showed that Implant II exhibited higher amplitude of peak to valley distances, compared to Implant I. Regarding the roughness parameters (Figure 2B), Implant II presented Ra values higher than Implant I (1.21 [SD 0.15] μm and 0.8 [SD 0.08] μm, respectively.) Also, increased values of Rz, Rt and Rsm were found on Implant II. Rsk was more negative on Implant II, but the two implants showed similar RKu values.

Observation of the three-dimensional-topometric images of the implants’ surface (Figure 3) showed that Implant I displayed a higher variation in the peak protuberances, compared to Implant II.

| Threaded Region | Implant I | Implant II |
|----------------|-----------|------------|
| Ra (μm)        | 0.80 ± 0.08 | 1.21 ± 0.15 |
| Rz (μm)        | 4.97 ± 0.68 | 7.08 ± 0.77 |
| Rt (μm)        | 6.25 ± 1.29 | 9.03 ± 1.71 |
| Rsm (μm)       | 0.02 ± 0.01 | 0.03 ± 0.00 |
| Rsk            | -0.11      | -0.22      |
| Rku            | 3.27       | 3.34       |

Significantly different from Implant I.

Figure 1. SEM images of the geometry (A, B) and surface topography at low (C, D) and high (E - H) magnification of the threaded regions of Implants I and II. Implants presented a similar geometry, but a distinct microtopography. Implant I presented an irregularly rough morphology with different sized cavities and flat areas and Implant II exhibited a relatively homogeneous microporous surface.

Figure 2. Roughness profiles (A) and parameters (B) of the threaded regions of Implants I and II. Implant II showed higher amplitude of peak to valley distances (A) and higher values of Ra, Rz, Rt, Rsm and Rsk (B).
Figure 3. Three-dimensional-optical profilometric profiles of the threaded regions of Implants I and II. Evidence of a more homogeneous rough surface was noticed for Implant II.

Table 1. Semi-quantitative XPS analysis of the Ti implants (at %)

|        | C 1s      | O 1s      | Ti 2p      | Al 2p      | N 1s      | Si 2p      | F 1s      | P 2p      | Ca 2p      |
|--------|-----------|-----------|------------|------------|-----------|------------|-----------|-----------|------------|
| Implant I | 45.13 (3.47) | 35.74 (2.21) | 8.28 (0.72) | 5.13 (0.72) | 3.14 (0.29) | 1.36 (0.15) | 0.55 (0.08) | 0.50 (0.07) | 0.17 (0.03) |
| Implant II | 60.66 (7.40) | 26.79 (4.93) | 8.06 (1.86) | 1.02 (0.56) | 1.79 (0.22) | 0.75 (0.15) | 0.21 (0.06) | 0.18 (0.02) | 0.56 (0.16) |

*Significantly different from Implant I (P ≤ 0.05).
M (SD) = mean (standard deviation).
Chemical composition

XPS representative spectra and results of the elemental atomic percentage (Figure 4 and Table 1, respectively) showed the presence of Ti (Ti 2p doublet, at ~459 eV), O (O 1s peak, at ~531 eV), C (C 1s, at ~285 eV) and N in higher percentages; other elements were present in significantly lower concentrations, i.e. Al, Si, F, P and Ca. Implants I and II had similar percentages of Ti and O (Table 1). Regarding the elements found in low percentages, the most evident difference between the two implants was the significantly lower percentage of Al on the surface of Implant II (Table 1). Pb was not detected. EDS spectra provided similar information regarding the qualitative composition of the implants (Figure 5).

