Early bone healing onto implant surface treated by fibronectin/oxysterol for cell adhesion/osteogenic differentiation: in vivo experimental study in dogs

Jung-Seok Lee1,†, Jin-Hyuk Yang1,†, Ji-Youn Hong2, Ui-Won Jung1, Hyeong-Cheol Yang3, In-Seop Lee4, Seong-Ho Choi1,*

1Department of Periodontology, Research Institute for Periodontal Regeneration, Yonsei University College of Dentistry, Seoul, Korea
2Department of Periodontology, Kyung Hee University School of Dentistry, Seoul, Korea
3Department of Dental Biomaterials Science, Dental Research Institute, Seoul National University School of Dentistry, Seoul, Korea
4Atomic-Scale Surface Science Research Center, Yonsei University, Seoul, Korea

Purpose: This study aimed to evaluate the effects of fibronectin and oxysterol immobilized on machined-surface dental implants for the enhancement of cell attachment and osteogenic differentiation, on peri-implant bone healing in the early healing phase using an experimental model in dogs.

Methods: Five types of dental implants were installed at a healed alveolar ridge in five dogs: a machined-surface implant (MI), apatite-coated MI (AMI), fibronectin-loaded AMI (FAMI), oxysterol-loaded AMI (OAMI), and sand-blasted, large-grit, acid-etched surface implant (SLAI). A randomly selected unilateral ridge was observed for 2 weeks, and the contralateral ridge for a 4-week period. Histologic and histometric analyses were performed for the bone-to-implant contact proportion (BIC) and bone density around the dental implant surface.

Results: Different bone healing patterns were observed according to the type of implant surface 2 weeks after installation; newly formed bone continuously lined the entire surfaces in specimens of the FAMI and SLAI groups, whereas bony trabecula from adjacent bone tissue appeared with minimal new bone lining onto the surface in the MI, AMI, and OAMI groups. Histometric results revealed a significant reduction in the BIC in MI, AMI, and OAMI compared to SLAI, but FAMI demonstrated a comparable BIC with SLAI. Although both the BIC and bone density increased from a 2- to 4-week healing period, bone density showed no significant difference among any of the experimental and control groups.

Conclusions: A fibronectin-coated implant surface designed for cell adhesion could increase contact osteogenesis in the early bone healing phase, but an oxysterol-coated implant surface designed for osteoinductivity could not modify early bone healing around implants in normal bone physiology.

Keywords: Cell adhesion, Dental implants, Fibronectins, Surface properties, Titanium.

INTRODUCTION

Since the dental implant was first introduced, various fixture designs and surface treatments have been developed for enhancement of osseointegration [1,2]. These designs aimed to achieve implant treatment with less time required for the healing period and longer-term clinical stability. Along the above-mentioned lines, machined-surface dental implants were replaced by those with rough surfaces [3]. Notably, sand-blasted, large-grit, acid-etched
dental implants not only provide extensively increased contact area with adjacent alveolar bone tissue [4], but also enhance cell attachment for de novo bone formation onto the implant surface [5].

Dental implants are supported by underlying bone tissue via direct bone contact (osseointegration); therefore, increasing the bone-to-implant contact area has been a main research topic in implant dentistry [6,7]. In previous studies, newly formed bone could be observed on the dental implant surface even in fatty marrow areas [8], in which increased bone density was observed around the dental implant compared to the marrow area. These findings can be explained by contact osteogenesis in Davies’ hypothesis [5,9]. The author described the mechanism of peri-implant bone healing, in which two types of healing are characterized: differential bone formation from recipient bone (distance) and attached cells onto the implant surface (contact osteogenesis). Therefore, cellular events, including attachment, proliferation, and differentiation on the dental implant surface have been suggested to be the most important factors for peri-implant bone healing and, additionally, Davies [5] proposed that surface topographies also affect contact osteogenesis. Many other studies have proven his hypothesis by demonstrating increased bone-implant contact around rough-surfaced implants [10,11], such as sandblasted, large-grit, and acid-etched surfaces, compared to those with machined surfaces [12,13].

