Dental implant treatment for two adjacent missing teeth in the maxillary aesthetic zone: a comparative pilot study and test of principle

Dental implants are increasingly being applied in the aesthetic zone; therefore, it is essential to be able to establish a predictable aesthetic result. According to the professionals’ opinion, dental implant crowns in the aesthetic zone are successful if a harmonious anatomical outcome has been established with the right dimensions of white and pink structures (Belser et al. 2004; Meijndert et al. 2007). On the other hand, regeneration of a soft-tissue contour with intact inter-proximal papillae and a gingival outline that is harmonious with the gingival silhouette of the adjacent teeth appears to be one of the major challenges (Den Hartog et al. 2008). In case of a single-tooth replacement, the presence of inter-proximal papillae is determined predominantly by the attachment level of the neighbouring teeth (Kan et al. 2003; Gündüz et al. 2005; Kourkouta et al. 2009), which favours the aesthetic outcome of single-tooth replacements in case of periodontally unaffected neighbouring teeth. However, the advantage of having neighbouring teeth on both sides of a single-tooth replacement is not present if two adjacent teeth are missing. As a result, the presence of a papilla between two implant crowns is predominantly dictated by the highest bone level between the implants (Kourkouta et al. 2009). Inter-implant distance appears to be another important factor in the preservation of bone height between two adjacent implants and should be at least 3 mm. In case of an inter-implant distance of < 3 mm, a loss of crestal bone height can be expected. This is caused by the lateral component of the peri-implant bone loss around implants. Overlap of both resorption areas between the adjacent implants will eventually result in vertical reduction of the inter-implant bone crest level (Tamow et al. 2000; Castaldo et al. 2004; Kourkouta et al. 2009). The reduced papilla height between two adjacent implants in comparison with single-tooth replacement complicates the aesthetic outcome. Only a maximum of 3 mm of inter-implant soft-tissue height should be expected instead of 3–5 mm of soft-tissue height between an implant and a natural tooth (Castaldo et al. 2004). To avoid black triangles and ensure that the distance between the contact point and the inter-implant bone

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crest is fully filled with soft tissue, the contact point of the two adjacent implant crowns should be positioned more apically to obtain a longer contact area. This technique is often used in case of compromised papilla presence, but impedes the manufacture of harmonious anatomically shaped crowns.

Therefore, the purpose of rehabilitation in the aesthetic zone should be to maintain bone around implants at an ideal level. Vertical and lateral bone loss around most implant systems at the interface of implant and abutment is up to 1.5 mm and is due to chronic irritation from bacteria products out of this interface (Hermann et al. 1997; Tarnow et al. 2000; Cardaropoli et al. 2006). This means that bone around implants must be at least 1.5 mm wide at the apical sides to ensure that the level of bone crest and thus the level of soft tissue remain stable. Sometimes, when the two missing adjacent teeth are an upper central incisor and a lateral incisor, there is lack of space to create enough distance between the implants and between the implants and their neighbouring teeth. Also, utilization of a smaller diameter implant in the region of the lateral incisor does not solve this problem. It is suggested that platform-switched implants, with less widespread lateral resorption, could have an effect (Rodríguez-Ciurana et al. 2009). Another option could be to place only one implant in the region of the central incisor and a prosthetic restoration consisting of an implant crown on this implant connected with a cantilever at the position of the lateral incisor. In this option, bone crest height is not affected by the lateral resorption of the adjacent implant. This option has not been evaluated so far in the literature. Therefore, the purpose of this prospective comparative pilot study was to evaluate hard and soft peri-implant tissue levels of patients.

