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Comparative assessment of BMP-9 levels and clinical implant parameters in immediately restored implants using bone compaction drilling versus conventional drilling: a randomized controlled study.

ABSTRACT

Background- This study aimed to compare the effects of bone compaction drilling and conventional drilling techniques on bone marker BMP-9 and clinical implant parameters.

Materials and Methods: Forty-two participants were randomized into two groups (n = 21 each): OD and CD. Randomization was computer-generated with allocation concealment, and both participants and outcome assessors were blinded. Over a 6-month follow-up, peri-implant crevicular fluid (PICF) levels of bone morphogenetic protein-9 (BMP-9), implant stability, and crestal bone loss were evaluated.

Results: At 2 weeks, the CD group exhibited significantly higher BMP-9 levels (177.67 ± 8.24 pg/mL) than the OD group (150.43 ± 4.96 pg/mL; $p < 0.05$). By 16 weeks, the OD group showed greater BMP-9 expression (440.90 ± 33.57 pg/mL vs. 423.62 ± 15.58 pg/mL). Implant stability was consistently higher in the OD group at all time points, with ISQ values initially declining at 2 weeks and increasing thereafter in both groups. The OD group also demonstrated significantly less crestal bone loss at each follow-up interval ($p < .05$).

Conclusion:

Bone compaction drilling or osseodensification using Densah burs demonstrated enhanced secondary healing, as indicated by increased BMP-9 levels, along with improved implant stability and reduced crestal bone loss compared to conventional drilling. These findings indicate a potential biomechanical advantage for implant placement.

Keywords: BMP-9;Crestal bone loss; Immediate restoration; Implant; Implant stability; Osseodensification

INTRODUCTION

Dental implants have completely transformed the realm of oral rehabilitation with a well-documented success rate typically ranging between 90%- 95% after 10 years of follow-ups.^{1,2} However in the maxillary arch, there's often a deficiency in both the quality and quantity of bone, because of which achieving successful osseointegration of implants is particularly challenging. Insufficient bone surrounding implants may adversely affect implant stability, percentage of bone-to-implant contact (BIC) and bone volume (BV), which consequently delays osseointegration.³

In the past, undersizing the osteotomy,⁴ osteotome technique by Summer's⁵ and implant site preparation using peizosurgery⁶ have been introduced to improve osseointegration in low bone density areas. However major drawbacks like increased mechanical strain on the bone leading to bone compression and ischemia, increased marginal bone loss and risk of overheating associated with ultrasonic devices reduced the clinical implications of these techniques.^{6,7,8} Also use of wider, longer implants with a reverse buttress, larger thread depth, narrow pitch and a self-tapping design were reported to be beneficial in compromised bone sites.⁹

Huwais in 2013¹⁰ developed an innovative osteotomy preparation method known as osseodensification (OD). This bone compaction drilling technique sparked a significant change in the methods used for implant site preparation, being a non-subtractive drilling technique. Osseodensification technique is indicated where there is insufficient quantity or quality of bone. Relying on the elastic and plastic properties of bone, the

47 osseodensification (OD) technique aids in preserving the bone bulk, increasing the density
48 at the implant preparation site and also accelerate the formation of new bone.¹¹

49 Various animal studies¹²⁻¹⁶ have reported that the use of osseodensification or bone
50 compaction drilling compared to conventional drilling has resulted in higher insertion and
51 removal torque values, improved primary and secondary implant stability, higher BV and
52 BIC. This favours and rationalizes the use of bone compaction drilling technique.

53 Although numerous studies on animal models exists, its clinical impact in humans during
54 bone healing remains limited. Also, our understanding of the molecular mechanisms
55 involved in the process and its impact on crestal bone loss is lacking.

56 Various bone turnover markers act on the regulation of molecular and biological events
57 leading to osseointegration. Bone morphogenetic proteins (BMPs) belong to the
58 transforming growth factor beta (TGF- β) family. They play a crucial role in the formation
59 of bone and differentiation of stem cells. BMP-9, alternatively referred to as Growth
60 Differentiation Factor 2 (GDF2), is recognized as one of the most potent osteogenic bone
61 morphogenetic proteins.¹⁷ The levels of BMP-9 during the early and late stages of peri-
62 implant bone healing may reflect the biological advantages conferred by osteogenic
63 differentiation.¹⁸ Additionally, implant stability and crestal bone loss (CBL) remain critical
64 indicators of long-term implant success.