According to the theory of contact osteogenesis, the microtexture of a rough surface could retain fibrin complex on the surface, and enable enhanced attachment and migration of undifferentiated cells, which are the initial steps of bone formation [9,14]. Since the clinical success of these rough surface implants [3,15], various other approaches have been developed for focusing on enhancing cellular events on the modified surfaces, that is, hydrophilic or anodized implant surfaces [16-19]. However, the separate effects of cellular events, such as cell attachment and differentiation, have not yet been elucidated.

The current development of implant surface technology has targeted the generation of a “biomimetic surface” on dental implants [1,20], in which biologic molecules have been applied onto the implant surface to stimulate osteogenesis and mimic each developmental step in the healing process. Extracellular matrix, peptides, and various growth factors are representative biologic molecules. Fibronectin is a major extracellular matrix that mediates attachment of cells to other cells or to other surfaces, such as the basement membrane. Recent studies have introduced a fibronectin-coated dental implant system in the research stage [21,22], and immobilization of fibronectin increased cell attachment [23] and osteoblastic protein expression at the in vitro level [24]. Another strategy in creating biomimetic implant surfaces is the use of growth factors [25,26] or their natural substitutes, such as oxysterol [27], for enhancement of osteogenic cell differentiation. Previous studies have demonstrated that oxysterols regulate differentiation of stem cells into osteogenic cells via the hedgehog pathway, and also prevent adipogenic differentiation in vitro [28,29]. Another in vivo study also found increased bone healing and augmentation in a spinal fusion model in animals [30].

Even though these two types of biomimetic implant surface developments showed increased cell attachment and differentiation in vitro [23,27], there was a lack of enhancement of osseointegration by using fibronectin or oxysterol in clinically-mimicking an in vivo animal model. Therefore, the present study aimed to evaluate the effects of fibronectin and oxysterol immobilized on machined-surface dental implants for the enhancement of cell attachment and osteogenetic differentiation, on peri-implant bone healing in the early healing phase using an experimental model in dogs.

MATERIALS AND METHODS

Animals

Five male mongrel dogs, aged 18–24 months and weighing approximately 30 kg, were used. All of the dogs had intact dentition and a healthy periodontium. Animal selection, management, and preparation, as well as the surgical protocol, followed the routine procedure approved by the Animal Care and Use Committee, Yonsei Medical Center, Seoul, Korea (2011-0072-1).

Implant preparation

Machined-surface implant (MI)

Cylindrical, threaded implants of commercially pure titanium (Ø3.4 mm, 10-mm length) with a machined surface were provided from the Research Institute of Dentium, Seoul, Korea.

Apatite-coated MI (AMI)

The MI was treated by calcium phosphate (CaP) nano-coating at a thickness of 500 nm, using ion beam-assisted deposition, as described previously [22]. For apatite formation on CaP-coated surfaces, samples were immersed into the solution, including Dulbecco’s phosphate-buffered saline (DPBS, Gibco-BRL, a division of Life Technology, Grand Island, NY, USA) and reagent grade CaCl₂ (100 mg/L). Apatite-coated samples were then rinsed with distilled water twice and dried at ambient temperature.

Fibronectin-loaded and AMI (FAMI)

FAMI samples were fabricated using the same method as for AMI, except that DPBS solution containing fibronectin was used instead of normal DPBS.

Oxysterol-loaded and AMI (OAMI)

CaP-coated, MIs were immersed in DPBS solution containing oxysterol for 2 days, and washed three times with distilled water and dried. The samples were immersed again in DPBS solution without oxysterol for one more day for additional apatite coating on the oxysterol/apatite coated surface, and then washed and dried.

Sand-blasted, large-grit, acid-etched surface implant (SLAI)

Commercially available dental implants (Implantium, Dentium Co., Seoul, Korea), which were treated by large grit sand-blasting
Effects of cell adhesion/osteoinduction on osseointegration

Histologic and histometric analysis

The animals were sacrificed with anesthesia drug overdose, and block sections, including segments of implants, were preserved and fixed in 10% neutral-buffered formalin. The specimens were dehydrated in ethanol, embedded in methacrylate, and sectioned in the mesio-distal plane using a diamond saw (Exakt, Apparatebau, Norderstedt, Germany). From each implant site, a central section was taken to a final thickness of about 30 μm, and the sections were stained with hematoxylin and eosin.