The surgical and prosthetic procedures of the future restoration in the ideal prospective position were performed. For the simultaneous augmentation procedures, an autogenous bone graft, collected during drilling or harvested from transparent acrylic resin (Vertex Castapress; Wolhusen, Switzerland) and overlaid with a Bio-Gide membrane [Bio-Oss®, Geistlich, Wolhusen, Switzerland] and overlaid with a Bio-Gide® GB membrane [Bio-Gide®, Geistlich]. The wound was closed primarily with sutures [Ethilon 5–0, Johnson & Johnson Health Care, Piscataway, NJ, USA]. For pain control, 600 mg ibuprofen [Bruden Bruis 600; Abbott BV, Hooldorp, the Netherlands] was prescribed, to be taken three times daily if needed. Following surgery, a temporary removable partial denture was adjusted to not exert pressure on the wound. Two weeks after implant surgery, the sutures were removed. Three months after implant

Methods

Patient selection

The patients selected for this study were referred to the Department of Oral and Maxillofacial Surgery [University Medical Center Groningen, University of Groningen, Groningen, the Netherlands] for implant therapy. To be included in this study, patients had to present with two missing adjacent teeth, a central and a lateral maxillary incisor. All patients had to be 18 years or older and were included in the study only after providing informed consent. The study was approved by the hospital medical ethical committee and written informed consent was obtained from all patients. Patients were selected on the basis of the following inclusion criteria:

- sufficient mesio-distal, bucco-lingual and interocclusal space available for the placement of two implant-retained restorations with the right anatomical design;
- sufficient bone available for the placement of two dental implants with a minimum inter-implant distance of at least 3 mm and a minimum tooth–implant distance of 1.5 mm [if required, a bone augmentation procedure was performed at least 4 months before implant placement] and
- implant site free from infection.

Exclusion criteria for this study were as follows:

- presence of medical and general contraindications for the surgical procedures;
- presence of active and uncontrolled periodontal disease;
- bruxism;
- smoking;
- history of previous dental implant therapy in the same region and
- history of local radiotherapy to the head and neck region.

The study population was divided into two groups:

1. “Implant–cantilever group”: Five patients to treat with one dental implant in the region of the central incisor (NobelReplace Groovy Regular Platform, Nobel Biocare AB, Göteborg, Sweden); prosthetic restoration will consist of an implant crown connected with a cantilever at the position of the lateral incisor.

2. “Implant–implant group”: Five patients to treat with two adjacent dental implants [NobelReplace Groovy Regular Platform at the position of the central incisor and NobelReplace Groovy Narrow Platform at the position of the lateral incisor]; prosthetic restoration will consist of two single-tooth implant crowns.

Surgical and prosthetic procedures

All patients were treated at the same department [Oral and Maxillofacial Surgery, University Medical Center Groningen, University of Groningen, Groningen, the Netherlands] by one experienced oral–maxillofacial surgeon and two experienced prosthodontists. Preoperatively, diagnostic casts were made with a diagnostic arrangement representing the future restoration in the ideal prosthetic position. Next, this ideal prosthetic position was transformed into a surgical guide from transparent acrylic resin (Vertex Castapress; Vertex-Dental BV, Zeist, the Netherlands). If it was clear that not enough bone was present to insert an implant with primary stability, a bone augmentation procedure was carried out with bone harvested from the retromolar region in a separate session. One day before implant placement, patients began using a 0.2% chlorhexidine mouthwash [Corysab, GlaxoSmithKline, Utrecht, the Netherlands]. One hour before implant surgery, patients started taking antibiotics (amoxicillin 500 mg, three times daily for 7 days). Under local anaesthesia [Ultracaine D-S Forte, Aventis Pharma Deutschland GmbH, Frankfurt am Main, Germany], the implants were placed, according to the procedure prescribed by the manufacturer, guided by the surgical guide. The implants used in this study were tapered and roughened to the top of the implants with a titaniumoxide surface (TiUnite, Nobel Biocare AB, Göteborg, Sweden). A mucoperiosteal full-thickness flap was raised to provide a clear view of the surgery area. The shoulder of the implants was placed at a depth of 2–3 mm apical to the buccal and cervical aspects of the future clinical crown to allow soft tissue to develop an adequate emergence profile. The implants were placed with an insertion torque of at least 45 N cm. If necessary, the osseous crest was contoured or slightly overcontoured to acquire a bone wall of at least 2 mm on the facial aspect of the implant. Furthermore, if part of the implant surface remained uncovered or if only a thin layer of labial bone was present, a local bone augmentation procedure was performed. For the simultaneous augmentation procedures, an autogenous bone graft, collected during drilling or harvested intra- orally, was combined with Bio-Oss® (Bio-Oss®, spongiosa granules [0.25–1 mm], Geistlich, Wolhusen, Switzerland) and overlaid with a Bio-Guide® GB membrane [Bio-Guide®, Geistlich]. The wound was closed primarily with sutures [Ethilon 5–0, Johnson & Johnson Health Care, Piscataway, NJ, USA]. For pain control, 600 mg ibuprofen [Bruden Bruis 600; Abbott BV, Hooldorp, the Netherlands] was prescribed, to be taken three times daily if needed. Following surgery, a temporary removable partial denture was adjusted to not exert pressure on the wound. Two weeks after implant surgery, the sutures were removed. Three months after implant
placement, the implants were uncovered and a healing abutment [NobelRepace healing abutment, Nobel Biocare AB] was placed.

