Immediate single-tooth implants in the anterior maxilla: 3-year results of a case series on hard and soft tissue response and aesthetics.

Jan Cosyn, Aryan Eghbali, Hugo de Bruyn, Kristiaan Collys, Roberto Cleymaet, Tim de Rouck

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<td>Cosyn, Jan; Ghent University, School of Dental Medicine, Department of Periodontology and Oral Implantology</td>
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CONFLICT OF INTERESTS AND SOURCE OF FUNDING

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ABSTRACT

**Purpose:** The objective of this prospective study was to assess the overall outcome of immediate single implant treatment in the anterior maxilla after a 3-year observation period.

**Material and methods:** Thirty consecutively treated patients with a thick gingival biotype, ideal gingival level/contour and intact socket walls at the time of tooth extraction were treated for single-tooth replacement in the aesthetic zone by two experienced clinicians. Treatment included minimal mucoperiosteal flap elevation, immediate implant placement (NobelReplace TiUnite®, Nobel Biocare, Göteborg, Sweden), insertion of a grafting material between the implant and the socket wall and connection of a screw-retained provisional restoration. The latter was replaced by a cemented crown 6 months thereafter. Patients were clinically and radiographically re-examined after 3 years to assess implant survival, complications and hard and soft tissue conditions. The aesthetic outcome was objectively rated using the Pink Esthetic Score (PES) and White Esthetic Score (WES) by a blinded clinician who had not been involved in the treatment.

**Results:** Twenty-five patients could be re-evaluated after 3 years. One early implant failure had occurred resulting in an implant survival rate of 96 %. Radiographic examination yielded on average 1.13 mm mesial, respectively 0.86 mm distal bone loss. The clinical conditions showed fairly low peri-implant plaque (18 %) and bleeding (24 %) and mean probing depth was 3.17 mm. Mean mesial/distal papilla shrinkage and midfacial soft tissue recession in reference to the pre-operative status accounted for 0.05 mm, 0.08 mm and 0.34 mm, respectively. Between the 1- and 3-year reassessment mesial papillae showed significant re-growth (0.36 mm; p = 0.015). Advanced midfacial recession (> 1 mm) was found in 2/25 (8 %) cases. Five (21 %) cases were aesthetic failures (PES < 8 and/or WES <6) and 5/24 (21 %) showed an (almost) perfect outcome (PES ≥ 12 and WES ≥ 9). The remainder (14/24 or 58 %) demonstrated acceptable aesthetics.

**Conclusions:** The proposed strategy seems a valuable and predictable treatment option for well-selected patients in the mid-long term as shown by almost full papillary re-growth and a low risk for advanced midfacial recession.
CLINICAL RELEVANCE

**Scientific rationale:** Ample studies have been published on immediate single implant treatment. However, no data have been reported on hard and soft tissue response and aesthetic outcome after a 3-year observation period.

**Principal findings:** One early failure occurred and mean bone loss was 1 mm. Papillae showed almost full regrowth and only 2/25 (8%) cases demonstrated advanced midfacial recession (> 1 mm). Nineteen (79%) cases showed acceptable to excellent aesthetics.

**Practical implications:** This strategy seems a valuable and predictable option for well-selected patients in the mid-long term. Main advantages include time gain, immediate aesthetics and comfort.
**Introduction**

Single implant treatment in the anterior maxilla is considered highly predictable and successful, at least in terms of implant survival and hard tissue remodelling following conventional implant surgery (Creugers et al. 2000, Berglundh et al. 2002, den Hartog et al. 2008, Jung et al. 2008). Hitherto, aesthetic aspects of therapy have only been sporadically reported even though these are more and more becoming the key for success in daily practice. The latter may be a reflection of an evolving society with more demanding patients expecting a restoration to be an exact copy of the original tooth and finalized within the shortest possible time span. Consequently, implant protocols have been reassessed ultimately resulting in immediate implant placement and restoration. Albeit this strategy offers obvious advantages such as time gain, immediate aesthetics and comfort, immediate implantation may not avert post-extraction remodelling (Botticelli et al. 2004, Araújo et al. 2005). As such, immediate single implant treatment may be a risky procedure in terms of soft tissue stability especially when patients are improperly selected and surgery is performed by inexperienced clinicians (De Rouck et al. 2008b). To reduce the risk for advanced midfacial soft tissue recession a number of prerequisites have been described (De Rouck et al. 2008b). At least in the short term, immediate implant placement has shown to be predictable under these conditions (Kan et al. 2003a, De Rouck et al. 2008a). The purpose of the present study was to document the overall outcome of immediate single implant treatment in the anterior maxilla in the mid-long term focusing on soft tissue dynamics and aesthetic aspects. The research hypothesis was that this treatment concept would demonstrate a low risk for advanced midfacial recession after 3 years of function.

