Research paper

Microrobotized blasting improves the bone-to-textured implant response.
A preclinical in vivo biomechanical study

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\textbf{Abstract}

This study evaluated the effect of microrobotized blasting of titanium endosteal implants relative to their manually blasted counterparts. Two different implant systems were utilized presenting two different implant surfaces. Control surfaces (Manual) were fabricated by manually grit blasting the implant surfaces while experimental surfaces (Microblasted) were fabricated through a microrobotized system that provided a one pass grit blasting routine. Both surfaces were created with the same \(\mu\text{m}\) average particle size alumina powder at \(310\) KPa. Surfaces were then etched with \(37\%\) HCl for \(20\) min, washed, and packaged through standard industry procedures. The surfaces were characterized through scanning electron microscopy (SEM) and optical interferometry, and were then placed in a beagle dog radius model remaining in vivo for \(3\) and \(6\) weeks. The implant removal torque was recorded and statistical analysis evaluated implant system and surface type torque levels as a function of time in vivo. Histologic sections were qualitatively evaluated for tissue response. Electron microscopy depicted textured surfaces for both manual and microblasted surfaces. Optical interferometry showed significantly higher \(S_a\), \(S_q\), values for the microblasted surface and no significant difference for \(S_d\) and \(S_{dr}\) values between surfaces. In vivo results depicted that statistically significant gains in biomechanical fixation were obtained for both implant systems tested at \(6\) weeks in vivo,
1. Introduction

The host to metallic implant response has been extensively studied and through implant system surgical hardware (relationship between implant bulk design and surgical instrument dimension) engineering, surface engineering at multiple length scales, and tissue engineering approaches, the host-to-implant response has significantly reduced the time required for the establishment of a biomechanically competent bone-implant interlocking (Coelho et al., 2015).

Although it has been recently shown that alterations in surgical hardware alter the osseointegration pathway of metallic devices (Coelho and Jimbo, 2014), surface engineering has been deemed equally important as an ad-hoc engineering design tool since the implant surface is the first component of the implant to interact with the host biomolecules and thus may modulate tissue and cellular events that may improve/accelerate tissue integration. Such surface engineering techniques have extensively been explored at the micrometer and nanometer scales (Coelho et al., 2015; Johansson et al., 2012), as well as surface coatings with bioactive ceramics (Jimbo et al., 2012). It has been reported that surface modifications promote earlier osseointegration, which has significantly contributed to the transition of the clinical modality. Currently, the majority of commercially available implant surfaces are physically and/or chemically modified and are alleged by industries to promote shorter healing periods. Evidence in the literature also indicates that surface modified implants are beneficial when utilized especially in clinically compromised situations (Khang et al., 2001; Pinholt, 2003; Stach and Kohles, 2003).

To date, continuous research in this field has focused in further enhancement of osseointegration and recent reports showed that the addition of biomolecules provided favorable outcomes (Coelho et al., 2014a; 2014b; Yoo et al., 2014). While promising, tissue engineering approaches are still far from reaching industrial scale due to challenges in product economic viability and large scale industrial surface engineering methods comprise grit blasting the implant surface through methods that provide little control of the process (i.e. manually). This is supposedly one of the reasons why implant surfaces, even within the same manufacturer and type, may vary in roughness depending on their batches. Although these batches all may be within the so-called moderately roughened surface roughness, which is known to present the strongest bone responses (Wennerberg and Albrektsson, 2009), the coarse differences in the surface roughness would result in different biologic outcomes. Due to the fact that cells and proteins respond sensitively to fractional alterations in the surface topography (Cooper, 2000), a slight deviation in surface roughness could result in adverse effects. Valverde et al. (2013) reported that implant surface roughness may be altered by the blasting media size, velocity, and surface coverage and suggested that the controlled blasting procedure could allow tailored surface topographical conditions. The microrobotization of the surface roughening procedure not only standardizes precision of the surface topography but also allows its modification by alteration of multiple parameters.

In this study, we have tested the bone/implant interface biomechanical competence of two different implant systems that were fabricated with the same materials and pressure parameters either manually or through a microrobotized process in a laboratory in vivo model. The objective was to test whether the microrobotized roughening process would present improved osseointegration to the implant surface as compared to the manually treated surfaces. It was hypothesized that the high precision and homogeneity of the micro-robotized surface roughening procedure would present higher osteoconductivity compared to the manually blasted controls.

