Clinical and histopathological analysis of zirconia intramucosal inserts used for improving upper dentures retention.

Running title: Safety of intramucosal zirconia inserts.

Fábio Gonçalves¹, Eliane Pedra Dias², Tania Mari Cestary³, Rumio Taga³, Raquel Virgínia Zanetti¹, Artemio Zanetti¹, José Mauro Granjeiro⁴,*

¹ Centro de Estudos de Implantes Osseointegrados da Universidade Cidade de São Paulo (CEIO-UNICID)
² Department of Pathology, Medical School, Fluminense Federal University, Niteroi, Brazil
³ Department of Biological Sciences, University of Sao Paulo, Bauru, Brazil;
⁴ Department of Cell and Molecular Biology, Fluminense Federal University, Niteroi, Brazil

Niterói, February, 23 2008

Corresponding author

* José Mauro Granjeiro, DSc (jmgranjeiro@vm.uff.br)
Cell and Molecular Biology Department, Biology Institute
Fluminense Federal University - UFF
Phone: 55 21 2629 2324 Fax: 55 21 3701 1617
Outeiro de São João Baptista, s/n
Campus do Valonguinho, Centro/Niterói – RJ
ZIP 24.020-150
Brazil
Abstract

Intramucosal inserts made of zirconia ceramic have been used to rehabilitation of edentulous patients. This study aimed to follow up on patients with complete dentures having intramucosal zirconia inserts and to perform histological analysis of biopsies of the epithelium around the inserts. Twelve subjects of both sexes, 31-66 years-old, received complete denture treatment having the intramucosal inserts in place. Detailed clinical, prosthetic and surgical procedures were described. Clinical exams were done on postoperative days 3, 7, 15, 120 and 360. After one year following the denture insertion, the subjects received local anesthesia and the tissues surrounding the insertion sites in the alveolar ridge were removed using a scalpel. Biopsies were fixed for up to 48 hours in 10% phosphate-buffered formaldehyde and sections of 5 µm thick were cut and stained with hematoxylin and eosin and Gomori trichrome. Immunohistochemistry was used to identify endothelium (anti-CD3) and T lymphocytes (anti-CD31). Removing and reinserting the denture was painful until day 15, but all patients reported a marked increase in the retention and stability of the complete denture with little or no discomfort after 30 days. The histopathologic analysis showed that zirconia inserts were well tolerated by the oral mucosa, presenting collagen fibers in the tissue around the insert, maintaining mild inflammatory response, and allowing reepithelialization, expressed by parakeratosis, epithelial hyperplasia and presenting granular layer. In conclusion, intramucosal zirconia inserts did not affect the health of oral mucosa and provide adequate retention and stability of the complete denture and comfort to the patients.

Key words: intramucosal insert, total prosthesis, histopathology, biocompatibility, dental rehabilitation
Introduction

The rehabilitation of edentulous patients continues to be a major challenge to dentistry. A number of techniques and materials have been investigated in order to improve retention and stability of the complete denture. The first study describing the use of intramucosal metal inserts associated to the complete upper denture was published in the 1940s (1), in which 30 patients were followed for 10 years. During the 1950s, adjustments in the insert size and the use of non-reactive metal inserts of two different sizes in the same prosthesis were proposed to increase stability and retention (2). Later in the 1970s and 1980s, the focus was on the design of the retention pin with the purpose of creating retention devices, such as precision attachments (3), that would reduce the need for relining the denture and prevent dysplastic changes in the epithelium around the inserts (4).

After the 1990s, there was a significant decrease in the number of reports on the use of intramucosal inserts, probably due to the advent of osseointegrated implants and their use in dentistry. However, a fundamental requirement for successful osseointegrated implants is the availability of sufficient quantity and quality of bone for dental implant placement (5). Bone-grafting procedures may be necessary for the reconstruction of the atrophic maxillae, increasing the complexity and morbidity of the procedure. However, oral rehabilitation using implants is not appropriate for a large number of patients due to restrictions such as local and systemic conditions that are contraindications to the use of implants, high-cost procedures and patients’ refusal.