**Material and Methods**

**Patient selection**

This prospective study was based on data from patients who had been treated for immediate single-tooth implants at the Dental Clinic of the Free University in Brussels (VUB). The 1-year results were previously published by De Rouck and co-workers (2008a). This paper addressed the outcome of the same study sample after 3 years of function focusing on soft tissue dynamics between the 1- and 3-year reassessment and aesthetic aspects. Patients were selected during a screening visit on the basis of inclusion and exclusion criteria.

Inclusion criteria were as follows:

1. At least 18 years old.
2. Good oral hygiene defined as full-mouth plaque score ≤ 25 % (O’Leary et al. 1972).
3. Presence of a single failing tooth in the anterior maxilla (15-25) with both neighbouring teeth present.
4. Ideal soft tissue level/contour at the facial aspect of the failing tooth in perfect harmony with the surrounding teeth.

5. Thick gingival biotype as determined by De Rouck et al. (2009b).

6. Adequate bone height apical to the alveolus of the failing tooth (≥ 5 mm) to ensure primary implant stability of at least 35 Ncm.

Exclusion criteria were as follows:

1. Systemic diseases.
2. Smoking.
5. Presence of active infection (pus, fistula) around the failing tooth.
6. Loss of the buccal bone crest after extraction of the failing tooth.

The study was conducted in accordance with the Helsinki declaration of 1975 as revised in 2000.

Surgical procedure

Following screening, comprehensive clinical and radiographic examination was performed by two experienced clinicians (JC/TDR). All patients consented to the planned treatment strategy. The surgical procedure can be found in detail in a previous paper (De Rouck et al. 2008a). Briefly, patients were advised to start antibiotic (Amoxicillin 500 mg) and analgesic therapy (Ibuprofen 600 mg) one hour pre-operatively and oral disinfection (Corsodyl®, GlaxoSmithKline, Genval, Belgium) was done just prior to surgery. Teeth scheduled for immediate replacement were removed as atraumatically as possible using periotomes after minimal mucoperiosteal flap elevation including the papillae of both adjacent teeth. Immediate implant placement (NobelReplace tapered TiUnite®, Nobel Biocare, Göteborg, Sweden) was performed if the buccal bone crest was intact. Special attention was paid to a correct selection and three-dimensional positioning of the implant as described by Buser et al. (2004). Following confirmation of the primary stability using a Torque Controller (Nobel Biocare, Göteborg, Sweden), implant impression was made. Deproteinized bovine bone particles (Bio-Oss® 0.25 - 1 mm, Geistlich Biomaterials, Wolhusen, Switzerland) soaked in blood were inserted to fill the void between the implant and the alveolus. Finally, an appropriate healing abutment was installed and the wound was closed by means of single sutures (Vicryl® 5/0, Johnson & Johnson, St-Stevens-Woluwe, Belgium). All surgical procedures were performed by one and the same experienced implant surgeon (JC).
Restorative procedures

For details on the restorative and technical procedures, we wish to refer to a previous paper (De Rouck et al. 2008a). Briefly, an individualized screw-retained provisional crown was fabricated in the dental laboratory using the implant impression taken at the time of surgery. An engaging titanium temporary abutment (Nobel Biocare, Göteborg, Sweden) served as a carrier for an appropriate hollowed denture tooth. Approximately 3 h following implant installation the temporary crown was placed and tightened by one and the same experienced prosthodontist (TDR) at 15 Ncm onto the fixture. The provisional restoration was adjusted to clear centric and eccentric contacts in order to avoid full functional load.

After 6 months, the provisional restoration was replaced by a permanent cemented restoration. A standard aesthetic titanium abutment (Esthetic Abutment, Nobel Biocare, Göteborg, Sweden) was used to connect the permanent metal-ceramic restoration. Cementation was performed using temporary cement (Temp-Bond NE®, Kerr, Scafati, Italy). All prosthetic procedures were conducted by one and the same experienced prosthodontist (TDR) and all permanent restorations were fabricated in one and the same dental laboratory (Dental Art, Zottegem, Belgium).

