

Surgical predictability of vertical GBR in the posterior mandible

Flap design, management and passivation of soft tissues as principal keys for success

By Drs. Marco Ronda and Claudio Stacchi

The effectiveness of guided bone regeneration (GBR), a technique used to promote horizontal or vertical bone regeneration, has been well-documented since the early 1990s.¹⁻⁴ The stability of the regenerated bone and its positive response in time, once functioning, has also been well-demonstrated.⁵⁻⁸

Vertical GBR is a technique with great potential but one that requires both the precise adherence to surgical protocols and application by operators with the appropriate knowledge and manual skills to ensure optimum management of soft tissues. In addition to achieving primary closure of the flaps, maintaining this closure during the entire period necessary for the formation and maturation of the new bone is a pre-requisite for the avoidance of membrane exposure, which inevitably leads not only to bacterial contamination but, nearly always, to the impairment of the surgical procedure of regeneration.^{9,10}

Numerous studies have described various clinical protocols regarding the management of soft tissues in both the upper and lower arches.¹¹⁻¹⁷

This retrospective analysis describes the surgical technique of the management of soft tissues applied during GBR with non-resorbable membranes in 127 cases of vertical defects of the posterior mandible and evaluates the clinical results obtained.

Materials and techniques

Between 2000 and 2012, a total of 127 cases of vertical bone defects in edentulous posterior mandibles were treated with the use of GBR with non-resorbable membranes.

The technique was applied by following a surgical protocol, which has undergone few variations during the years.

From 2000 to 2008, expanded polytetrafluoroethylene (e-PTFE)



Fig. 1



Fig. 2

Photos/Provided by Drs. Ronda Marco and Claudio Stacchi

titanium-reinforced non-resorbable membranes (Gore-Tex TR9, W.L. Gore & Associates, Flagstaff, Ariz.) were used as a barrier device in 72 cases (Fig. 1).

From 2009 to 2012, high-density polytetrafluoroethylene (d-PTFE) titanium-reinforced non-resorbable membranes (Cytoplast TI250XL, Osteogenics Biomedical, Lubbock,

Texas) were used as a barrier device in 55 cases (Fig. 2).

All the membranes were fixed mesially and distally on the lingual side with the use of titanium pins (Helmut Zepf Medizintechnik, Seitingen, Germany) or mini-screws (Pro-Fix, Osteogenics

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ICOI heats up spring in Vegas

Group's annual symposium takes over the Bellagio Hotel with 'Maxilla'-focused sessions

By Sierra Rendon, Implant Tribune

More than 1,200 attendees, including 700 doctors and 250 auxiliaries, laboratory technicians, students and industry personnel, hit the Bellagio Hotel on the Las Vegas Strip for this year's Spring Symposium.

This event included an in-depth, challenging focus on "The Maxilla: Single Tooth to Full Arch Reconstruction" and welcomed main podium lecturers such as Dr. Jaime Lozada, Dr. Giuseppe Cardaropoli, Dr. Joseph Kan and Dr. Michael Sonick.

For implant doctors or team members who came looking to stock up on supplies or to look for products to bring home to the office, the exhibit hall was brimming with new technology and other treats. More than 100 exhibitors brought the industry's latest and greatest options for implantologists to use in their practices.

See pages B10 and B11 for more scenes and stories from the ICOI's Spring Symposium.



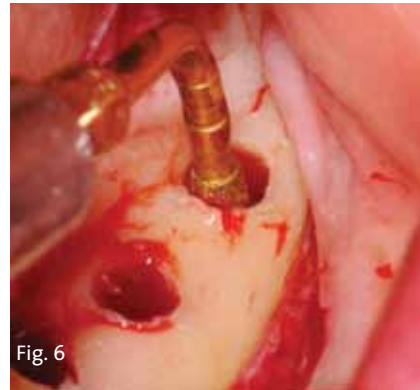
Dr. Jaime Lozada provides the first main podium lecture on the first day of the ICOI's Spring Symposium in Las Vegas. Lozada's session focused on 'Extraction Site Management' and, more specifically, the IDR technique. Photo/Sierra Rendon, Managing Editor

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Biomedical, Lubbock, Texas) (Fig. 3).