Recently, intramucosal inserts made of zirconia (ZrO$_2$ or zirconium oxide) ceramic have become available (6). This ceramic material has high strength (7), is inert and biocompatible (8, 9) and being use as abutments (10). The zirconia inserts have a highly polished surface that minimizes the formation and accumulation of biofilms (11). In spite of the well-known biocompatibility of zirconia, no clinical or histological studies have been performed to evaluate the effect of the use of intramucosal inserts made of zirconia ceramic on the epithelium in contact with such devices.

The objective of this study is to follow up on patients with complete dentures having intramucosal zirconia inserts, as well as to perform histological analysis of biopsies of the epithelium around the inserts.

Material and Methods

Zirconia intramucosal inserts

The intramucosal inserts (MUCHOR® (Dyna Dental Engineering B.V.) used were made of zirconium oxide (ZrO$_2$, Y$_2$O$_3$ Ytrium stabilized) very well polished. Figure 1 shows the insert composed of two parts: one with elliptical shape, which remains in
contact with the mucosa (3.2 mm diameter and 5.2 mm height) and the second one, which will be retained into the denture acrylic (4.0 mm diameter).

Sample

Twelve subjects of both sexes, aged between 31 and 66 years, received complete denture treatment in the Center for the Study of Osseointegrated Implants at the University City of São Paulo (UNICID/CEIO). This research followed the principles of the Declaration of Helsinki and the free and informed consent was obtained from all patients after a clear and complete explanation of the study was provided.

In this study, eligibility criteria included patients who were experiencing problems due to lack of retention of complete dentures, alveolar bone loss and maxillary atrophy with loose mucosa and pneumatized maxillary sinuses. Patients were excluded if they had chronic systemic disorders such as uncontrolled diabetes and hypertension, were not interested in the procedure and refused to cooperate, or did not fulfill the clinical and radiographic criteria required for the placement of osseointegrated implants. Eligibility and exclusion criteria were verified by anamnesis, clinical examinations and panoramic x-ray. Table 1 summarizes the characteristics of the subjects selected for the study.

Initially, the subjects underwent surgical procedures to prepare the mouth for complete denture, with the removal of anatomical or even pathological elements that could contribute to the lack of stability of the complete upper denture. The hyperplastic mucosa was removed by cryosurgery, followed by the use of tissue conditioners to promote tissue repair.

In the construction of complete dentures, upper and lower dental impressions were taken, casts were prepared and mounted in the articulator using the Zanetti technique (12), intermaxillary registration was performed, the denture teeth best suited for each patient were selected and the dentures were inserted. Adjustments for overextension and relieve of pressure areas, as well as occlusal adjustments were made during the first 15 days following insertion of the denture.

After being adapted and balanced, each complete denture was relined with zinc oxide-eugenol cement and sent to the dental laboratory. Also, an acrylic guide plate was made to mark the positions for the denture inserts.

A dental stone cast was made from the relined denture, and an adequate number of intramucosal inserts, required to meet the needs of each patient, were placed on it using a surveyor to assure a proper insertion path, thus further facilitating the construction of the denture. An acrylic surgical guide was made from the stone cast having the intramucosal inserts in place in order to assure a uniform and correct
positioning of the preparation sites in the mucosa. The denture was then invested, and after polymerization of the acrylic resin, it was removed from the mold and polished.

With the surgical guide in place, the injection sites of local anesthesia were identified. The anesthesia was strictly limited to the locations for the insert preparation sites in the mucosa, thus avoiding excess of anesthesia or anesthesia of the posterior vestibule, which could result in local or peripheral edema that would interfere with the full insertion of the denture. Preparation sites 5 mm deep were made using a special drill with low speed of rotation with water-cooling, provided by the manufacturer of the inserts. After the preparation sites were made and adjusted, the denture was inserted and checked for fitness, assuring full insertion. A small amount of dental adhesive paste was applied to the palatal area of the denture to increase stability and initial retention, contributing to tissue repair of the mucosa around the intramucosal inserts.

