Surgical management of peri-implantitis

Specialist Periodontist, Dr Jeremy Vo, explains how he has embraced AIR-FLOW® technology in the management of Peri-Implantitis

By Dr Jeremy Vo

Surgical intervention is often required in the treatment of advanced peri-implantitis lesions. Peri-implantitis is defined as an inflammatory process around an implant, with soft tissue inflammation and loss of supporting marginal bone.

The aim of surgical therapy is to allow access for the decontamination of implant surfaces which have been exposed to oral biofilms. Several approaches for implant decontamination have been described and can be broadly categorised to include mechanical, chemical and laser instruments.

Mechanical removal of hard and soft deposits can be achieved with rubber cups, curettes, and/or ultrasonic devices. Curettes of different materials have been manufactured, specifically for the debridement of implant surfaces. These materials include steel, titanium, carbons fibre, Teflon and plastic. Ultrasonic devices with polyetheretherketone (PEEK) coated tips are also specific for implant surfaces. A more aggressive approach has been proposed which involves intentional removal of the implant threads. This is known as ‘implantoplasty’ and the aim is to produce a polished, smooth collar which better supports oral hygiene compared with the original rough surface of the implant.

Chemical decontamination is aimed at disinfecting the implant surface by direct application during surgery. Following elevation of the soft tissues, the implant surface can be rinsed with several different substances including chlorhexidine, sodium chloride, hydrogen peroxide and citric acid. Unfortunately, no chemical agent has shown superior results when compared with others.

Laser decontamination - including the use of Er:YAG and CO₂ lasers - have also been utilised during surgery in an attempt to improve clinical outcomes. The evidence is, however, weak and has not shown significant improvement when compared with conventional mechanical therapy.

There are 3 main approaches for surgical intervention including:

**Access surgery**

The primary aim of access surgery is to decontaminate the implant surface. Commonly, intrasulcular incisions will allow the conservation of the soft tissues around the implant once the mucoperiosteal flaps are elevated. Inflamed peri-implant tissues are degranulated and the implant surface is decontaminated. A clinical study with 5 years follow up reported complete resolution of advanced peri-implantitis lesions in 42% of implants and 63% survival of implants.

**Resective surgery**

This surgical technique allows implant decontamination to take place, but rather than conserving soft tissues, a reverse bevelled incision combined with osteoplasty reduces the pocket depths around the implant. As a result, the neck of the implant is usually left exposed to the oral cavity and therefore, this technique is only suggested for implant defects in non-aesthetic areas. The 2-year outcome of resective peri-implantitis surgery found complete resolution of clinical signs of disease in almost 60% of implants.

**Regenerative surgery**

Regenerative surgery is aimed at improving hard tissue integration around the implant (re-osseointegration) as well as remineralising recession of the peri-implant mucosa. Following mucoperiosteal flap elevation, the implant surface is decontaminated and the intrabony defect is degranulated. Various approaches to bone grafting have been described. Bone substitute materials such as Bio-Oss® (Geistlich Biomaterials) can be used to fill the intrabony defect which is then covered with a resorbable or nonresorbable membrane. A 4-year clinical study found significant reductions in probing pocket depth and radiographic defect fill with a regenerative technique involving Bio-Oss® and Bio-Gide®.

The clinical approach

The development of biofilms on the implant surface plays a significant role in the initiation and progression of peri-implant disease. The bacterial microflora is composed predominantly of Gram-negative anaerobes and is similar to microflora found around teeth with severe periodontitis. Unfortunately, in the management of peri-implantitis, no definitive gold standard has been identified for implant decontamination. Implant surface roughness and irregularities can enhance bacterial attachment and prevent adequate instrumentation. The tips of the curettes are often too large to reach the deeper parts of the thread.

Recently, a powered air-abrasive system utilising Erythritol (a sugar substitute) has been proposed as an effective method of biofilm removal from the implant surface that is safe on hard and soft tissues (EMS AIR-FLOW® EL-308/A; Electro Medical Systems (EMS)).” The abrasiveness of Erythritol is low and it does not cause extensive damage to the surface topography of the implant compared with the use of conventional steel curettes or ultrasonics. Furthermore, in vitro data suggests that it has an antimicrobial effect.

Once the implant surface has been decontaminated, the morphology of the bony defect may help determine the most suitable surgical approach. Generally, if the defect is circumferential with intact bony walls, or has an intrabony component, the use of a regenerative approach will provide improved clinical outcomes. On the other hand, if the defect is supra-bony or the implant presents with some degree of buccal dehiscence, an apically repositioned flap would be indicated in these non-aesthetic areas. **Figure 1. Endographic assessment at baseline.**

**Figure 2a-c. Pre-operative clinical photographs.**

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Surgical therapy is the cornerstone of peri-implantitis therapy.
The following case study illustrates a protocol that was used to treat advanced peri-implantitis. The case was treated successfully with a 6-month follow-up. Success was defined by a reduction in probing pocket depths (PPIs), along with a reduction of soft tissue redness and bleeding on probing.

**Case Study: Regenerative approach for treatment of peri-implantitis**

A 70-year-old female was referred for advanced peri-implantitis in the mandible. She presented complaining of pain and she also noted discharge from one of the anterior implants. Her medical history was non-contributory and she was a non-smoker.

Clinical examination revealed 5 implants in the mandible supporting a fixed full arch reconstruction. Probing pocket depths were of the order of 8-9 mm around 3 of the anterior implants. The distal implants had normal probing depths. CBCT imaging revealed an intrabony component of 6.5 mm for the implant in the 41 position, 4.2 mm at the 41 implant and 3.3 at the 33 implant.

A preoperative phase was carried out, including assessment of oral hygiene and non-surgical implant decontamination in 1 session. After 6 weeks, the patient underwent surgical treatment. This comprised of full thickness mucoperiosteal flaps being raised and the chronic inflammatory tissue removed from the defects around the 3 implants with the use of tetrol curettes. The implant surface was then decontaminated using EMS AIR-FLOW® technology with very fine erythritol powder (EMS AIR-FLOW® PLUS Powder). The implants were also irrigated and cleansed with saline-soaked cotton swabs.

A crater shaped defect was present around all the implants at the buccal and lingual surfaces, however the implants had a dehiscence on the buccal aspect. The craters were filled with BioOss granules (Geistlich) and Bio-Gide was placed to cover the defects. Lastly, the flaps were repositioned and secured with mattress and sling sutures. Systemic antibiotics were administered postoperatively. The full arch prosthesis was reissued at the completion of surgery.

Clinical parameters and radiographic examinations were performed at 3 and 6 months. At both intervals, there was resolution of the clinical parameters for all 3 implants, including plaque index, bleeding on probing and probing pocket depth. At these visits, non-surgical maintenance was carried out, including oral hygiene reinforcement and removal of biofilm via EMS AIR-FLOW® technology and EMS AIR-FLOW® PLUS Powder.

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