Three-year results of All-on-4 in daily practice

Abstract

Objective

All-on-4 treatment concept is widely applied for complete-arch rehabilitations. Nevertheless, minor technical and biological complications can occur. The objective of this study was to evaluate the three-year clinical and radiographic data of complete-arch fixed dental prostheses supported by four implants according to the All-on-4 protocol.

Materials and methods

Thirty consecutive edentulous patients or patients with terminal dentition (18 females and 12 males; mean age of 67.4 years), with a preference for implant-supported complete-arch screw-retained fixed dental prostheses, were enrolled and treated according to the All-on-4 protocol between January 2008 and December 2011. The outcomes evaluated were implant and prosthesis survival and success rates, any technical and biological complications, periimplant marginal bone loss and patient satisfaction.

Results

One hundred and twenty regular platform implants were placed. No patients dropped out. One implant failed two months after placement, resulting in a cumulative implant survival rate of 99.2%. No definitive prostheses failed. Eight technical and three biological complications were reported in 11 patients during the entire follow-up period. At the three-year examination, the mean marginal bone loss was 1.52 ± 0.41 mm.

Conclusion

Within the limitations of the present study, the All-on-4 protocol was deemed a viable treatment concept for the complete-arch rehabilitation of both jaws in the medium term. Further long-term prospective studies are needed to confirm these results.
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Introduction

Complete edentulism is associated with decreased masticatory function, as well as unfavorable esthetics due to the loss of support for the facial musculature, decreased vertical dimension and speech impairment. The conventional method for treating edentulous patients is to rehabilitate them with a complete removable denture. However, the denture must be adjusted over time to compensate for the progressive tissue changes associated with denture wearing. Advances in implant dentistry have allowed a shift from a complete removable denture to an implant-supported overdenture for the oral rehabilitation of edentulous patients. The McGill consensus statement in 2002 and some independent studies state that mandibular two-implant overdentures are the gold standard for edentulous patients. However, technical and biological complications can occur.

In two pilot retrospective studies, Maló et al. presented a planning protocol for the rehabilitation of the edentulous mandible and maxilla using four implants (All-on-4, Nobel Biocare, Kloten, Switzerland) to overcome anatomical limitations in the mandible that make it challenging to treat without the use of more complex techniques. The two most anterior implants are placed axially, and the posterior implants are placed in an angled position to maximize implant length and avoid anatomical structures (i.e., the mental nerve and anterior border of the maxillary sinus). The All-on-4 treatment concept seems to be a safe, effective and efficient surgical and prosthetic procedure on both jaws after ten years in function. However, as confirmed in a recent systematic review, few independent uncontrolled prospective and retrospective studies have confirmed these preliminary results.

The purpose of this prospective observational study was designed to evaluate patients treated according to the All-on-4 protocol in the mandible or maxilla. The patients were selected and treated in two private centers in Rome and Sassari, between January 2008 and December 2011.

The inclusion criteria were a healthy patient aged 18 years or older at the time of implant placement, able to give her or his informed consent for participation, with a residual alveolar crest, distal to the first premolar, of ≤ 5 mm in height and ≤ 4 mm in width, assessed by computed tomography (CT) or scans, and refusal of a conventional bone augmentation procedure; partially edentate patient with hopeless dentition based on multiple risk factors, including endodontic and periodontal criteria, prosthetic restorability of the tooth, cost–benefit ratio and patient preference in terms of refusing any major bone augmentation procedures.

The exclusion criteria were general medical (American Society of Anesthesiologists Physical Status Class III or IV) and/or psychiatric contraindications; pregnancy or nursing; any interfering medication, such as steroid therapy or bisphosphonate therapy; alcohol or drug abuse; heavy smoking (> 10 cigarettes/day); radiation therapy to the head or neck region within five years; high or moderate parafunctional activity; untreated periodontitis; poor oral hygiene and motivation, defined as full-mouth bleeding on probing and a full-mouth plaque index of ≥ 25%; known allergic or adverse reactions to the restorative material; and unavailability for regular follow-ups.

This investigation was conducted according to the principles embodied in the Declaration of Helsinki of 1975 for biomedical research involving human subjects, as amended in 2008. All of patients were duly informed about the nature of the study and gave their written consent.