All patients were instructed not to remove the denture for 72 hours, the time taken for the initial healing of the mucosa around the intramucosal inserts. After the third day, the patients could remove the denture only for cleaning and then return it to place.

Clinical Evaluation
The subjects returned to the study center (CEIO-UNICID) on postoperative days 3, 7, 15, 120 and 360. At these follow-up visits, the clinical aspect of the mucosa in contact with the insert was subjectively evaluated, as well as the retention and the stability of the denture, subject’s comfort level when eating, removing and reinserting the prosthesis, and also changes in speech.

Biopsy and histopathological evaluation
After a minimal period of one year following the denture insertion, the subjects received local anesthesia and the tissues surrounding the insertion sites in the alveolar ridge were removed using a scalpel (n° 10 blade). All biopsies were fixed for up to 48 hours in 10% phosphate-buffered formaldehyde (pH 7.2) and sent to the Laboratory of Pathology of the Antônio Pedro University Hospital, Fluminense Federal University, RJ, Brazil. After macroscopic analysis, the biopsies were dehydrated in increasing concentrations of ethanol, cleared in xylol, and immersed in liquid paraffin at 60° C. After obtaining the paraffin blocks, sections 5 mm thick were cut and stained with hematoxylin and eosin (H&E) and Gomori trichrome. Histological evaluation was conducted by an experienced pathologist. The endothelium and T lymphocytes were identified by immunohistochemistry using anti-CD3 and anti-CD31 antibodies, respectively, as described before (13). Soon after, sections 6 µm thick were cut from the paraffin-embedded tissues and mounted on silane-coated glass slides (Super Frost
Plus; Erviegas, São Paulo, Brazil). The immunohistochemical characterization of cells was performed using the avidin–biotin–peroxidase complex (ABC) method. Slides were deparaffinized in xylol and ethanol, and incubated in 3% hydrogen peroxide diluted in phosphate-buffered saline (PBS) (pH 7.2) for 40 min at ambient temperature. Then, the slides were immersed in citrate buffer (pH 6.0) (P4809; Sigma, Saint Louis, MO) for 20 min at 950 °C for antigen retrieval. Soon after, the slides were incubated in 3% normal rabbit serum diluted 1:100 in distilled water for 20 min at room temperature, and were sequentially incubated with rabbit anti-human CD3 and CD31 polyclonal antibodies diluted 1:100 in 1% PBS- bovine serum albumin (BSA) overnight at 40 °C. Following the incubation period, the slides were washed with PBS and incubated with biotinylated goat anti-rabbit IgG antibody diluted 1:100 in 1% PBS-BSA for 60 min at room temperature. Then, the slides were incubated in avidin-biotin complex (K0492; Dako) for 45 min at room temperature, followed by incubation in a solution of 5 mg of 3,3'-diaminobenzidine (D4293; Sigma) diluted in 10 mL of PBS containing 180 µL of 20% hydrogen peroxide for 20 to 30 min at room temperature. After washing with distilled water, the slides were counterstained with Harris hematoxylin for 5 min. Tissue samples from giant cell tumors of bone were used as positive controls. Negative controls were obtained by substituting the primary antibodies with 1% PBS-BSA and normal rabbit serum (X0903; Dako).

Results

Clinical Evaluation

On postoperative day 3, the patients reported pain when removing and reinserting the denture, which had been held in place by dental adhesive paste. A yellowish cicatricial tissue was present at the insertion sites between postoperative days 3 and 7.

On postoperative day 15, patients were still on a pureed diet and showed some improvement in the retention of the denture, but still reported pain on its removal and reinsertion.