After positioning the graft material around the implants, which were left protruding from the crest (Fig. 4), the membranes were also stabilized on the buccal side with the same fixation devices (Fig. 5). Preparation of the implant sites, for the most coronal portion of the osteotomy, involved the use of twist drills and, for the most apical portion, near the mandibular nerve, a piezoelectric OT4 insert (Piezosurgery, Mectron, Carasco, Italy) (Fig. 6).

Implants (Spline Twist and Tapered Screw-Vent, Zimmer Dental, Carlsbad, Calif.) were inserted leaving their most coronal portion protruding from the crest for a length equivalent to the vertical bone regeneration planned. In certain cases — those in which it was not possible to obtain adequate primary stability in low quantities of residual bone — the



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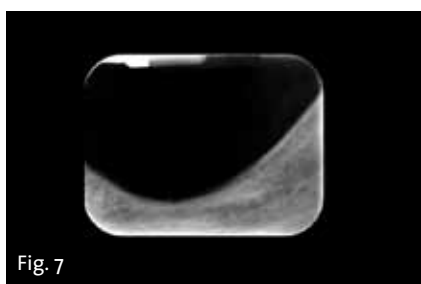


Fig. 7

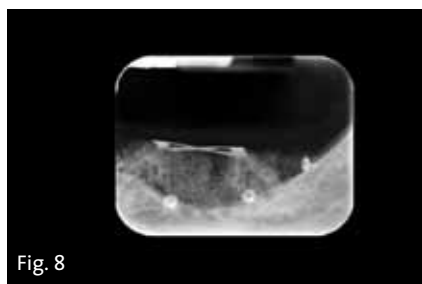


Fig. 8



Fig. 9



Fig. 10

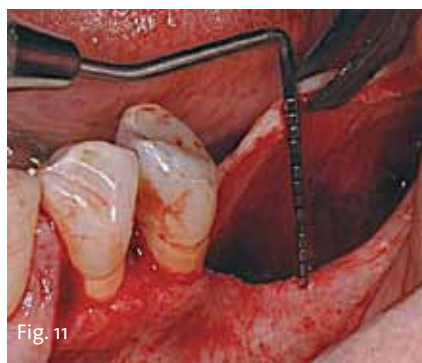


Fig. 11

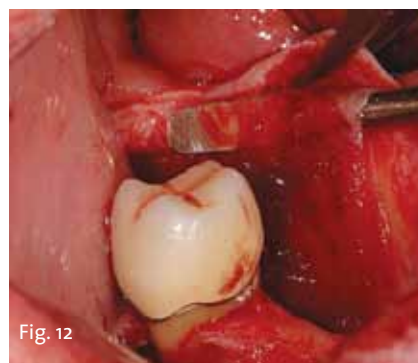


Fig. 12

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vertical bone regeneration preceded the positioning of the implants (Figs. 7, 8).

Multiple cortical perforations, which created openings for osteopromotion, were then made with a piezoelectric OP5 insert (Piezosurgery, Mectron, Carasco, Italy) in order to stimulate blood and cell migration from the bone marrow spaces to the regeneration area.^{18,19}

During the period of time analyzed, various graft materials, alone or combined, were used together with the membranes: autologous bone; tricalcium phosphate; DBM (Dynagraft, Keystone Dental, Burlington, Mass.); MFDBA (Puros, Zimmer Dental, Carlsbad, Calif.); or combinations of mineralized and demineralized allograft bone (MFDBA & DFDBA, enCore, Osteogenics Biomedical).

Surgical management of soft tissue

All surgeries as well as postoperative care are carried out by a single operator. For each patient, treatment includes the analysis of a diagnostic wax-up and CT or CBCT scan performed with a template. The objective is not only to position the implants where the quantity of residual bone allows but to position their platforms on the ideal line situated approximately 2 mm under the cement-enamel junction of the adjacent teeth.