Histologic preparation

Ten experimental groups were allocated according to the type of implant surface (MI, AMI, FAMI, OAMI, and SLAI) and observational period (2 and 4 weeks). Five types of implants were installed in a randomly selected unilateral edentulous ridge, and the same types of implants were installed on the contralateral side at 2 weeks after the first surgery. The order of installation sites was rotated in five animals for even distribution of the experimental site. After allowing differential healing periods (2 and 4 weeks) following implant installation surgery, the animals were sacrificed for histological analysis.

Twelve weeks before implant installation surgery, all premolars and first molars were extracted at both mandibles under general anesthesia and sterile conditions in an operating room using 0.05 mg/kg atropine (subcutaneous injection), 2 mg/kg xylazine (Rompun, Bayer Korea, Seoul, Korea), and 10 mg/kg ketamine hydrochloride (Ketalar, Yuhan Co., Seoul, Korea) intravenously. The dogs were placed on a heating pad, intubated, administered 2% enflurane, and monitored with an electrocardiogram. After disinfecting the surgical sites, 2% lidocaine HCl with epinephrine 1:100,000 (Kwangmyung Pharm, Seoul, Korea) was administered by infiltration at the surgical sites. Implant installation operations were also performed under the same conditions as the tooth extraction procedure. A midcrestal incision was made, and mucoperiosteal flaps were carefully reflected on the buccal and lingual aspects. The edentulous ridge was carefully flattened with a surgical bur under sterile saline irrigation in order to obtain a widened ridge to accommodate a standardized ridge shape. Five prefabricated implants were installed in rotational order from anterior to posterior sites in individual animals with an even distribution of installed sites in each group, and the same order of installed implants was applied at both sides of the edentulous ridge in the same animal. Implant site preparation was performed by sequential drilling, and the flaps were sutured after implant installation using 5–0 resorbable suture materials (Vicryl 5/0, Polyglactin 910, Ethicon, a division of Johnson, Somerville, NJ, USA). The sutures were removed after 7–10 days, and a soft diet was provided throughout the study period.

Histologic and histometric analysis

Histologic and histometric analyses were performed using incan-
ven bone lining newly formed on the implant surface without any relationship with the recipient bone tissue (yellow asterisks in Fig. 2C and E). Most of FAMIs and SLAs showed the latter type of bone tissue, suggesting contact osteogenesis in the healing processes of osseointegration.

In the specimens from all groups at 4 weeks, newly formed bone tissue increased around the surface and in the space between the implant threads (Figs. 3 and 4). The proportion of newly formed woven bone around the implant was reduced at 4 weeks compared to 2 weeks.
Histometric analysis

The results of histometric analyses (mean, standard deviation, and 95% confidence interval for the mean) were presented in Tables 1 and 2 and Fig. 5. Statistical analyses using a linear mixed model revealed that the BICs were significantly different between groups, according to the observational period ($P<0.01$) and type of surface treatment ($P<0.01$). At the 2-week observational period, the BIC averaged $54.08\% \pm 9.50\%$, $59.94\% \pm 5.49\%$, $68.54\% \pm 11.42\%$, $58.65\% \pm 8.84\%$, and $77.14\% \pm 7.38\%$ for the MI, AMI, FAMI, OAMI, and SLAI groups, respectively. The SLAI group, as a positive control, showed the highest BIC among all groups, and there were significant differences from the MI ($P<0.01$), AMI

Figure 3. Representative photomicrographs of all experimental and control groups at 4 weeks after implant installation. The woven bone area had decreased in all of the specimens at the 4-week observational period. Most of the spaces between threads were filled with lamellated bone rather than woven bone, and increased, direct contact of bone to dental implants could be found along the whole length of the dental implants. There were still no significant differences in bone healing visible at the bone-to-implant interface in low magnification views. (A) Machined-surface implant (MI), (B) apatite-coated MI (AMI), (C) fibronectin-loaded and AMI, (D) oxysterol-loaded and AMI, and (E) sand-blasted, large-grit, and acid-etched surface implant. The scale bars in all panels represented 1 mm.