2. Materials and methods

2.1. Materials

Sixty commercially pure grade 2 titanium alloy, threaded endosseous implants were utilized in the study. Half of these implants were control surfaces (Manual) that were fabricated by manually grit blasting while the other half were experimental surfaces (Microblasted) that were fabricated through a microrobotized system that provided a one pass grit-blasting routine. For both processes the implants were mounted with customized fixtures in a rotating column. Both surfaces were created with the same ~50 μm average particle size alumina powder at ~310 KPa. Both surfaces were then subjected to a 37% HCl etching procedure for 20 min and were washed and packaged through standard industry procedures. Information regarding both processing parameters are presented in Table 1.

Thirty implants (Duocon System, Signo Vincles, Campo Largo, PR, Brazil) of 3.8 mm diameter by 10 mm length presented a microthreaded cervical component with a large thread pitch along the body of the implant (Fig. 1a). The other thirty implants (Slim System, Signo Vincles, Campo Largo, PR, Brazil) presented dimensions of 3.5 mm diameter by 10 mm length presented with a large thread pitch throughout its
length from apical to cervical. Three implants of each system and surface were utilized for surface characterization, and twelve implants of each system and surface were utilized in the in vivo animal study.

2.2. Surface texture characterization

For surface texture characterization, three implants per surface condition were utilized. Scanning electron microscopy (SEM, Philips XL 30, Eindhoven, The Netherlands) was performed at various magnifications under an acceleration voltage of 15 kV. Verification of particle embedding composition was performed by energy dispersive spectroscopy (EDS) at various accelerating voltages (PGT IMIX with PRISM light element detectors). Surface roughness was evaluated in the different surface implants by optical interferometry (IFM, Phase View 2.5, Palaiseau, France) at the flat region of the implant cutting edges (three measurements per implant). $S_a$ (arithmetic average high deviation), $S_q$ (root mean square), $S_{ds}$ (density of summits), and $S_{dr}$ (developed surface ratio) parameters were determined. A filter size of 250 $\mu$m x 250 $\mu$m was utilized for a total of 15 measurements per surface of each implant type. IFM parameter statistical analysis was performed by one-way ANOVA. Statistical significance was indicated by p-levels less than 5%.

2.3. In vivo laboratory model

For the laboratory in vivo model, 6 adult male beagle dogs approximately 1.5 years old were acquired after the approval of the Ethics Committee for Animal Research at Ecole National Veterinaire D’Alfort.

Prior to general anesthesia, intramuscular atropine sulfate (0.044 mg/kg) and xylazine chlorate (8 mg/kg) were administered. A 15 mg/kg ketamine chlorate dose was then used to achieve general anesthesia.

![Fig. 1](left) the Duocon and (right) Slim implant geometries.

<table>
<thead>
<tr>
<th>Blasting parameter</th>
<th>Manual</th>
<th>Microblasted</th>
</tr>
</thead>
<tbody>
<tr>
<td>Distance from nozzle</td>
<td>5 mm</td>
<td>1.5 mm</td>
</tr>
<tr>
<td>Nozzle angulation</td>
<td>1st Pass: perpendicular to implant long axis, 2nd pass: $+45^\circ$ relative initial position, 3rd pass: $-45^\circ$ from initial position</td>
<td>Perpendicular to implant long axis</td>
</tr>
<tr>
<td>Implant rotation velocity</td>
<td>220 rpm</td>
<td>105 rpm</td>
</tr>
<tr>
<td>Blasting time</td>
<td>$\sim$60 s</td>
<td>$\sim$20 s</td>
</tr>
</tbody>
</table>
The central region of the radius diaphysis was used for the surgery. After hair shaving, skin exposure, and antiseptic cleaning with iodine solution at the surgical and surrounding area, approximately a 5 cm length incision to access the periosteum was performed and a flap was thereafter reflected for bone exposure.

Four implants were placed along the radius from proximal to distal in an alternated distribution of system and surface type in every radius. This approach resulted in balanced surgical procedures that allowed the comparison of the same number of implant surfaces per time in vivo, limb, surgical site (1 through 4), and animal. The implants remained in vivo for either 3 or 6 weeks (right and left radius provided samples that remained in vivo for 3 and 6 weeks, respectively). The implants were placed at distances of 1 cm from each other along the central region of the bone.