After performing local anesthesia, (articaine hydrochloride 4 percent with epinephrine 1:100.000, Septanest, Ognà, Muggiò, Italy), a horizontal, mid-crestal, full thickness incision is performed in keratinized tissue. The incision extends from the distal margin of the last tooth adjacent to the treatment area to the ramus of the mandible, ending with a releasing incision on its buccal surface.

In the second molar area, to preserve the integrity of the lingual nerve, the scalpel should be inclined at an approximately 45 degree angle with the tip in vestibular direction, and the blade should touch the external oblique line while the incision is made in distal and buccal direction.

In the proximal vestibular zone, the incision continues intrasulcularly involving the last two teeth adjacent to the area to be treated and concludes with a vertical hockey stick releasing incision.

Lingually, the incision continues intrasulcularly until the gingival zenith of the last tooth and continues along the crest of the ridge for approximately 1 cm in the thickness of the keratinized gingiva. Full thickness flaps is then elevated and the mental nerve is isolated. The mobilization and release of the buccal flap is obtained with a horizontal periosteal incision performed with a new blade for the entire length of the flap, from the distal to the mesial release.

This longitudinal incision is performed approximately 5 mm apically from the crestal incision and should only affect the periosteal fibers. The passivation of the vestibular flap, thus obtained, allows for a mean coronal elevation of the flap of approximately 20 mm: this is the sum of the amount of tissue present above the periosteal line of incision (5 mm) and the stretching of the flap following the periosteal incision (15 mm) (Figs. 9, 10).

The lingual flap is also full thickness elevated until the mylohyoid line is reached. This maneuver allows for the obtaining of a mean coronal elevation

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of approximately 15 mm (Fig. 11). At this point, following the technique previously described by Ronda and Stacchi¹⁷, the mylohyoid muscle insertion on the inner surface of the lingual flap is identified, approximately 5 mm apically from the crestal line of incision.

This insertion, with the use of a blunt instrument, is first isolated (Fig. 12), and then separated from the flap by applying light tensile force. This maneuver allows for the near doubling of the lingual flap passivation and brings the coronal elevation from approximately 15 mm to approximately 30 mm (Figs. 13, 14).

The flaps thus passivated can be sutured covering the membrane without tension, using two different suture lines: one horizontal mattress suture with 3-0 PTFE approximately 5 mm apically from the crestal line of incision (Cytoplast Suture, Osteogenics Biomedical) and a series of interrupted sutures with 4-0 PTFE to complete the flap closure. The releasing incisions are closed with resorbable sutures (6-0, 7-0) (Serafit, Serag Wiessner, Naila, Germany).

The sutures are removed after approximately 12-15 days and, during this period, the patient uses a chlorhexidine 0.2 percent mouthrinse twice a day for one minute. In addition, antibiotics (amoxicillin/clavulanic acid 875+125mg) and NSAIDs (ibuprofen 600 mg) are prescribed for one week.

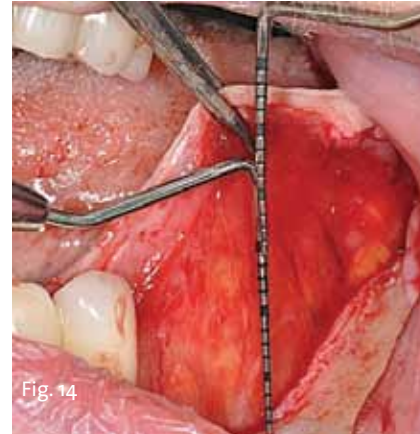
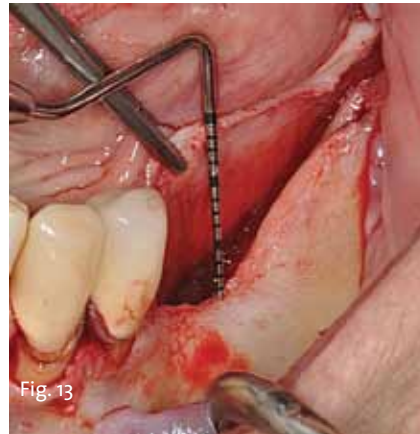
After a period of approximately six months, during which new bone formation is obtained and completed, the patient undergoes a second procedure for the removal of the membrane and fixation system, completing soft-tissue management (Figs. 15, 16).

