NovaBone® Dental Putty

Clinical Publications
NovaBone Dental Putty is a Calcium-Phosphosilicate bone substitute that uses a next-generation uni-dose cartridge system engineered for enhanced handling and improved performance.

- Available in pre-filled syringes and a unique proprietary cartridge delivery system.
- NovaBone is available in disposable uni-dose cartridges.
- The diameter of the cannula is 2.8 mm (inner) and 3.0 mm (outer), designed to be ideal for dispensing of the graft, especially in minimally invasive techniques such as gaps in immediate implant and crestal approach sinus lifts.
- Cartridges are available in various sizes and are used in conjunction with the cartridge dispenser.
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Placement of Implants Simultaneously With Direct Sinus Lift in Ridges With Minimum Remaining Bone: Lack of Association Between Residual Bone Height and Primary Implant Stability

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Background
In case of minimum residual bone height, direct sinus augmentation is utilized prior to implant surgery aiming to provide adequate ridge dimensions. Most clinicians consider 5 mm of residual bone height to be the threshold for simultaneous sinus lift and implant placement.1 In cases of ridges with minimum residual bone height, a staged approach can be a lengthy procedure that increases treatment time until final restoration. The present clinical investigation was conducted to assess the potential benefit of utilizing a putty bone graft with a viscous consistency2 to ascertain primary implant stability during simultaneous implant placement in ridges with minimum residual bone height.

Aim
The aim of this retrospective study was to evaluate the clinical outcomes following placement of implants simultaneously with lateral window sinus augmentation with a calcium-phosphosilicate (CPS) putty bone substitute (NovaBone Dental Putty, NovaBone Products Alachua, FL) in ridges with minimum residual bone height.

Materials & Methods
Seventeen healthy, adult patients with less than 5 mm of vertical bone height in at least one posterior maxillary site, underwent sinus floor elevations according to a modification of the out fracture osteotomy technique.3 Neighboring sites that had greater than 5 mm distance from the floor of the sinus were included in the analysis as a separate subgroup to evaluate the effect of residual bone height in primary implant stability. Following elevation of the Schneiderian membrane, the CPS putty bone substitute was delivered pre-mixed using a cartridge delivery system and was directly injected into the prepared sinus cavity.4 A total number of 30 implants (Internal Tapered, BioHorizons, Birmingham, AL, USA) were inserted in the severely resorbed ridges (<5 mm) and their neighboring sites (≥5 mm) and primary implant stability for each subgroup was recorded. Primary stability was recorded as the maximum insertion torque (MIT) achieved using a torque wrench for the placement of the implant in its final position.2 Two distinct torque values were used as reference points (20 N/cm², 35 N/cm²) and the MIT for each implant was recorded as greater, equal or lesser than the corresponding reference.2

Fig 1:
A. Intra-operative view of the prepared bony window on the lateral wall of the sinus with the aid of a piezotome.
B. The out fractured bone wall was kept in saline throughout the procedure. Sinus curettes were utilized to perform careful elevation of the Schneiderian membrane across the floor of the sinus.
C. The putty bone substitute was delivered into the sinus using a cartridge delivery system that simplified the grafting procedure. An implant in the second premolar position was placed in the native bone and thus excluded from the analysis.
D. An implant in the 1st molar position was placed in the augmented sinus. The out fractured window was replaced in the lateral wall of the sinus.
Results

- No patients experienced any complications associated with the sinus surgery, or the implant placement. 0% sinus perforations.
- The preoperative ridge height ranged from 3.32 mm to 8.14 mm with 61.29% of the sites exhibiting less than 5 mm of native bone height.
- In all cases (100%) at least 20 N/cm² of MIT were achieved.
- Logistic regression analysis failed to show that the initial bone height of the residual ridges was associated with the MIT achieved during implant placement (P>0.05).
- At the second stage, appointment of the mean vertical bone height was 13.32 mm (±1.82 mm) as seen on post-operative radiographs. All implants successful osseointegrated (100% success). The difference between the initial vertical bone height and the bone height at the time of loading displayed a highly statistically significant difference (P<0.001).

Conclusion

Within the limitations of this study, we conclude that:

- The placement of implants simultaneously with direct maxillary sinus lift utilizing the proposed technique for adequate primary stability may be a viable treatment option in severely resorbed posterior maxillary sites.
- The diminished preoperative vertical dimensions of the residual ridges did not seem to negatively influence the osseointegration of implants placed in this study.

References


Poster Presented at
The Academy of Osseointegration Annual Meeting, Mar 6 – Mar 8, 2014, Seattle, WA
Use of a Viscoelastic Putty Alloplast for Minimally Invasive Indirect Sinus Elevation

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†Advanced Education Program in Periodontology, University of Minnesota, MN, ††Private Practice, Mumbai, India ‡Private Practice, New Delhi, India

Background
Use of percussive forces with osteotomes for indirect sinus elevation (ISE) is a successful modality despite incidence of Schneiderian membrane (SM) perforation as high as 26% and reported benign paroxysmal positional vertigo (BPPV). Transalveolar sinus floor elevation (TSFE) employing hydraulic pressure has been advocated for ISE to minimize complications associated with osteotomes. This study used viscoelastic calcium-phosphosilicate alloplastic putty (CPS) [NovaBone Dental Putty®, NovaBone LLC, Alachua, FL] for TSFE. The viscoelastic biomaterial acts as a “protective cushion”, yet permits hydraulic pressure to lift the SM while minimizing risks of 1) BPPV or 2) mechanical perforations of SM associated with osteotomes. The purpose of this case series is to assess 6 months post-loading clinical, radiographic and patient-related outcomes following CPS use for TSFE with simultaneous implant placement.

Methods
Twenty patients (12 male, 8 female) exhibiting an edentulous maxillary posterior region with residual ridge height > 5 mm were enrolled. Implant osteotomies were intitiated to ~ 1 mm below sinus floor, ~ 0.2 cc of CPS putty injected to base of osteotomy, and sinus floor in-fractured via osteotome and mallet. Perforation of SM was assessed via palpable resistance with a sinus probe [Sinus Crestal Approach Kit, Zimmer Dental, Carlsbad, CA]. Additional 0.5 cc increments of CPS were injected in osteotomy so implant insertion produced hydraulic pressure to elevate sinus floor and SM. Periapical radiographs obtained at right-angle projection were digitally calibrated using reference points on adjacent teeth to compensate for potential foreshortening measurements and Sinus Graft Remodeling Index (SGRI), respectively at 4-months post-surgery and 6-months post-loading.

