Introduction

The dental rehabilitation of partially or totally edentulous patients with oral implants has become common practice with reliable long-term results. However, unfavourable local conditions of the alveolar ridge due to atrophy, periodontal disease, trauma sequel, malformation or neoplasia may cause insufficient bone volume, which may complicate the therapy of the masticatory function with dental implants. When alveolar ridges lack the appropriate bone volume, additional surgical reconstructive procedures are required.

The use of autologous bone grafts with dental implants was described originally by Brånemark et al. in 1975, and today is a well-accepted procedure in oral and maxillofacial rehabilitation. Insertion of an endosseous implant requires sufficient bone volume for complete bone coverage. Physiologically, an ideal bone grafting material should provide osteogenicity, osteoinductivity and osteoconductivity for new bone formation. Despite some recent advances in bone-substitute technology, autogenous bone grafts remain the “gold standard” in reconstructive surgeries because of their osteoinductive, osteoconductive and non-immunogenic properties. Guided bone regeneration (GBR) is an alternative technique to onlay grafting for localised alveolar ridge augmentation prior to dental implant placement. The clinical potential of membrane techniques for bone regeneration was recognised by Nyman et al. They demonstrated that membranes act as a physical barrier when applied over bone defects, preventing the ingrowth of competing, non-osteogenic cells into the membrane-protected space. Space provision, such as guided tissue regeneration, was shown to be effective in regenerating new bone on atrophied alveolar ridge, either vertically or horizontally, with the use of a membrane. Similar to onlay bone graft, which also serves as a space maintainer, GBR may incur similar complications that pertain to the use of onlay graft. Complications related to GBR may come from membrane exposure, miniscrew exposure and contamination. Exposed membranes or fixation screws often cause local inflammation with decreased bone formation.
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The significance of early membrane exposure on the regenerative outcome has been somewhat controversial in guided tissue regeneration and GBR procedures. Several studies have shown better responses when the membranes remained submerged than when they became exposed during healing. However, other studies failed to show such differences. It must be pointed out that patients affected by partial edentulism do not easily accept major surgical procedures that may imply hospitalisation or general anaesthesia. These disadvantages, together with the fact that dental implants do not demand a large amount of bone, lead to the growing use of autogenous block bone grafts from intraoral sources rather than from extraoral. The use of the mandible as a donor site is said to be less invasive, to save surgical and anaesthetic time and to be accomplished in the outpatient operatory. Harvesting of bone grafts from the retromolar region has been reported several times before. In the repair of localised alveolar defects, bone grafts from the retromolar region offer several benefits: a) the proximity of donor and recipient sites that reduces operative and anaesthetic time; b) conventional surgical access; and c) making them ideal for outpatient implant surgery. Minimal discomfort and decreased morbidity are also reported for this type of bone grafting. This technique can be combined with impacted third molar extractions.

The purpose of the present retrospective study was to evaluate the surgical success and to assess the rate of complications that arise from harvesting retromolar bone grafts in a group of partially edentulous patients prior to implant placement. We used a two-stage technique. In the first surgical stage, one or more cortico-cancellous bone blocks harvested from the retromolar region were fixed with osteosynthesis titanium screws to the recipient site as onlay grafts to achieve a horizontal and/or vertical augmentation of the ridge volume. In the second procedure, three to six months later, the screws were removed and implants were placed. The results regarding bone augmentation, donor and recipient site morbidity, bone graft stability and resorption prior to implant placement were recorded during the postoperative period and healing phase. Complications associated with this procedure mostly involve infection, incision line opening, nerve dysfunction, wound dehiscence, loss of portion of the bone graft, and graft mobilisation. A short review of the literature is presented in Table 2.