Figure 4. High magnification views of representative experimental and control samples at 4 weeks. All of the spaces between threads were filled with newly formed bone with high density. (A, B, D) Newly formed bone appeared in these spaces, approaching the implant surface from the recipient bone. The newly formed bone was partially in contact with the surface. (C, E) Thickened rims of newly formed bone on the surface could be found at most of the installed implant surface area, and connected with recipient bone tissues or regenerated bone sprouting from the lamellated bone around the dental implants. (A) Machined-surface implant (MI), (B) apatite-coated MI (AMI), (C) fibronectin-loaded and AMI, (D) oxysterol-loaded and AMI, and (E) sand-blasted, large-grit, and acid-etched surface implant. Scale bars in all panels were 100 μm.
Table 1. Results of histometric analyses in bone-to-implant contact proportion.

| Group   | 2 Weeks | 4 Weeks |
|---------|---------|---------|
|         | Mean ± SD | 95% CI  | Mean ± SD | 95% CI  |
| MI      | 54.08 ± 9.50<sup>a</sup> | 42.29–65.88 | 70.53 ± 7.77<sup>b</sup> | 60.88–80.17 |
| AMI     | 59.94 ± 5.49<sup>a</sup> | 52.72–66.36 | 71.10 ± 11.75<sup>c</sup> | 56.51–86.89 |
| FAMI    | 68.54 ± 11.42 | 50.37–86.70 | 73.95 ± 9.74 | 61.86–86.04 |
| OAMI    | 58.65 ± 8.84<sup>d</sup> | 47.66–69.63 | 74.94 ± 5.55<sup>d</sup> | 68.04–81.83 |
| SLAI    | 77.14 ± 7.38 | 67.98–86.30 | 84.34 ± 4.06 | 79.30–89.38 |

SD: standard deviation, CI: confidence interval, MI: machined-surface implant, AMI: apatite-coated MI, FAMI: fibronectin-loaded and AMI, OAMI: oxysterol-loaded and AMI, SLAI: sand-blasted, large-grit, and acid-etched surface implant. <sup>a</sup>Significantly different between measurements in MI and SLAI groups (P = 0.004). <sup>b</sup>Significantly different between measurements in AMI and SLAI groups (P = 0.031). <sup>c</sup>Significantly different between measurements in OAMI and SLAI groups (P = 0.022). <sup>d</sup>Significantly different between at 2- and 4-week period measurements in the same group (P = 0.019, *P = 0.042, *P = 0.015).

Table 2. Results of histometric analyses in bone density.

| Group   | 2 Weeks | 4 Weeks |
|---------|---------|---------|
|         | Mean ± SD | 95% CI  | Mean ± SD | 95% CI  |
| MI      | 29.94 ± 11.80 | 15.18–44.50 | 46.92 ± 9.58<sup>e</sup> | 25.02–58.82 |
| AMI     | 33.99 ± 4.57 | 28.31–39.67 | 50.82 ± 11.89<sup>f</sup> | 36.06–65.58 |
| FAMI    | 33.78 ± 9.34 | 18.91–48.65 | 42.62 ± 3.96 | 37.71–47.53 |
| OAMI    | 34.23 ± 8.99 | 23.07–45.39 | 51.91 ± 2.86<sup>f</sup> | 48.36–56.46 |
| SLAI    | 42.74 ± 3.42 | 38.49–46.98 | 50.18 ± 7.75 | 40.55–59.80 |

SD: standard deviation, CI: confidence interval, MI: machined-surface implant, AMI: apatite-coated MI, FAMI: fibronectin-loaded and AMI, OAMI: oxysterol-loaded and AMI, SLAI: sand-blasted, large-grit, and acid-etched surface implant. <sup>e</sup>Significantly different between at 2- and 4-week period measurements in the same group (P = 0.014, *P = 0.017, *P = 0.010).

Figure 5. Results of histometric analyses of the proportion of bone-to-implant contact (%) and bone density (%). Asterisks (*) indicate significant differences as shown by statistical analyses. MI: machined-surface implant, AMI: apatite-coated MI, FAMI: fibronectin-loaded and AMI, OAMI: oxysterol-loaded and AMI, SLAI: sand-blasted, large-grit, and acid-etched surface implant.