Illustrations
Easy access into the sinus through the crestal floor for sinus augmentation with Osteotome technique can be achieved in 4 simple steps. Step 1: Prepare the osteotomy to less than 1 mm from the sinus floor. Step 2: An osteotome is then used to gently fracture the bone in the area. Step 3: The canula from the cartridge tip can be pressed against the surface of the bone and the putty is injected into the area resulting in membrane elevation with hydraulic pressure from Putty delivery. Step 4: An implant can then be placed in the augmented area.

Representative Case 1

Fig A: Tracing of incisions
Fig B: Reflection of full thickness mucoperiosteal flap
Fig C: Application of an osteotome following placement of a small quantity of putty that will function as a “cushion” during percussion of the sinus floor
Fig D: Adaptation of the cartridge in the osteotomy site and ejection of the CPS putty
Fig E: Insertion of a Tapered Internal implant in the osteotomy
Fig F: The implant achieved optimal primary stability
Fig G: Pre-operative radiographic view
Fig H: Post-loading radiographic view showing a radiopaque line delineating the new floor of the sinus
Results
Twenty-eight implants were placed in conjunction with TSFE utilizing CPS putty. All sites healed uneventfully. Median time to loading was 4 months. Mean residual ridge height was 7.34 mm (SD ±1.41) (intra-observer r – 0.89). Incidence of intraoperative SM perforations and/or BPPV was zero. At implant insertion a mean 2.71 mm (SD ±1.36) of linear radiopaque CPS graft was apical to the implant while original sinus floor lamina dura was still recognizable (SGRI-score 1). At mean 16 months follow-up (range 7-31 months) CPS was reduced to 1.71 mm (SD ±1.26). A “new sinus floor outline” (SGRI-score 3) was noted in 81.25% of patients. Mean linear radiographic bone gain was 4.95 mm (SD ±1.99). All implants were osseointegrated at 6-months postloading for 100% survival rate. No biological or restorative complications noted. All patients indicated treatment outcome as “highly satisfied.”

Discussion
The presented technique utilizes a simplified minimally invasive TSFE technique based on the application of hydraulic pressure by a viscous bone graft that is designed to act as an incompressible fluid. The continuous flow of the viscous biomaterial via a cartridge system results in the gradual and continuous elevation of the membrane. The use of the presented technique for sinus floor elevation may offer a conservative approach for localized augmentation of the sinus floor while minimizing postoperative complications. The consistency of the putty helps in minimizing membrane perforations and associated adverse events during percussion with osteotomes.

Conclusion
Use of viscoelastic CPS putty for TSFE aims to provide a clinical option for simultaneous placement of implants with minimal complications or post-operative morbidity.

References

Implants placed simultaneously with lateral window sinus augmentation using a putty alloplastic bone substitute for increased primary implant stability: a retrospective study.

Kher U, Mazor Z, Stanitsas P, Kotsakis GA.

Introduction: The aim of this retrospective study was to evaluate the primary stability of implants placed in significantly pneumatized maxillary sinuses with minimum residual bone height.

Methods: Seventeen patients who had been treated with simultaneous implant placement in sites with <5 mm of vertical bone height using a modified direct sinus lift technique were included. Implants placed in adjacent sites with at least 5 mm of bone height were included as quasi-controls.

Results: A total of 30 implants were inserted with a maximum insertion torque number >20 N/cm. Logistic regression analysis failed to show any association between residual bone height and primary implant stability. Implant survival was 96.67% (29/30) during a mean follow-up of 15.74 months postloading.

Conclusions: The diminished preoperative vertical dimensions of the residual ridges did not seem to negatively influence the osseointegration of implants placed in this study. The prerequisite for simultaneous sinus augmentation and implant placement is an adequate primary stability of the implant and not a fixed minimum bone height level.


A simplified approach to the minimally invasive antral membrane elevation technique utilizing a viscoelastic medium for hydraulic sinus floor elevation.

Kotsakis GA, Mazor Z.

Purpose: Maxillary sinus augmentation surgery is frequently employed to provide adequate vertical bony dimensions in posterior maxillary sites. When significant gain in bone height is sought for, an invasive lateral-window approach is routinely used to achieve sinus floor elevation. The minimally invasive antral membrane elevation technique was initially conceived as a surgical improvisation that has been shown to lead to up to, or exceeding, 10 mm of bone height, while enhancing the profile of the transalveolar sinus augmentation technique. This approach is based on the use of hydraulic pressure that is applied to the Schneiderian membrane via a saline-inflatable balloon. Even though this technique has been shown to be a treatment modality, the need for specialized equipment, training, and corresponding costs may hinder its widespread application. The purpose of this clinical paper is to introduce a simplified approach to the minimally invasive antral membrane elevation technique.

Methods: The simplified minimally invasive antral membrane elevation technique is based on the application of hydraulic pressure by a viscous bone graft that acts as an incompressible fluid. The specific clinical steps of this technique will be demonstrated to illustrate how grafting of the maxillary sinus is achieved simultaneously with the atraumatic elevation of the Schneiderian membrane, thus resulting in even less operative time.

Conclusions: This simplified technique may make the minimally invasive antral membrane elevation technique more accessible to implant surgeons as it eliminates the need for purchase of specialized equipment and aids in further decrease of intra-operative time accomplished with the original technique.

Sinus elevation with an alloplastic material and simultaneous implant placement: a 1-stage procedure in severely atrophic maxillae.

Jodia K, Sadhwani BS, Parmar BS, Anchlia S, Sadhwani SB.

**Aim & Objective:** The aim of the study is to evaluate clinically and radiographically the long term success of one-stage direct (lateral) sinus lift procedure using alloplastic bone graft material and bio-absorbable membrane in conjunction with two stage implant placement in atrophic partially edentulous posterior maxilla.

**Methods:** One stage direct maxillary sinus lift in conjunction with two stage implant placement was carried out in 12 patients at 13 sites. All the patients were partially edentulous with posterior maxillary alveolar ridge height of >5 mm and were in the age group of 20-50 years. Bioactive glass putty, bio-absorbable collagen membrane and 3.75 × 11.5 mm implants were used. Loading of implants was done 6 months after placement of implants. Patients were evaluated clinically and radio-graphically 6, 18, 30 months after placement of implants to assess increase in residual ridge height, peri-implant condition (marginal bone loss, plaque and gingival index) and implant stability.

**Results:** Maxillary first molar was the most common site (69.23 %) for sinus lift and implant placement. Caries was the most common cause (76.92 %) for loss of tooth. Increase in residual ridge height ranged from (71.43 to 133.33 %) as measured by Denta-Scan. Implant survival rate was 100 %. Marginal bone loss ranged from 0.68 to 1.22 mm. Implant stability was measured by periotest (-2.7 to -3.6). Only one patient had perforation of sinus membrane, but it was sealed satisfactorily by bio-absorbable membrane.