Before implant placement, all of the patients underwent CT or CBCT scan according to a double-scan protocol. In the case of immediately post-extraction implants, a previously reported two-piece radiographic guide was used for the diagnostic study and for virtual implant planning. If the operator decided to wait for healing of the post-extraction socket, the implant sites had to heal for at least three months before radiographic examination (Figs. 1–2).

DICOM data of the two sets of scans were transferred to a 3-D software planning program (NobelGuide, Nobel Biocare) and matched to each...
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other. Four implants were planned for the rehabilitation of each participant in post-extraction or healed sockets, according to the All-on-4 protocol. Planning data of the patients who required template-guided surgery were sent to a milling center (NobelProcera, Nobel Biocare), where stereolithographic surgical templates with hollow metallic cylinders to guide implant placement in the virtually planned position were fabricated. Patients received professional oral hygiene prior to the surgery and were instructed to rinse with a 0.2% chlorhexidine mouthwash for 1 min, starting two days prior to the intervention.
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Surgical protocol

An antibiotic (2 g of amoxicillin and clavulanic acid or 600 mg clindamycin if allergic to penicillin) was administered 1 h prior to surgery and continued for six days (1 g amoxicillin and clavulanic acid or 300 mg clindamycin b.i.d.) after surgery. Local anesthesia was induced using a 4% articaine solution with 1:100,000 epinephrine (Ubistesin; 3M Italia, Milan, Italy). Implants were placed in the planned anatomical sites either conventionally or using a fully guided approach. A flapless or a flap approach was performed in order to maintain an adequate residual band of keratinized mucosa around the implants. When the alveolar crest was too thin (knife edge) to place the implant, the alveolar crest was remodeled using piezoelectric bone surgery under copious irrigation with sterile saline to obtain a flat bony crest. Each patient received four NobelSpeedy Groovy implants (Nobel Biocare), featuring a flat-to-flat matched implant-abutment interface with a 0.7 mm tall external hexagonal prosthetic connection and a rough, highly crystalline and phosphate-enriched titanium oxide surface (TiUnite, Nobel Biocare). All of the implants were placed according to the surgical and prosthetic protocols recommended by the manufacturer (IFU 73494 Manual 2/All-on-4 and IFU 71286), with no deviations from the original protocol. The drilling sequence was chosen according to the manufacturer’s instructions in relation to the bone quality, achieving an insertion torque at implant placement ranging from 35 to 45 N cm in the mandible or from 35 to 55 N cm in the maxilla, measured using a surgical unit (OsseoCare Pro Drill Motor Set, Nobel Biocare). In the post-extraction sites, the gaps between the implants and the surrounding socket walls were filled with 0.25–1 mm granules of deproteinized bovine bone (Geistlich Bio-Oss or Geistlich Bio-Oss Collagen, Geistlich Pharma, Wolhusen, Switzerland), hydrated using the patient’s blood mixed with antibiotic solution (Rifocin 250 mg/10 mL, Sanofi-aventis, Milan, Italy).

Seventeen- or thirty-degree angled multi-unit abutments (Nobel Biocare) were immediately connected to the distal implants for better orientation of the screw access hole. Straight multi-unit abutments (Nobel Biocare) were used in the anterior implants if needed.

A prefabricated screw-retained acrylic resin provisional restoration without any cantilever was delivered immediately after surgery (Figs. 3–5). All of the patients received oral and written recommendations regarding medication, oral hygiene maintenance and diet. Post-surgical analgesic treatment was provided with ibuprofen 600 mg, administered every 8 h for two days after the surgery, and later on if needed. The patients were instructed to rinse the mouth with a 0.2% chlorhexidine mouthwash t.i.d. without brushing the implant area (Figs. 6 & 7).

Prosthetic protocol

After three to four months of healing, a definitive impression was taken at the implant or abutment level according to a previously reported protocol.27 Definitive prostheses with titanium or zirconia frameworks, fabricated using CAD/CAM technology (NobelProcera), were screwed on at either the implant or abutment level according to the manufacturer’s instructions three to five months after implant placement (in the mandible and the maxilla, respectively). The definitive implant-supported complete-arch FDP was designed with (hybrid design) or without pink material in the cervical region (crown design) and veneered with ceramic, acrylic or composite according to the patients’ needs (Figs. 8a–c). Clinical accuracy of the framework (strain-free screwing and absence of an open margin upon clinical and radiographic examination) was evaluated before prosthesis delivery.28–30 The occlusion was adjusted avoiding any premature contacts. Mutually protected occlusion with anterior guidance or balanced occlusion was used in cases of opposing natural dentition or an FDP and complete removable denture, respectively. Follow-up visits were scheduled at one and six months and then annually up to five years of function. The patients underwent a professional cleaning by a dental hygienist every four to six months (Figs. 9a–c). Panoramic and periapical radiographs were obtained annually after definitive prosthesis delivery (Fig. 10).