Results

The goal of this study was to describe the results and complications that occurred both during and after surgery in 127 cases of vertical GBR with non-resorbable membranes, until their removal. Certain complications in a considerable percentage of cases can lead to the failure of the entire regenerative procedure. In order to list and analyze them, the classification proposed by Fontana et al. (2011)²⁰ was used.

Beyond the normal sequelae associated with surgery (edema, blood extravasation and hematoma), neurological complications (B, Fontana 2011) occurred in three cases (2.4 percent). Paresthesia is believed to have been related to the release and elevation of the vestibular flap, which most likely caused the stretching of mental nerve fibers. In all three cases, the symptoms of paresthesia subsided one month after the surgery.

During the healing period, no membrane exposure occurred in any of the cases (no Class I, II or III complications, Fontana 2011). In nine cases (7.1 percent), graft sepsis occurred in the absence of membrane exposure (Class IV, Fontana 2011). All Class IV complications occurred during the first month after the regenerative procedure.



Discussion

The objective of this retrospective analysis is to focus on the complications associated with the surgical technique of vertical regeneration with non-resorbable membranes in order to evaluate the level of surgical predictability associated with this procedure in view of the complexity and difficulty in augmenting the posterior ridge.

From the analysis of the results described, the general percentage of failure was 7.1 percent.

However, it is evident that with the application of conventional passivation techniques, and the introduction of the new lingual flap management technique, the extent of coronal displacement of the flaps guarantees the specialist a sufficient quantity of tissue to perform a tension-free suture above the regeneration area.

This is confirmed by the fact that no membrane exposure occurred in the 127 cases analyzed. The primary cause of failure of this technique, from the analysis of our data, is the bacterial contamination of the graft-membrane-implant complex in its entirety.

Contamination can already occur during surgery (inappropriate handling of surgical instruments, graft contamination as a result of bacteria present in saliva) or during the postoperative phase (failed primary closure of the flaps or early exposure of the membrane). As seen, the appropriate management of soft tissue allows for an entirely passive and hermetic primary closure of the flaps, as well as its maintenance, for the entire duration of the healing period.

The problem yet unresolved is that of the cases in which graft sepsis occurs, despite flap closure being perfectly maintained.

In this situation, which always mani-

fest itself during the first month after the procedure, intra-operative graft contamination plays a fundamental role. Given the difficulty in keeping the surgical area completely isolated from salivary contamination during the GBR procedure (above all, in the posterior mandible), the reduction of surgical time is one of the keys for minimizing the risk of infection.

In this regard, it could be useful to harvest autologous bone from a donor site, which is not from the actual area of regeneration, prior to the GBR procedure (with an inevitable increase in morbidity), or the use of commercial bone grafts alone, with the objective of entirely eliminating both autologous bone harvesting and the risk of infection associated with prolonged operating times.²¹

Conclusions

The current flap passivation techniques available to the specialist have significantly reduced the percentage of failure associated with early exposure of the membrane.

Therefore, we can surmise that vertical GBR is a realistically feasible solution in regard to surgical success (treatment results' stability over time has already been extensively demonstrated), despite the technique being considered highly "operator-sensitive."

The fact that vertical GBR is a difficult procedure is not, by any means, to be underestimated. It requires extensive knowledge and should be carried out after appropriate training, which must enable the specialist to acquire a complete theoretical and practical knowledge both in the fields of periodontology and implant dentistry.

References are available upon request from the publisher.

About the authors



DR. MARCO RONDA graduated with a degree in medicine from the University of Verona. A one-year course in advanced surgery, taught by Dr. Massimo Simion, and a Masters course in Regenerative Surgical Techniques at the University of Pennsylvania are among the many specialization courses he has attended. Ronda periodically gives lectures and provides practical training courses in implantology and bone-regeneration techniques at his practice in Genoa. He is also invited to speak at many national and international meetings and cooperates with several Italian universities including Milan, Trieste, Modena, Genoa and Pisa, and he is an adjunct professor at Bologna University.