65 This study aims to compare bone compaction drilling and conventional drilling techniques
66 using both molecular (BMP-9 expression) and clinical (implant stability, CBL) parameters.
67 The null hypothesis posits that there is no significant difference between the two
68 techniques.

69 MATERIALS AND METHODS

70 A two year study was carried out in a tertiary hospital setting. Ethical approval (XIV-
71 PGTSC-IIA/PII) was obtained and registration in the Clinical Trial Registry of India
72 (ICMR-NIMS) (CTRI/2023/03/050409) was done prior to the commencement of the study.
73 The study followed a randomized controlled design (computer-generated random numbers)
74 with participants and outcome assessor blinding following the CONSORT (Consolidated
75 Standards of Reporting Trials) guidelines.¹⁹ Allocation of treatment was concealed using
76 opaque sealed envelope. Sample size of 21 participants in each group was calculated based
77 on the minimum difference $d = \max(\sigma_1, \sigma_2)$, considered to be clinically significant. Type I
78 error $\alpha = 5\%$ corresponding to 95% confidence level, Type II error $\beta = 10\%$ for detecting
79 results with 90% power of study.¹⁷

80 Participants visiting prosthodontics clinic for rehabilitation of a single edentulous space in
81 the maxillary arch that could be restored with an implant supported single unit crown were
82 screened as per predetermined inclusion and exclusion criteria (Figure 1). Participants
83 aged 18-60 years of both genders, capable of comprehending and signing an informed
84 consent, with a bounded edentulous space in the maxillary arch post-extraction for at least
85 3 months were included. Keratinized tissue >2 mm from mid crest to mucogingival
86 junction, simplified oral hygiene index of 0-3 indicating good to fair oral hygiene and
87 adequate bone for optimal implant placement and a safe distance (≥ 2 mm) from vital
88 tissues and an opposing dentition with a stable occlusion were assessed and included in the
89 study.

90 Participants with history of systemic conditions or under medications, presence of any
91 local risk factor, history of treated periodontitis, smoking, any parafunctional habits,
92 pregnant or lactating women were excluded. Also participants fulfilling any criteria either
93 group 1 or 2 according to second ITI (International Team of Oral Implantology) Consensus

94 report were excluded from the study.²⁰The participants underwent comprehensive clinical
95 examination, laboratory investigations, and radiological assessment before being enrolled
96 in the study.

97 All patients were prescribed prophylactic antibiotics before implant surgery. Drilling
98 technique for bone compaction drilling or osseodensification group (OD) was using

38 99 Densah Burs (Versah International, USA) (Figure 2 and 3) and for CD group was using the
100 conventional drills supplied by the manufacturer corresponding to the implant (Figure
101 2 and 4). MIS implants (MIS Implants Technologies Ltd, Israel) were used which have an
15 102 SLA treated surface with V shaped design, pitch of 1.0 mm, thread depth of 0.5 mm and an
29 103 internal hexagonal connection. Implants were placed 0.5 mm subcrestal into the prepared
104 osteotomy with the rate of 30 Ncm, and tightened using a torque wrench. Following
105 implant placement with either technique, immediate restoration was done with provisional
106 poly methyl methacrylate (PMMA) crown fabricated using CAD/CAM within 1 week
107 with prosthesis held out of occlusion.²¹ A follow-up program of six months was designed for
108 all participants. Final implant prosthesis was fabricated after appropriate healing time (6
109 months) abiding by the principles of implant protected occlusion.

110 Samples for the assessment of BMP-9 levels were collected at 2 weeks, 8 weeks and 16
111 weeks post-implantation. Samples were collected prior to clinical measurements to
112 minimize blood contamination. Area around implant was isolated with the help of cotton
113 roles and any layer of supragingival plaque, if present was removed carefully. After air
114 drying the area, sterile absorbent paper points were inserted in the sulcus until light
115 resistance was felt and held steady for 30 seconds. For each implant site, four samples
116 were collected per time point and were transferred into a single Eppendroff tube containing
117 100µl of phosphate buffer solution. (Figure 5A). Visibly contaminated samples with blood
12 118 were discarded and recollected after ensuring haemostasis. Samples were centrifuged at