The primary outcome measures were as follows:

- An implant was classified as “successful” when the following criteria31 were fulfilled: did not cause pain or suppuration, did not show any mobility, did not show any signs of radiolucency, and did not show periimplant bone loss of > 1.5 mm during the first year and then > 0.2 mm yearly.
- An implant was classified as “surviving” when the implant remained in the jaw and was stable after the prosthesis was removed, even though all of the success criteria were not fulfilled.
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Figs. 8a–c
Lateral (a & c) and frontal (b) intra-oral photographs of the carbon-fiber-reinforced definitive prosthesis screwed onto the implants.

Figs. 9a–c
Lateral (a & c) and frontal (b) intra-oral photographs of the definitive prosthesis taken three years after implant placement.

Fig. 10
Panoramic radiograph three years after implant placement.

An implant-supported complete-arch FDP was defined as “successful” when the dental prosthesis remained in function and the esthetic evaluation, assessed by the dentist and the patient, was satisfactory during the study period.

An implant-supported complete-arch FDP was considered as “surviving” when the dental prosthesis remained in function, even though all of the success criteria were not fulfilled.

The secondary outcome measures were as follows:

- Any technical (fracture of the framework and/or the veneering material, screw loosening, etc.) and/or biological (pain, swelling, suppuration, etc.) complications were considered.

- The distance from the most coronal margin of the implant collar and the most coronal point of bone-to-implant contact was taken as the marginal bone level. This level was evaluated on intra-oral digital radiographs taken with the paralleling technique using a film holder (Rinn XCP, DENTSPLY, Elgin, Ill., U.S.) at implant placement (baseline) and then yearly up to three years of function. Radiographs were accepted or rejected for evaluation based on the clarity of the implant threads. All readable radiographs were displayed in image analysis software (Digora for Windows 2.8, SOREDEX, Tuusula, Finland) that was calibrated for every single image using the known measure of the implant thread pitch. Measurements of the mesial and distal bony crest level adjacent to each implant were made to the nearest 0.1 mm and averaged at the patient level.

- Patient satisfaction was evaluated with a questionnaire one month after delivery of the final prosthesis and at the three-year follow-up examination, provided by independent and blinded outcome assessors. The assessor asked the following questions: Are you satis-
fied with the function of your implant-supported prosthesis? Are you satisfied with the esthetic outcome of your implant-supported prosthesis? Would you undergo the same therapy again?

An independent assessor evaluated the implant and prosthetic survival and success rates. Complications were assessed and treated by the same clinicians. The marginal bone loss was evaluated by an independent radiologist.

Statistical analysis

Patient data were collected in an Excel spreadsheet (Microsoft). Finally, all of the data were exported into IBM SPSS Statistics for Macintosh (Version 22.0; IBM, Armonk, N.Y., U.S.) for statistical analysis. A bio-statistician with expertise in dentistry analyzed the data using PASW Statistics for Windows (Version 18.0; SPSS, Chicago, Ill., U.S.). Descriptive analysis was performed for numeric parameters using mean ± standard deviation (median; 95% CI). Dichotomous and continuous outcomes were compared using the chi-squared test and one-way analysis of variance, respectively. Differences in the proportions of patients with implant failures, prosthesis failures and complications (dichotomous outcomes) were compared between the subgroups using the Fisher exact test. Patient was the statistical unit of the analyses. All statistical comparisons were conducted at a 0.05 level of significance.