On postoperative day 30, all patients reported a marked increase in the retention and stability of the complete upper denture with no need for dental adhesive, were able to eat a normal diet, and felt little or no discomfort when removing and reinserting the denture. The appearance of the mucosa at the insertion sites was identical to the palatal mucosa. During this time, no patient developed any local infectious process related to the insertion sites.

At quarterly follow-up visits, there was little need for occlusal adjustment or adjustment of denture borders. On postoperative day 360, only one subject required the removal of two inserts from the denture to reduce its retention due to difficulties in
removing the prosthesis. Aspect of normal oral mucosa was found after three years (Figure 2).

**Histopathological Evaluation**

The macroscopic examination revealed mucosa fragments varying in size from 0.4x0.4x0.2 to 0.8x0.7x0.4 cm, with an average volume of 0.08 cm³, brown color, elastic, and showing central hole lined with white mucosa. The microscopic analysis (Figures 3A-B) revealed two patterns in the analyzed slides: (a) oral mucosa of the surgical margin: mild keratinization, moderate epithelial hyperplasia, keratinocytes with perinuclear halos, moderate spongiose, and mononuclear exocytosis; and (b) periorificial oral mucosa (in contact with the insert): parakeratosis, mild to severe epithelial hyperplasia, minimal to extensive granular layer; lamina propria with dense collagen bands (Figure 3C), fading of the loose subepithelial connective tissue, few mononuclear leukocytes (scar). The focal mononuclear infiltrate, which was composed predominantly of T lymphocytes, was confirmed by anti-CD3 (Figure 3D). Sparse new formed vessels, well evidenced by anti-CD31 (Figures 3E-F).

**Discussion**

The histopathological observations carried out in this study indicate that polished zirconia ceramic is very well tolerated by the mucosal tissue, leading to the accumulation of collagen fibers in the area around the insert, maintaining mild inflammatory response, and allowing reepithelialization, which is expressed by parakeratosis, epithelial hyperplasia and presenting granular layer. No granulomatous reaction or important inflammatory foci was observed. Microscopic observations suggest that zirconia ceramic can be an effective alternative material for use in intramucosal inserts. This is in agreement with the very good results obtained from clinical evaluations, which showed that the use of inserts leads to a marked increase in the retention of the denture, thus providing comfort and confidence to patients. The data reported here are in strong accordance to previous literature (14, 15).

This technique was suggested by Hans Nordgren and developed by Gustav Dahl at the end of the first half of the 20th century (1). It made use of intramucosal metal inserts, which according to histopathological analysis, promoted the formation of a densely fibrous connective tissue around the insert, showing sclerosed vessels, metabolically active basal layer and parakeratosis (16). After some time, the insertion site in the palatal mucosa may become hyperkeratinized or show histological evidence of mild chronic inflammation; however, to date there are no reports of patient complaints. Clinically, the insertion site was healthy without edema or other signs of inflammation, showing pink mucosa of normal appearance. Dahl (1), a pioneer in this
area, reports the use of stainless steel or chrome-cobalt alloy inserts, mushroom shaped, with sizes proportional to the thickness of the mucosa, thereby facilitating their insertion and improving the retention of the denture. Another important aspect of this technique is to place the insertion sites slightly to the palatal side of the alveolar ridge, where the submucosa is thicker.

Along the years, the most important modifications of this technique have been focused on the shape of the insert, which had always been made of metal alloys. In the present study, we report for the first time the use of highly polished zirconia ceramic as the insert. Here, the clinical cases studied, it was observed that the replacement of metal by zirconia did not produce noticeable changes in the cellular response to the inserts. Previous work has being shown the safety, biocompatibility and stability of zirconia ceramics (17-19), and the lack of mutagenic and carcinogenic effect (20).

The laboratory and surgical procedures are very simple and easily performed by the general clinician. The observed clinical responses reflect the healing process of the oral mucosa after surgical procedures; the healing process was completed within 30 postoperative days. The cooperation of the patient is extremely important during this stage for the patient must keep the denture in place as long as possible, even during periods of discomfort.