One week after abutment connection, an open tray impression was made at the implant level using an impression post [Impression Coping Implant Level Open Tray for NobelReplace, Nobel Biocare AB], a custom acrylic resin impression tray [Lightplast base plates; Dreve DenTamid GmbH, Unna, Germany] and a polyether impression material [Impregum Penta; 3M ESPE, St Paul, MN, USA]. In the dental laboratory, a screw-retained provisional restoration was fabricated, consisting of a temporary abutment (NobelReplace temporary abutment Engaging; Nobel Biocare AB) against which a veneering composite (Solidex; Shofu, Inc., Kyoto, Japan) was modelled. In the implant–cantilever group, the lateral incisor was modelled as a cantilever. A metal reinforcement was placed at the palatal side at the connection between the two composite crowns. The plaster cast was prepared in such a way that the lateral incisor could be overcontoured in the region of contact with the mucosa. In this way, the illusion was created that the cantilever crown emerged out of the mucosa. In the implant–implant group, two solitary screw-retained provisional restorations were fabricated. The provisional crowns were contoured so that the peri-implant soft tissue was optimally supported. Extra care was given to the inter-proximal areas: the inter-proximal papillae were provided enough space to regenerate. The cantilever crown was cleared from heavy contacts; only light contact was allowed. With excursion, all contact was avoided. One week after the impression was made, healing abutments were removed and the provisional implant crowns were placed and torqued to 32 N cm. For 3 months, the patients visited the prosthodontist once a month for examination. Three months later (6 months following implant placement), another implant-level impression was made for the fabrication of a definitive restoration. In the dental laboratory, a soft-tissue cast was prepared. First, a waxing of the definitive restoration was made on a temporary abutment (NobelReplace Temporary Abutment Engaging; Nobel Biocare AB). After that, the waxing was cut back to the desired form and scanned for fabrication of custom-made zirconia abutments (Procera, Nobel Biocare AB). If the screw access hole was located at the mid-palatal side, the porcelain was added directly to the abutment to create a screw-retained crown. If the access hole was not located at the mid-palatal side, a custom-made zirconia abutment was fabricated together with a full ceramic cement-retained restoration. Again, in the implant–cantilever group, the lateral incisor was modelled as a cantilever with a zirconia base connected to the centrally located restoration. The cantilever crown was cleared from heavy contacts; only light contact was allowed. With excursion, all contact was avoided. In the implant–implant group, two solitary restorations were fabricated.

Fig. 1. Implant crown with a cantilever as a lateral incisor, dental implant located at the central incisor.

Fig. 2. Intraoral radiograph of dental implant and neighbouring teeth from the implant–cantilever group.
pellet alone. Cement-retained restorations were fastened with Fuji Plus cement (GC, Alsip, IL, USA) [Fig. 1].