**Conclusions:** One stage lateral sinus lift procedure with alloplastic bone graft material in combination with 2 stage implant placement has a predictable outcome in patients with severe resorption of posterior maxilla.


Clinical and histologic comparison of two different composite grafts for sinus augmentation: a pilot clinical trial.


**Background & Objectives:** Many different techniques as well as bone graft regimens have been suggested for performing this procedure. It was the goal of this study to compare, clinically and histologically, two different composite grafting regimens used for sinus augmentation.

**Materials & Methods:** Five patients, needing a bilateral sinus augmentation to allow implant placement, were recruited for this study. Right sinuses were grafted with cortical bone (collected from overlying the sinus membrane) and bovine hydroxyapatite (HA), while the left side sinuses were grafted with overlying autologous bone plus a bioglass (BG) material. Bone core biopsies were taken at 6 months after sinus graft or at the time of implant insertion. A waiting period of 6 additional months was granted to allow healing, before prosthetic restoration and functional loading. The level of peri-implant bone was evaluated 12 months after loading. A comparative histomorphometric analysis was conducted and a statistical analysis was performed.

**Results:** All implants in both groups were functional after a 12-month loading period. No bone loss was observed radiographically or clinically in both groups. Histologic analysis revealed that both composite grafts had a high biocompatibility. In the bovine HA-containing group, minimal xenogenic graft absorption was noted. In contrast, BG group samples presented a high absorption rate with some remaining particles imbedded in new normal bone.

**Conclusions:** Sinus augmentation using a combination of autogenous bone plus either bovine HA or BG is a predictable technique.

Histological and Clinical Evaluations of NovaBone Dental Putty vs. Dynablast in Extraction Sockets

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†University at Buffalo, School of Dental Medicine, Department of Restorative Dentistry, Buffalo, NY, ††University of Kentucky, College of Dentistry, Department of Restorative Dentistry, Lexington, KY.

Background
Alveolar bone loss following tooth loss is an ongoing process due to the lack of functional stimulation. The rate and degree varies among individuals, but is greatest (40% — 60%) during the first few months or years after tooth loss (1). Prosthetic reconstruction and replacement of teeth (and associated structures) becomes more difficult and often unsatisfactory when anatomic deformities in the jaw bone ridges occur. Available evidence suggests that alveolar bone ridge resorption may be attenuated or prevented, at least initially, by reconstructive surgical techniques employed at the time of tooth removal. Several materials have been promoted for ridge preservation and include socket filling materials such as non-resorbable hydroxylapatites (particulate or solid block) and polymers (such as HTR Synthetic Bone), resorbable materials such as autogenous bone, particulate freeze-dried bone allograft (demineralized or non-demineralized), natural hydroxylapatite (bovine and equine derived), synthetic “resorbable” hydroxylapatites, and tricalcium-phosphate; and both non-resorbable (e-PTFE) and resorbable (collagen or polyactic acid-type) guided tissue regeneration barriers (2). Furthermore, the bone graft materials have been marketed primarily in form of particles as well putty.

The purpose of this clinical study was to assess the clinical and histological outcomes of two marketed putty graft materials used for socket preservation following dental extractions, one being an allograft, one an alloplast. This is a clinical and histological study with limited data submitted at this time.

Materials and Methods
For this study, an Alloplast material in putty form, NovaBone (NovaBone Products) was used in comparison with an allograft in putty form, Dynablast. Upon institution’s IRB approval, generally-healthy males and females volunteers adult between the age of 18 and 75, were scheduled for extraction of single-rooted teeth and socket preservation; upon enrollment, subjects were randomly assigned to either NovaBone Dental Putty or Dynablast at time of tooth-extraction; reason for extraction was non restorability of the involved teeth (1A-2A).

Surgical Procedure
The extraction procedures followed the method previously described by Kutkut et al (3). Briefly, following gaining anesthetic effect with local infiltration of anesthetic agents, the sulcus of the teeth was incised (1B-2B). Periotomes were used to gently luxate the tooth to gain mobility, then extraction forceps used to remove the tooth (1C-2C). The sockets were then degranulated with Bone curettes, and then grafted. The graft materials were inserted in the sockets up to reach the alveolar bone crest (1E, 2E). The grafted sockets were covered with a collagen membrane (Conform, ACE) positioned under the unsupported free gingival margin, without elevation of any flap (1F, 2F). The margins of the socket were sutured with Cytoplast 3.0 (Cytoplast). During the suturing procedure, the collagen membrane was protected with a small perioseal elevator, with the intent of not engaging the barrier with the suture (1G, 2G). Clinical measurements at baseline and 5 months included: presence of attached gingiva (AG), ridge width (RW) at the crest, ridge width at 10 mm apically to the crest (1D-, 2D). Membrane exposure was also measured following suturing and at 1 and 2-weeks post-op to monitor the wound closure.

Bone biopsies were harvested at 5-7 months (1H-1I, 2H-2I) when implants were placed (1J, 2J) with primary stability at 35 Ncm. Bone cores were immersed in 10% buffered formalin and processed for undecalcified histological analysis. At this time, preliminary clinical data from 8 subjects, 4 for each group, are presented. However, histological data for 7 subjects (NovaBone n=3; Dynablast n=4) are presented.

Socket / Ridge Regeneration
Results

Mean values for AG were 3.75 mm (±1.89) at baseline and 5.25 mm (±1.25) at 5 months for NovaBone (n=4); 5.75 mm (±2.75) at baseline and 6.0 mm (±1.41) at 5 months for Dynablast. Differences between the 2 groups were not statistically significant. Mean values for RW at the crest were 8.25 mm (±2.36) at baseline and 6.75 mm (±2.5) at 5 months for NovaBone; for Dynablast, values were 7.75 mm (±0.95) at baseline and 7.63 mm (±2.05) at 5 months. Measurements of the RW 10 mm apical to the crest were 13.87 mm (±3.37) at baseline and 13.5 mm (±2.38) at 5 months for NovaBone; for Dynablast, 12.5 mm (±1.73) and 12.5 mm (±2.38). There were no statistically significant differences between the 2 groups. For NovaBone, collagen exposures mean values were approximately 38.07 mm² (±7.43) after extraction to 17.2 mm² (±3.68) at week 1 and 5.29 mm² (±3.58) at week 2; for Dynablast, 27.66 mm² (±14.62) to 11.96 mm² (±11.14) at 1 week and 7.85 mm² (±5.28) at 2 weeks.

