Current knowledge about the hydrophilic and nanostructured SLActive surface

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Abstract: This review summarizes the present documentation for the SLActive surface, a hydrophilic and nanostructured surface produced by Straumann Company in Switzerland, and covers the results from 15 in vitro, 17 in vivo, and 16 clinical studies. The SLActive surface is a development of the large grit-blasted and acid-etched SLA surface, and is further processed to a high degree of hydrophilicity. In general, the in vitro and in vivo studies of the SLActive surface demonstrate a stronger cell and bone tissue response than for the predecessor, the SLA surface, produced by the same company. However, in most studies, this difference disappears after 6–8 weeks. In the clinical studies, a stronger bone response was reported for the SLActive surface during the early healing phase when compared with the SLA surface. However, the later biological response was quite similar for the two surfaces and both demonstrated very good clinical results.

Keywords: SLActive, surface, in vitro, in vivo, clinical results

Introduction

We have attempted to summarize current published evidence for the SLActive implant from the Straumann company in Switzerland. The SLActive surface is a relatively recent innovation, with papers published over the last 5 years. This implant design has been launched by Straumann, based on it having a high surface energy, and being more hydrophilic than the previously presented SLA implant from the same company. High surface energy was once suggested by Baier et al1,2 to be an important surface characteristic, resulting in stronger bone responses, although one previous attempt to confirm this hypothesis proved unsuccessful,3 possibly due to the then tested high surface energy implant losing its physical state when in contact with the tissues. Another explanation may be the fact that surface energy exists at different levels, and the surface energy in this latter study was not high enough. Allegedly, the SLActive surface is supposed to overcome the tissue contamination problem by keeping the implant in a fluid container after surface alteration. The SLA surface is, in principle, produced by coarse grit-blasting with 0.25–0.5 mm corundum grit at 5 bar, followed by acid-etching.4 SLActive surfaces are produced with the same sandblasting and acid-etching, but they are rinsed under nitrogen protection to prevent exposure to air and then stored in a sealed glass tube containing isotonic NaCl solution.4 Schwarz5 claimed that this specific preparation process leads to a hydroxylated/hydrated surface, able to retain the high surface energy by reducing the adsorption of potential contaminants from the atmosphere (eg, hydrocarbons and carbonates).
A computerized search as of April 15, 2011 for scientific papers related to the SLActive surface yielded 15 in vitro studies, 14 in vivo studies in animals, three experimental studies in humans, and 16 clinical studies that will be summarized under these headings. A number of the cited studies investigated what was referred to as a “modified SLA surface” that did not always explicitly state that this modified surface was identical to the sometimes later commercialized SLActive surface. It is not uncommon for commercial companies to test different variations of surfaces before launching one type of surface. The present authors are unaware whether all modified surfaces tested actually are identical to the SLActive surface. In our literature overview, we have decided to use the same terminology as that used by the respective author(s), which is why the investigated implant is termed either SLActive or modified SLA surface in our text.

**In vitro studies**

Several in vitro studies have been performed to investigate cellular responses to SLActive surfaces. Many kinds of cells, including human osteoblast-like MG63 cells, human alveolar osteoblasts, and human mesenchymal stem cells from bone marrow, and human periodontal ligament cells, have been cultured on titanium disks with either an SLActive or a modSLA surface and compared with cells cultured on SLA surfaces. Only one study showed an increase of cell adhesion on the SLActive surface within 3 hours, while most of the studies showed a similar attachment of cells on both SLA and SLActive surfaces. However, a more differentiated phenotype of cells grown on SLActive was noticed compared with SLA, attested by a remarkable increase of alkaline phosphatase activity, osteocalcin, and osteoprotegerin production. In addition, the production of higher levels of local factors responsible for a bone-forming microenvironment, such as a 10-fold increase of prostaglandin E2 and a 2.5-fold increase of transforming growth factor-beta on SLActive compared with SLA surfaces, meant a significant enhancement of numerous gene expressions, associated with transforming growth factor-beta and bone morphogenetic protein signaling pathways.

ModSLA surfaces seem to provide a local noninflammatory microenvironment, since dendritic cells cultivated on modSLA maintain an immature phenotype compared with dendritic cells cultured on SLA surfaces. Moreover, SLActive surfaces were suggested to enhance angiogenesis induced by osteoblasts, because higher levels of angiogenesis growth factors, ie, vascular endothelial growth factor-A, fibroblast growth factor-2, and epidermal growth factor were found on cells cultivated on SLActive compared with SLA surfaces. Furthermore, gene expression of several angiogenesis-related factors was found to be increased.