Data collection
Data were collected starting pre-operatively (Tpre), directly after implant surgery (Tpost), directly (within a month) after placement of the definitive implant crown (T0) and 1 year after placement of the definitive restoration (T1).

The following parameters were assessed:

- implant loss during the entire evaluation period;
- pocket probing depth at Tpre (only neighbouring teeth), T0 and T1: the depth was measured to the nearest millimetre at three locations around the implants and the neighbouring teeth [mid-buccally and at both approximal sides];
- papilla index according to Jemt (1997) at T1;
- marginal bone level and bone crest level: the most coronal bone peak of the bone crest and the first bone to tooth level, the vertical distance between line a and the first bone to tooth level, the most coronal bone peak of the implant and the abutment was used as a reference line (line a) from which all distances were measured, (2) the first bone to implant level: the vertical distance between line a and the first bone to implant level, measured at the implant side facing the adjacent implant and at the implant side facing the neighbouring tooth, (3) the bone level of the neighbouring tooth: the vertical distance between line a and the first bone to tooth level, the most coronal bone peak of the inter-implant bone crest and the most coronal bone peak of the bone crest between the implants and their neighbouring teeth and
- a subjective appreciation of the final result of the treatment was assessed using a patient questionnaire modified from the one used by Meijndert et al. [2007]. The questionnaire comprised an overall satisfaction score [range 0–10], two questions concerning the implant-supported restoration and two questions concerning the peri-implant mucosa [possible score 0–4].

Statistical analysis
Because of the setting being a pilot study, statistical analysis has been restricted to means, median and standard deviation.

Results
Mean age in the implant–cantilever group was 33 years (range 20–43) and two males and three females were present in this group. The mean age in the implant–implant group was 28 years (range 18–49), and four males and one female were present in this group. The reason for tooth loss was trauma for all patients in both groups. All 10 patients could be evaluated during the 1-year evaluation period. No implants failed in any group during the 1-year follow-up. Mean and median pocket probing depths are listed in Table 1. These pocket probing values of the implants are in line with each other in the two groups. Pocket probing depths are larger around the implants than around the natural neighbouring teeth. Papilla indices are listed in Table 2. Scores are relatively low, pointing towards a

<table>
<thead>
<tr>
<th>Location</th>
<th>Implant-cantilever</th>
<th>Implant-implant</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Tpre</td>
<td>T0</td>
</tr>
<tr>
<td>Central incisor</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Midbuccally</td>
<td>2.2</td>
<td>2.2</td>
</tr>
<tr>
<td>Proximal side facing adjacent tooth</td>
<td>2.2 (0.84)</td>
<td>2.2 (0.84)</td>
</tr>
<tr>
<td>Midbuccally</td>
<td>3.2</td>
<td>3.2</td>
</tr>
<tr>
<td>Proximal side facing adjacent tooth</td>
<td>3.2 (1.1)</td>
<td>3.2 (1.1)</td>
</tr>
<tr>
<td>No implant/lateral implant</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Proximal side facing adjacent tooth</td>
<td>3.2 (1.64)</td>
<td>3.2 (1.64)</td>
</tr>
<tr>
<td>Midbuccally</td>
<td>2.8</td>
<td>2.8</td>
</tr>
<tr>
<td>Proximal side facing adjacent tooth</td>
<td>2.8 (1.45)</td>
<td>2.8 (1.45)</td>
</tr>
<tr>
<td>Cuspid</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Midbuccally</td>
<td>1.2</td>
<td>1.2</td>
</tr>
<tr>
<td>Proximal side facing adjacent tooth</td>
<td>1.2 (0.71)</td>
<td>1.2 (0.71)</td>
</tr>
</tbody>
</table>

NA, not applicable; Tpre, evaluation visit before implant surgery; T0, evaluation visit directly after placement of definitive restoration; T1, evaluation visit 1 year after placement of definitive restoration.
compromised papilla presence. The frequency distributions of the scores of both groups are more or less the same. The mean marginal bone level, the bone crest level and changes during the evaluation period are listed in Table 3. Marginal bone loss occurs, yielding similar results for both groups. Patients' opinion is listed in Table 4. Patient satisfaction is more or less the same in both groups and is very high, with a mean overall satisfaction score of 8.8 for the implant–cantilever group and 9.2 for the implant–implant group.