Histomorphometry

A 2.7 mm inner diameter trephine was used to obtain the core from the center of the grafted area upon re-entry at 5 months to evaluate the quality and quantity of bone. The cores were preserved in 10% formalin and sent to the Hard Tissue Laboratory at University of Minnesota for histological & histomorphometric analysis. Upon receipt, specimens were dehydrated with a graded series of alcohols for 9 days. Following dehydration, the specimens were infiltrated with a light-curing embedding resin (Technovit 7200 VLC, Kulzer, Wehrheim, Germany). Following 20 days of infiltration with constant shaking at normal atmospheric pressure, the specimens were embedded and polymerized by 450 nm light. Specimens were prepared in an apico-coronal direction and were cut into a thickness of 150 μm on a cutting/grinding system (EXAKT Technologies, Oklahoma City, OK). The cores were polished to a thickness of 45-65 μm using a micro-grinding system. The slides were infiltrated with constant shaking at normal atmospheric pressure, 7200 VLC, Kulzer, Wehrheim, Germany). Following 20 days of infiltration, the specimens were embedded and polymerized by 450 nm light. Specimens were prepared in an apico-coronal direction and were cut into a thickness of 150 μm on a cutting/grinding system (EXAKT Technologies, Oklahoma City, OK). The cores were polished to a thickness of 45-65 μm using a micro-grinding system. The slides were stained with Stevenel’s blue and Van Gieson’s picro fuchsin and cover-slipped for histologic analysis using bright-field and polarized microscopy.

<table>
<thead>
<tr>
<th>NovaBone</th>
<th>Dynablast (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Attached Gingiva (AG) (mm)</td>
<td>3.75 ± 1.89</td>
</tr>
<tr>
<td>Ridge Width (RW) – Crest (mm)</td>
<td>8.25 ± 2.36</td>
</tr>
<tr>
<td>RW– 10 mm apical to crest (mm)</td>
<td>13.87 ± 3.37</td>
</tr>
</tbody>
</table>

Table 1: Clinical Measurement Data at baseline and 5 months post-operative

Fig 3A: NovaBone 40x magnification at 5 mos  
Fig 3B: Dynablast 40x magnification at 5 mos

Discussion

Several materials can be used as graft immediately following tooth extraction, to preserve alveolar ridge dimensions, with the aim of future simple implant placement. Putty materials have been used, particularly allografts and alloplasts, such as bioactive phosphosilicate putty. The results of this study indicates that both materials can be used, and that clinician or patient’s choice can be satisfied with either material. In the present study, approximately 5-7 months following extraction, osteotomies prepared in either graft material sockets, and implants placed, resulted in implant primary stability. The data confirms that bioactive calcium-phosphosilicate can be used as bone graft for socket preservation (4).

Conclusion

Although these are preliminary data and no final conclusions can be made, a trend is observed indicating that synthetic putty bone graft performs as well as human demineralized putty bone graft material in socket preservation procedures. However, the synthetic putty bone graft seems to favor soft-tissue healing. Histologically, the percentage of residual graft material for the human demineralized putty material is higher than the synthetic material tested in this study.

References


Poster Presented at

The Academy of Osseointegration

Annual Meeting, Mar 7 – Mar 9, 2013, Tampa, FL
Socket Grafting with Alloplastic Bone Putty and Xenograft Particulate: A Histological Comparative Evaluation

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Background
Ridge preservation procedures aim in the timely regeneration of bone in the extraction socket and the preservation of the alveolar ridge dimensions during the healing period to provide a favorable site for implant placement.1, 2 Different biomaterials exhibit individualized absorption rates and lead to different percentages of vital bone formation in the healed sockets.3, 4 The aim of this study was to histologically evaluate and compare bone regeneration in post-extraction sockets utilizing either a putty calcium-phosphosilicate alloplastic bone substitute (CPS Putty) (NovaBone Dental Putty, NovaBone Products, Alachua, FL) or a particulate bovine xenograft (BO) (Bio Oss, Osteohealth, Shirly, NY) for ridge preservation.

Methods
Twenty-two patients with a mean age of 51 years old scheduled to undergo tooth extractions and ridge preservation were included in this study. Willing participants signed the consent form and were enrolled in the study. The study was conducted in accordance with the Helsinki Declaration of 1975, as revised in 2000. Patients were recalled after 4 or 6 months according to a randomization list to evaluate healing and to obtain a core for histological analysis during implant placement.

Exclusion criteria:
• Medical history that contraindicates surgical treatment
• Medications that alter soft or bone tissue healing (e.g., biphosphonates, chronic steroid intake, etc)
• Pregnancy for females
• Acute periodontal or odontogenic disease
• Destruction of more than 50% of the buccal plate at the time of extraction

Surgical Procedure
Ridge preservation was performed by utilizing the socket-plug technique as previously described by Kotsakis et al. (2012)5. Briefly, at the surgical appointment, care was taken to extract the teeth atraumatically in an attempt to preserve the remaining bone structure. Following extraction, the sockets were meticulously debrided and any inflammatory tissue was removed. Either a calcium-phosphosilicate alloplastic putty (CPS Putty) was placed into the sockets using a cartridge delivery system without any prior mixing or further manipulation, or particles of an anorganic bovine xenograft were mixed with sterile saline and packed into the socket. A collagen plug was utilized to occlude the sockets and a horizontal mattress suture was used to stabilize it. No pre- or post-operative antibiotics were administered, and all patients were placed on 0.12% chlorhexidine oral rinse post-operatively. Analgesics were prescribed on a need basis (Ibuprofen, 400 mg). Postoperative evaluations were performed at 7 and 14 days and at 4 weeks to assess wound healing and record any adverse events.

Histomorphometry
Following either 4 (subgroup 1) or 6 months (subgroup 2) of healing, surgical reentry was performed in the healed ridges for implant placement. Prior to the initiation of the osteotomies, a trephine bur with a 2.7 mm – 3.0 mm internal diameter was used to obtain a bone core from each site. The cores were preserved in 10% formalin and sent to the Hard Tissue Laboratory at University of Minnesota for histological & histomorphometric analysis. Upon receipt, specimens were dehydrated with a graded series of alcohols for 9 days. Following dehydration, the specimens were infiltrated with a light-curing embedding resin (Technovit 7200 VLC, Kulzer, Wehrheim, Germany). Following 20 days of infiltration with constant shaking at normal atmospheric pressure, the specimens were embedded and polymerized by 450 nm light. Specimens were prepared in an apico-coronal direction and were cut into a thickness of 150 μm on a cutting/grinding system (EXAKT Technologies, Oklahoma City, OK). The cores were polished to a thickness of 45-65 μm using a micro-grinding system. The slides were stained with Stevenel’s blue and Van Gieson's picro fuchsin and cover-slipped for histologic analysis using bright-field and polarized microscopy.

