Saucerization of osseointegrated implants and planning of simultaneous orthodontic clinical cases

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The field for Orthodontics has seen significant expansion with the advent of new diagnostic and therapeutic approaches in all specialties, such as medical and dental implantology, sleep medicine, orthognathic surgery, computed tomography, gerodontology, etc. This requires the mastery of new concepts and technical terms typical of the jargon used by each specific area. Such mastery plays a key role in discussions about diagnosis and planning of clinical cases with professionals from other specialties.

Dental osseointegrated implants, for example, completely changed the practice and scope of dentistry in the last 20 years. Many adult orthodontic patients have already had one or more osseointegrated implants installed or may be planning, or need to do so. Many young orthodontic patients have also had osseointegrated implants installed because of tooth loss caused by trauma or partial anodontia.

Osseointegrated implant saucerization is a phenomenon worthy of recognition and consideration in orthodontic planning to establish functional and aesthetic prognosis. With this insight in mind, we intend to discuss the concept of saucerization, with the specific purpose of answering a few important questions. Given the occurrence of saucerization, should special care be given to teeth located in the neighborhood of osseointegrated implants when moving teeth and finishing orthodontic cases?

The concept of osseointegration is a peculiarity of the teeth and implants in our bodies: The importance of cervical soft tissues

Osseointegration allows the direct anchorage of an implant through bone tissue formation around the implant without the growth or development of fibrous tissue at the bone-implant interface.

Teeth are the only body structures that traverse or penetrate an epithelial lining or coverage (Figs 1, 2 and 3). By extension, dental implants also have this feature and the anchorage provided by osseointegration is a prerequisite for implant stability. Long-term implant survival depends on the adhesion of the epithelium and connective tissues to the titanium surface since a complete soft tissue cervical sealing protects the bone from the highly contaminated oral environment.

The marginal gingiva and peri-implant mucosa share many clinical and microscopic characteristics. The gingival mucosa around
Successful implants usually display no inflammatory lesions. When lesions do occur, they are small and located adjacent to the junctional epithelium. Clinically, a healthy or slightly inflamed gingiva, as well as the peri-implant mucosa, if proper oral hygiene is performed, exhibit inflammatory infiltrates at similar locations and with similar extension. Several studies have shown similarities between the peri-implant mucosa and the gingiva in terms of their epithelial and connective structures. However, the absence of root cementum on the surface of the implants change the orientation plane and the adhesion of the fibers between teeth and implants. The importance of sealing the soft tissue at implant sites to achieve functional success has not been completely or thoroughly evaluated.

Studies on the topography of periodontal tissue vasculature revealed that the gingiva and connective tissue above the bone crest of the tooth are nourished by supraperiosteal vessels that originate in the alveolar process and periodontal ligament. In the soft and hard peri-implant tissues the mucosa region is nourished by terminal branches of wide vessels originating from the periosteum of the bone implant site. In both cases the vessels built a “plexus clevicular” lateral to the junctional epithelium. All natural teeth in the connective portion above the crest showed a rich vasculature, unlike the implant sites as very few vessels were observed in this region. This finding reinforces the suspicion that the peri-implant soft tissue may have a slightly decreased ability to defend itself against external aggression compared to the natural periodontal tissues (Fig 1).

The mechanical resistance between the gingiva and the peri-implant mucosa was tested in

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**FIGURE 1** - In the normal periodontium, at A, the collagen fibers are highlighted, extending from the gingival alveolar bone (AB) crest to the cementum (C), gingiva and periodontal ligament (P) to form a cross-hatch pattern at the connective attachment. The rich blood vascular (V) and fibroblastic (F) components can be seen, to a lesser extent in the cervical peri-implant connective tissue (CT). B shows schematically that the bundles of collagen fibers in the peri-implant cervical connective attachment tend to run parallel to the surface of the intermediate prosthesis (IT). GE = gingival epithelium; JE = junctional epithelium; IJE = implant junctional epithelium; D = dentin; M = marrow space; IP = implant.
FIGURE 2 - The tooth is the only structure of the body that crosses the lining epithelium and interacts with the internal environment. Layout of the periodontal structures relative to the biological distances: dentin (D), cementum (C), alveolar bone (B), periodontal ligament (PL), junctional epithelium (JE), gingival epithelium (GE) and gingival connective tissue (GCT). The junctional epithelium has 15-30 cell layers and as it proliferates in the apical direction it enables the contact of EGF molecules with bone cells, thereby stimulating bone resorption and maintenance of the biological distances. In the human body, between the epithelium and the bone, there is always connective tissue interposition due to the presence of EGF in the underlying epithelial and connective tissues. EGF is released by the Epithelial Rests of Malassez and keeps the alveolar bone away from the cementum through the same mechanism and thus prevents dentoalveolar ankylosis.