Osteoblastic cytocompatibility

Cell adhesion and proliferation

Human osteoblastic bone marrow cells were cultured over the surface of the implants, and the colonized implants were characterized for proliferation and differentiation events (Figures 6 - 8). In both implants, low magnification SEM images (Figure 6A, B) showed adhered cells 3h after plating, whereas at high magnification (Figure 6C, D), cells exhibited a round central shape with long cytoplasmic extensions and were able to adapt to the underlying irregular surface topography. At day 28, SEM images showed that the implants’ surface was completely covered by a cell layer (Figure 6E, F). However, a thicker cell layer

![Figure 5](https://www.ejomr.org/JOMR/archives/2012/1/e4/v3n1e4ht.htm) | J Oral Maxillofac Res 2012 (Jan-Mar) | vol. 3 | No 1 | e4 | p.6
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![Figure 6](https://www.ejomr.org/JOMR/archives/2012/1/e4/v3n1e4ht.htm)
was seen over Implant II; in addition, the cell layer exhibited a more complex organization with a fibrilar matrix, compared with that observed over Implant I, which showed mostly a continuous cell layer (Figure 6G, H).

**Alkaline phosphatase (ALP) activity**

The time-dependent ALP activity, normalized to the total protein content (Figure 7), showed an increase with culture time in both colonized implants. ALP activity was higher over Implant II, attaining a statistically significant difference at day 21; in addition, maximum values were observed earlier, at day 21, compared to colonized Implant I (maximal values, at day 28).

**Matrix mineralization**

SEM images of 28-day colonized implants showed the presence of mineralized globular deposits in close association with the cell layer in both implants, but the cell layer growing over Implant II exhibited a more complex fibrilar structure and a higher abundance of the globular structures (Figure 8A, B). Figure 8C shows a representative high magnification image of the mineralized deposits integrated in the fibrilar matrix, and Figure 8D displays the X-ray spectrum of the globular deposits, showing the presence of Ca and P peaks. Ca\(_i\) levels in the culture medium (Figure 8E) were relatively constant until approximately day 21 (Implant I) and day 16 (Implant II), decreasing significantly afterwards.

**DISCUSSION**

It is well established that surface topography of Ti implants is of particular interest for the osseointegration process, especially within the areas confined by single threads that are representative of the implants microtopography. This study compares two sandblasted and acid-etched implant systems, from Eckermann Laboratorium S.L. (Alicante, Spain). According to the manufacturer, the system patented as Eckeyte® (Implant II) has improved surface properties compared to the Implant system I, launched previously. Implants I and II presented a similar geometry, namely thread spacing, distance peak-valley and valley shape. However, they exhibited a distinct microtopography, which was produced by sandblasting with Al\(_2\)O\(_3\) followed by an etching process involving HCl, H\(_2\)SO\(_4\), and HF solutions, differing on the experimental protocols. In the acid-etched process, factors such as acid concentration, relative proportions of the acids used, and time of treatment played important roles in determining the final surface properties. These differences in surface morphology were expected to influence cell behavior and mineralization processes.

![Figure 7. ALP activity, expressed as nmol/min/mg protein, of human bone marrow osteoblastic cells grown over Implants I and II for 28 days. ALP activity increased throughout the culture time, and colonized Implant II exhibited earlier maximal values, compared to colonized Implant I.](http://www.ejomr.org/JOMR/archives/2012/1/e4/v3n1e4ht.htm)