Poster Presented at
The Academy of Osseointegration Annual Meeting, Mar 7 – Mar 9, 2013, Tampa, FL
Results
Post-operative healing was uneventful in all sockets from both the test groups. Histomorphometry revealed that the residual graft values were significantly higher in the xenograft group compared to the alloplast group at the 6 month observation period. For cores harvested at 4 months post-extraction the amount of residual bone graft in the xenograft group was higher as well, but the difference did not reach statistical significance. Table 1 displays vital bone in defects filled with CPS Putty that ranges from 36% - 57% (average 47.15% ± 8.5%). Residual bone graft (RBG) was found to range from 30% at 4 months to a minimum of 3% at 6 months and an average of 17.4% ± 9.4%. The vital bone values in the BO group (Table 1) ranged between 17% - 27% (average 22.2% ± 3.5%). The difference in vital bone volume between the two groups was found to be statistically significant in favor of the CPS putty group P<0.05. Specimens in the BO group showed a range of RBG of a maximum of 38% at 4 months to a minimum of 18% at 6 months with an average of 25.7% ± 5.9%.

Table 1: Vital Bone & Residual Graft (RBG) values of both CPS & BO Product at varying time periods

<table>
<thead>
<tr>
<th>Tooth #</th>
<th>Months</th>
<th>% Bone</th>
<th>% RBG</th>
<th>Tooth #</th>
<th>Months</th>
<th>% Bone</th>
<th>% RBG</th>
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Table 2: Comparative evaluation of the RBG % between two study groups at two time intervals

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<th>BO</th>
<th>Months</th>
<th>% Bone</th>
<th>% RBG</th>
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<tr>
<td>4 months</td>
<td>23.67 ± 5.57</td>
<td>35.00 ± 4.24</td>
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<tr>
<td>6 months</td>
<td>8.03 ± 4.08**</td>
<td>23.14 ± 3.44</td>
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<td>All Samples</td>
<td>17.40 ± 9.39</td>
<td>25.60 ± 5.89</td>
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Discussion
Historically, the function of biomaterials has been to replace damaged and missing tissues. In contemporary implant dentistry, the aim of ridge preservation is to provide a healed site with adequate dimensions and favorable bone quality for the successful placement of an implant in the most timely manner. The quality and quantity of the regenerated bone should allow for the placement of an implant surrounded by bone tissue able to withstand functional loading. Results of histological analysis in our study revealed bone formation in sockets implanted with both biomaterials, with sockets grafted with CPS putty displaying consistently more bone regeneration than those in the bovine xenograft group. A similar percentage of residual bone graft was found at 4 months post-extraction in sockets of both groups with the those in the CPS putty group exhibiting a slightly smaller percentage in comparison to those in the bovine xenograft group. At the 6 month interval, sockets in the CPS putty group exhibited a statistically significant smaller amount of residual bone graft demonstrating the continued bone turnover in the CPS putty group, whereas in the xenograft group, the amount of reduction in the percentage of residual bone particles was not statistically significant between 4 and 6 months, implying that this graft requires a protracted period of healing.

Conclusion
The higher percentage of residual bone graft in the xenograft group in comparison to the CPS putty indicates an increased graft turnover rate in the latter group. Results from this study demonstrate that CPS putty may be the material of choice in comparison to anorganic bovine bone xenograft in ridge preservation procedures when timely implant placement is the desired outcome. Further randomized clinical trials are required to investigate whether the increased amount of new bone formation in the CPS putty group has a clinical impact on implant survival.

References
A Randomized, Blinded, Controlled Clinical Study of Particulate Anorganic Bovine Bone Mineral and CPS Putty Bone Substitutes for Socket Preservation.

Kotsakis, GA, Salama M, Chrepa V, Hinrichs JE, Gaillard P.

Introduction: The purpose of this study was to compare an anorganic bovine bone graft particulate to that of a calcium-phosphosilicate putty alloplast for socket preservation.

Methods: Thirty teeth were extracted from 24 patients. The sockets were debrided and received anorganic bovine bone mineral (BOV, n=12), calcium-phosphosilicate putty (PUT, n=12), or no graft (CTRL, n=6). The sockets were assessed clinically and radiographically 5 months later. Eight sockets in the BOV group and nine in the PUT group received implants 5 to 6 months post-grafting. The maximum implant insertion torque (MIT) was measured as an index of primary implant stability. The data were analyzed with the Mann-Whitney test.

Results: Both test groups had statistically significantly less reduction in mean ridge width (BOV: 1.39±0.57 mm; PUT: 1.26±0.41 mm) in comparison to the control group (2.53±0.59 mm). No statistically significant difference was identified between the test groups. MIT for PUT was 35 N/cm (MIT grade 4) for seven of the nine implants. MIT values in the BOV group ranged from grade 1 (10 to 19 N/cm) to grade 4, which was statistically significantly lower than for the PUT group. The overall implant success rate was 94.1% (16 of 17 implants were successful). No implants were lost in the PUT group; one implant failed in the BOV group.

Conclusions: Both tested bone substitutes can be recommended for preservation of alveolar ridge width following extraction. PUT might be more suitable for achieving primary stability for implants placed at 5 to 6 months postextraction.


Histomorphometric evaluation of a calcium-phosphosilicate putty bone substitute in extraction sockets.

Kotsakis GA, Joachim FP, Saroff SA, Mahesh L, Prasad H, Rohrer MD.

Objective: The objective of this study was to evaluate bone regeneration in 24 sockets grafted with a calcium-phosphosilicate putty alloplastic bone substitute.

Methods: A core was obtained from 17 sockets prior to implant placement for histomorphometry at 5 to 6 months post extraction. Radiographic analysis during the same postextraction healing period showed radiopaque tissue in all sockets.

Results: Histomorphometric analysis revealed a mean vital bone content of 31.76% (± 14.20%) and residual graft content of 11.47% (± 8.99%) after a mean healing period of 5.7 months.

Conclusions: The high percentage of vital bone in the healed sites in combination with its timely absorption rate suggest that calcium-phosphosilicate putty can be a reliable choice for osseous regeneration in extraction sockets.

Clinical Evaluation of 262 Osseointegrated Implants Placed in Sites grafted with Calcium-Phosphosilicate Putty: A Retrospective Study.

Babbush CA, Kanawati A.