**Discussion**

Reporting no implant failures of a study group with solitary implant crowns in the aesthetic region, conventional healing and a follow-up period of at least 1 year is not uncommon. Palmer et al. [1997], Jemt & Lekholm [2003], Cardaropoli et al. [2006] and Zarone et al. [2006] all reported a 100% survival rate. In general, survival rates of implants are very high in this region [Den Hartog et al. 2008]. However, survival rates of implants supporting a crown with a cantilever has not been reported so far. It must be noted that the cantilever crown was cleared from heavy contacts; only light contact was allowed. However, higher forces may have an impact on the cantilever during biting, thus exerting moment forces on the implant.

Mean pocket probing values around the implants of both groups were comparable, the presence of a cantilever and possible moment forces on the implant apparently has no or negligible negative effect on the pocket probing depth. Mean pocket probing depths were larger around the implants than around the natural neighbouring teeth. The observed values and difference between implants and natural teeth is in agreement with other studies [Bragger et al. 1997; Hultin et al. 2000; Meijndert et al. 2008]. This is due to the biological width being different around natural teeth compared with implants.

### Table 2. Frequency distribution of scores of papilla index 1 year after placement of the definitive crown

<table>
<thead>
<tr>
<th>Score</th>
<th>Implant-cantilever</th>
<th>Implant-implant</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Central-implant</td>
<td>Cantilever-cusp</td>
</tr>
<tr>
<td>0</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>1</td>
<td>0</td>
<td>3</td>
</tr>
<tr>
<td>2</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>3</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>4</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

Score 0, no papilla formation; score 1, less than half of the papilla; score 2, at least half of the papilla is present; score 3, papilla fills whole approximate space; score 4, abundance of papilla.

### Table 3. Mean, SD and Median of marginal bone level, bone crest level and changes during the evaluation period in mm

<table>
<thead>
<tr>
<th>Location</th>
<th>Implant-cantilever</th>
<th>Implant-implant</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>T1 Mean (SD) Median</td>
<td>T1 Mean (SD) Median</td>
</tr>
<tr>
<td></td>
<td>Tpost</td>
<td>Tpost–T1</td>
</tr>
<tr>
<td>Central incisor</td>
<td>Marginal bone level facing the adjacent central implant</td>
<td>–1.63 (0.84) –1.3</td>
</tr>
<tr>
<td>Bone crest</td>
<td>Bone crest level between central incisor and central implant</td>
<td>–2.2 (0.56) –2.1</td>
</tr>
<tr>
<td>Central implant</td>
<td>Marginal bone level facing the adjacent central incisor</td>
<td>–0.02 (0.04) 0</td>
</tr>
<tr>
<td>Bone crest</td>
<td>Marginal bone level facing no implant/lateral implant</td>
<td>0.06 (0.14) 0</td>
</tr>
<tr>
<td>Bone crest</td>
<td>Bone crest level between central implant and no implant/lateral implant</td>
<td>–1.19 (0.9) –1.7</td>
</tr>
<tr>
<td>No implant/ lateral implant</td>
<td>Marginal bone level facing the adjacent central implant</td>
<td>–1.32 (1.67) –0.5</td>
</tr>
<tr>
<td>Bone crest</td>
<td>Marginal bone level facing the adjacent cuspid</td>
<td>0.34 (0.46) 0.3</td>
</tr>
<tr>
<td>Cuspids</td>
<td>Marginal bone level facing the adjacent no implant/lateral implant</td>
<td>0.43 (0.48) 0.4</td>
</tr>
</tbody>
</table>

Positive numbers in the column Tpost and T1 mean a bone level apically of the reference line (+ = microgap).

