Survival of allogenic corticocancellous bone blocks

Horizontal alveolar process augmentation for implant placement

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Introduction

Loss of mastication or aesthetics that is to be restored by dental implants requires sufficient volume and quality of alveolar bone.1–3 It is important for the primary stability and the long term success of any dental implant treatment. The famous golden standard remains to be the autologous bone block4,5 as it is not involved in any immunological concerns, and contains vital cells. However, the vitality of the graft is highly dependent on the perioperative storage of the graft.6 It is generally accepted that class IV and V, according to the Cawood and Howell classification,7 need block augmentation before implant placement. The use of osseous allograft blocks for alveolar process augmentation is not very well documented in the literature.

Antonio Barone et al.8 published his study in 2009 and showed a good success with the osseous allogenic block, 24 blocks were used to augment the maxilla in 13 patients. Five blocks were used for ver-
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Figs. 6–8: Six months after bone augmentation. Out of these 24 blocks, two were a failure due to soft tissue exposure and thus completely removed. The remaining blocks were loaded with 38 implants at later stages and all implants achieved good primary stability. Contar et al. also published their paper in 2009. A total of 34 osseous allogenic blocks were used in 15 patients, one block had an early exposure. A number of 51 implants were placed into the grafted area with sufficient primary stability. None of the implants were lost within an observation period between 24 to 35 months. Carinci et al. published a paper in 2010 where implants placed in the resorbed maxilla, which had been grafted with osseous allogenic blocks and reported a survival rate of 98.3% over a mean follow-up of 26 months. This study showed results comparable to same areas augmented with autologous iliac crest bone. In 2015, Krasny et al. published an article in which 21 patients were treated with 26 grafts. In two grafts, there were complications with soft tissue, and one augmentation had to be redone because ofiatrogenic causes. After three to six months of healing, 33 implants were placed. Within an average observation time of 36 months (28–50), no implant had been lost. Araujo et al. published a systematic review on the same matter in 2013, in which a total of 253 osseous allogenic blocks were placed in 194 patients with a mean follow-up of twelve months (3–66). All studies showed good success from 95% to 100%.

Materials and methods

A total of 15 Patients were treated by the same surgeon in Godt Smil Odense from November 2013 to March 2015. Nine Patients were male, six were female. The youngest patient is 26 and oldest patient is 78-years-old. These 15 patients received 19 allogenic bone blocks to horizontally augment the atrophic alveolar process both in the mandible and in the maxilla prior to implant placement. All patients were in good general health, one was a smoker. All patients underwent periodontal therapy, if needed, before surgical intervention.

Surgical procedure

Premedication is 2,000 mg Imadrax (Amoxicillin), 1,000 mg Pinex (Paracetamol) and 400 mg Ibuzetin (Ibuprofen) 60 minutes before treatment. All Patients were instructed to rinse their mouth with 0.05% Chlorhexidine solution twice for one minute. The same strength of chlorhexidine solution was used for the perioral skin using a chlorhexidine-impregnated gaze. Local anaesthesia was administered as infiltration buccally and palatally/lingually (Xyloplyin® dental adrenalin 20 mg/ml + 12.5 microgram/ml lidocain hydrochlorid + adrenalin, DENTSPLY). Venous blood was sampled with a so-called butterfly (Vacuette® Greiner bio-one). The blood was collected in 10 ml
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tubes (A-PRF®+) and centrifuged according to Choukroun’s protocol.

A mucoperiosteal trapezoid flap was raised exposing the defect area. Neighbouring teeth were cleaned of any debris. Allogenic bone blocks weather, J bone or iliac crest, were customised chairside and fixed to the recipient site with osteosynthesis screws. The block was further adjusted after fixation has taken place, making sure there were no sharp edges and keeping the graft at least 1 mm away from any tooth surface. A PRF was placed over the block and healing was done by primary intention. Sutures were Resilon (Glycolon 5-0). Postoperative medication was Imadrax (Amoxicillin), 1,000 mg twice a day for three days, Ibumetin (Ibuprofen, 400 mg) in combina-

Results

None of the patients reported any problems during healing. All 19 bone blocks were integrated to the recipient site and bleeding at osteotomy after six months of healing gave a 100% result. There was great variation in resorption that was measured from the head of the screw: nine cases showed no resorption from the head of the screw, the remaining cases showed resorption ranging from 0.5 to 1.5 mm from the head of the osteosynthesis screw. Peripheral areas of the blocks, however, can exhibit a higher degree of resorption, but they were not measured in any way in this study.

Discussion

It must be noted that none of the grafts were used for vertical augmentation. This sample is a part of a bigger sample that involves more complex bone augmentation with use of more types of biomaterials such as spongious allogenic bone, prefabricated or not. In the bigger sample, there were failures that were not present when the augmentation was strictly horizontal and done by corticospongious blocks alone, indicating that this is a very predictable procedure when done like described in this case series.

The surgeon’s opinion is that resorption is related to width of the augmentation, thickness of the cortex on the corticospongious allogenic bone block after modification in the mouth, and area of the mouth, whereas the mandible showed more resorption. However, the surgeon’s opinion is per se of low evidence, and these topics should be investigated more in well-planned studies.

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Literature

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