119 1000 rpm for 10 minutes and stored at -80°C. While PICF volume was not measured
 120 directly, standardization was ensured by consistent sampling time and PBS dilution. BMP-
 121 9 levels were quantified using sandwich ELISA technique (E0051Hu, Bioassay
 122 Technology Laboratory, Zhejiang, China) and expressed in picogram/millilitre.
 123 Primary implant stability was assessed immediately after implant placement. Smart Peg
 124 (MIS Implants Technologies Ltd, Israel) specific to the implant system and the restorative
 125 platform diameter was utilized and subsequently resonance frequency analysis (RFA) was
 126 conducted using Osstell (Integration Diagnostics, Savedalen, Sweden)). Average of three
 127 readings was recorded in terms of implant stability quotient (ISQ). Secondary stability was
 128 recorded in similar way in terms of ISQ values assessed at 2 weeks, 8 weeks and 16 weeks.
 129 (Figure 5B).

130 Standardized periapical radiographs were captured using the long-cone paralleling
 131 technique, with individualized sensor holders employed for consistent and reproducible
 132 positioning. The acquired images were analysed using ImageJ software (National Institutes
 133 of Health, Bethesda, MD, USA). Dimensional calibration of the images was performed
 134 using the known length of the inserted implant to correct for any magnification errors.
 135 Crestal bone loss was measured by subtracting the baseline measurements recorded at the
 136 time of provisional prosthesis delivery from measurements taken at 2 weeks, 8 weeks, 16
 137 weeks, and 6 months postoperatively (Figure 5C). Radiographic analysis was conducted
 138 independently by two calibrated investigators. To assess intra- and inter-examiner
 139 reliability, 20% of the radiographs were randomly selected and re-evaluated after a two-
 140 week interval. Intra-class correlation coefficients (ICCs) were calculated, and values above
 141 0.85 were considered acceptable, indicating high reliability. To determine the method
 142 error, duplicate measurements were analysed using Dahlberg's formula, and the standard
 143 error of measurement (SEM) was calculated. This ensured that any reported differences in

bone loss between groups exceeded the magnitude of measurement error, thereby supporting the validity of the statistical findings.

Sample size was calculated based on the variation in BMP-9 levels among the study groups, using the appropriate formula which is

$$n = k \frac{(z_{\alpha} + z_{\beta})^2 (\sigma_1^2 + \sigma_2^2)}{d^2}$$

Where n = number of samples to be collected

$\sigma_1 = 300$, The SD of BMP-9 in first group with reduced torque, $\sigma_2 = 250$, The SD of

BMP-9 in second group with conventional torque.³³ The data were analysed with a

statistical software package (IBM SPSS Statistics, v26.0; IBM Corp) ($\alpha = .05$).²² Unpaired t test, repeated measures ANOVA followed by Bonferroni post hoc analysis and Pearson's correlation were the statistical tools used.

RESULTS

The OD group had notably lower levels of BMP-9 after 2 weeks compared to the CD

group (OD group: 150.43 ± 4.96 and CD group: 177.67 ± 8.24) which was statistically

significant ($P < 0.05$). However, the levels of BMP-9 were higher at 8 weeks (OD group:

274.19 ± 12.16 and CD group: 267.24 ± 14.50) and 16 weeks (OD group: 440.90 ± 33.57 and

CD group: 423.62 ± 15.58) in the osseodensification group with the difference being

statistically significant at 16-week ($P < 0.05$). There was a consistent upward trend in BMP-

9 levels from the 2nd week to the 16th week in both the groups indicating increased

formation of bone during this period. (Table 1)

The mean ISQ values at baseline (OD group: 75.19 ± 4.92 and CD group: 64.90 ± 6.80), at 2

weeks (OD group: 69.24 ± 7.74 and CD group: 61.43 ± 8.26), at 8 weeks (OD group:

72.76 ± 5.12 and CD group: 62.95 ± 8.40) and at 16 weeks (OD group: 74.62 ± 4.57 and CD

group: 64.57 ± 4.17) indicated that the implant stability of the OD group was more than the CD group which was statistically significant ($P < 0.05$) at every time point. There was a decrease in the ISQ values at 2 weeks when compared to the baseline and subsequently showed a gradual increase in both the groups. (Table 2)