Implant survival and complications

As performed during the first year of function, patients were evaluated for implant survival and complications after 3 years. A distinction was made between biologic and technical complications.

Hard tissue parameters

As performed during the first year of function, a peri-apical radiograph using the long-cone paralleling technique was taken after 3 years. Each X-ray holder (XCP Bite Block®, Dentsply Rinn, Elgin, IL, USA) had been individualized with an occlusal jig (Futar D Fast®, Kettenbach Dental, Eschenburg, Germany) in order to standardize the procedure. Changes in marginal bone levels at the mesial and distal aspect of the implant were determined following digitalization (SprintScan 35 Plus®, Polaroid, Cambridge, MA, USA) and by using the appropriate software (Vixwin 2000 v1.11®, Dentsply Gendex, Lake Zurich, Switzerland). Details can be found in a preceding paper (De Rouck et al. 2008a).

Soft tissue parameters

As performed during the first year of function, the clinical condition of the implant restoration was evaluated after 3 years by means of the following parameters:
1. **Plaque score.** A dichotomous score was given (0= no visible plaque at the soft tissue margin; 1= visible plaque at the soft tissue margin) at four sites per implant (mesial, midfacial, distal, palatal).

2. **Probing depth.** It was measured to nearest 0.5 mm at four sites per implant (mesial, midfacial, distal, palatal) using a manual probe (CP 15 UNC, Hu-Friedy®, Chicago, USA).

3. **Bleeding on probing.** A dichotomous score was given (0= no bleeding; 1= bleeding) at four sites per implant (mesial, midfacial, distal, palatal).

**Soft tissue dimensions** were measured as follows:

1. **Papilla levels.** The levels were recorded by means of an acrylic stent provided with direction grooves. Papilla level (mesial and distal) was defined as the distance from the top of the groove to the top of the papilla measured to the nearest 0.5 mm using a manual probe (CP 15 UNC, Hu-Friedy®, Chicago, USA).

2. **Midfacial mucosa level.** The level of the peri-implant mucosa at the midfacial aspect of the tooth/restoration was measured using the same acrylic stent provided with a central direction groove. The midfacial level was defined as the distance from the top of the groove to the first contact with the peri-implant mucosa measured to the nearest 0.5 mm using a manual probe (CP 15 UNC, Hu-Friedy®, Chicago, USA). **Given the aforementioned research hypothesis, midfacial mucosa level was considered the primary outcome variable of the study.**

All soft tissue dimensions were recorded by two clinicians (JC/TDR). The results on the inter-examiner reliability can be found in an earlier paper (De Rouck et al. 2008a).

**Aesthetic outcome**

All aesthetic evaluations relating to the soft tissues and implant crowns were performed by one clinician (AE) who had not been involved in any treatment. This clinician was calibrated prior to the study on the basis of 20 single implant cases in the anterior maxilla. Per case a frontal and occlusal colour slide was available and each case was scored twice with an interval of one week. The 20 single implant cases were also scored by another clinician (JC). The results on the intra- and inter-examiner reliability can be found in a recent paper (Cosyn et al. 2010).

The **Pink Esthetic Score (PES)** by Fürhauser et al. (2005) was used to evaluate the aesthetic outcome of the peri-implant soft tissues. This index includes 7 variables: mesial papilla, distal papilla, midfacial level, midfacial contour, alveolar process deficiency, soft tissue colour and soft tissue texture. Each parameter is assessed with a 0-1-2 score with 2 being the best and 0 being the worst score. Thus, a maximum score of 14 can be reached.
Papillae are evaluated for completeness; the other variables are assessed by comparison with a reference tooth which is the contralateral tooth for incisor and cuspid replacements and the neighbouring premolar for premolar replacements. The authors set the threshold for clinical acceptance arbitrarily at 8/14. A score of 12/14 or more was considered (almost) perfect.

