Implant rehabilitation of extremely atrophic mandibles (Cawood and Howell Class VI) with a fixed-removable solution supported by four implants: One-year results from a preliminary prospective case series study

Abstract

Objective

The objective of this study was to report one-year preliminary data on Cawood and Howell Class VI patients rehabilitated with a fixed-removable solution.

Materials and methods

Completely edentulous patients, aged 18 years or older, presenting with severely atrophic mandibles (Class VI according to Cawood and Howell) were enrolled and treated using four implants, a CAD/CAM titanium bar and a low-profile attachment system to support an implant-supported overdenture. Outcome measures were success rates of the implants and prosthesis, complications, marginal bone level changes, bleeding index, plaque index and patient satisfaction (Oral Health Impact Profile).

Results

A total of 16 Osstem TSIII implants were placed in four consecutive edentulous participants. All of the treated patients were female with an average age of 71.5 (range: 64–82). Patients were followed up for a mean period of 13.8 months (range: 12–16) after loading. No participants dropped out, and no deviation from the original protocol occurred. At the one-year follow-up, no implants or prosthesis had failed and no biological or technical complications had occurred. At the one-year follow-up, the mean marginal bone loss was 0.23 ± 0.07 mm. The Oral Health Impact Profile summary scores demonstrated a significant decrease throughout the study, from 66.5 ± 3.7 to 19.3 ± 2.8. At the one-year follow-up, the bleeding index was 1.6% and the plaque index was 4.7%.

Conclusion

Within the limitations of this study, an overdenture fully supported by four implants and a CAD/CAM titanium bar with a low-profile attachment system, can be considered an effective and predictable option for patients with Cawood and Howell Class VI atrophic mandibles. Minimum marginal bone remodeling, good periodontal parameters and patient satisfaction can be expected.

Keywords

Implant, atrophic mandible, CAD/CAM titanium bar, overdenture.
Introduction

Prosthetic rehabilitation on implants in severely atrophic patients is a challenge. In a Cawood and Howell Class VI case, the mandible presents with a depressed ridge form, inadequate in height and width, evident basilar bone loss, and alveolar nerve exposure.1 A mandibular implant overdenture is a viable treatment option for edentulous mandibles, improving overall patient satisfaction compared with a removable complete denture.2 These studies document successful treatment outcomes and better oral health-related quality of life as compared with wearing of complete dentures.3 The number of implants to be placed and the type of retention have been controversially discussed.4–6 The implant survival rate of mandibular overdentures is high regardless of the number of implants.7 Therefore, two single standing implants with ball attachments have sometimes been considered a risk and some investigators suggest using four implants with a splinting bar.6, 8–10

In patients with an alternated skeletal maxillomandibular relationship, a fixed-removable solution may be a viable option for soft- and hard-tissue reconstruction and for the All-on-4 concept for the rehabilitation of patients presenting with extremely atrophic mandibles (Cawood and Howell Class VI).1, 11 A fixed-removable solution may be a feasible option to overcome the technical complications of other treatment options.4, 12 Moreover, hygienic maintenance of the prosthesis can be challenging when extensive prosthetic flanges are needed to provide adequate lip and check support to overcome esthetic problems typical of aging.13

The purpose of this preliminary case series study was to report one-year preliminary data on Cawood and Howell Class VI patients rehabilitated with a fixed-removable solution. This study followed the Strengthening the Reporting of Observational Studies in Epidemiology reporting guidelines.14

Materials and methods

This preliminary investigation was designed as a prospective study conducted according to the principles embodied in the Helsinki Declaration of 1975, as revised in 2008. Completely edentulous patients, aged 18 years or older, presenting with severely atrophic mandibles (Class VI according to Cawood and Howell)1 were enrolled and treated in consecutive order after being informed about the nature of the study and providing their written informed consent. All of the surgical and prosthetic procedures were performed in a private center in Rome, Italy, by a certified implantologist (MT) between September 2015 and February 2016. Exclusion criteria were general contraindications to oral surgery, pregnancy or nursing, intravenous bisphosphonate therapy, alcohol or drug abuse, heavy smoking (≥ 10 cigarettes/day), radiation therapy to the head or neck region within the last five years, parafunctional activity, untreated periodontitis, full-mouth bleeding on probing, and a full-mouth plaque index of ≤ 25%, and allergy or adverse reactions to the restorative materials.