Abstract
Along with the widespread use of dental implants, regenerative procedures have become an indispensable tool for implant surgeons in managing residual ridges and the surrounding bone. Putty bone grafts have significantly superior handling characteristics in comparison to particulates. These include ease of placement, enhanced particle containment, and a viscous consistency that has allowed for unique delivery systems to be developed. The aim of this study was to report the clinical efficacy of calcium phosphosilicate (CPS) putty in a wide variety of indications related to implant reconstruction and to report the survival rate of implants placed in these grafted sites. The CPS putty was used as the graft material of choice. Treatments were categorized into following groups: extraction graft, extraction with immediate implant placement, all-on-four concept, peri-implantitis treatment, bone augmentation before implant placement, implant replacement graft, and grafting around implant placed in resorbed ridges. Included in the analysis were 65 patients (36 men, 29 women) with a mean age of 63 ± 12 years. In total, 262 implants were placed. Four implants were diagnosed with peri-implantitis and were treated as described in category 4, for a total of 266 grafted sites. Two implants from the extraction graft category and 3 implants from the all-on-four group were lost and replaced with successfully osseointegrated implants during a mean study follow-up period of 12.24 ± 2.32 months. The implant success rate at 1 year was 98.1% (257/262). Based on results of this large-scale, retrospective study we conclude that (1) the use of putty bone grafts can simplify bone-grafting procedures and reduce intraoperative time in various grafting indications, (2) this study verified the efficacy of a CPS putty bone graft biomaterial in a large array of implant-related surgical indications, and (3) implants placed in sites grafted with CPS putty yield very high survival rates.


Socket Grafting with Calcium-Phosphosilicate Alloplast Putty: A Histomorphometric Evaluation

Mahesh L, Salama MA, Kurtzman GM, Joachim FP.

Background: Socket grafting with a bone graft substitute immediately after extraction is essential to preserve the ridge architecture for implant placement. Several bone graft substitutes have been tested for their ability to regenerate osseous tissue in the sockets. Evidence suggests that socket bone typically regenerates during a period of 6 to 8 months or longer, depending on several factors including the original ridge dimensions, type of graft, and the overall systemic health of the individual. The purpose of this study is to histologically evaluate the bone regeneration potential of a novel synthetic calcium-phosphosilicate putty (CPS) graft substitute.

Methods: After extraction of the involved teeth, CPS putty graft was placed, and the sockets were covered with a collagen plug. Cores were taken from 20 patients for histological evaluation prior to implant placement. Ten cores were processed decalcified with hematoxylin and eosin (H&E) stain and the remaining 10 were processed undecalcified. Histomorphometric data obtained from both sets is presented.

Results: Histomorphometric analysis revealed an average vital bone content of 49.5 (±20.7). A residual graft content of 4.3% (± 7.8) was observed following a healing time of 4.9 (± 0.8) months.

Conclusions: Clinical and histomorphometric data suggests that CPS putty is a good choice for socket bone regeneration.

Guided Bone Regeneration Simultaneously with Implant Placement Utilizing an Alloplastic Putty

Alejandro Kovacs†, Georgios A Kotsakis†, Tarun Kumar††
†Resident, Advanced Education Program in Periodontology, University of Minnesota, MN, ††Professor and Head, Department of Implantology, Bapuji Dental College & Hospital, Davangere, India.

Background
In edentulous sites where the residual ridge exhibits significant reduction in the orofacial plane that compromises the placement of an implant surrounded by adequate bone volume, ridge augmentation procedures should be employed. When guided bone regeneration (GBR) is performed simultaneously with implant placement to compensate for horizontal ridge deficiencies that leave implant threads exposed, it can significantly reduce healing time and eliminate the need for a second surgery, thus increasing patient acceptance. The aim of this case series is to report the clinical outcome of GBR with a calcium-phosphosilicate (CPS) alloplastic putty bone substitute performed simultaneously with implant placement.

Methods
Twelve patients presenting with Class I Seibert defects in 14 edentulous sites were treated with GBR using a CPS putty with a collagen membrane (CM) or titanium mesh (TiM) following implant placement. In order to be included in the study, at least one implant thread had to be exposed on the facial aspect of the implant following implant placement. During 1st stage surgery, the distance from the most apical level of the bone crest on the facial aspect of the implant to the platform of the implant was estimated. The same measurement was retaken during second stage surgery. All patients were followed-up at least 6 months after delivery of the implant-supported restorations.

Representative Case 1
A 22 yr old male patient presented with an edentulous area in the lower left first molar area. The ridge width as measured at the crest was 3 mm. A 4.2 mm x 11.5 mm MIS Seven Implant was placed in the area. A 4 mm dehiscence defect was noted after implant placement. The area was decorticated to facilitate bleeding and a macroporous titanium mesh (0.1 mm) was placed over the implant and contoured to adjacent bone envelope. The mesh was stabilized with titanium tacks. CPS Putty was injected inside the mesh through the sides using the unique cartridge delivery system. Tissues were approximated to get complete primary closure. There was no exposure of the titanium mesh during healing and second stage re-entry was performed at 4.5 months for mesh removal. Upon mesh removal, complete ridge regeneration in the area was noticed. The cover screw had to be exposed by scraping the newly formed bone using Piezo surgery. The implant was restored with PFM crown.

Representative Case 2
A 30 yr old female patient presented with edentulous area in the lower left first molar region. The buccolingual width was measured at 3 mm at the crest of the ridge. A MIS Seven 4.2 mm x 10 mm implant was placed in the area. A dehiscence defect exposing approximately 4 mm of the implant was noticed. The area was decorticated to facilitate bleeding and a titanium mesh was placed and contoured to adjacent bone. The mesh was stabilized with titanium tacks and CPS putty was injected underneath the mesh through the sides and complete primary closure was achieved. Titanium mesh exposure was noticed 4 months post operatively and re-entry showed complete bone regeneration in the area. The cover screw was exposed by scraping the bone with Piezo instruments. The area was restored with PFM crowns.
Results
In this study 6/14 sites were augmented with CPS putty and TiM while 8/14 were augmented with CPS putty and a CM. The average intra-operative defect measurement during 1st stage implant surgery was 4.57 mm (±2.50). After a median healing time of 5 months (range: 4-7 months) patients were scheduled for 2nd stage surgery. Intra-operative measurements showed the average defect distance to be 0.14 mm (±0.36), (P<0.01). Implants were not surrounded by hard tissue, resistant to probe penetration only in two sites that accounted for the above mentioned mean defect value measured at 2nd stage surgery. In one of the two cases with a remaining defect, a CM that was prematurely exposed had been used, while in the second case, a TiM had been utilized. In total, premature membrane exposure was noted in 2/6 cases with TiM and in 1/8 cases with CM. New tissue was almost completely covering the osseointegrated implant in the cases where premature TiM exposure occurred. All implants were successfully osseointegrated (14/14) at the time of 2nd stage and maintained successful osseointegration after a follow-up of at least 6 months post loading for a cumulative success rate of 100%.