**In vivo studies**

The first data on osseointegration of modSLA titanium implants was reported by Buser et al. The authors compared the percentage of bone to implant contact of SLA and modSLA implants after 2, 4, and 8 weeks of healing, using six miniature pigs receiving 46 implants (23 SLA and 23 modSLA) in a split-mouth design. They found that modSLA implants showed a significantly greater mean percentage of bone to implant contact at 2 (49.30% vs 29.42%; \( P = 0.017 \)) and 4 weeks (81.91% vs 66.57%; \( P = 0.011 \)) of healing, while no differences were observed at 8 weeks. Similar results were also reported by Bornstein et al , who placed 30 SLA and 30 SLActive implants in the mandibles of five dogs and reported a significantly higher percentage of bone to implant contact for modSLA implants at 2 weeks of healing (\( P < 0.05 \)), but this difference was no longer evident after 4 weeks. Further evidence was provided by a study performed using the tibia of sheep and involving 30 implants, 15 SLA and 15 SLActive, retrieved after 3 and 6 weeks. The percentage of bone to implant contact was significantly higher for modSLA implants at 3 weeks, but the difference was not statistically significant after 6 weeks of healing.

Schwarz et al performed several studies to investigate the performance of SLActive implants, compared with SLA implants. They placed 64 implants, 32 with an SLA surface and 32 with modSLA surface (SLActive) in the upper and lower jaws of four foxhound dogs (16 implants per animal) and assessed histologically, immunohistochemically, and histomorphometrically the bone formation, the peri-implant tissue reaction, angiogenesis, and osteocalcin activity, and the percentage of bone to implant contact after days 1, 4, 7, and 14 of healing. They observed that blood clots around the modSLA surfaces seemed to be stable at 1 day of healing and were substituted by a dense connective tissue after 4 days. Moreover, at day 14, the modSLA implants seemed to be surrounded by firmly attached, mature, parallel-fibered woven bone. In addition, they noticed that after 14 days of healing, modSLA implants were surrounded by well vascularized loose connective tissue exhibiting collagen fibers, that had started to extend and attach in part perpendicularly to the surface, with no clear separation into an inner and an outer zone. A further investigation confirmed these outcomes in a study performed in 15 beagle dogs involving a total of 240 implants.
The interfacial biomechanical properties of the SLActive implant surface were compared with the SLA surface in an animal experiment by Ferguson et al.26 They placed 162 implants, ie, 81 SLA and 81 modSLA, in the maxilla of 27 miniature pigs in a split-mouth study and analyzed the removal torque and bone-implant interfacial stiffness after weeks 2, 4, and 8 of healing (nine pigs/54 implants per healing time) and found the removal torque values to be on average 8%–21% higher for modSLA ($P = 0.003$) and the interfacial stiffness values on average 9%–14% higher for modSLA implants ($P = 0.038$) at each healing time.

It was hypothesized that the hydrophilic properties noted for modSLA-surface implants may also improve bone formation at buccal dehiscence defects, probably due to stabilization of the blood clot. Schwarz et al.27 created acute-type buccal dehiscence defects (height 3 mm, width 3 mm) during the placement of 24 implants, ie, 12 SLA and 12 modSLA, in the upper and lower jaws of four beagle dogs and evaluated the bone regeneration after 2 and 12 weeks of healing. They found that modSLA implants exhibited a complete defect fill at 12 weeks following implant placement, while wound healing at SLA implants was merely characterized by the formation of dense connective tissue, without any increase in bone formation. However, it is not clear to what extent a submerged healing procedure may have prevented the collapse of the mucoperiosteal flap into the defect area.