The International Journal of Periodontics & Restorative Dentistry has published his study regarding a new surgical technique of lingual flap management that has been proven to increase bone volume in all cases. He is also the author of an article that was published in the Clinical Oral Implants Research journal that compares expanded PTFE and dense PTFE in guided bone regeneration. He may be contacted at mronda@panet.it.



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'The current flap passivation techniques available to the specialist have significantly reduced the percentage of failure associated with early exposure of the membrane.'



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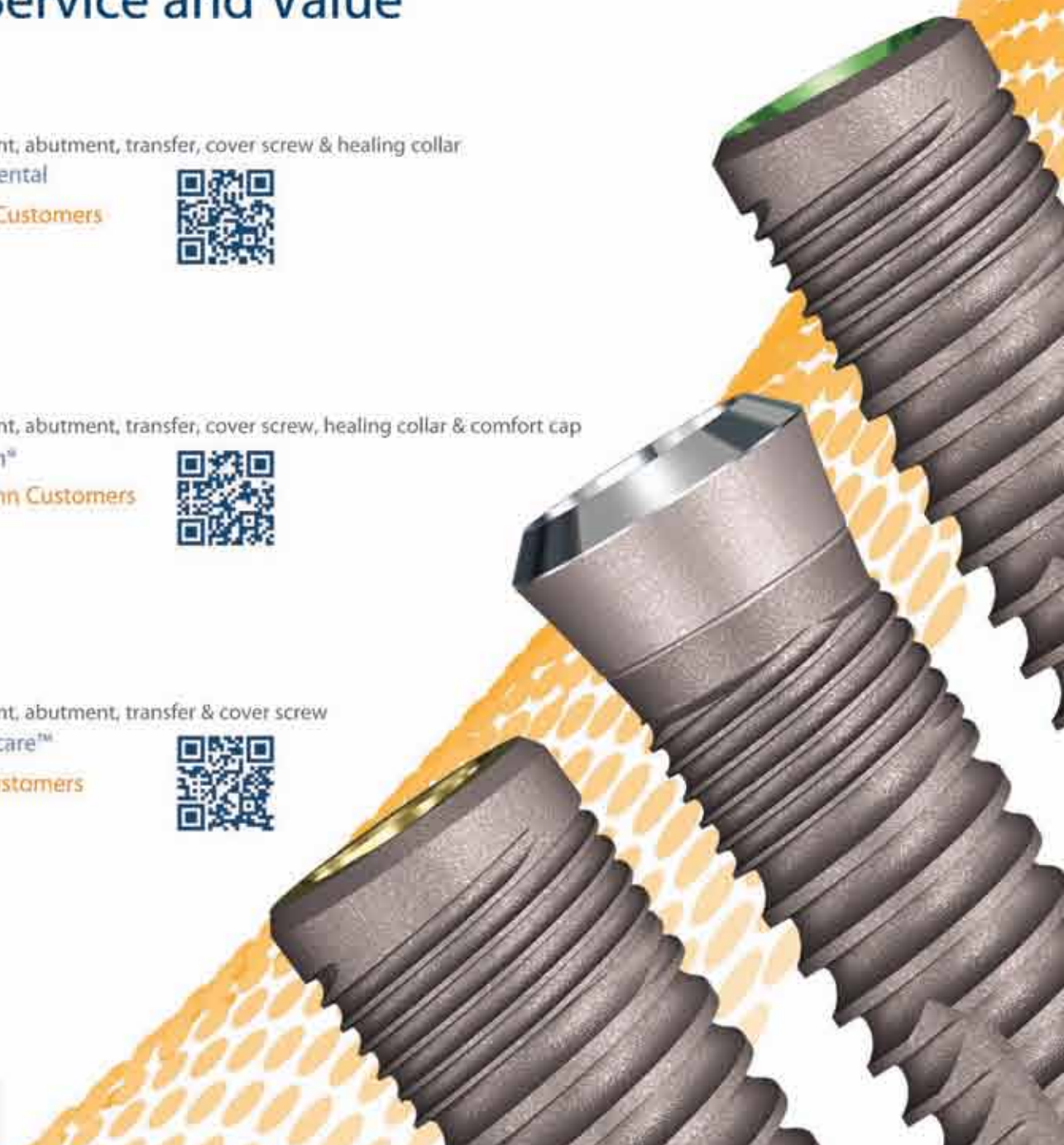
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Implant position in the esthetic zone