![Figure 8. Matrix mineralization of human bone marrow osteoblastic cell cultures grown over Implants I and II for 28 days. High magnification SEM images: A, B and C = formation of globular mineralized structures closely associated with the cell layer. D = representative X-ray spectra of the mineralized deposits, showing the presence of Ca and P peaks. E = levels of ionized calcium (Ca\(_i\)) in the culture medium throughout the 30-day culture time, showing the consumption of Ca\(_i\) from the medium due to the formation of calcium containing deposits in the cell layer.](http://www.ejomr.org/JOMR/archives/2012/1/e4/v3n1e4ht.htm)
temperature and time of exposure are variables that are worked out to create different topographies [3]. A variety of studies reported that specific acid-etching protocols produce microporous on Ti surfaces with sizes ranging from 0.5 to 4 μm in diameter [3,8], which appear similar to that seen on the surface of Implant II. This type of microporous surface has been shown to greatly improve osseointegration [3,19]. Regarding the different surface features of Implants I and II, a previous work performed in our laboratory, in titanium surfaces submitted to different acid-etching protocols, showed that etching treatments based mostly on H2SO4 solutions provided a surface with a topography similar to that seen on Implant I, and treatments with HF based solutions were very effective in smoothing irregular rough titanium surfaces creating an homogeneous rough topography [20]. Ti is very reactive to fluoride ions, forming soluble TiF4 species, and it is reported that this type of treatment creates a microrough topography and, also, fluoride incorporation, that appears to enhance osteoblastic differentiation [21] and osseointegration in \textit{in vivo} experimental studies [22]. However, these considerations are merely speculative in which concerns the conditions used to produce the topography of Implants I and II, because such details were not disclosed by the manufacturer.

The surface of the tested implants differs also on amplitude, spacing and hybrid roughness parameters, as expected from their different topography. The average surface roughness (Ra) was higher on Implant II, compared to Implant I. However, the Ra values of both systems are within the range considered to yield improved implant performance [5]. It is worth mentioning that Ra gives only a good overall description of height variations (as it is the arithmetic average of the absolute deviations from the mean line over a sampling length), and it is not sensitive to small changes in the surface profile [23]. There are other parameters that complement surface description [23]. Rz, the arithmetic average of the five highest profile peaks and the five lowest profile valleys over the entire measurement trace, was higher on Implant II. The same was observed for Rt, the maximum peak to valley of the entire measurement trace, and for Rsm, the arithmetic average spacing between the falling flanks of the peaks on the mean line measured over the sampling length. However, Implants I and II have similar Rku values, a parameter that describes the probability density sharpness of the profile. On the other hand, Rsk, defined as skewness, which measures the symmetry of the deviation from a mean plan, was more negative on Implant II than on Implant I, meaning that, comparatively, Implant II has more valleys than peaks, which is in line with that observed on the high magnification SEM images. Also, the three-dimensional topography images showed evidence of a more homogeneous rough surface on Implant II.

The chemical composition and changes on the surface of Ti implants differ, depending on their bulk composition and surface treatments. These are relevant issues as they affect the initial biological behaviour following implantation, such as protein adsorption and cell attachment, conditioning the subsequent proliferation and differentiation of bone cells at the implant surface [3]. The present results showed that Implants I and II presented some differences in the chemical composition of the surface. XPS analysis revealed the presence of Ti, but also O, C and N, as expected, as these elements are commonly adsorbed on Ti surface [3]. The Ti 2p doublet and the O 1s peak reflected of titanium oxide (TiO2) layer [3], and the C peak indicates an organic contamination (presence of a carbon overcoat on the implant) that is normally associated with implant handling (during packaging, for example) [24]. Al 2p peak is due to alumina residual particles on the surface, resulting from the blasting process with Al2O3. Alumina is insoluble in acid and is hard to remove from the Ti surface. These particles might be released into the surrounding tissues and cause deleterious effects on the implant osseointegration [3]. Also, the chemical heterogeneity of the implant surface may decrease the excellent corrosion resistance of Ti in physiological environment [25]. Implant II showed significantly lower levels of Al, suggesting differences on the blasting and/ or the acid-etching protocols. Regarding this, a previous work reported that HF solutions were more effective than H2SO4 solutions in eliminating Al contamination arising from the blasting process [20]. The implants presented also residual amounts of F, resulting from the presence of HF in the acid-etching treatment. Si, P and Ca were found in very low levels, reflecting an inorganic contamination, normally associated with implant handling, as referred above [24]. It should be noted that the organic and inorganic pollution normally found on the implant surface is typically inhomogeneous across the implant, and should not be mistaken for controlled chemical or biochemical modifications [24]. Implant osseointegration requires the recruitment of osteoprogenitor cells and their proliferation and differentiation into functional osteoblasts that are able to produce a mineralized extracellular matrix at the interface. Bone cells can recognize and respond to substratum structures both \textit{in vivo} and \textit{in vitro} [3], and, in the present work, human bone marrow cells, cultured in experimental conditions that favour osteoblastic differentiation [26,27], were seeded over Implants I and II, and the colonized implants were evaluated for the elicited cell response. Results showed that both