Preoperative photographs, radiographs (Figs. 1a–c) and model casts were produced for initial screening and case evaluation. A radiographic guide was made by duplicating the relined pre-existent removable complete mandibular denture, if judged viable from an esthetic and functional perspective; otherwise, a new radiographic guide was made according to the functional and esthetic requirements. A cone beam computed tomography (CBCT) scan (CRANEX 3Dx, Soredex, Tuusula, Finland) was taken of each enrolled patient wearing the radiographic guide and a bite index in centric occlusion with an extraoral volume transfer element (Evobite, 3DIEMME, Cantù, Italy), fixed using a dedicated silicone material (3DIEMME). Then, the radiographic guide and the bite index were repositioned in the master cast and optical scanning was performed. Radiographic and prosthetic data were imported into a dedicated diagnostic and medical imaging software (3Diagnosys 4.2, 3DIEMME). The digitalized model and radiographic guide were accurately superimposed over the reconstructed bone volume by CBCT, based on the volumetric elements, present in both the CBCT volume and the optical scan (Evobite). Four implants per patient were planned in the anterior area of the mandible, according to the prosthetic setup. After careful functional and esthetic evaluation and final verification, the prosthetic-driven plan was approved, and a stereolithographic surgical template was fabricated with a newer rapid prototyping technology (New Ancorvis, Bargellino, Italy).

One hour before implant placement, patients received a single dose of an antibiotic (2 g of amoxicillin or 600 mg of clindamycin if allergic to penicillin) and professional hygiene therapy. Prior to the start of surgery, the patients rinsed
**Figs. 1a–c**
Preoperative radiographs (a & b) and intraoral photograph (c).

**Figs. 2a & b**
Clinical view (a) after placement of the four implants (Osstem TSIII, 4.5 × 10 mm) according to a one-stage protocol. Radiograph after implant placement (b).

**Fig. 3**
Definitive impression taken using plaster as both splinting and impression material.
with 0.2% chlorhexidine for 1 min. The surgical template (New Ancorvis) was fitted in the patient’s mouth, then local anesthesia was administered with a 4% articaine solution with 1:100,000 epinephrine (Ubistein, 3M ESPE, Milan, Italy). The implants were placed in the planned anatomical sites according to a one-stage approach (Figs. 2a & b)\textsuperscript{15} using the surgical template. Each patient received four Osstem TSIII bone level implants (Osstem TSIII, Osstem, Seoul, South Korea), placed either without a flap or with a minimally invasive flap, according to the drilling protocol recommended by the manufacturer (OsstemGuide Kit). After surgery, the existing removable complete denture was relined chairside (Sofreliner Tough Soft, Tokuyama Dental, Montecchio Precalcino, Vicenza, Italy) to accommodate the healing of the hard and soft tissue, thereby ensuring no pressure on the healing abutments. Occlusion was checked during osseointegration of the implants.

After implant placement, all of the patients received oral and written recommendations on medication, oral hygiene maintenance and diet. Analgesics (500 mg of paracetamol plus 30 mg of codeine, or 600 mg of ibuprofen) were administered as needed. Eight weeks later, a definitive impression was taken using plaster (Snow White Plaster No. 2, Kerr, Orange, Calif. U.S.) as both splinting and impression material (Fig. 3)\textsuperscript{16} A complete mounting technique was used to articulate the opposite arch cast (KaVo PROTAReo 7, KaVo Dental, Biberach, Germany). Then, esthetics and function of the final occlusal vertical dimension, tooth position and interocclusal record were verified and approved by both the clinician and the patient at the try-in appointment. Afterward, the master cast and the try-in were digitalized with an optical scanner (Identica T500, Medit, Seoul, South Korea).