Results

Thirty-two patients were screened for eligibility. Two patients were not enrolled in the trial, because of refusal to sign the informed consent. A total of 120 NobelSpeedy Groovy implants were placed using either computer-assisted template-guided (n = 15) or conventional freehand surgery (n = 15) in 30 consecutive patients (18 females and 12 males) with a mean age of 67.4 ± 6.9 (range of 51–87). Thirty CAD/CAM screw-retained implant-supported complete-arch FDPs (18 in the mandible and 12 in the maxilla) were delivered. No patients dropped out of the study within three years after implant placement and no deviation from the original protocol occurred. All of the patients were followed up for a minimum period of three years (mean of 53.8 months; range of 36–84 months). Data collected were included in the statistical analysis. The main patients’ and interventions’ characteristics are summarized in Table 1.

At the three-year follow-up examination, one out of 120 implants (0.8%) had failed, resulting in a cumulative implant survival rate of 99.2%. The only implant failure occurred in one patient two months after placement, before delivery of the final prosthesis. The affected implant had an infectious etiology (pain, swelling and suppuration) and it was 11.5 mm long in position 16, placed using guided surgery in a healed site. The implant was replaced three months after bone healing. The temporary prosthesis was shortened, but not replaced by a conventional complete removable denture, and the patient was adequately informed and instructed to pay attention and follow a soft diet. At the three-year follow-up examination, no definitive prostheses had failed, resulting in a cumulative prosthesis survival rate of 100%.

Eleven patients experienced one technical or biological complication each, resulting in eight technical and three biological complications reported during the entire follow-up period. Six technical and one biological complication were reported during the healing period with temporary prostheses, while two technical and two biological complications were reported after definitive prosthesis delivery. All of the complications were successfully resolved.

Three prosthetic screws loosened in the temporary prostheses (in three patients) during the healing period, and this was resolved by retightening the screws, stabilizing the occlusion and advising the patients not to overload the prostheses (not to ingest food that may require significant masticatory effort). Three fractures of the provisional acrylic prostheses occurred (in three patients) during the healing period. The temporary prosthesis was adjusted chairside, the occlusion was stabilized and a night guard was delivered for each patient. Fracture of the composite veneering material of the definitive implant-supported cross-arch FDP occurred in one patient two years after loading, most likely due to occasional parafunctional habits. These situations were resolved by adjusting the definitive prosthesis chairside, stabilizing the occlusion and delivering a night guard.

The first biological complication was reported six weeks after implant placement in an 11.5 mm long implant placed in position 45, using guided surgery in a healed site. The patient reported pain and swelling without suppuration. The temporary abutment was replaced with a healing abutment. The temporary prosthesis was shortened to the...
right canine. The implant was left to heal for four months according to a conventional loading protocol. The other two complications were periimplantitis, consisting of a mean mesiodistal peri-implant bone loss of 2.6 and 2.8 mm, reported at the one-year follow-up examination. The first case of periimplantitis developed around a 15 mm long implant in position 35, placed using conventional freehand surgery in a healed site. The second case developed around a 13 mm long implant in position 32, placed using guided surgery in a healed site and immediately loaded. No other technical or biological complications occurred during the entire follow-up period.

After an initial mean marginal bone loss of 1.16 ± 0.40 mm (1.06 mm; 95% CI: 0.92–1.20), all of the implants lost a mean of 0.21 ± 0.11 mm (0.20 mm; 95% CI: 0.16–0.24) between the one-and two-year follow-ups, and 0.16 ± 0.07 mm (0.15 mm; 95% CI: 0.13–0.17) between the two-and three-year follow-ups. At the three-year follow-up, the mean marginal bone loss was 1.52 ± 0.41 mm (1.42 mm; 95% CI: 1.27–1.57). The radiographic data are shown in Table 2.

All of the patients were fully satisfied with the function and esthetics of their definitive prostheses, and all of the patients declared that they would undergo the same treatment again.

**Maxilla versus mandible**

Twelve patients were treated in the maxilla, while 18 patients were treated in the mandible. There were no statistically significant differences between centers for the number of patients who had failed implants (1/12 vs. 0/18; risk ratio = NA; \( p = 0.399 \)) or complications (3/12 vs. 8/18; risk ratio = 0.5625; 95% CI: 0.19–1.70; \( p = 0.442 \)). At the three-year follow-up examination, the mean marginal bone loss was 1.49 ± 0.34 mm (1.42 mm; 95% CI: 1.23–1.61) in the maxilla versus 1.54 ± 0.46 mm (1.38 mm; 95% CI: 1.17–1.59) in the mandible (\( p = 0.756 \)). The radiographic data are shown in Table 3.