http://www.ejomr.org/JOMR/archives/2012/1/e4/v3n1e4ht.htm
implants allowed the initial cell adhesion and spreading and the subsequent cell proliferation, as assessed by SEM observation at an early (3h) and a later (28 days) culture time. In addition, ALP activity increased significantly during the culture time, suggesting an osteoblastic differentiation pathway [26]. ALP has a determinant role in the onset of the matrix mineralization, by providing phosphate ions from the hydrolysis of organic phosphates (in vitro, from the added β-glycerophosphate) that, together with calcium ions (in vitro, present in the culture medium), are used in the formation of the cell mediated calcium phosphate mineralized matrix. Accordingly, 28-day colonized implants showed the presence of calcium phosphate deposits, closely associated with the cell layer, as observed by SEM and coupled EDS. Concentration of Ca\textsubscript{i} in the culture medium was determined throughout the culture time, in order to get information on the time-dependent profile of the mineralization process. The measured Ca\textsubscript{i} levels reflected changes occurring between every medium change, so that the values were not cumulative. The decreased Ca\textsubscript{i} levels observed at later culture times reflected the consumption of Ca\textsubscript{i} from the culture medium, that is being used in the formation of calcium phosphate deposits in the collagenous matrix. Matrix mineralization is the last event of the osteoblast differentiation, and the observed behaviour is representative of the in vivo osteoblast phenotype [26]. This cell response is expected, as it is in agreement with the known biocompatibility of Ti [3], and also considering that the surface roughness of both implants is within the range reported to provide a good biological performance [2]. Compared to Implant I, Implant II showed a thicker fibrilar cell layer, an earlier peak of ALP activity and onset of matrix mineralization (as evident by the time-profile of the Ca\textsubscript{i} consumption) and a more abundant mineralized matrix. The better performance of Implant II is related to its surface properties, and surface microtopography certainly plays an essential role in this behaviour. Homogeneous rough microporous surfaces are reported to promote the adhesion of fibrin improving the osteoconductivity and providing contact guidance for the osteogenic cells migrating along the implant surface, compared to irregularly rough surfaces [3,28]. Also, this type of surface topography appears to provide a higher bone-to-implant contact, as described in in vivo studies [9,10]. Some differences were also observed on the surface chemical composition, and this might also affect biological response [5]. It is worthwhile to note that, as referred above, Implant II contains a significantly lower percentage of Al, which is a positive feature, considering the deleterious effects of this element on cell behaviour [3].

The main limitation of this study is the lack of information in commercial or scientific literature on the details of the manufacturing process of the two implant systems, which is not surprising as it is a trade secret of the manufacturer. Thus, it is not possible to correlate specific surface properties with the observed osteoblastic cell response. Nevertheless, the homogeneous rough surface microtopography of Implant II fills many of the features reported to contribute to an improved bone cell response and better clinical outcome, as referred above.

CONCLUSIONS

Implants I and II presented a similar geometry, but a distinct microtopography, and also some differences in the surface roughness parameters and chemical composition. However, the homogeneous rough and microporous surface of Implant II is most probably a main factor for its improved osteoblastic performance, reflected by the presence of a thicker fibrilar cell layer and an earlier and more abundant matrix mineralization. The lower Al content of Implant II also deserves to be noted, due to the known cytotoxicity of this element. The earlier osteoblastic differentiation that appears to occur on Implant II might be a relevant factor during the first stages of bone apposition, which in turn may speed up and enhance the osseointegration process anticipating a better clinical outcome.

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