Establishing a treatment plan is paramount

By Siamak Abai, DDS, MMedSc

Since the advent of modern root form osseointegrated implant dentistry in 1952, clinicians have strived for improvements in implant positioning in the esthetic zone to achieve predictable restorative and esthetic results.

Years of clinical experience in congruence with controlled clinical studies have helped establish parameters as a guide for these results. Establishing a treatment plan and clinical protocol prior to implant placement is paramount.

Treatment planning traditionally begins with comprehensive medical and dental evaluations, articulated diagnostic casts, radiographs, cone-beam computed tomography (CBCT) scans and a diagnostic wax-up. Patient demands must be taken into consideration prior to surgery, and pre-surgical mockups may be necessary to convey the information to the patient.

The advancement of CBCT technology has led dentistry into a new realm of dimensional accuracy. In combination with the use of a surgical or guided stent, proper 3-D positioning of an implant has led to more accurate clinical results.

The importance of the implant position can be manifested in the four dimensionally sensitive positioning criteria: mesiodistal, labiolingual and apico-coronal location, as well as implant angulation.¹ The ultimate goal is not only to avoid sensitive structures, but to respect the established biological principles to achieve esthetic results.

Mesiodistal criteria

Correct implant position in a mesiodistal orientation allows the clinician to avoid damaging adjacent critical structures. A minimum distance of 1.5 mm between implant and existing dentition prevents damage to the adjacent teeth and provides proper osseointegration and gingival contours²⁻⁴ (Fig. 1a).

Distances of less than 3 mm between two adjacent implants leads to increased bone loss and can reduce the height of the inter-implant bone crest. A distance of more than 3 mm between two adjacent implants preserves the bone, giving a better chance of proper interproximal papillary height (Fig. 1b).

Labiolingual criteria

An implant placed too far labially can cause bone dehiscence and gingival recession while an implant placed too far lingually can cause prosthetic difficulties. A thickness of 1.8 mm of labial bone is critical in maintaining an implant

soft-tissue profile⁵ (Fig. 2).

Labially oriented implants compromise the subgingival emergence profile development, creating long crowns and misalignment of the collar with respect to the adjacent teeth.⁶

Apico-coronal criteria

Peri-implant crestal bone stability plays a critical role in the presence of interdental papilla.⁷ Implants placed too shallow may reveal the metal collar of the implant through the gingiva. Countersinking implants below the level of the crestal bone may give prosthetic advantages but can lead to crestal bone loss.

The ideal solution would be the placement of an implant equicrestal or subcrestal to the ridge. However, the existing microgap at the implant abutment junction leads to bone resorption because of peri-implant inflammation.⁸ It is suggested an implant collar be located 2 mm apical to the CEJ of an adjacent tooth if no gingival recession is present⁹ (Fig. 3).

Implant angulation

Implant angulation is particularly important in treatment planning for screw-retained restorations. Implants angled too far labially compromise the placement of the restorative screw while implants angled too far lingually can result in unhygienic and unesthetic prosthetic design.