Discussion
Guided Bone Regeneration is a very technique sensitive surgical procedure that requires the right combination of surgical techniques and biomaterial selection. Results from this 14 ase series indicated that CPS Putty in combination with either a CM, or a Ti-mesh produced excellent bone regeneration in GBR surgeries. CPS Putty is a third generation synthetic bone graft substitute. The material is cohesive and provides adequate retention at the defect site. Various studies indicate a capability possessed by CPS particles to stimulate differentiation towards cell lineage with therapeutic potential in tissue engineering.1,2 This unique phenomenon (osteostimulation) occurs exclusively with CPS based substitutes and has been shown to be superior to conventional osteoconduction. Recently several publications have been published on the bone regeneration efficacy of CPS Putty in a variety of indications.3,5

Conclusion
Calcium-phosphosilicate putty can be a successful scaffold for new bone growth in GBR procedures. Both a collagen membrane and a titanium mesh can be employed in conjunction with CPS putty for bone regeneration around implants. TiM may exhibit a more frequent exposure rate, which was not found to compromise the clinical outcome in this study. Further randomized, controlled clinical trials are required to validate our results.

References

Poster Presented at
The Academy of Osseointegration
Annual Meeting, Sept 29 – Oct 1, 2013, Philadelphia, PA
Horizontal Augmentation Utilizing Ridge-Split Technique & CPS Putty for Expansion of Alveolar Ridge in the Anterior Maxilla

Taran Kumar†, Sudhindra Kulkarni ††, Raison Thomas†, Dhoom Mehta†
†Bapuji Dental College, Davangere, India, ††SDM Dental College, Dharwad, India.

Background
The alveolar bone deficiencies are caused due to various factors that include: trauma, faulty extraction, socket compression, and long term edentulism. At times, the quantity of the bone may be adequate for implant placement, but may not be in the proper location for an appropriate maxillo-mandibular relation and thus necessitate procedures to augment the ridge. Edentulous ridges can be split and expanded to widen a narrow ridge to accept a standard-sized implant. Alveolar ridge expansion with Piezo powered saw and subsequent expansion of the split bone with motorized expanders can create space for Immediate Implant Placement. The Piezoelectric Saw is ideal for performing precise bone cuts while offering better control on the expansion. The aim of this study was to place the implant immediately by ridge split using the Piezo unit and Calcium-Phosphosilicate (CPS) for expanding the alveolar ridge.

Methods
A total of 15 patients who reported to the Department of Implantology and needed the correction of the deficient anterior maxilla were included in the study. All the patients were informed of the procedure and consent was taken. All the patients who were included in the study underwent a preoperative assessment that included models and wax-up to determine the desired tooth/implant positions. Afterwards, all the patients underwent a radiographic evaluation which included Periapical IOPAR and OPG. The following parameters were recorded during the study: 1. Pre-operative ridge width, 2. Post-operative ridge width during second stage while connecting healing caps, 3. Primary stability of the implants as measured in terms of insertion torque, 5. Implant survival and success.

Surgical Procedure
A Mid-crestal incision design was followed for all the cases with two vertical release incisions on the mesial and distal sides, and a full thickness flap was reflected. Using the piezo surgery, the sagittal bone cuts were made initially at the crest leaving at least 1 mm of margin at the palatal bone. The mesial and distal vertical cuts were made at a distance of 1 mm from the adjacent teeth. The implant sites were then designated as per the pre-operative plan with a pilot drill. Further expansion of the site was done with bone expanders (MIS expander screw kit) to the planned width. All the sites irrespective of implant placement were grafted with a calcium-phosphosilicate alloplastic bone substitute, performed and covered with either collagen membrane or titanium mesh over the split, and the flaps were released to achieve primary closure.

Representative Case 1
A 45 year old female patient had lost her teeth due to trauma 4 months back and desired replacement of the missing teeth. Clinical examination revealed missing teeth no. 11, 12 and ridge deficiency on the buccal aspect. The ridge preoperatively at the crest was 2.5 mm. CPS putty was injected between the split ridge. 4x13 mm Neobiotec ISII implants were placed in the region of 11 and 12 utilizing the surgical stent. RCM collagen membrane was utilized to cover the grafted site. 4.5 months later, an impression was made and PFM cemented crowns were delivered to the patient.
Representative Case 2
A 32 year old male patient had many missing teeth due to periodontal disease which were extracted 2-4 years back. The patient desired replacement with implant supported full arch restoration. Clinical examination revealed all the teeth were missing except canines and premolars. The anterior ridge in the region of the central incisor and lateral incisor was deficient on the buccal aspect. The radiographic findings and the implant planning revealed that the width was adequate and needed to be displaced buccal for optimal implant position. Pre-operative ridge width was 3 mm. After splitting the ridge as per the protocol followed, CPS putty was injected between the split ridge. 3.8 mm x 12 mm Biohorizons laserlok implants were placed in the region of 11 and 21 utilizing the surgical stent. RCM collagen membrane was utilized to cover the grafted site. 4.5 months later, an impression was made and hybrid screw retained was restoration delivered to the patient.

Results
The mean pre-operative ridge width was 2.9 mm (range 2.4 mm) and the postoperative width was 7.1 mm (range 5.5-8 mm). The smallest implant diameter was 3.5 mm and the maximum was 4.6 mm diameter (average of 3.8 mm). The insertion torque was 25-35 ncm (average of 24.6 ncm) in the single-staged cases, whereas in the two-staged case, the torque at placement was 35-40 ncm. 8 of the implants were restored with individual and remaining 22 with splinted crowns or bridge. None of the implants displayed any crestal bone loss, and the bone levels remained stable throughout the study.

Discussion
The alveolar ridge split procedure in combination with immediate implant placement has been described for more than 10 years. This procedure discards the need for onlay grafts. Moreover, it avoids the use of a secondary surgical site that exhibits postoperative morbidity associated with bone harvesting. In the present study we report a total of 15 cases where piezosurgical expansion of the maxillary anterior areas was done and 30 implants were placed. The total amount of expansion gained was 4.15 mm which facilitated placement of implants. The grafting procedure was carried out to contour augment the buccal plate to give long-term stability to both hard and soft tissues and also to avoid accidental buccal bone resorption, which would happen when the split buccal bone is thin. There are several advantages of a one stage split and placement, including but not limited to reduction in overall treatment time by an approximate 12-14 weeks and better soft-tissue control as the implant and the split heals at the same time. The soft-tissue from the buccal side that gets displaced to close the split in the first surgery can be recovered, reestablishing an adequate zone of keratinised tissue on the buccal aspect.

Conclusion
It can be concluded that the piezoelectric alveolar ridge splitting technique is a very successful alternative for extended grafting procedures if the ridge shows adequate height. Implants can be placed simultaneously after the ridge split and expansion with adequate insertion torques and achieve 100% survival and success. There is enhanced ease of grafting using CPS putty in between split ridges before implant placements without any voids left behind. Significant ridge width enhancement can be achieved with the use of this technique. Bone regenerated with collagen membrane along with CPS putty graft is comparable to augmenting the site with bioactive glass graft and a titanium mesh. Simultaneous placement of the implant with the split reduced the total treatment time.