In another experiment,28 192 implants (96 with an SLA surface and 96 with an SLActive surface), were placed in 192 sites with standardized buccal dehiscence defects (height 4 mm, width 3 mm) and then randomly assigned to a submerged or nonsubmerged healing procedure in the upper and lower jaws of 12 beagle dogs (16 implants per dog). They found that SLActive titanium surfaces promoted bone regeneration (significantly higher percentage of bone to implant contact, new bone height, and new bone fill area) in acute-type buccal dehiscence defects at every stage of healing, and a submerged healing procedure additionally improved the outcome of bone regeneration at follow-up of 1–8 weeks. Schwarz et al.29 performed a study with the same design to compare bone regeneration in dehiscence-type defects around modSLA surfaces (SLActive) and acid-etched surfaces with a calcium phosphate nanometer particle modification (NanoTite®), and found that bone filling and area of mineralized tissue was comparable for the two groups. However, modSLA implants revealed significantly higher mean bone to implant contact and new bone height at weeks 2 and 8 of healing. In contrast, Al-Hamdan et al.30 found similar bone responses to SLActive and Nanotite implants in a beagle dog experiment with a follow-up of 2–8 weeks.

Lai et al.31 placed 36 implants, ie, 18 with an SLA surface and 18 with a modSLA surface, in the mandibles of six beagle dogs and left them to heal nonsubmerged for 2, 4, and 8 weeks. One-third of the implants were placed with a 0.5 mm and one-third with a 1 mm bone gap around the coronal 5 mm of the implants, while the remaining one-third was inserted without a gap. Histological observation and histomorphometric analysis showed a significantly greater percentage of bone to implant contact, newly formed bone, and defect filling with modSLA implants compared with SLA implants at weeks 2 and 4 of healing, but no significant differences were found between the two groups at week 8. No differences in healing patterns were found for the two defect sizes.

Liñares et al.32 analyzed the effect of loading time on modSLA implants. They placed 24 implants with a modSLA surface after tooth extraction in the mandibles of six miniature pigs (four implants in each animal) according to a split-mouth design, ie, the implants were loaded after four weeks of healing in one side of the mandible, while on the contralateral side, the implants were loaded immediately after placement. After 8 weeks of loading, the animals were sacrificed and 21 implants were histologically and histomorphometrically analyzed, with the percentage of bone to implant contact and the mean distance of the implant shoulder to the first bone to implant contact being similar for both groups.

Mardas et al.33 operated on 36 New Zealand white rabbits, comprising 12 with induced osteoporosis, 12 with osteoporosis treated with alendronate, and 12 healthy controls, and evaluated bone formation histomorphometrically after 30 and 120 days. A guided bone regeneration procedure involving titanium domes with two different surfaces, ie, one SLA and one modSLA (SLActive), were placed over the prepared bone defects. The investigators reported that newly formed total bone, newly formed mineralized bone, and bone to implant contact were higher at the modSLA-treated sites during all the experimental periods. Therefore, the use of modSLA may further promote extraskeletal bone formation and osseointegration in healthy and osteoporotic-like conditions.

Three papers34–36 reported experimental in vivo studies in humans, and investigated for the first time the sequence of events during early osseointegration in human volunteers. Lang et al.34 placed 49 solid screw-shaped implants with a diameter of 2.8 mm and a length of 4 mm, either with an SLA surface or an SLActive surface in the mandibular retromolar area of 28 human volunteers and retrieved them after days 7, 14, 28, and 42 of submerged healing. Although all the devices healed uneventfully, only 30 (61%) specimens were available...
for analysis due to the difficulty in harvesting. At both days 7 and 14 of healing, no differences were observed between the two surfaces and the percentage of bone to implant contact increased from 6% to 12.2% for SLA and from 6% to 14.8% for SLActive. The 28-day specimens showed a significant increase of bone to implant contact of at least three-fold from the 7-day examination, with a bone to implant contact of 32.4% for SLA and 48.3% for SLActive, representing a statistically significant difference between the two surfaces and in favor of the SLActive surface. Six weeks after insertion, the percentage of bone to implant contact had further increased, reaching 62% for both implant surfaces, without any difference between them.

The same cohort was included in another study in which the new bone formation and the role of bone debris and bone particles during the early healing stages were assessed. New bone apposition was higher on SLActive screws after weeks 2 and 4, but the difference did not reach statistical significance. Furthermore, with the same protocol, 18 solid screw-type cylindrical titanium implants, 4 mm long and 2.8 mm wide, either with an SLA or with an SLActive surface, were inserted in the mandibles of nine volunteers. The authors found that genes had the same pattern of expression around the two different surfaces at day 4, while at day 7 several functionally important gene categories were overrepresented in the SLActive specimens, including genes involved in bone mineralization/ossification, vascular endothelial growth factor signaling, focal adhesion and mitogen-activated protein kinase signaling, as was the category of extracellular space localization genes (osteogenesis and angiogenesis-associated gene expression). At day 14, the gene expression profile was associated with more mature wound healing on both surfaces, with bone morphogenetic protein signaling upregulated also on SLA surfaces.