The clinical results obtained for the treated patients confirmed a marked increase in the retention of the complete upper denture in all cases. All 12 patients reported improvement in their diets due to an increased ability to chew semisolid and solid foods. As a consequence, the patients’ self-esteem and confidence increased, thus improving the quality of their social interactions. However, it is important to point that the development of the technique to obtain the described results never neglected the principles of the construction of a complete upper denture. In other words, although the use of intramucosal inserts increase the retention of complete dentures, it is important not to overlook the principles of occlusal balance and stability, neither stop the search for methods to achieve better casting accuracy in recreating anatomical features. It is important to note that we are still dealing with a complete denture and precise measurements of the vertical dimension and intermaxillary relations are also required for complete dentures with intramucosal inserts.

Conclusion

Based on both clinical and histopathological observations we can conclude that intramucosal zirconia inserts did not affect the mucosal health and provide adequate retention and stability of the complete denture and comfort to the patients. Patients did not report any complaints after wearing the denture with zirconia inserts for more than
one year, and no alterations were observed in the insertion sites that could impact the mucosal health.

Reference List


Figure Captions

**Figure 1.** Photography showing the shape and dimensions of zirconia intramucosal inserts.

**Figure 2.** Clinical aspect of the maxillary alveolar ridge three years after surgery showing normal oral mucosa around the insertion sites (A, B).

**Figure 3.** Histopathological features. A. Periorificial oral mucosa with parakeratosis (*), granular layer, epithelial hyperplasia (arrow) and fibrosis (arrow head, HE). B. Detail: granular layer (arrows), parakeratosis, and mild subepithelial inflammatory infiltrate (HE). C. Fibrosis (arrows, Gomori trichrome). D. Immunohistochemistry: T lymphocytes stained with anti-CD3. E – F Immunohistochemistry: endothelial cells stained with anti-CD31 (arrow). Original magnification: A. 20x, B. 100x, C-E. 400x, F. 100x.
Table 1. Characteristics of the subjects using intramucosal inserts made of zirconia ceramic and postoperative follow-up visits where biopsies of the mucosa around insertion sites were performed.

<table>
<thead>
<tr>
<th>Case</th>
<th>Name</th>
<th>Sex</th>
<th>Age (years)</th>
<th>Follow-up visit (months)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>MAT</td>
<td>F</td>
<td>60</td>
<td>12</td>
</tr>
<tr>
<td>2</td>
<td>LAD</td>
<td>M</td>
<td>54</td>
<td>12</td>
</tr>
<tr>
<td>3</td>
<td>AN</td>
<td>M</td>
<td>58</td>
<td>12</td>
</tr>
<tr>
<td>4</td>
<td>MLB</td>
<td>F</td>
<td>62</td>
<td>12</td>
</tr>
<tr>
<td>5</td>
<td>DAFS</td>
<td>F</td>
<td>31</td>
<td>12</td>
</tr>
<tr>
<td>6</td>
<td>LGR</td>
<td>F</td>
<td>55</td>
<td>36</td>
</tr>
<tr>
<td>7</td>
<td>JP</td>
<td>M</td>
<td>57</td>
<td>12</td>
</tr>
<tr>
<td>8</td>
<td>ST</td>
<td>F</td>
<td>66</td>
<td>18</td>
</tr>
<tr>
<td>9</td>
<td>CT</td>
<td>M</td>
<td>55</td>
<td>15</td>
</tr>
<tr>
<td>10</td>
<td>FMP</td>
<td>F</td>
<td>37</td>
<td>18</td>
</tr>
<tr>
<td>11</td>
<td>CA</td>
<td>M</td>
<td>55</td>
<td>12</td>
</tr>
<tr>
<td>12</td>
<td>RCP</td>
<td>F</td>
<td>49</td>
<td>12</td>
</tr>
</tbody>
</table>

1 All study subjects were Caucasians.

2 All retention devices were inserted in the mucosa covering the maxillary alveolar ridge.