The *White Esthetic Score* (WES) by Belser et al. (2009) was used to evaluate the aesthetic outcome of the visible part of the implant restoration. This index includes 5 variables: tooth form, tooth volume, tooth colour including the assessment of hue and value, tooth texture and translucency. Again, each parameter is assessed with a 0-1-2 score with 2 being the best and 0 being the worst score. Thus, a maximum score of 10 can be reached. All variables are assessed by comparison with a reference tooth which is the contralateral tooth for incisor and cuspid replacements and the neighbouring premolar for premolar replacements. The authors set the threshold for clinical acceptance arbitrarily at 6/10. A score of 9/10 or more was considered (almost) perfect.

The *overall aesthetic outcome* was assessed by combining the results of the PES and WES. If $\text{PES} \geq 12$ and $\text{WES} \geq 9$, the treatment was considered (almost) perfect. If $\text{PES} < 8$ and/or $\text{WES} < 6$, the result was considered a failure.

**Statistical analysis**

Data analysis was performed using the patient as the experimental unit. For all parameters mean values were calculated, if applicable. Descriptive statistics also included frequency distributions for papillae and midfacial mucosa level. The changes between the 1- and 3-year reassessment were examined using the Wilcoxon signed ranks test. The level of significance was set at 0.05.

**Results**

From the 32 patients who had been scheduled from May 2005 to June 2006, 30 (14 men, 16 women; mean age of 54 with a range from 24 - 76) were treated for single-tooth replacement in the aesthetic zone by means of an immediate implant (Nobelreplace tapered TiUnite®, Nobel Biocare, Göteborg, Sweden). Two patients had to be excluded during surgery as loss of the buccal bone crest had occurred after tooth extraction. For details on the reasons for tooth loss, implant locations and dimensions, we wish to refer to an earlier paper (De Rouck et al. 2008a). During the 3-year observation period, 1 and 3 patients were lost to follow-up after 3 and 12 months, respectively. One of them moved and 1 could not be contacted even after several attempts. The other 2 patients agreed to come in for evaluation but did not show up in the end.
Implant survival and complications

At one month follow-up, one of the implants was found to be mobile causing pain and discomfort (tooth location 21; diameter 5 mm - length 16 mm). Besides this one early failure, all implants remained well-integrated resulting in a 96 % implant survival rate after 3 years of function. With respect to complications, one permanent crown lost retention at 8 months of follow-up and was re-cemented. There were no other technical, nor biologic complications.

Hard tissue parameters

Table 1 shows mean bone loss from baseline (implant installation) at the 1- and 3-year reassessment. Bone loss significantly increased between these intervals ($p \leq 0.038$). After 3 years of function radiographic examination yielded on average 1.13 mm mesial, respectively 0.86 mm distal bone loss. Hence, the mean overall bone loss was 1.00 mm.

(HERE APPROXIMATELY TABLE 1 PLEASE)

Soft tissue parameters

In table 2 the clinical conditions of the implant restorations are shown. Throughout the study period, mean plaque levels remained low (< 20 %) indicating good oral hygiene. Between 1 and 3 years, a significant reduction in probing depth from 3.46 mm to 3.17 mm ($p = 0.015$) occurred coinciding with a significant bleeding on probing drop from 41 % to 24 % ($p = 0.002$).

(HERE APPROXIMATELY TABLE 2 PLEASE)

Table 3 depicts the dimensional changes of the soft tissue outline around the implant restorations in relation to the status prior to tooth extraction. Mesial papillae showed a significant re-growth between 1 and 3 years ($p = 0.015$) pointing to a mean loss of only 0.05 mm from the pre-operative status at study termination. A similar trend was found for distal papillae ($p = 0.117$) resulting in a final mean loss of 0.08 mm. At 3 years follow-up severe mesial papilla loss (> 1 mm) was found in 1/25 (4 %) and severe distal papilla loss (> 1 mm) in 4/25 (16 %) cases. In 13/25 (52 %) patients mesial papillae regained at least their original height. In 14/25 (56 %) patients distal papillae regained at least their original height.
The midfacial mucosa level did not alter significantly between the 1- and 3-year reassessment ($p = 0.135$). At study termination a mean recession from the pre-operative status of 0.34 mm was found (table 3). At 3 years follow-up advanced midfacial recession (> 1 mm) was found in 2/25 (8 %) cases. In 10/25 (40 %) patients the midfacial mucosa regained at least its original level.

A case is shown in figure 1 illustrating papillary re-growth and stability of the midfacial mucosa level between the 1- and 3-year reassessment.