(\(P = 0.04\), and OAMI (\(P = 0.03\)) groups. Although there was no significant difference in BIC between the FAMI and SLAI groups, the FAMI group had a wider confidence interval for the mean BIC than the SLAI group; a similar maximum level of confidence interval was found in FAMI (86.70%) and SLAI (86.30%), but the corresponding values for the minimum levels of intervals were 50.37% and 67.98%, respectively. The other experimental groups (MI, AMI, and OAMI) showed a significantly increased BIC at 4 weeks compared to the same experimental group at 2 weeks (\(P = 0.03\), \(P = 0.04\), and \(P = 0.02\), respectively). In addition, there were no significant differences in BIC between any of the groups at the 4-week observational period, when the BIC averaged 70.53% ± 7.77%, 71.10% ± 11.75%, 73.95% ± 9.74%, 74.94% ± 5.55%, and 84.34% ± 4.06% for the MI, AMI, FAMI, OAMI, and SLAI groups, respectively.

Significantly increased bone density was found from the 2-week to 4-week observational periods in the linear mixed model (\(P < 0.01\)), but no effects were found according to the type of surface treatment (\(P = 0.134\)). Bone density at the 2-week observational period averaged 29.84% ± 11.80%, 33.99% ± 4.57%, 33.78% ± 9.34%, 34.23% ± 8.99%, and 42.74% ± 3.42% for the MI, AMI, FAMI, OAMI, and SLAI groups, respectively. The corresponding values for bone density at 4 weeks were 46.92% ± 9.58%, 50.82% ± 11.89%, 42.62% ± 3.96%, 51.91% ± 2.86%, and 50.18% ± 7.75%, respectively; there were significant differences from 2 weeks to 4 weeks in MI (\(P = 0.02\), AMI (\(P = 0.03\), and OAMI (\(P = 0.01\)).

**DISCUSSION**

This study was based on our two previous studies that demonstrated an increase in cell adhesion on fibronectin and patite-coated titanium surfaces [23] and enhancement of osteoblastic differentiation by oxysterol and apatite-coated titanium surfaces [27]. Since the development of dental implants using titanium, various surface treatments have been studied to make a bioactive titanium surface through the concept of mimicking cellular events in bone formation and remodeling processes [1,20]. The two above-men-
tioned types of implant surfaces were also intended to transform the surface from "biotolerant" to "bioactive" titanium: The fibronectin and apatite coating was intended to promote cellular adhesion onto the surface as a first step in the healing process [23], and the oxysterol or apatite coating was intended to promote differentiation of the adhered cells [27]. However, most dental clinicians still prefer rough surface implants without any biomimetic surface modification, even though various types of biomimetic surface implants have been commercially available. A systematic review of the Cochrane Collaboration reported that there was no clinical evidence of superiority in any particular type of dental implant [31].

The present study aimed to determine whether biomimetic implant surfaces that showed greater cell attachment and differentiation at in vitro level [23,27] could enhance the histologic parameters of the bone-to-implant interface in experiments in vivo mimicking clinical situations. In the present experimental model of normal bone without any defects, two types of bone healing patterns were observed around the dental implant surface: (1) a thin rim of woven bone deposited onto the implant surface (yellow asterisks in Fig. 2); and (2) bony trabecula sprouting from recipient bone tissue and contacting the implant surface (black asterisks in Fig. 2). These correspond to distance osteogenesis and contact osteogenesis in Davies' hypothesis [5], respectively. In a previous study, Davies described contact osteogenesis, in which de novo bone formation occurred on the implant surface; this pattern of healing was also demonstrated in the present results [5,9]. At the 2-week observational period, mineralized tissue could be seen on the implant surface contacting connective tissue fibers. Furthermore, the thickness and degrees of mineralization were varied, which indicated the bone formation process originated from the implant surface rather than the recipient bone tissue.