After insertion, each implant received its proprietary cover screw to avoid tissue overgrowth. The soft tissue was sutured in layers according to standard procedures, with the periosteum and muscle layers sutured with Vicryl 4-0 (Ethicon, Johnson & Johnson, Miami, FL) and the skin with 4-0 nylon (Ethicon).

Fig. 2 – Scanning electron micrographs for the (a, c, e) manual and (b, d, f) for the microblasted surfaces at various magnifications. Note the presence of embedded alumina particles for the manual and the absence of particle embedding for the microblasted surface (arrows). Representative micrographs obtained from Duocon implants, similar characteristics were also observed for Slim implants.
Postoperative antibiotic and anti-inflammatory medications included a single dose of benzyl penicillin benzatine (20,000 UI/kg) intramuscularly and ketoprofen 1% (1 ml/5 kg). The dogs were euthanized by an anesthesia overdose, and the limbs were retrieved by sharp dissection. The soft tissue was carefully removed by surgical blade, and an initial clinical evaluation was performed to determine implant stability. If an implant was clinically unstable, it was excluded from the study.

For the torque testing, the radii were adapted to an electronic torque machine equipped with a 250 Ncm torque load cell (Test Resources, Minneapolis, MN, USA). Custom machined tooling was adapted to each implant internal connection and the bone block was carefully positioned to avoid specimen misalignment during testing. The implants were torqued in the counter clockwise direction at a rate of \(-0.196 \text{ rad/min}\), and torque to interface fracture was recorded. The torque machine was set to automatically stop the measurement when a torque drop of 10% from the highest recorded torque was detected. The rationale for this procedure was to minimize interface damage prior to histologic procedures, subsequently allowing qualitative histologic assessment. As such, careful biomechanical testing allowed each block to be used for both biomechanical and histologic evaluation (Coelho and Lemons, 2009; Granato et al., 2009, Granato et al., 2011).

The bones were reduced to blocks containing the implants and surrounding bone, and were immersed in 10% buffered formalin solution for 24 h. The blocks were then washed in running water for 24 h, and gradually dehydrated in a series of alcohol solutions ranging from 70% to 100% ethanol. Following dehydration, the samples were embedded in a formalin solution for 24 h. The blocks were then washed in running water for 24 h, and gradually dehydrated in a series of alcohol solutions ranging from 70% to 100% ethanol.

Table 2 presents the statistical summary for the different surfaces.

<table>
<thead>
<tr>
<th>Surface</th>
<th>$S_a$ (μm)</th>
<th>$S_q$ (μm)</th>
<th>$S_{ds}$ (summits/mm²)</th>
<th>$S_{dr}$ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Microblasted</td>
<td>0.66</td>
<td>0.80</td>
<td>4677.50</td>
<td>35.00</td>
</tr>
<tr>
<td>Standard dev</td>
<td>0.13</td>
<td>0.17</td>
<td>1290.85</td>
<td>2.90</td>
</tr>
<tr>
<td>Manual</td>
<td>0.33</td>
<td>0.42</td>
<td>6433.17</td>
<td>38.67</td>
</tr>
<tr>
<td>Standard dev</td>
<td>0.09</td>
<td>0.10</td>
<td>2330.29</td>
<td>5.41</td>
</tr>
<tr>
<td>p-values</td>
<td>0.001</td>
<td>0.002</td>
<td>0.21</td>
<td>0.90</td>
</tr>
</tbody>
</table>

Preliminary statistical analyses showed no effect of implant site (i.e., there were no consistent effects of positions along the radii) on all measurements. Therefore, site was not considered further in the analysis. Further statistical evaluation of torque at 3 and 6 weeks employed a mixed-model ANOVA where the animal was considered the statistical unit ($n=6$). Statistical significance was indicated by $p$-levels less than 5%, and post-hoc testing employed the Fisher LSD test.

Fig. 3 depicts representative scanning electron micrographs for both the manual and microblasted surfaces at different magnifications. From a qualitative perspective, both surfaces presented similar textured morphology (Fig. 2) due to the grit-blasting followed by the acid-etching procedures. Relative to the microblasted surfaces, the manually prepared implant surfaces presented evidence of alumina particles embedded within the textured titanium surface.

3. Results

Table 2 presents the $S_a$, $S_q$, $S_{ds}$, and $S_{dr}$ mean and standard deviations, as well as the $p$ values obtained for the different surfaces’ comparisons. Significant differences favoring the microblasted surface were detected in the $S_a$ and $S_q$ values and no differences were observed in $S_{ds}$ and $S_{dr}$ values (Table 2).