(HERE APPROXIMATELY TABLE 3, FIGURE 1 PLEASE)

**Aesthetic outcome**

Table 4 shows the results of all 7 criteria of the PES. Midfacial level and soft tissue colour were most satisfying showing a perfect match with the corresponding tooth in 17/25 (70 %) cases. Unfavourable results were most prevalent for the alveolar process showing severe deficiency in 5/25 (20 %) cases.

The mean PES was 10.48 (SD 2.47; range 5 - 14). Figure 2 shows the cumulative percent of the PES. Dotted lines indicate the upper limit for an unacceptable result (PES = 7) and a favourable, yet imperfect result (PES = 11). Four out of 25 (16 %) cases showed an unfavourable outcome and 9/25 (36 %) an (almost) perfect result.

(HERE APPROXIMATELY TABLE 4 & FIGURE 2 PLEASE)

Table 4 shows the results of all 5 criteria of the WES. Since 1 patient refused to replace the provisional restoration, the results on the WES were based on 24 cases.

Tooth texture was most satisfying indicating an ideal result in 21/24 (88 %) cases. Unfavourable results were most prevalent for tooth colour with a severe mismatch in 5/24 (21 %) and a perfect result in only 9/24 (38 %) cases.

The mean WES was 8.17 (SD 1.52; range 5 - 10). Figure 3 shows the cumulative percent of the WES. Dotted lines indicate the upper limit for an unacceptable result (PES = 5) and a favourable, yet imperfect result (PES = 8). Two out of 24 (8 %) cases showed an unfavourable outcome and 12/24 (50 %) an (almost) perfect result.

(HERE APPROXIMATELY FIGURE 3 PLEASE)
The overall aesthetic outcome was assessed by combining the results of the PES and WES. Five out of 24 (21%) single implant treatments showed an (almost) perfect result (PES ≥ 12 and WES ≥ 9). An acceptable result (PES: 8 – 11 and WES: 6 – 8) was found for 14/24 (58%) cases. The aesthetic outcome was unfavourable for 5/24 (21%) single implant treatments. Three of them (13%) were considered unfavourable because of a PES < 8, another one (4%) because of a WES < 6. One case (4%) showed a PES < 8 and WES < 6 and could be regarded as a complete aesthetic failure.

Discussion

In the present study 96% of the implants survived and mean bone loss was 1 mm after 3 years of function. These data correspond well with the existing knowledge on survival and bone remodelling of conventionally installed single TiUnite® (Nobel Biocare, Göteborg, Sweden) implants after a comparable observation period (Turkyilmaz et al. 2007, De Bruyn et al. 2009, Eghbali et al. 2010). Hence, the timing of implant placement relative to tooth extraction does not seem to be decisive for implant survival or bone remodelling, which has also been demonstrated by at least 4 controlled clinical studies using various implant systems (Lindeboom et al. 2006, Palatella et al. 2008, Block et al. 2009, Van Kesteren et al. 2010).

As described in a systematic review on single implants in the anterior maxilla (den Hartog et al. 2008), few reports included data on clinical parameters. In our study plaque levels were fairly low and comparable between the 1- and 3-year reassessment. In contrast, probing depth and bleeding on probing significantly reduced within this time frame. This trend had already started in the first year (De Rouck et al. 2008a). Similar observations have been described in other short- (Proussaefs et al. 2002) and long-term studies (Apse et al. 1991).

Even though a number of reports are available on immediate single implant treatment, few prospective studies have been published with data on soft tissue dynamics (Kan et al. 2003a, De Rouck et al. 2008a, Palatella et al. 2008, Block et al. 2009, Van Kesteren et al. 2010, Raes et al. 2011). However, these papers all described short-term results with observations up to 2 years. The results of the present 3-year study showed that papillae had not fully remodelled after 1 year of function. Significant papillary re-growth was observed especially at the mesial aspect and at the 3-year reassessment papillae had basically regained their original height. These findings indicate that the presence of papillae may not be the key issue following immediate single implant treatment providing these were intact at the time of tooth loss. Similar observations have been described following conventional implant surgery (Jemt 1997, Choquet et al. 2001, Kan et al. 2003b, Henriksson & Jemt 2004,
Cardaropoli et al. 2006). In these studies the bone peak at the adjacent tooth was considered the pivotal factor in maintaining papilla height between a single implant and tooth.

