This is an author produced version of *Rehabilitation of oncology patients with hard palate defects Part 2: Principles of obturator design*.

White Rose Research Online URL for this paper:
http://eprints.whiterose.ac.uk/90417/

**Article:**
Restorative Dentistry

Rehabilitation of Oncology Patients with Hard Palate Defects.

Part 2: Principles of Obturator Design

Rahat Ali*,
BSc(Hons), BDS, MSc, MFGDP (UK), MFDS RCS, PGC (HE),
Specialist Registrar in Restorative Dentistry,

Asmaa Altaie*,
BDS, MSc,
Clinical Teaching Fellow/PhD student in Restorative Dentistry,

Brian Nattress*,
BChD, FDS RCS, MRD RCS, PhD,
Senior Lecturer/Honorary Consultant in Restorative Dentistry,

*Leeds Dental Institute, Clarendon Way, Leeds, LS2 9LU
Rehabilitation of Oncology Patients with Hard Palate Defects

Part 2: Principles of Obturator Design

Abstract
The first part of this series on the conventional rehabilitation of oncology patients with hard palate defects discussed the dental challenges posed by oncology patients and the surgical/restorative planning interface for conventional dental rehabilitation. This article will describe Aramany’s classification of hard palate defects, Brown’s classification of palatal defects and focus on the basic principles of obturator design which need to be appreciated when prosthetically rehabilitating a patient with a hard palate defect.

Clinical relevance statement
A good understanding of basic removable prosthodontic theory relating to denture design, dental materials science and head and neck anatomy is an absolute prerequisite when designing an obturator for a patient.

Learning Objective
This article will describe: (a) how hard palate defects can be classified and (b) the basic principles of obturator design which need to be appreciated when rehabilitating a patient with a maxillectomy defect.
Introduction

Resecting a palatal tumour will result in a surgical defect. The resulting defect can be managed by primary surgical closure, reconstruction with a surgical flap or prosthetic obturation. The decision as to which modality of treatment is chosen should be made within an oncology multidisciplinary team with input from surgeons, clinical oncologists, radiologists and restorative dentists (1). With modern surgical techniques, it is encouraging to see that more surgeons are offering reconstructions for head and neck oncology patients post resection. The use of microvascular flaps to reconstruct head and neck cancer patients has increased over the years, with deep circumflex iliac artery (DCIA) and radial forearm free grafts (RFF) being commonly used to reconstruct maxillary defects (2). It is encouraging to see that the use of dental implants to rehabilitate oral cancer patients has also increased since 1995 (2). There are however a cohort of patients who have not been reconstructed and may be unsuitable for or may not be interested in implant based rehabilitation. Such patients with acquired hard palate defects will need conventional dental rehabilitation with an obturator. For these patients, the obturator will help to separate the oral and nasal cavities, help to restore normal speech and swallowing and provide support for the lip and cheek.

This article is the second paper in a series of three. It will focus on the conventional, non-implant based prosthetic obturation of oncology patients with surgically acquired hard palate defects. It will describe Aramany’s classification of hard palate defects, Brown’s Classification of palatal defects and focus on the basic principles of obturator design.
Principles of Obturator Design

Defects have been successfully managed with obturators (Latin: “obturare” meaning “to block up”) for many years. Ambroise Pare was probably the first individual to artificially close a palatal defect in the 1500’s (3). The early obturators were used primarily to close congenital defects as opposed to acquired defects. The pioneering works of these prosthodontists set the foundation for the current generation of maxillofacial prosthodontists.

Aramany classified partially dentate maxillectomy patients into six groups (4). This classification is based on the relationship of the defect to the remaining abutment teeth and represents a systematic means of describing maxillectomy patients and potentially framework designs for an obturator (Figure 1). Although it is a very useful classification, it only considers the defect in the horizontal plane and gives no indication of the extent of the defect in the vertical axis.

Brown et al suggested a modified classification of maxillary defects (5). It describes the defect in both the vertical and horizontal planes for both dentate and edentulous maxillectomy patients. The vertical classification is as follows:

Class 1: Maxillectomy without an oral/antral communication.

Class 2: Low Level Maxillectomy not involving the orbital floor or contents.

Class 3: High Level Maxillectomy involving the orbital contents.

Class 4: Radical Maxillectomy involving orbital exenteration.

Classes 2, 3 and 4 are further classified in the horizontal plane as either

a: Unilateral Alveolar Maxillectomy

b: Bilateral Alveolar Maxillectomy

c: Total Alveolar Maxillary Resection.
For any removable prosthesis to be successful, it must be retentive, stable and have good support. These prosthetic principles are especially applicable to an obturator.