**Clinical studies**

Randomized controlled clinical trials comparing SLA and SLActive are still few. One is the pilot study of Oates et al., in which the new bone formation and the role of bone debris and bone particles during the early healing stages were assessed. New bone apposition was higher on SLActive screws after weeks 2 and 4, but the difference did not reach statistical significance. Furthermore, with the same protocol, 18 solid screw-type cylindrical titanium implants, 4 mm long and 2.8 mm wide, either with an SLA or with an SLActive surface, were inserted in the mandibles of nine volunteers. The authors found that genes had the same pattern of expression around the two different surfaces at day 4, while at day 7 several functionally important gene categories were overrepresented in the SLActive specimens, including genes involved in bone mineralization/ossification, vascular endothelial growth factor signaling, focal adhesion and mitogen-activated protein kinase signaling, as was the category of extracellular space localization genes (osteogenesis and angiogenesis-associated gene expression). At day 14, the gene expression profile was associated with more mature wound healing on both surfaces, with bone morphogenetic protein signaling upregulated also on SLA surfaces.

In another investigation, 22 patients with bilateral tooth loss in the same jaws were rehabilitated with 96 dental implants, 48 SLA and 48 SLActive, such that all patients received at least one SLA and one SLActive implant. Prosthetic loading was initiated after 8 weeks of healing in the mandible and after 12 weeks in the maxilla. No patients were lost to follow-up and only one SLActive implant failed after 3 weeks of placement, giving an overall survival rate 1 year after loading of 100% for SLA and 97.91% for SLActive (difference not significant). Resonance frequency analysis was performed at surgery, and weeks 1, 3, and 6 later, but there was no statistically significant interaction between SLA and modSLA implants. Mean bone loss at the loading stage (baseline, implant insertion), measured for 93 implants and only by means of panoramic X-rays, was found to be 0.22 ± 0.06 mm around SLA and 0.18 ± 0.05 mm around SLActive, and after 1 year of function, with a marginal bone loss of 0.46 ± 0.07 mm around SLA and 0.43 ± 0.11 mm around SLActive implants.

In a clinical prospective study, 20 patients, who had been treated at least 6 months previously with radiochemotherapy in the oral region, were rehabilitated with 102 implants, 50 SLA and 52 SLActive, randomly assigned according to a split-mouth design. The prosthetic load was carried out after 6 weeks in the mandible and 10 weeks in the maxilla. There was one patient dropout (five implants) due to recurrence of cancer and therefore 97 implants were followed for an average time of 14.4 months. Survival rate was 96% for the SLA, because two implants were lost before loading, and 100% for SLActive. The amount of bone loss was evaluated only by orthopanoramic radiography and reported as
a mean only, ie, 0.4 mm medially and 0.4 mm distally for SLA implants and 0.3 mm medially and 0.3 mm distally for SLActive implants, although whether the resolution level of mean orthopanoramic radiography is satisfactory for a statistical analysis can be seriously debated.

Ganeles et al41 placed 383 SLActive implants (197 immediately nonocclusively loaded and 186 loaded with nonocclusively restoration after 28–34 days) in 266 patients. Change in bone levels was assessed for 323 implants (missing data for 60 implants) by mean of standardized periapical radiographs, taken at baseline, at time of permanent restoration (20–23 weeks post-surgery) and after 5 and 12 months. The 5-month results42 revealed a mean bone level change from baseline (baseline not specified) that was 0.81 ± 0.89 mm and 0.56 ± 0.73 mm in the immediate and early loading group, respectively, with a statistically significant difference. The mean bone level change at 12 months was 0.90 ± 0.90 mm for immediate loading and 0.77 ± 0.93 mm for the early loading, showing a significant difference between treatment groups. However, implantation depth was different, with immediately loaded implants placed, on average, 0.3 mm deeper than later loaded implants. The frequency of bone loss >2 mm was low overall, but three-fold higher in the immediately loaded group. Bone gain was noted for 16% of the implants. Bone loss was significantly correlated with treatment center, implant length and implant position. Implant survival at 12 months was claimed to be 98% in the immediate group (four implants lost) and 97% in the early group (six implants lost), with no implant failures reported in type IV bone (where 7.3% of implants were placed). However, the dropout rate was not reported.