The removal torque statistical summary is presented in Fig. 3. At 3 weeks, increased levels of removal torque were observed for both implant systems with the microblasted surface relative to the control. However, statistical significance was only reached for the Slim system ($p<0.02$). At 6 weeks, the microblasted surface condition in both implant systems presented significantly higher removal torque values ($p<0.01$) (Fig. 3).
Qualitative histologic sections depicted that healing chambers were formed along the body of both implant systems due to their macrogeometry and surgical instrumentation dimensions geometric relationship. At 3 weeks, an osteogenic tissue typical of intramembranous-like bone healing pathway was observed occupying the healing chambers for both the manually blasted (Fig. 4a) and microblasted (Fig. 4b) surfaces. However, higher degrees of bone formation and osteoid were observed for the microblasted surfaces relative to their manually blasted counterparts. At 6 weeks, the healing chambers presented substantially more bone filling relative to three weeks (Fig. 4c-d).

4. Discussion

This study observed the effect of microrobotized surface blasting procedures on osseointegration. Implant surfaces that were treated manually with the same setups were considered as controls. Although treated with the same aluminum oxide (alumina) particles and similar parameters, the results of the surface topographical measurements clearly indicated that the 2 blasting procedures generated 2 different surfaces. It was evident from the amplitude parameters that implant surfaces treated by the microblasted (microrobotized) procedure resulted in significantly higher $S_a$ and $S_q$ values than that of the manually blasted procedure. Morphologically, remnants of the alumina particles could evidently be captured with the SEM, whereas the surface for the microblasted group presented a homogeneous morphology. Apparently, these differences resulted in higher osteogenic responses to the implant surface for the microblasted implant group compared to the manually blasted implant group. Specifically, the removal torque (RTQ) values were significantly higher for both implant types at 6 weeks for the microblasted surfaces and at 3 weeks, the RTQ values were significantly higher for one implant type (Duocon) with microblasting and for the Slim group the mean values were higher for the microblasted group (no statistical difference). These results suggested that osteogenic cells sensitively responded to the rougher and homogeneous microstructures, however at the same time, suggested that the interplay between the macrogeometry is of some importance. The macrogeometry of the Slim group generates a healing chamber situation between the implant and the wall of the osteotomy for the entire length of the implant even at the cortical regions, where for the Duocon group, the neck (micro-threaded) portion was engaged in a press-fit situation. These differences may have resulted in the biologic differences since the healing chamber situation has been suggested to promote rapid osteoconduction due to the intramembranous-like healing pathway (Coelho and Jimbo, 2014).

Fig. 4 – At 3 weeks, (a) manual and (b) microblasted surface implants presented a healing chamber filled with osteogenic tissue where initial bone formation observed in higher amounts for the microblasted surface relative to its manually blasted counterpart. At 6 weeks, (c) manual and (d) microblasted surface implants presented high degrees of bone filling relative to three weeks. The gaps observed in the micrographs between tissue and implant were likely generated by the mechanical testing performed prior to histologic sectioning. Representative micrographs of the Slim implant.
The reason why remnants of alumina were evident on the manually blasted surface remain largely speculative, however, the irregular blasting patterns of the manual blasting procedure may have generated more embedding of the alumina particles. Although the negative chemical effect on the bone formation by the residual alumina has been discarded in some studies (Piattelli, 2003; Wennerberg et al., 1996a; Wennerberg et al., 1996b), some other studies indicate that Al ions that may be released in the vicinity of the implant interface, which may inhibit normal bone deposition and mineralization (Capdevielle et al., 1998; Stea et al., 1992; Thompson and Puleo, 1996). However, in clinical reality, the commercially available implants, which possess alumina particles on their surfaces have successfully been in function for decades (Jimbo and Albrektsson, 2015; Pjetursson et al., 2012). Within the current existing evidence, it can be suggested that the effect of the remaining alumina on the implant surface may have had some influence, however, not to the extent that the effects could be considered negative. Hypothetically, the promotion of osseointegration was mainly due to the effect of the successful surface roughening procedure, which was nearly 150% of an increase in average roughness and root square mean roughness.

In this study, microblasting the implant surface significantly promoted the biomechanical stability of the implants placed in the beagle dog model after 3 and 6 weeks in vivo. It would be of interest to investigate whether the effect of a controlled surface blasting procedure influences extended healing periods.

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