Support can be defined as the resistance to movement of prosthesis towards the underlying tissue. For an obturator, support can be derived from the residual hard palate, alveolar ridge, the remaining teeth and from within the defect itself. According to the recommendations of Desjardins (6) a flat vaulted hard palate and a broad ridge are better adapted to provide support than a high tapering palate and a narrow tapering alveolar ridge. To maximise support for the patient’s new denture, maximum palatal coverage of the hard palate should be employed (Figure 2a and 2b). If the remaining teeth are to be used to provide support, they should be loaded with rest seats and used in sufficient numbers to minimise movement of the prosthesis towards the underlying mucosa. They should be located as close to the defect as possible and not place excessive torque on the teeth (6) (Figure 2b).

Given the demands that will be placed on the abutment teeth to provide adequate support, they should be periodontally sound.

The presence of the defect allows structures within it to be used to obtain support for the prosthesis. This support can be achieved by contact of the prosthesis with any anatomical structure that provides a firm base, and includes the floor of the orbit, pterygoid plate and nasal septum (6). Using the orbital floor increases the vertical height of the appliance and can create difficulties with insertion and removal. Care must also be taken to avoid pressure on the sensitive turbinate nasal bones. The nasal septum is also not ideal to provide support as it is partly cartilage and has very little surface area that can be utilised. The most common place to use for support (within the defect) is the pterygoid plate. Positive contact on to the plate can tripodize and stabilise the device, and minimise its rotation into the defect.

Retention of a prosthesis refers is the resistance to vertical displacement. As with support, retention can be gained from the residual maxillary structure, or anatomical structures within the defect. Implants within the residual alveolar ridge, the surgical defect or the zygomatic bones can also be used to help retain the obturator. Any remaining teeth can be an asset for retaining the prosthesis as they can be engaged (with clasps). Movement of the obturator may generate tremendous torquing forces, which in turn will be dissipated by the retaining abutment teeth. It is recommended that to maximise direct retention, clasps are located as close to and as far from the
defect, with at least one clasp in between these two extreme points (6). The preparation of guide planes to produce one single path of insertion can also aid in retention (7). If the abutment teeth have good bony support and no signs of active periodontitis, a cast clasp and rest seat can be prescribed to augment retention and support for the prosthesis (Figure 2b). For Aramany Class IV defects, retention can be maximised by placing retentive clasp arms on the remaining premolar and molar teeth, if they are periodontally sound and have adequate root support (8). For Class Aramany II defects, clasps can be prescribed for the anterior and posterior teeth (7). If the teeth have a guarded long term prognosis, wrought gold or stainless steel clasps should be used to minimise torquing forces on the abutments (Figure 2a).

Retention can also be obtained from the defect itself. Naturally occurring undercuts within the residual hard/soft palate can be engaged by the prosthesis to augment retention. Extension of the obturator into the area superior to the lateral scar band, created by the use of a split skin graft at the time of surgery, provides an ideal undercut to aid retention of the appliance (Figure 3). Extension of the obturator onto the nasopharyngeal surface of the soft palate will also improve retention (9). Therefore at the surgical planning stage, it is imperative that the surgeon tries to preserve as much palatal tissue as possible, without compromising resection of the tumour. Patient satisfaction with an obturator will significantly decrease if more than one quarter of the hard palate or one third of the soft palate is resected (10).

So far, the concept of retention has been discussed with respect to vertical displacement of the appliance. An obturator is unique in that it can also rotate out of the defect due to the force of gravity and function. It is difficult to eliminate this and patients may use adhesive to prevent the obturator from dislodging. To minimise this, indirect retention can be incorporated into prosthesis by extending the connector beyond the axis of rotation onto any residual teeth that are present, assuming that they are periodontally sound. For Class II Aramany defects, this could involve extending the metal framework/indirect retainer onto to the opposite side of the defect and onto to the remaining canine and premolar teeth (8).

Stability refers to the resistance to dislodgement by functional forces. If an obturator is designed to gain optimal support and retention from the remaining tissues (eg. maximal lateral extension into the defect/extension of the bracing components onto the abutment teeth) the need for stability will also be appeased. In particular, Sharma
and Beumer discussed the need to maximally extend the bung up the lateral wall of the defect to improve retention, stability and provide support for the lip and cheek (11). Desjardins (1978) has further recommended that the posterior part of the obturator is lower (in height) than the anterior portion. This design feature will encourage downward movement of any fluid into the nasopharynx (6) 

There are essentially two principle styles of obturator: (a) The fully extended hollow bulb (usually rigid) and (b) the open top which may be designed with either a rigid or flexible rim (12). The use of either design is dictated by the requirements of individual cases.