FIGURE 3 - The form of the alveolar bone crest, with its rhomboidal aspect, corresponds to the morphology of the junctional epithelium (JE) which fosters the steady release of EGF, depicted by the arrows. The collagen fibers of the connective attachment (CA) perpendicular to the cementum (C) can help limit the effect of EGF on bone cells. The cementoblasts (Cb) on the root surface have receptors for EGF and other mediators of bone turnover, which ultimately protect teeth from resorption. D = dentin; PL = periodontal ligament; B = alveolar bone, E = enamel; Ob = osteoblasts.

dogs and revealed that probe penetration was greater in implants than in teeth: 2 mm and 0.7 mm, respectively.14 In peri-implant soft tissues, the probe displaced the junctional epithelium and connective tissue on the implant’s adhesion surface interface and stopped at the bone crest. Occasionally, bleeding occurred due to vessel rupture. In the teeth, the probe stopped at the apical portion of the junctional epithelium, identifying the bottom of the gingival sulcus. The bleeding was minimal, in contrast with that of the implants.14

The effects of dental bacterial plaque after three weeks and after three months in the gingiva and peri-implant tissues were comparatively evaluated.6 Both tissues exhibited inflammatory lesions identical in size and composition features. Within three months the bleeding was similar and both inflammatory infiltrates had the same characteristics, but the apical extent was more pronounced in the peri-implant mucosa than in the gingiva. This finding implies that the defense mechanisms of the gingiva are more efficient than those of the peri-implant tissues in preventing future spreads of sulcus microbiota.6 However, the neck of an osseointegrated dental implant tends to display normal function and aesthetics, provided that adequate oral hygiene is maintained. This also applies to normal teeth.
Saucerization of osseointegrated implants: Concept and Mechanism

Saucerization occurs in all osseointegrated implants, regardless of their design, surface type, platform, connection type, commercial brand or patient conditions (Fig 12). Although the speed with which it occurs can vary, its occurrence seems to be part of the integration of implants with epithelium and gingival connective tissue.

The cervical region of osseointegrated implants, when exposed to the oral environment, usually exhibits some degree of bone resorption (Figs 4-11), of approximately 0.2 mm depth.4,5,11 The plane of the resorbed osseointegrated bone surface forms an open angle with the implant’s cervical region on nearly all of its surfaces. Three-dimensionally, this cervical bone resorption—observed in all types of osseointegrated implants—is in the shape of a saucer, i.e., it is shallow and superficial, hence "saucerization." This process can be extended over time,

FIGURE 4 - The gingival stratified squamous epithelium (GE) is juxtaposed with its normal thickness soon after the placement of healing caps or intermediate prosthesis and crown. The ulcerated epithelium has its cell membranes exposed to mediators that interact with their receptors. Under stress the cells increase the production of mediators. The EGF (arrows) of the epithelial cells themselves stimulates peri-implant epithelial proliferation and initiates the formation of the peri-implant junctional epithelium. EGF from saliva (S) probably participates in this process because it is greatly increased during oral surgery.

FIGURE 5 - The peri-implant junctional epithelium (JE) produces new cell layers and assumes a conformation similar to the junctional epithelium of natural teeth. This new conformation of the peri-implant junctional epithelium brings it closer to the osseointegrated surface, increasing the local concentration of EGF and, as a result, accelerating bone resorption and starting saucerization.