A CAD/CAM titanium bar was virtually designed (Figs. 4a & b) according to the ridge and prosthetic contours and implant position in order to enhance a vertical path of insertion of between 4 and 6°. Then, a one-piece titanium bar was manufactured (Fig. 5) from a homogeneous solid block of a medical titanium alloy (Ti6Al4V, New Ancorvis). Four to five threadable OT Equator attachments (Rhein 83, Bologna, Italy) were placed along the implant bar. The fit of the implant bar was clinically and radiographically tested in the patient’s mouth according to a previously published protocol.\textsuperscript{17, 18} A cast cobalt–chromium alloy metal framework (Vitalium, DENTSPLY International, York, Pa., U.S.)
was conventionally fabricated on to the CAD/CAM titanium bar as a counterpart (Fig. 6). Finally, the overdenture was finished, sealing the borders to minimize food impaction and saliva and air leakage. The titanium bar was screwed at the abutment level according to the manufacturer’s instructions, and the fixed-removable solution was delivered (Figs. 7a–c). All of the patients were then enrolled in a standard implant recall program. Oral hygiene maintenance was checked and radiographs were taken early after final prosthesis delivery. Occlusion was checked at every appointment (Figs. 8a & b).

**Outcome measures**

The primary outcome measures were the following:

- **Success rates of the implants and prosthesis:** An implant was considered a failure if it presented with any mobility, assessed by tapping or rocking the implant head with the metallic handles of two instruments, progressive marginal bone loss or infection, and any mechanical complications rendering the implant unusable, although still mechanically stable in the bone. A prosthesis was considered a failure if it needed to be replaced with another prosthesis.

- **Complications:** Any biological (pain, swelling, suppuration, etc.) and/or mechanical (screw loosening, fracture of the framework and/or the veneering material, etc.) complications were evaluated.

- **Marginal bone levels:** The levels were assessed using intraoral digital periapical radiographs (Digora Optime, SOREDEX; photostimulable phosphor imaging plate, size 2, pixel size of 30 μm, resolution of 17 lp/mm) at implant placement (baseline) and one year after loading. Intraoral radiographs were taken with the paralleling technique by means of a periapical radiograph with a commercially available film holder (Rinn XCP, Dentsply Rinn, Elgin, Ill., U.S.). The radiographs were accepted or rejected for evaluation based on the clarity of the implant threads. All readable radiographs were uploaded to an image analysis software package (DiW 2.8, SOREDEX) that was calibrated using the known length or diameter of the dental implants and displayed on a 24 in. LCD screen (iMac, Apple, Calif., U.S.) and evaluated under standardized conditions (ISO 12646:2004). The marginal bone levels were determined from linear measurements performed by an independent calibrated examiner on each periapical radiograph, from the mesial and distal margin of the implant neck to the most coronal point where the bone appeared to be in contact with the implant.

- **Patient satisfaction with function and esthetics** was assessed using a scale of 1–10, where
Figs. 7a–c
Clinical view of the overdenture at the time of prosthesis delivery (a & b) and radiographic view early after final prosthesis delivery (c).

Figs. 8a & b
One-year follow-up of a mandibular CAD/CAM titanium bar screwed on to implants (a). Dental panoramic tomogram taken one year after loading, showing the perfect fit of the bar on the implants (b).
Results

A total of 16 Osstem TSIII implants (14 regular platform and two mini platform) were placed in four consecutive edentulous participants. All of the participants were followed up for a minimum of one year (mean: 13.8 months; range: 12–16) after definitive loading. All of the treated patients were female with an average age of 71.5 (range: 64–82). The main patient and implant characteristics are shown in Table 1. No participants dropped out, and no deviation from the original protocol occurred. At the one-year follow-up, no implants or prosthesis had failed, resulting in cumulative implant and prosthetic survival rates of 100%. No biological or technical complications occurred during the follow-up, resulting in cumulative implant and prosthetic success rates of 100%. At the one-year follow-up, the mean marginal bone loss was $0.23 \pm 0.07$ mm. The OHIP summary scores demonstrated a significant decrease ($P = 0.0002$) throughout the study, from $66.5 \pm 3.7$ to $19.3 \pm 2.8$. At the one-year follow-up, the bleeding index was 1.6% and the plaque index was 4.7%. All of the data are summarized in Table 1.

Discussion

This prospective study was designed to evaluate the one-year clinical and radiographic outcomes and patient satisfaction of Cawood and Howell Class VI patients treated with a fixed-removable overdenture supported by four implants, placed using guided surgery, and a CAD/CAM titanium bar. Because it was designed as a single-cohort, proof-of-concept study, the main limitations were the lack of a control group and a small sample size. Hence, this investigation should be considered as a pilot for future multicenter randomized clinical trials with control group comparison.