**Guided versus conventional freehand surgery**

Fifteen patients were treated using computer-assisted template-based surgery and 15 with conventional freehand surgery. There were no statistically significant differences between centers for the number of patients who had failed implants (1/15 vs. 0/15; risk ratio = NA; \( p = 0.999 \)) or complications (4/15 vs. 7/15; risk ratio = 0.5714; 95% CI: 0.21–1.55; \( p = 0.449 \)). At the three-year follow-up examination, the mean marginal bone loss was 1.48 ± 0.47 mm (1.33 mm; 95% CI: 1.09–1.57) in the guided surgery group versus 1.55 ± 0.34 mm (1.46 mm; 95% CI: 1.29–1.63) in the conventional freehand surgery group (\( p = 0.365 \)). The radiographic data are shown in Table 3.

**Discussion**

The present study reported data on 30 implant-supported restorations delivered according to the All-on-4 protocol and followed for at least three years after implant placement. Because it was designed as a single-cohort prospective study, the main limitation of the present research was the lack of a control group and the small sample size. Another limitation of the present study was the variability within the cohort of patients. In the present study, the three-year implant (99.2%) and prosthesis success rates (100.0%), as well as the mean bone loss of 1.52 ± 0.41 mm, indicate that the All-on-4 treatment concept is a promising treatment modality. Furthermore, the results of this prospective observational study are consistent with other studies investigating the same topic.

A recent systematic review by Patzelt et al., which included 4,804 implants, demonstrated a mean cumulative implant and prosthesis survival rate at three years of 99.0 ± 1.0% and 99.9 ± 0.3%, respectively. At the three years amounted to 1.3 ± 0.4 mm. However, 12 out of the 13 included studies were considered to be highly biased. Most of the studies included (69%) in the systematic review derived from a limited number of investigators in Italy and Portugal, which may limit the generalizability of the findings, and only 31% of the studies reported a complete follow-up period of three years.

Malo et al. retrospectively reported a cumulative patient-related success rate of 93.8% up to ten years of follow-up in the mandible and a prosthesis survival rate of 99.2%. In the maxilla, a five-year survival rate of 93% was reported, and the survival rate of the prostheses was 100%. The mean marginal bone loss was 1.52 ± 0.30 mm after three years. Similar results were reported by Browaeys et al., who highlighted unacceptable ongoing bone loss in 49.2% of the patients.

Balshi et al. retrospectively analyzed the outcomes of 200 arches (800 implants) treated
### Table 1
Patients' and interventions' characteristics.

<table>
<thead>
<tr>
<th>Males</th>
<th>12 (40.00%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Females</td>
<td>18 (60.00%)</td>
</tr>
<tr>
<td>Mean age at implant placement</td>
<td>67.4 ± 6.9</td>
</tr>
<tr>
<td>Smokers (&lt; 10 cigarettes/day)</td>
<td>5 (16.67%)</td>
</tr>
<tr>
<td>Patients treated in the maxilla</td>
<td>12 (40.00%)</td>
</tr>
<tr>
<td>Patients treated in the mandible</td>
<td>18 (60.00%)</td>
</tr>
<tr>
<td>Patients treated using guided surgery</td>
<td>15 (50.00%)</td>
</tr>
<tr>
<td>Patients treated using conventional surgery</td>
<td>15 (50.00%)</td>
</tr>
<tr>
<td>Post-extraction implants</td>
<td>19 (15.83%)</td>
</tr>
<tr>
<td>10.0 mm implant length</td>
<td>14 (11.67%)</td>
</tr>
<tr>
<td>11.5 mm implant length</td>
<td>42 (35.00%)</td>
</tr>
<tr>
<td>13.0 mm implant length</td>
<td>62 (51.67%)</td>
</tr>
<tr>
<td>15.0–16.0 mm implant length</td>
<td>2 (1.67%)</td>
</tr>
<tr>
<td>Patients with prosthesis failures</td>
<td>0</td>
</tr>
<tr>
<td>Patients with implant failures</td>
<td>1 (3.33%)</td>
</tr>
<tr>
<td>Patients with complications</td>
<td>11 (36.67%)</td>
</tr>
</tbody>
</table>