FIGURE 6 - The peri-implant junctional epithelium (JE) conformation is similar to the junctional epithelium of natural teeth. It derives structural balance from the peri-implant connective attachment to stabilize its proliferative activity. On the bone surfaces resorption decreases, approaching normal bone turnover. Thus, the peri-implant bone surface undergoes corticalization, indicative of process stabilization.
FIGURE 7 - During the removal of the healing caps or intermediate prosthesis there occurs the formation of the peri-implant junctional epithelium (JE) that covers the surface interface with the mucosa, including the gingival tissue. When it is still thin and disorganized, the peri-implant junctional epithelium tends to show a reddish appearance and can bleed if touched, given its frailty (A). When organized and mature, the peri-implant junctional epithelium appears pink, resembling the epithelium of the adjacent mucosa. Occasionally, the underlying microcirculation (B) can be seen as the JE becomes transparent.

Consuming on average 0.1 mm of peri-implant cervical bone tissue each year.⁴,⁵,¹¹ In a personal communication, Albrektsson reported that this cervical bone loss tends to decrease over the years to a level even lower than that recorded in previous studies, and that these results would soon be reported in the literature.

Many theories and explanations have been provided to account for saucerization but almost all have had difficulty explaining some of its features. One of these theories attributes saucerization to the occlusal masticatory load that implants have to sustain. However, when osseointegrated implants are out of occlusion or are fitted only with the gingival healing caps for many months or even years, without ever coming into occlusion, saucerization is also present (Fig 13). On the other hand, when implants remain submerged for a few months/years, the bone moves toward the more cervical surface and may even grow over the cover screws (Fig 12). This bone gain requires osteotomy maneuvers in order to place healing caps or an intermediate prosthesis.

Shortly after the placement of healing caps, or directly from the intermediate prosthesis and crown, the stratified squamous epithelium

FIGURE 8 - After saucerization, the peri-implant bone surface normalizes, with corticalization (arrows) indicative of stabilization of the pericervical bone remodeling process (toluidine blue, 10X).
of the oral mucosa is juxtaposed to the surface with its normal thickness (Fig 4). When an epithelium is ulcerated their cell membranes are exposed to mediators in order to interact with their receptors, in the same manner as in oral ulcers and surgical wounds, including in the peri-implant region.

The epidermal growth factor (EGF) in the saliva and in the epithelial cells stimulates peri-implant epithelial proliferation, thereby triggering the formation of the peri-implant junctional epithelium. The peri-implant junctional epithelium produces new cell layers and assumes a conformation similar to the junctional epithelium of natural teeth (Fig 5). This new conformation of the peri-implant junctional epithelium brings it closer to the osseointegrated surface, increasing the local concentration of EGF and, as a result, accelerating bone resorption and starting saucерization (Fig 5). Two recent papers have reviewed EGF functions and history.12,13

A few weeks or months after the peri-implant junctional epithelium and saucerization are formed they start moving away from each other. A stable biological distance is then established between the implant-integrated cervical bone and the peri-implant junctional epithelium, as occurs with natural teeth. From this stage, saucerization balance and stabilization are in place, allowing the bone on the cervical surface to resume corticalization (Figs 6, 8-11). It is probably due to this stabilization over the years that bone loss resulting from cervical saucerization diminishes its rhythm,14,11 provided that the conditions of hygiene and periodontal health are close to ideal. This situation has been noted in clinical cases that were followed up for many years after placement of osseointegrated implants (Figs 10 and 11).

The reestablishment of the junctional epithelium in the peri-implant oral mucosa may be due to stimulation by the EGF of the mucous

FIGURE 9 - Clinical case of implant in the upper lateral incisor region after six years, highlighting saucerization with regular bone surface and osseointegration.
epithelium itself through what is known as the autocrine effect. Although it probably takes place throughout the mucosa, it is particularly active in ulcerated areas where this autocrine effect is compounded by salivary EGF. As a result, a considerable increase occurs in cell layers to the extent that the peri-implant junctional epithelium is formed. Once the epithelium-implant integration occurs, salivary EGF penetration ceases or is drastically reduced and the process of cell-renewal epithelial proliferation goes back to normal.

The thickness of the gingival tissue appears to have a considerable effect on alveolar crest bone loss. When this thickness is 2 mm or smaller, the cervical bone loss tends to be significantly greater.21
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These results could probably be explained in light of the EGF. The thickness of the gingival tissue at the time of implant placement is commensurate with the distance from the implant junctional epithelium to be formed relative to bone tissue, i.e., EGF molecules rise to the bone surface in lower concentration.