For every millimeter of lingual inclination, the implant should be placed an additional millimeter apically to create an optimal emergence profile.¹⁰ In general, implant angulation should mimic angulation of adjacent teeth (Fig. 4). Furthermore, maxillary anterior regions require a subtle palatal angulation to increase labial soft-tissue bulk.¹¹

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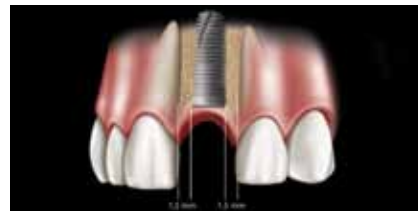


Fig. 1a: Minimum distance of 1.5 mm between implant and existing dentition. Photos/Provided by Glidewell Laboratories

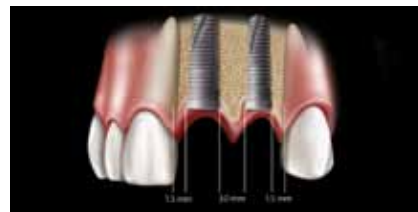


Fig. 1b: Minimum distance of 3 mm between two adjacent implants.



Fig. 2: Proper labiolingual placement with 1.8 mm thickness of labial bone.

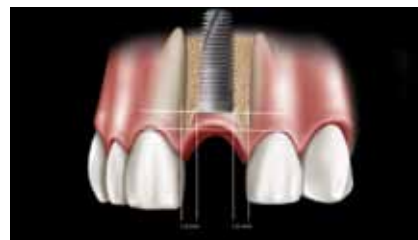


Fig. 3: Lateral view of implant placed with the collar at the level of crestal bone with adjacent teeth CEJ 2 mm coronal to the collar of the implant.

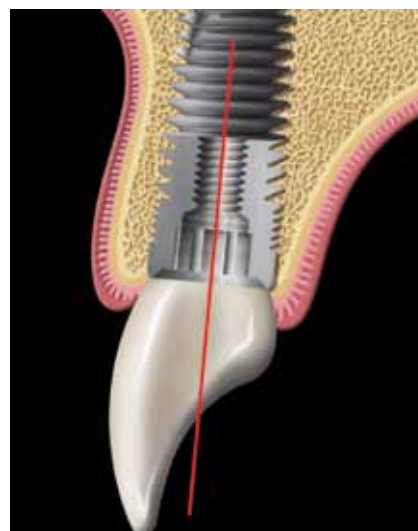


Fig. 4: Proper implant angulation with screw access in the cingulum area.



Fig. 5a: Inclusive Tapered Implant at placement.



Fig. 5b: Inclusive custom healing abutment in place.



Fig. 5c: Contoured soft-tissue sulcus after healing.



Fig. 5d: Screw-retained IPS e.max[®] crown (Ivoclar Vivadent; Amherst, N.Y.) in place.



Fig. 5e: PA to verify seating of crown.



Fig. 5f: Buccal view of final restoration at delivery.

Controlled tissue contouring for immediate loading including BioTemps® Provisionals

INCLUSIVE® Tooth Replacement System



A 3.7 mm X 11.5 mm Inclusive® Tapered Implant was placed in the surgical site for tooth #7, as well as tooth #10.



BioTemps bridge #7-10 was seated, allowing for nice tissue healing prior to the final impressions.



After four months of integration, the BioTemps and patient-specific abutments were removed, revealing very healthy pink tissue with trained tissue contours.



The custom impression copings match the patient-specific soft tissue contours of the custom temporary components.



The final zirconia abutments were fabricated by the dental laboratory and torqued into place.



The final implant-retained bridge was cemented into place.

Clinical dentistry by Timothy F. Kosinski, DDS, MAGD

\$695* Single Tooth Replacement
(Case as shown: \$1,646*)

System includes: implant, drills, custom healing abutment, custom temporary abutment, BioTemps Provisional, custom impression coping, final custom abutment and ceramic crown.

*Price does not include shipping or applicable taxes.

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All Inclusive implants, abutments and components are manufactured in our Irvine, Calif., facility.



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Premium Products - Outstanding Value

The Inclusive Tooth Replacement System is a convenient and effective method of promoting and preserving the ideal tissue contours throughout the implant treatment process. It includes treatment options for these clinical situations:

- ◆ Immediate loading with custom temporary abutment and BioTemps® Provisional
- ◆ Single-stage implant placement with custom healing abutment