The sites receiving FAMI showed histologic evidence of contact osteogenesis along a larger area of the implant surface compared to the other experimental groups at 2 weeks of healing. The first step in contact osteogenesis would be cell adhesion onto the implant surface, followed by further cellular events, including proliferation and differentiation [14]. Fibronectin binds to cell adhesion molecules (integrin) on the cell surface, and increases/stabilizes cell-cell/cell-stratum adhesion [32,33]; therefore, these in vivo results might be caused by the function of fibronectin, like in vitro results. This healing pattern could also be demonstrated in sites receiving SLAI as a positive control group. Numerous previous studies have already found increased bone-to-implant contact in SLA-surfaced implants compared to MIs, which resulted from increased surface energy by microroughness of the surface [13,34,35]. In the histometric results, the SLAI group showed a higher BIC compared to the other three experimental groups (MI, AMI, and OAMI), and there was no significant difference between the FAMI and SLAI groups. However, the FAMI group also did not show any differences from the other groups, unlike the SLAI group. While the SLAI group showed a narrow range of histometric results, sites receiving FAMI showed a relatively wide range of results with regard to the BIC proportion; 95% confidence intervals for the mean were 67.98–86.30 in the SLAI group and 50.37–86.70 in the FAMI group. Rough surface implants can provide increased contact surface and mechanical interlocking with the recipient bone tissue [11,36], and these might increase the stability of SLAI during the early healing period. Increasing the stability of implants could also stabilize the in vivo healing process, which can be influenced by not only cellular events, but also the various environments of the recipient site (bone quality, biologic responses by cytokines, and immune responses). For this reason, FAMI might show a less uniform increase in the BIC compared to the SLAI, despite enhancement of cell adhesion.

Oxysterol is an oxidized derivative of cholesterol, and its novel characteristic of regulating differentiation into osteogenic cells has been the subject of increasing research attention. In previous studies, oxysterol induced in vivo osteogenesis like bone morphogenetic protein–2 (BMP–2), or synergistically increased the osteoinductive effects of BMP–2 [30,37,38]. Since complications of BMP–2 application have emerged in research and clinical fields [39,40], oxysterol has received attention as a substitute or regulating factor for BMP–2. Our previous study also showed an increase in alkaline phosphatase with oxysterol surface coating, and simultaneous apatite and oxysterol coating could synergistically enhance in vitro osteogenesis [27]. In the present study, the authors hypothesized that osteoinduction by oxysterol could enhance bone formation by osteoblasts adhered to the implant surface, and increase the BIC and bone density in the early healing phase. However, oxysterol coating on the implant surface failed to enhance the BIC and bone density in the space between threads, and these findings are in agreement with a previous study [26] that demonstrated that implant surface coating with recombinant human BMP–2 did not increase the BIC or bone density in a nondefect, normal bone, despite successful vertical augmentation in the supra-alveolar defect area without any graft materials [25,26]. Osteoinductive molecules might not accelerate or enhance bone healing at the bone-to-implant interface in normal bone, even though they can increase bone formation in the area with unfavorable defects. However, to confirm these hypotheses, further studies should be performed in various defect models.

This study aimed to determine the effects of enhancement of cell adhesion onto the implant surface and osteogenic differentiation by surface coating techniques on bone healing processes, especially at the bone-to-implant interface, in the early healing phase. Two types of implants focusing on cell adhesion increased the BIC at 2 weeks after implant installation: sand-blasted/acid-etched and fibronectin-coated surface implants. However, surface coating with osteoinductive molecules did not modify the BIC or bone density around any of the implant surfaces. Cell adhesion onto the implant surface may enhance contact osteogenesis around the implant in the early healing phase, when the implant is installed in normal bone without any defect.
CONFLICT OF INTEREST

No potential conflict of interest relevant to this article was reported.

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ORCID

Jung-Seok Lee http://orcid.org/0000-0003-1276-5978
Jin-Hyuk Yang http://orcid.org/0000-0002-8238-1374
Ji-Youn Hong http://orcid.org/0000-0003-1040-7077
Ui-Won Jung http://orcid.org/0000-0001-6371-4172
Hyeong-Cheol Yang http://orcid.org/0000-0001-9420-7894
In-Seop Lee http://orcid.org/0000-0002-0407-0431
Seong-Ho Choi http://orcid.org/0000-0001-6704-6124

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