Saucerization timing and orthodontic treatment

In natural teeth, the union of the junctional epithelium to the cervical enamel and surface is performed by means of several kinds of union structures, which effect an efficient sealing for salivary EGF (Figs 1, 2 and 3) in the peri-implant
FIGURE 13 - Osseointegrated Implants submerged from A to D. In this situation saucerization does not occur. Bone repair fosters partial overlap of implant coverage (as at B, C and D) because there is no formation of peri-implant junctional epithelium that would provide EGF molecules (arrows) in the vicinity of the bone surface. As soon as the healing caps are fitted, the formation of the peri-implant junctional epithelium (PJE) begins and so does saucerization (E). GE = gingival epithelium; GCT = gingival connective tissue; B = alveolar bone. (C: toluidine blue, 10X).
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FIGURE 14 - Implant installed in the region of tooth 12. The periapical radiograph (A) shows the proximity of the roots of teeth 11 and 13 due to the missing lateral incisor, which renders implant placement impossible; B shows the fixed orthodontic appliance for separation of the roots and crowns of teeth 11 and 13, thereby creating adequate space, suitable for implant installation in the region of tooth 12.

FIGURE 15 - The same clinical case of the previous figure with abutment mounted on the implant (A). Periapical radiograph (B) showing adequate intraradicular space between 11 and 13, which allowed the installation of the implant in the correct position. C shows the prosthetic crown cemented onto the abutment.
junctional epithelium. This sealing, however—provided by the epithelium-implant junction—is less efficient and supposedly allows a constant salivary EGF input which, in conjunction with the EGF of the junctional epithelium and mucosa, sets in motion a process of slow and steady approach to the cervical bone (Figs 1, 4, 5, 6, 9).

After an osseointegrated implant has been placed, peri-implant saucerization can normally be expected to occur, regardless of implant type (Figs 14 and 15). So what is the average distance that should be maintained by orthodontists between the cervical regions of neighboring natural teeth—when using osseointegrated implants—so that the cervical bone level of these implants is not affected by neighboring saucerization?

This concern may be even greater in upper anterior teeth such as, for example, lateral incisor implants (Figs 10, 11, 14, 15) in cases of partial unilateral or bilateral anodontia. Or, again, in cases of incisors and canines lost by accidental injury. The aesthetic and functional implications of the gingiva should be considered in planning and installing implants, such as the shape and size of the papillae, as well as the maintenance of a harmonious smile line.

Can saucerization, eventually, adversely affect the cervical hard and soft tissues of teeth located in the neighborhood of implants in patients treated orthodontically and whose teeth were harmoniously aligned with the implants? What special orthodontic care would be required to avoid or reduce the undesirable long-term consequences of osseointegrated implant saucerization occurring in the neighborhood of natural teeth?

The more we succeed in clarifying the phenomena related to cell and tissue saucerization, the more we will be able to learn about the care, and the aesthetic and functional nuances involved. Additional refinement and details concerning the evolution of the operative and restorative procedures of dentistry as a whole come to light every day, dissolving boundaries or obstacles between the most diverse specialties.

**Final considerations**

Orthodontists should increasingly familiarize themselves with the jargon of other clinical specialties, including implantology, as well as their concepts and more specific issues. This need stems from increased transdisciplinary actions undertaken by professionals in the joint planning of clinical cases involving multiple specialties, and whose ultimate goal is to rehabilitate the patient’s mouth.

Bone saucerization around osseointegrated implants is one such concept that forms a specific part of the implantology jargon. Orthodontists should consider the occurrence of this peri-implant bone phenomenon while simultaneously placing osseointegrated implants and moving the other teeth, realigning or relocating them harmoniously, many a time with such proximity to the cervical region that the condition should be carefully evaluated for its risks and aesthetic and functional benefits.

Further research is probably needed to answer the following question: Given the occurrence of saucerization, what are the special needs and care required by teeth located in the neighborhood of osseointegrated implants when moving teeth and finishing orthodontic cases?
REFERENCES


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