Bornstein et al41 inserted 56 SLActive implants in the posterior mandible of 39 patients and loaded 54 of the implants (96.4%) after 21 days of healing, while two were left posterior mandible of 39 patients and loaded 54 of the implants (96.4%) after 21 days of healing, while two were left healing for another 3–4 weeks, since they were considered “spinners” and were given 3 weeks of additional healing, and one implant in one patient was lost to follow-up. After 24 months, marginal bone loss from the baseline of implant placement, assessed by mean of periapical radiographs, was 0.2 mm, with 25 implants having a loss of 0.3–1.0 mm, no implant with a bone loss >1 mm, and 39 implants with a bone gain.

Roccuzzo et al46 placed 35 implants in the maxillary posterior region of 35 consecutively enrolled patients. Of these, 29 implants were loaded with a provisional prosthesis in occlusion after 21 ± 2 days, while the six remaining implants were left healing for another 3–4 weeks, since they rotated slightly at day 21. After 1 year of follow-up, there were no patient dropouts and no implant losses. Bone loss from baseline (abutment connection), derived from measurements on periapical radiographs, was reported only as a mean value of 0.22 ± 0.35 mm.

Rossi et al47 placed 40 SLActive short (6 mm) implants in 35 consecutive patients, 10 of whom were heavy smokers, and loaded them after 6 weeks of healing with single crowns and followed them up for 2 years. The dropout rate was not revealed. They found a 95% survival rate after 2 years, due to two implants being lost before loading, with a mean resonance frequency analysis value of 74.8 ± 6.1 at the 6-week evaluation. Bone loss from implant placement was noticed on periapical standardized radiographs for 35 implants and was 0.75 mm, reported only as a mean of 0.34 ± 0.38 mm before loading and increased by 0.23 ± 0.33 in the first year of loading, and by another 0.21 ± 0.39 mm in the second year of loading.

In another paper,48 11 patients (nine edentulous and two partially edentulous) were treated with bilateral sinus floor augmentation, performed with synthetic biphasic calcium phosphate on one side and with deproteinized bovine bone on the other side (randomly selected), and the placement of 62 SLActive implants after 8 months of healing. Time of the loading was not reported. The overall survival rate (no dropouts) was 96.8%, since two implants were lost, one made of each material, and the success rate was 91.7% for biphasic calcium phosphate and 95.7% for deproteinized bovine bone. The mean marginal bone loss from baseline (time of the delivery of the fixed prosthesis) to the 1-year follow-up was reported to be 0.29 ± 0.10 mm for implants placed in deproteinized bovine bone (23 implants), 0.43 ± 0.20 mm for implants placed in biphasic calcium phosphate (24 implants), and 0.52 ± 0.27 mm for implants placed in the native bone (15 implants).
In a case report, Marchetti et al. treated a 59-year-old patient with a severely atrophied maxilla using a Le Fort I osteotomy and interpositional iliac bone graft, and 13 weeks later, with the placement of seven immediately-loaded SLActive implants. After 20 months of follow-up, they found no implant failure and minimal bone loss, measured on periapical radiograms (baseline implant placement), as an average of 0.78 (range 0–1.75) mm.

Stoker et al. placed 248 SLActive implants in the intraforaminal area of atrophic mandibles in 124 patients and loaded them immediately with an implant-retained overdenture and evaluated them for a mean period of 2 years (12–40 months). The dropout rate was not reported and the survival rate at the end of the evaluation period was 98.8%, with only three implants lost. During the healing phase, the implant stability quotient value increased significantly.

Al-Nawas et al. investigated whether small diameter (3.3 mm) implants made from titanium-zirconium alloy performed similarly to titanium Grade IV small diameter implants, with the same SLActive surface. Ninety-one patients were enrolled and received 182 small diameter implants, one for each type, in the interforaminal region of the mandible and were rehabilitated with implant retained overdentures after 6–8 weeks. Of these patients, 87 were available for the study period of 12 months. Three implants were lost and two implants (in the same patient) had recurrent peri-implant infection. The mean peri-implant bone level from the day of the surgery was 0.34 ± 0.54 mm for titanium-zirconium and 0.31 ± 0.56 mm for titanium Grade IV after 12 months, but most of the changes occurred within the first 6 months. Nevertheless, because the two types of implants were made of different alloys, the authors of the present paper are uncertain whether the two surfaces are similar to or differ from SLActive surfaces produced on other materials.