There was evidence of significant difference ($P < 0.05$) in the crestal bone loss of both the

OD and CD groups at all the time intervals. The mean values of crestal bone loss in mm at 2 weeks (OD group: 0.07 ± 0.01 and CD group: 0.10 ± 0.03), at 8 weeks (OD group: 0.24 ± 0.03 and CD group: 0.29 ± 0.05), at 16 weeks (OD group: 0.37 ± 0.05 and CD group: 0.45 ± 0.07) and at 6 months (OD group: 0.58 ± 0.09 and CD group: 0.67 ± 0.08) indicated

that the crestal bone loss of the CD group was significantly more than the OD group. (Table 3)

Pearson's correlation analysis showed that there was a significant ($P < 0.05$) correlation between BMP-9 and implant stability in the OD group and between BMP-9 and crestal bone loss in the both the OD and CD group (Supplementary Table 1 and 2)

DISCUSSION

Over the past two decades, significant advancements in implantology have positioned endosseous implants as the preferred method for replacing missing teeth. Despite their widespread success, implant failures do occur, with rates reaching 8.16% and 4.93% in the

maxillary and mandibular arch respectively.²² Thiebot et al²³ noted that in most cases of

implant failures, the bone involved was classified as type III or IV. Ko et al²⁴ showed that the cortical bone thickness is least in the anterior and posterior maxilla, elucidating why

83% of the failures identified in Thiebot et al's study occurred in the maxilla, as also

observed in Kern et al's 5-year follow-up investigation.²⁵

The osseodensification technique, introduced by Huwaisin 2013¹⁰ utilizes specially

designed Densah burs to preserve and compact bone rather than removing it. This method promotes dense autografted bone formation around the implant, improving bone-implant contact and increasing insertion torque, thereby ensuring primary implant stability.¹¹

The demographic analysis confirmed no significant influence of age or gender between the groups, ensuring homogeneity. The central biological marker evaluated in this study was

bone morphogenetic protein-9. BMP-9 plays a pivotal role in osteoblast differentiation and bone tissue formation by activating the SMAD-dependent signalling pathway.²⁶ Studies by

Nieet al²⁷, Kawecki F²⁸, Haimov H²⁹ and others have enumerated the recent applications of BMP-9 ranging from alveolar bone healing to coatings on titanium implants.

BMP-9 levels increased progressively across all time points in both groups, with significantly higher levels observed in the OD group at the 8- and 16-week follow-ups.

These findings align with the known osteoinductive properties of BMP-9, which peak during the late stages of bone remodelling.³⁰ The lower levels observed in the OD group at 2 weeks may reflect the delayed inflammatory and resorptive phase due to early bone compaction.¹⁴ By 8 and 16 weeks, healing chambers created by OD likely served as active sites for osteogenesis, accounting for elevated BMP-9 expression.^{31,32}

Implant stability values both primary and secondary were significantly higher in the OD group compared to the CD group. These results which are in alignment with other animal and human studies,^{13,15,33,34, 35} reiterate that OD promotes a biomechanical environment conducive to immediate and long-term implant success.

Additionally, crestal bone loss was significantly lower in the OD group at all follow-ups.

This can be attributed to increased bone volume, higher bone density, and reduced micromovement due to superior primary stability. Although some previous studies have

reported no statistically significant difference in crestal bone loss between drilling

217 techniques, they were often limited by smaller sample sizes or restricted anatomical
218 locations.^{36, 37}

219 Importantly, this study found a positive linear correlation between BMP-9 levels and
220 implant stability, which was statistically significant in the OD group. This supports the
221 biological rationale that increased osteogenic activity, as signalled by BMP-9, contributes
222 to enhanced implant anchorage and stability over time.

223 The study also found a positive correlation between BMP-9 and crestal bone loss, which
224 may appear contradictory given BMP-9's role in bone formation. However, this
225 relationship likely reflects a reactive biological response rather than a direct causative one.
226 As bone remodelling intensifies (indicated by higher BMP-9 levels), localized bone
227 turnover may transiently elevate, especially in regions subject to biomechanical stress,
228 surgical trauma, or inflammatory response. Given the multifactorial nature of crestal bone
229 loss, which includes biological, biomechanical, and surgical influences, this correlation
230 should not be interpreted as BMP-9 causing bone loss. Instead, it suggests that elevated

3 231 BMP-9 may co-occur with active bone remodelling processes, some of which could
232 contribute to marginal bone alterations.