Negative numbers in the column Tpost–T1 mean bone loss during the evaluation period.

Tpost, evaluation visit directly after implant surgery; T1, evaluation visit 1 year after placement of definitive restoration.
of the implants occurred in the period from the placement of the implants to 1 year after placement of the definitive crowns. Marginal bone level was, at placement of the implants, more or less at the level of the top of the implant. This phenomenon of resorption of bone in the vicinity of the microgap has been described as a result of a chronic irritant, such as bacteria, coming from the implant–abutment interface. A resorption of 1.5–2 mm has been reported (Hermann et al. 1997; Tarnow et al. 2000). On the other hand, the mean marginal bone loss at the side of the implants facing the cantilever tended to be slightly larger in comparison with the other approximal implant sides of the implant–cantelever group and the implant–implant group. Mean bone crest resorption distally of the central implant in the implant–cantelever group was comparable with the mean inter-implant bone crest resorption between the central implant and the lateral implant in the implant–implant group. Mean bone crest resorption distally of the central implant in the implant–cantelever group is 1.1 mm. Mean bone crest resorption between the central implant and the lateral implant in the implant–implant group is 1.4 mm. Although the inter-implant distance is more than 3 mm, there could still be an effect of a lateral resorption area. Considerably large standard deviations were observed for mean changes in the marginal bone level and the crestal bone level. Similar observations were reported in other studies (Palmer et al. 2000; Small & Tarnow 2000; Steveling et al. 2001; Karoussis et al. 2003; Tawil & Younan 2003; Meijndert et al. 2008). The large standard deviations suggest a considerable variability in changes in the marginal bone level between individual patients, making a reliable prediction of the expected changes in hard and soft peri-implant tissues for an individual patient rather difficult. Variations in the distance from the contact point to the approximal crestal bone and variations in the level of the marginal approximal bone of the adjacent might be the basis of the variation in individual changes of the approximal peri-implant tissues [Tarnow et al. 1992; Kan et al. 2003; Grunder et al. 2005].

Patients’ opinion was listed in Table 4. Patient satisfaction was very high, without differences between the groups. It appears from the papilla index scores that the presence of papillae, especially between the implant crown and cantilever and the adjacent implant crowns, is compromised. This disagreement has been described before by Meijndert et al. (2007). Also, in this study, patients were less critical than one might expect.

Conclusions
In this 1-year prospective comparative study, no significant differences in hard- and soft-tissue levels were observed between patients with a missing central and lateral upper incisor treated with either one implant and an implant crown with a cantilever or two implants with solitary implant crowns. Thus, a prosthetic solution wherein patients with a missing central and lateral upper incisor are treated with only one implant restored with an implant crown and cantilever could serve as an alternative for treatment with two implants restored with solitary implant crowns.

References

Table 4. Frequency scores of the satisfaction questionnaire about the appearance of the implant crown, the appearance of the mucosa and the mean and SD of the overall appearance of the implant crowns

<table>
<thead>
<tr>
<th>Score</th>
<th>Implant-cantilever group (n = 5)</th>
<th>Implant-implant group (n = 5)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0 1 2 3 4</td>
<td>0 1 2 3 4</td>
</tr>
<tr>
<td>Shape of the crown</td>
<td>0 0 0 1 4</td>
<td>0 1 0 0 4</td>
</tr>
<tr>
<td>Colour of the crown</td>
<td>0 0 0 0 5</td>
<td>0 0 0 1 4</td>
</tr>
<tr>
<td>Shape of the mucosa</td>
<td>0 0 3 0 2</td>
<td>0 1 0 1 3</td>
</tr>
<tr>
<td>Colour of the mucosa</td>
<td>0 1 0 1 3</td>
<td>0 0 0 2 3</td>
</tr>
</tbody>
</table>

Mean (SD)

Overall score (range 0–10) 8.8 (0.8) 9.2 (0.8)

Overall score: scale 0, completely dissatisfied to score 10, completely satisfied.

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References