### Table 2
Mean marginal bone loss ± standard deviation (mm) (95% CI) between follow-up examinations.

<table>
<thead>
<tr>
<th>Follow-up period</th>
<th>Baseline to 1 year</th>
<th>1–2 years</th>
<th>2–3 years</th>
<th>Baseline to 3 years</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.16 ± 0.40 (0.92–1.20)</td>
<td>0.21 ± 0.11 (0.16–0.24)</td>
<td>0.16 ± 0.07 (0.13–0.17)</td>
<td>1.52 ± 0.41 (1.27–1.57)</td>
<td></td>
</tr>
</tbody>
</table>

### Table 3
Three-year marginal bone loss ± standard deviation (mm) according to the jaw location and type of surgery.

<table>
<thead>
<tr>
<th>Location</th>
<th>Mandible (n = 18)</th>
<th>Maxilla (n = 12)</th>
<th>P-value†</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.54 ± 0.46</td>
<td>1.49 ± 0.34</td>
<td>P = 0.756</td>
<td></td>
</tr>
<tr>
<td>Type of surgery</td>
<td>Guided (n = 15)</td>
<td>Freehand (n = 15)</td>
<td>P-value†</td>
</tr>
<tr>
<td>1.48 ± 0.47</td>
<td>1.55 ± 0.34</td>
<td>P = 0.365</td>
<td></td>
</tr>
</tbody>
</table>

† Unpaired t-test assuming normal distribution.
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Grandi et al. analyzed 47 patients treated with 188 immediately loaded implants placed in the mandible according to the All-on-4 protocol.34 At the 18-month follow-up, no implant had failed and all of the restorations were stable. However, three patients experienced fracture of the provisional restoration. No significant differences in bone loss were found between axially placed and tilted implants at the 18-month follow-up.

Babbush et al. retrospectively examined 165 patients treated according to the All-on-4 protocol.35 The cumulative implant survival rate was 99.6% (99.3% in the maxilla and 100.0% in the mandible) for up to 29 months of loading. The definitive prosthesis survival rate was 100%. Recently, the same authors retrospectively analyzed the patient-centered outcomes, including the cost of treatment, length of the treatment period and comfort provided by the provisional restoration, in patients treated according to the All-on-4 protocol and compared these results to a historical control group, which included complete-arch FDPs supported by natural teeth or implants and implant-supported overdentures.36 This study demonstrated that the cost, length of treatment and comfort provided by the provisional restoration significantly favored the All-on-4 treatment modality.

In the present study, the overall percentage of complications experienced was large (36.6%). Nevertheless, this result did not differ from those normally encountered in oral rehabilitation in which implants are used as support for an FDP (33.6% at five years).37 Moreover, most of these complications were reported on the temporary restoration, during healing. The clinicians who carried out the procedures addressed all of the complications chairside. Furthermore, the condition of the patients remained stable up to the completion of the three-year follow-up period. In order to minimize the incidence of complications, dental clinicians should exert great effort in selecting patients, respecting the original protocols, and choosing reliable components and materials for implant-supported complete-arch FDPs.

Patients with untreated periodontitis were not included in the study. Implant therapy in patients with a history of chronic periodontitis and generalized aggressive periodontitis might be considered a viable treatment with similar survival outcomes to those reported for healthy patients. Periodontally compromised patients were included after being treated to reduce the inflammation and halt the disease progression, before tooth extraction and implant placement. According to Donos et al., it is necessary to treat and control the periodontal disease, regardless of its progression pattern and subtype, before implant therapy is initiated in order to improve the overall implant success and achieve a more favorable bone resorption pattern.38 Nevertheless, a comprehensive implant maintenance program has to be encouraged and continued in order to identify perimplant bone loss early on, particularly in patients with a history of periodontal disease.

Conclusion

Within the limitations of the present study, the All-on-4 concept is a predictable and minimally invasive treatment concept for the complete-arch rehabilitation of both jaws, regardless of jaw location and type of surgery. It may decrease the overall treatment time and re-establish adequate function in a cost-effective way. Further long-term prospective data (five years and more) and outcomes beyond cumulative survival rates are needed.

Competing interests

The authors declare that they have no competing interests. This study was completely self-financed and no funding was sought or obtained, not even in the form of free materials.
References