Bergkvist et al. investigated the relationship between bone mineral density assessed by computed tomography scans before implant placement, the implant stability measured at implant placement, and also the marginal bone around immediately loaded implants. Twenty-one patients were sequentially enrolled and 137 SLActive implants were placed and immediately loaded both in the maxilla and mandible, and were followed for 1 year. At 1-year examination, the survival rate was 100%, but the patient dropout rate was not reported. The change in mean marginal bone level from baseline was 1.18 ± 0.98 mm in the maxilla and 1.15 ± 1.23 mm in the mandible, with almost 80% of the surfaces (medial and distal) having a marginal bone loss <1 mm.

Discussion
Characteristics of SLActive surface that may explain tissue reactions
The SLActive implant is marketed on the basis of its high surface energy. However, in reality, there are several differences between SLA and SLActive surfaces that are just as likely as high surface energy to explain experimental or possible clinical differences between the two surfaces, eg, the differences in microroughness and nanoroughness between SLA and SLActive. Whereas SLA surfaces have changed from older batches with an SA (average height deviation from a mean plane) of about 1.5 µm and an Sdr (surface enlargement due to surface topography as compared with a flat reference area) % of 34 over to an Sa of 1.75 µm and an Sdr% of 97% , the SLActive surface has an even greater Sdr% of 143, if with a similar Sa value of 1.78 µm. In this case, the surface difference indicates that SLActive has a much greater number of peaks/valleys across the surface compared with SLA (Figure 1A and B). Furthermore, a distinct difference in nanoroughness between SLA and SLActive was noticed, with the latter surface
having a very clear nanoroughness pattern in contrast with the former surface\(^5\) (Figure 2A and B).

**Experimental findings of in vitro/in vivo studies**

Most in vitro and in vivo studies report a stronger short-term bone response to SLActive implants compared with SLA implants, but the difference disappears in many in vivo cases during the first 6–8 weeks.

**Findings from human experimental studies**

Three human experimental studies performed under the guidance of a university ethics committee documented a stronger bone response to SLActive compared with SLA implants at weeks 2 and 4 of follow-up, but no differences between the two implants at 6 weeks of follow-up. In our opinion, these human experimental studies failed to provide strong support for the alleged clinical advantage of SLActive implants, due to the differences having already leveled out at 6 weeks.

**Findings from clinical studies**

Some of the clinical studies measured bone loss only with orthopantograms which is a clear methodological shortcoming. Furthermore, the baseline for radiography in many studies was the time of loading, indicating that any bone loss preceding the time of loading is neglected. Naturally, the later the loading, the larger the potentially hidden bone loss. However, it must be noted that there is no consensus as yet about the most appropriate time for baseline radiography, hence an underestimation of true bone loss figures are not necessarily more common in clinical SLActive reports than for other clinical materials. There were other clinical papers on the SLActive implant where only mean bone loss was reported, which is a severe shortcoming of those papers because the minimal requirement is to present either standard deviations or separate those implants with >2 mm and >3 mm of bone loss from the rest of the materials, given that such implants may not be deemed successful.

**Results from comparative clinical studies**

Some clinical studies were properly randomized and the measured outcomes were either similar between SLA and SLActive or some early advantages at short-term follow-up were noticed for SLActive implants. However, reported differences in bone loss were very small and only reported by a couple of studies.

**Conclusion**

There are many published experimental and clinical studies of the SLActive implant. Most experimental studies indicate stronger bone responses to the SLActive surface compared with the SLA surface, at least in the first few weeks after implant placement. From a clinical viewpoint, many studies have been published where SLA and SLActive surfaces have been compared with one another and are quite positive. However, the clinical differences noted between the two types of surfaces are very small, and it is the opinion of the present authors that, based on the collective evidence from our present review, there are few indications supporting one implant type over another. Having said this, within the limitations of the maximal

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**Figure 2** (A) An SLA implant, flank area. (B) An SLActive implant, flank area. The image demonstrates a complex surface topography with several wavelengths included.
three-year observation period, SLActive surfaces seem to perform adequately and may, like SLA surfaces, be regarded as representing a well performing modern oral implant system.

Disclosure

The authors report no conflicts of interest in this work.

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