Hollow bulbs obturators improve speech resonance and decrease the weight of the obturator. This helps to minimise any downward displacement of the prosthesis due to gravity or function. It was reported that 29% of obturator patients experience significant oral/nasal leakage (13). Therefore the bulb must be carefully manufactured to produce an adequate seal and partition between the oral and nasal cavities. Hollow bulbs can also gain support from structures within the defect. Patients may have problems with insertion, and therefore a two-part design may have to be considered to overcome this problem, especially if the patient has significant trismus (Figure 4a and 4b). If a two part obturator is used, magnets can be used to unify the segments into one prosthesis. Magnets have been used to stabilize maxillofacial prostheses for over half a century (14) and can be useful adjuncts for the Restorative Dentist. Typically a magnet is embedded in the bung which the patient locates first into the defect. A separate magnet is on the denture portion of the appliance, which the patient inserts second. Such a design allows each segment of the obturator to have a separate path of insertion. Given that the magnets are self-seeking, it is easy for the patient to unite the denture with the bung.
A method of producing a hollow bulb obturator is shown in (Figure 5). The technician makes the hollow bulb in wax, before packing, flasking and processing it in acrylic resin. The hollow bulb is subsequently filled with Plaster of Paris and pumice. A lid is then waxed over the pumice/plaster filled bulb before processing the obturator for a second cycle. Once complete, the pumice/plaster is carefully removed from the bulb via a small cavity in the superior aspect of the bung. This opening is then restored with self-cured acrylic resin.

Using a flexible bung (Figure 4a and 4b) can atraumatically engage undercuts within the defect more effectively than hard acrylic. They are typically made out of resilient lining material. Retention of the silicone bung is excellent when adequate tissue undercuts remain around the soft palate and on the buccal aspect of the defect (9). However silicone rubbers flex under load. Consequently the seal on the surgical site may be compromised when the patient masticates on the resected area. As a result, patients often chew on the unresected site. There are further problems with flexible bungs made of resilient lining material. They will need regular replacement and are difficult to adjust (15). Adjustment of the silicone bung can produce a rough surface which is difficult to clean and may encourage fungal growth (9). Therefore where possible, the authors favour the use of hard acrylic. Open top designs tend to be smaller and lighter. The bulb can be carefully moulded to render the appliance more retentive (Figure 6a and 6b). However, they clearly will accumulate unpleasant nasal secretions and will need to be cleaned regularly.
Discussion

Given the variability in the different types of patient/defect/residual denture bearing anatomy which the clinician may encounter, it is not possible to discuss specifics of obturator design. Therefore the authors have attempted to discuss the general principles which underpin appliance design in this paper. The works of Desjardins (6) and Aramany (4) have been heavily quoted whilst describing the design features of an obturator. Their recommendations are not based on the results of any clinical, prospective, randomised control trials. Rather they are based on their experiences having provided numerous prostheses for oncology patients. Readers are encouraged to study their published works to gain further insight in this complex and challenging field. Furthermore a good understanding of basic prosthodontic theory relating to denture design, dental materials science and head and neck anatomy is an absolute prerequisite for any one managing head and neck resection patients. These fundamental principles need to be understood when designing an obturator. The next and final part of this series will describe the clinical stages involved in the construction of an acrylic, one part, hollow bulb obturator.
Acknowledgements

The authors acknowledge and thank the Medical Illustration Department at the Leeds Dental Institute for taking the photos shown in this manuscript. The authors also acknowledge the excellent technical support provided by all the Prosthetics Technicians at the Leeds Dental Institute, especially Paul Bartlett, Terry Haite, Richard Jones, Steve Scupham and Deborah Cook. The authors also thank the comments and contributions to this manuscript made by Richard Hardcastle, Chief Anaplastologist at St. Luke’s Hospital, Bradford.
References

Figures and figure legends

Figure 1: Aramany’s Classification of maxillectomy defects. Class I = midline resection; Class II = unilateral resection; Class III = central resection; Class IV = bilateral anterior-posterior resection; Class V = posterior resection & Class VI = anterior resection.

Figure 2 (a): A palatal defect that was obturated with an acrylic obturator. To minimise any torquing forces on the UR6, a flexible wrought gold clasp was used.
Figure 2 (b): A patient with an extensive defect where a swing-long prosthesis was provided. Note the use of multiple cast clasps and rests, the extension of the metal connector onto the UL3 for support/indirect retention and maximal palatal coverage.

Figure 3: A patient where a split thickness skin graft was prescribed for the lateral aspect of the resection. It formed an undercut that was recorded on the master impression and engaged by the obturator.

Figure 4 (a): A two-part, rigid hollow bulb obturator connected by magnets.
Figure 4 (b): A patient who had their entire hard and soft palate resected. She was rehabilitated with a two-part obturator connected by magnets. The bung was made of resilient soft lining material.

Figure 5: Production of a Hollow Bulb for an Oburator.
Figure 6 (a): A two-part, acrylic obturator with a flexible open lid bung.

Figure 6 (b): A one-part, acrylic obturator with a rigid open lid bung.