The results of the present one-year preliminary prospective case series study reported implant and prosthetic survival and success rates of 100% and greater patient satisfaction, indicating that patients with extremely atrophic mandibles (Cawood and Howell Class VI) may be rehabilitated using this fixed-removable solution.

Complete maxillary and mandibular dentures have been the conventional standard of care for edentulous patients. However, most patients report significant problems adapting to their mandibular dentures owing to a lack of comfort, retention and stability and to the inability to chew and eat properly, resulting in pain and discomfort. Recent evidence from studies carried out over the past decade has determined that the two-implant overdenture is considered...
Fixed-removable solution for atrophic mandibles

Table 1

<table>
<thead>
<tr>
<th></th>
<th>Age (years)</th>
<th>Sex</th>
<th>Smoking</th>
<th>Implants 7 mm length</th>
<th>Implants 8.5 mm length</th>
<th>Implants 10 mm length</th>
<th>Implants ≥ 4 mm wide</th>
<th>Failed implant</th>
<th>Failed prosthesis</th>
<th>MBL (mm)</th>
<th>OHIP T0</th>
<th>OHIP T1</th>
<th>BI</th>
<th>PI</th>
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<tr>
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<td>0</td>
<td>4</td>
<td>0</td>
<td>4</td>
<td>0</td>
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<td>71.0</td>
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<td>F</td>
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<td>4</td>
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<td>2</td>
<td>0</td>
<td>0</td>
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<td>63.0</td>
<td>21.0</td>
<td>1/16</td>
<td>0/16</td>
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<td>Patient 3</td>
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<td>4</td>
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<tr>
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<td>0</td>
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<td>64.0</td>
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<td>2</td>
<td>4</td>
<td>10</td>
<td>2</td>
<td>14</td>
<td>0</td>
<td>0.23 ± 0.07</td>
<td>66.5 ± 3.7</td>
<td>19.3 ± 2.8</td>
<td>1/64</td>
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<tr>
<td>Mean ± SD</td>
<td>71.5 ± 7.7</td>
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<td></td>
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<td>0.23 ± 0.07</td>
<td>66.5 ± 3.7</td>
<td>19.3 ± 2.8</td>
<td>1/64</td>
<td>3/64</td>
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</table>

MBL = marginal bone loss; OHIP = Oral Health Impact Profile; T0 = baseline; T1 = one year after definitive prosthesis delivery; BI = bleeding index; PI = plaque index; SD = standard deviation.

the first alternative treatment for the completely edentulous mandible.19, 20 Nevertheless, the placement of at least four implants of standard length may allow the delivery of an overdenture supported by a CAD/CAM titanium bar and a low-profile attachment system,21 avoiding any bearing area on the soft tissue and reducing the denture base extension.22 The OT Equator for bars exists in two types, castable and prefabricated (threadable). In the present study, the prefabricated shape was used. This type of attachment is initially of higher cost, but it is highly wear resistant, its surface being of titanium nitride. Furthermore, it is easy to replace, if needed.

A fixed dental prosthesis on four implants may be a possible alternative to a mandibular overdenture on four implants and supported by a CAD/CAM titanium bar with a low-profile attachment system. Nevertheless, it is associated with higher marginal bone loss, high frequency of complications and poor plaque control, particularly in extremely atrophic patients.23–26 In the present study, a trend of minimum marginal bone loss and good periodontal parameters was observed within the one-year follow-up, demonstrating that a good level of hygiene can be expected using this fixed-removable solution. Similar to with a fixed dental prosthesis, patient satisfaction significantly improves owing to an improvement in esthetics and masticatory function. Moreover, the prosthetic flanges of a fixed-removable solution allow for full lip and cheek support.

Conclusion

Within the limitations of this study, a mandibular overdenture on four implants and supported by a CAD/CAM titanium bar with a low-profile attachment system can be considered an effective and predictable option for patients with Cawood and Howell Class VI atrophic mandibles. Minimum marginal bone remodeling, good periodontal parameters and patient satisfaction can be expected.

Competing interests

MT is the Research Project Manager at Osstem AIC, Italy. However, this study was self-supported. Hence, the authors declare no conflicts of interest.
References