**Materials and methods**

A consecutive retrospective study was conducted on patients who underwent retromolar onlay bone grafting from January 2008 until January 2011. Files of 86 patients (77 males and 9 females) reporting 104 bone graft operations were reviewed. Patients ranged in age from 20 to 58 years (average 37.9 years). From the current study, patients were excluded if their data covered: a) grafting of bone defects caused by tumour resections, osteoradionecrosis and bisphosphonate-associated osteonecrosis, b) grafting of bone defects in syndrome patients with craniofacial involvement and with congenital malformations, such as cleft patients, c) grafting of extraction sockets and intraalveolar defects simultaneously with immediate implant placements and d) augmentations including the application of distraction osteogenesis. Medical history, cause of tooth loss and smoking status at the time of operation were recorded. All patients underwent primary clinical and radiographic examinations and were diagnosed as having an inadequate quantity of bone for implant placement.
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Table 3 shows the frequency of causes for the tooth loss. All the patients were informed in advance that bone reconstruction might be necessary prior to implant placement, since the need to augment the alveolar ridge can be evaluated correctly using panoramic radiographs only when there is vertical resorption of the ridge. Conventional radiographic examination provides little or no information about ridge thickness. Because of this, all the patients of the study underwent, except a conventional panoramic radiograph of the jaws, CT scans reformatted with Dentscan software. In disagreement with other authors, we believe that reformatted CT images always provide a precise treatment guide when the decision to graft or not to graft has to be made in critical cases. A total of 104 alveolar segments were treated: 22 procedures involved the maxilla and 82 the mandible. In six of the augmented areas on the maxilla posterior, a Sinus elevation was also performed. Fifteen patients included in the study were treated in separate procedures for augmentation of different alveolar sites. Each augmented site was studied. 67 procedures were carried out under local anaesthesia and 37 under general anaesthesia. All sites were treated in a similar fashion. The number of bone blocks, donor sites and number of implants inserted in each augmented site were also recorded. The choice of donor site, left or right, was determined preoperatively, based on defect morphology and recipient site location. When the augmentation was planned in the posterior mandible, a single surgical field was needed, thus reducing patient discomfort. The recipient site was healed completely prior to graft surgery. The proposed recipient site for the graft was exposed prior to graft harvest in all cases. In this manner, the dimensions and morphology of the bony defect were measured, and minimal time elapsed between graft harvest and placement (Figs. 1 et 2). Figure 3 presents the preoperative situation. The recipient site was prepared and recontoured with the Safescraper (C.G.M. S.p.A., Divisione Medica META, Italy) by pushing the end of the devise toward the bone surface and simultaneously pulling the devise backward. Collection of 2–3 ml of bone was feasible with a mean surgical time of five minutes for harvesting (Fig. 4). The collected bone was preserved in a sterile environment until grafting. To access the ramus area, the concavity formed by the border between the ascending ramus and the external oblique ridge was identified and used as a starting point for the mucosal incision.

The incision was made medially to the external oblique ridge and extended mesially toward the buccal aspect of the second molar. Care was taken to ensure that the incision was not extended too far lingually, preventing damage to structures on the lingual aspect of the mandible. A mucoperiosteal flap was elevated superiorly, exposing the lateral aspect of the ramus. The osteotomy was carried out with an osteotomy kit for Piezosurgery® (Mectron, Deutschland Vertriebs GmbH) and was started anteriorly to the coronoid process at the

_Surgical protocol_

Stage 1 surgery

The bone harvesting procedure was performed using a standardised surgical technique. The anaesthesia of all patients was carried out with Ultracain™ D-S (Hoechst Marion Roussel Deutschland, Frankfurt, Germany) containing 1:200,000 epinephrine at the donor and recipient sites. A single shot of 2,2 g amoxicillin and clavulanic acid (Augmentan®, GlaxoSmithKline Consumer Healthcare GmbH & Co. KG) or, if penicillin allergic, 600 mg Clindamycin (Clinda-saar®, MIP Pharma GmbH) as well as 250 mg Prednisolon (Solu-Decortin®, Merck Pharma GmbH) was administered intravenous to patients a few minutes prior to surgery.

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Postoperatively, patients were instructed to rinse their mouth with chlorhexidine 0.2% for two to three weeks twice a day. After this period, the sutures were removed. Removable provisional prostheses were adjusted generously. Patients were instructed to use their prostheses for cosmetic appearance and for eating rather than function for the whole period of healing, i.e. three months. At that time, patients were scheduled for implant surgery. No antibiotic therapy was continued after surgery and patients were instructed to use non-steroidal anti-inflammatory drugs (Ibuprofen®, Docpharm® Arzneimittelvertrieb GmbH & Co. KGaA) only if pain was present.

Stage 2 surgery

After a healing period varying between three to six months after the grafting procedure, clinical and radiographic evaluations were performed and implants were placed in a routine fashion using the special program for guided implant surgery CoDiagnostiX® (IVS Solutions AG). All patients underwent CT scan before the implant placement to assess new bone formation and plan the accuracy of the implant position. A crestal incision and subperiosteal dissection of the alveolus were performed and the fixation screws were removed. Implant site preparation was performed with guidance from the laboratory-made splint and the implants were positioned (Figs. 12–18). We used bone-level type titanium implants. The most common in use were Straumann® Bone Level implants (Institut Straumann AG, Switzerland), followed by Astra® (Astra Tech Inc.) and Camlog® (CAMLOG Vertriebs GmbH, Germany). In total, 155 implants were positioned, 39 in the maxilla and 116 in the mandible. Three to six months later, the prosthetic work was started.

Clinical appointments were performed after surgery to evaluate any complication at the donor and recipient site, such as dehiscence, infection, swelling, sensory disturbances or haemorrhage. Graft loss and graft removal were defined as failure; swelling, wound dehiscence, incision line opening, infection with pus or temporary paresthesia were defined as complications. Buccal nerve damage from incision along the external oblique ridge as well as the damage of inferior alveolar nerve is possible. Graft incorporation was evaluated following the removal of the fixation screws. Statistical analysis included descriptive statistics using IBM SPSS Software for Windows. The significance level of p ≤ 0.05 for all statistical tests was definite. The residents of our department operated and followed upon all patients, and the results were analysed in percentage and presented in tables and diagrams.

Disclosure

The authors do not have any financial interests, either directly or indirectly, in the products listed in the study.

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