233 This study rejects the null hypothesis and confirms that bone compaction drilling enhances
32 234 peri-implant bone healing, increases implant stability, and reduces crestal bone loss,
235 supported by the levels of bone marker and clinical parameters.

6 236 Limitations of this study include the relatively small sample size, limited follow-up
237 duration, and the unicentric design, which may introduce Berksonian bias. Future research
238 with broader populations and longer observation periods is needed to validate these
239 findings further. Additionally, more detailed exploration of the temporal relationship
240 between BMP-9 expression and specific bone remodelling phases may provide deeper
241 insights into peri-implant healing dynamics.

CONCLUSION

Within the limitations of the present study, it can be concluded that the bone compaction drilling technique demonstrated a superior biological response compared to conventional drilling, as evidenced by elevated BMP-9 levels. Additionally, clinical parameters, including improved implant stability and reduced crestal bone loss, further support the efficacy of osseodensification as a favourable technique for implant placement.

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372 TABLES

373 Table 1. Comparison of levels of BMP-9 in OD and CD groups

BMP Levels	Group	N	Mean (pg/ml)	Std. Deviation	Std. Error Mean	p-value
2 wk	OD group	21	150.43	4.96	1.083	0.000 ^a
	CD group	21	177.67	8.24	1.798	
8 wk	OD group	21	274.19	12.16	2.653	0.100
	CD group	21	267.24	14.50	3.164	
16 wk	OD group	21	440.90	33.57	7.326	0.039 ^a
	CD group	21	423.62	15.58	3.400	

374 Values are presented as mean±standard deviation

375 CD- conventional drilling, OD- osseodensification drilling

376 ^a)Statistically significant difference

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Table 2. Comparison of implant stability in OD and CD groups

Implant Stability	Group	N	Mean (ISQ)	Std. Deviation	Std. Error Mean	p-value
Baseline	OD group	21	75.19	4.92	1.074	0.000 ^a
	CD group	21	64.90	6.80	1.484	
2 weeks	OD group	21	69.24	7.74	1.688	0.003 ^a
	CD group	21	61.43	8.26	1.802	
8 weeks	OD group	21	72.76	5.12	1.116	0.000 ^a
	CD group	21	62.95	8.40	1.834	
16 weeks	OD group	21	74.62	4.57	0.996	0.000 ^a
	CD group	21	64.57	4.17	0.909	

Values are presented as mean±standard deviation

CD- conventional drilling, OD- osseodensification drilling

^a)Statistically significant difference

Table 3. Comparison of crestal bone loss in OD and CD groups.

Crestal bone loss	Group	N	Mean (mm)	Std. Deviation	Std. Error Mean	p-value
2 weeks	OD group	21	0.071	0.011	0.0025	0.000 ^a
	CD group	21	0.104	0.026	0.0057	
8 weeks	OD group	21	0.244	0.028	0.0060	0.001 ^a
	CD group	21	0.287	0.049	0.0107	
16 weeks	OD group	21	0.367	0.055	0.0120	0.000a
	CD group	21	0.469	0.070	0.0152	
6months	OD group	21	0.584	0.094	0.0205	0.002 ^a
	CD group	21	0.675	0.080	0.0174	

Values are presented as mean±standard deviation

CD- conventional drilling, OD- osseodensification drilling

^a)Statistically significant difference

Figure 1: CONSORT flow diagram.

Figure 2. Preoperative Computed Tomography scan. (A) Osseodensification (OD) group.
(B) Conventional drilling (CD) group.

Figure 3. Osseodensification (OD) group. (A) Preoperative site. (B) Implant site
preparation using Densah bur drills. (C) Prepared osteotomy.

Figure 4. Conventional drilling (CD) group. (A) Preoperative site. (B) Prepared
osteotomy using conventional drilling technique.

Figure 5. Assessment of outcomes. (A) BMP-9 using PICF (peri-implant crevicular fluid).
(B) Implant stability using RFA (Resonance Frequency Analysis). (C) Crestal bone loss
using IOPA (intraoral periapical radiographs).

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