References

Poster Presented at
The Academy of Osseointegration
Annual Meeting, Mar 6 – Mar 8, 2014, Seattle, WA
Clinical results of implant placement in resorbed ridges using simultaneous guided bone regeneration: a multicenter case series.
Konstantinidis I, Kumar T, Kher U, Stanitsas PD, Hinrichs JE, Kotsakis GA.

Objectives: The purpose of this case series was to evaluate the new bone formation following guided bone regeneration (GBR) with a calcium-phosphosilicate (CPS), alloplastic bone putty at peri-implant dehiscence defects and to assess survival rate of implants placed in the augmented sites after 12 months of function.

Methods: Implants were placed in patients exhibiting Seibert class I ridge defects resulting in peri-implant dehiscence defects. The defects were treated following GBR principles with the use of a CPS alloplastic bone graft putty in combination either with a collagen membrane or a titanium mesh. The height of each bony dehiscence was clinically measured at the time of implant placement and again during second-stage surgery. The percentage of complete defect coverage, frequency of adverse events, and risk factors for residual defect were determined.

Results: Thirty-six implants were placed in 26 patients. Twenty-seven of the 36 sites employed a collagen membrane in conjunction with the CPS while the remaining nine sites utilized a titanium membrane. Mean gain in bone height was 3.23 ± 2.04 mm, with 75 % of the peri-implant defects achieving complete regeneration. A negative correlation was identified between patient age and complete coverage of the peri-implant defect (p = 0.026). The implant survival rate at 12 months was 97.22 %.

Conclusions: Use of CPS bone putty during delayed implant placement at peri-implant dehiscence sites either in combination with a collagen membrane or a titanium mesh results in predictable defect coverage. The handling characteristics of CPS putty may simplify GBR protocol. Implants placed in conjunction with GBR have a very good survival rate after 1 year of follow-up.


Management of Fenestration Defects During Flapless Immediate Implant Placement in the Esthetic Zone.
Kher U, Tunkiwala A, Umrigar Z.

Abstract: The success of flapless immediate implant placement is dependent on the bony architecture on the buccal aspect of the socket. The presence of a fenestration defect in the buccal cortical plate may jeopardize the esthetic outcome, especially if the clinician does not undertake adequate soft-and hard-tissue augmentation procedures.

This article describes the use of an esthetic buccal flap design to deal with fenestration defects created during anterior implant placement immediately after extraction. This technique allows hard-tissue grafting of the fenestration defect around the implant in a postextraction socket.

Clinical and cone beam computed tomography comparison of NovaBone Dental Putty and PerioGlas in the treatment of mandibular Class II furcations.

Asmita, Gupta V, Bains VK, Singh GP, Jhingran R

Objective: The objective of this study is to compare the putty form of bioactive glass (NovaBone Dental Putty) and particulate form (PerioGlas) in the resolution of Class II furcation defects.

Background: Use of bone regeneration materials is becoming common in periodontal surgeries including furcation defects and the promising role of bioactive allograft materials has encouraged their presentation in different morphologic forms with their own advantages and disadvantages giving the operator ample of choices in his/her periodontal armamentarium.

Materials & Methods: A total of 28 patients with 40 Class II furcation defects were enrolled in the study and were randomly allocated to two groups with 20 sites in each group. Measurement of defects was done using clinical and cone beam computed tomography (CBCT) methods. The patients were followed-up at 6 months. Intergroup comparisons were done using Mann-Whitney U-test.

Results: There was no significance between group differences in clinical parameters and defect size at the baseline. After 6 months, particulate form showed a mean resolution of 50.48 ± 16.47% and 51.11 ± 9.48%, respectively for vertical defect and horizontal defect while putty form showed a mean resolution of 43.48 ± 9.33% and 42.88 ± 11.09%, respectively. Mean resolution in furcation width was 40.15 ± 13.00% for particulate form as compared with 36.27 ± 11.41% in putty form. Statistically, there was no significant difference between two groups except for the horizontal defect fill where PerioGlas showed statistically better results.

Conclusions: Putty form was comparable to particulate form for resolution of Class II furcation defects.


* PERIOGLAS Putty and NOVABONE Dental Putty are manufactured by NovaBone Products, LLC. PERIOGLAS®, NovaBone® and Smart Science™ are trademarks of NovaBone Products, LLC. NovaBone Putty is marketed as PERIOGLAS in China.

Use of the NovaBone augmentation material in the treatment of chronic periodontitis. Preliminary communication.

Slezák R, Paulusová V.

Abstract: The pilot study deals with the clinical results of the regenerative therapeutical method in intraosseous periodontal defects done by surgical augmentation procedure using a material of a new generation prepared on the basis of a bioactive glass.

Methods: A group of 10 individuals were treated properly for chronic periodontitis. Some of treated infraalveolar periodontal lesions, both periodontal pockets and interradicular defects, persisted or recurred in posterior teeth. In order to eliminate them they were indicated for advanced periodontal surgery or regenerative surgical therapy stimulating healing processes of supportive tissues, mostly alveolar bone around treated teeth.

Results: Relevant clinical parameters, i.e. values of the pocket depth, gingival recession, and loss of attachment were evaluated before and after the periodontal surgery and compared with each other. The therapy led to significant improvement of clinical parameters in terms of the reduction of pocket depth and loss of attachment values.

List Of Publications

Sinus Augmentation

Socket Preservation

GBR / Ridge Augmentation

Perio Defects
There are several ways to access the sinus via crestal approach and elevate the sinus membrane prior to augmentation: A) Osteotome Technique B) Piezo Technique C) Neurosurgical Bur D) Densah Burs, etc. Bone graft delivery into the sinus may be difficult and challenging. The NovaBone Cartridge System may simplify the delivery of graft into the sinus when accessed through the crest of the ridge. The tip of the cartridge is 2.8 mm in diameter and was designed specifically to deliver the graft seamlessly into the sinus. Putty consistency may help prevent membrane tears. The delivery of the putty graft may elevate the membrane with minimal instrumentation and hydraulic pressure.

Easy access into sinus through crestal floor for sinus augmentation with Osteotome technique can be achieved in 4 simple steps.

**Step 1**: Prepare the osteotomy to less than 1 mm from the sinus floor.

**Step 2**: An osteotome is then used to gently fracture the bone in the area.

**Step 3**: The canula from the cartridge tip can be pressed against the surface of the bone and the putty is injected into the area resulting in membrane elevation with hydraulic pressure from Putty delivery.

**Step 4**: An implant can then be placed in the augmented area.