An issue that gained a lot of attention in recent studies relates to the midfacial mucosa level following immediate single implant treatment (Lindeboom et al. 2006, Chen et al. 2007, Juodzbalys & Wang 2007, Kan et al. 2007, De Rouck et al. 2008a, Evans & Chen 2008, Palatella et al. 2008, Block et al. 2009, Chen et al. 2009, Redemagni et al. 2009, Van Kesteren et al. 2010, Raes et al. 2011). According to a review article immediate implants show an increased risk for advanced midfacial recession (> 1 mm) (Chen & Buser 2009). However, only 2/25 (8 %) of our cases demonstrated more than 1 mm recession and mean loss was only 0.34 mm after 3 years without deterioration between the 1- and 3-year reassessment. In addition, all the criteria of the PES midfacial level was most satisfying. On the basis of these observations, the risk for advanced midfacial recession was clearly low in our patients, hereby confirming the aforementioned research hypothesis. Interestingly, our findings contrast the conclusion of a review article by Chen & Buser (2009), yet seem to be in agreement with 3 controlled clinical studies describing a low risk for advanced midfacial recession following immediate single implant treatment (Palatella et al. 2008, Block et al. 2009, Van Kesteren et al. 2010). This contradiction is interesting and may be explained by disparities in study design (prospective versus retrospective), recording procedure (using a stent or standardized digital slides with fixed reference points or not), case selection (thin-scalloped biotype cases excluded or not; buccal bone crest intact or not), surgical aspects (surgeon’s experience; implant type; orofacial implant positioning; hard and/or soft tissue grafting or not) and restorative aspects (immediate provisionalisation or not). Given the complexity of this aspect of treatment outcome, a thorough systematic review would be valuable specifically comparing the risk for advanced midfacial recession between immediate and conventional single implant treatment.

Aesthetic aspects relating to the soft tissues and implant crown are more and more becoming the key for success. Although ample reports have been published on immediate implantation and restoration, few have documented the aesthetic outcome using objective criteria (Juodzbalys & Wang 2007, Chen et al. 2009, Raes et al. 2011). Since these papers all described short-term results with observations up to 1 year we believe this 3-year paper may add relevant information. When scrutinizing the results on the PES, 16 % of all cases showed an unfavourable outcome (PES ≤ 7), which is slightly higher when compared to earlier findings in the short term (0 % - 11 %) (Juodzbalys & Wang 2007, Chen et al. 2009, Raes et al. 2011). On the other hand, 36 % of our cases showed an (almost) perfect outcome (PES ≥ 12), which resembles quite well with available data (26 % - 39 %) (Juodzbalys & Wang 2007, Chen et al. 2009, Raes et al. 2011).
To our knowledge only 4 case series have been published documenting the aesthetic characteristics of single implants crowns (Belser et al. 2009, Buser et al. 2009, Cosyn et al. 2010, Raes et al. 2011). In the present study 8% of the cases could be considered failures in this respect (WES ≤ 5), which falls within the range described in the literature (0% - 21%) (Belser et al. 2009, Buser et al. 2009, Cosyn et al. 2010, Raes et al. 2011). In addition, 50% of our cases showed an (almost) perfect result which is in agreement with others (Buser et al. 2009, Cosyn et al. 2010, Raes et al. 2011), yet in contrast with Belser et al. (2009) showing perfection in only 18% of the cases. In this regard it must be emphasized that patients had been sent back to the referring dentist for restorative treatment.

Of particular importance is the overall aesthetic outcome combining the results of the PES and WES. As such, 21% showed perfection (PES ≥ 12 and WES ≥ 9) which is quite modest, yet in agreement with the current knowledge on single implant treatment (7% - 35%) (Belser et al. 2009, Buser et al. 2009, Cosyn et al. 2010, Raes et al. 2011). Similarly, 21% of our cases could be considered aesthetic failures (PES < 8 and/or WES < 6) which also falls within the range of what has been published (5% – 34%) (Meijndert et al. 2007, Belser et al. 2009, Buser et al. 2009, Cosyn et al. 2010, Raes et al. 2011). Clearly, optimal aesthetics seem difficult to achieve and failures are quite prevalent in spite of the fact that patients had been selected on the basis of stringent criteria and treated by experienced clinicians. It seems wise to warn patients for this relatively high risk.

In conclusion, the results of this 3-year prospective study indicate that immediate single implant treatment may be considered a valuable and predictable option in terms of implant survival and hard and soft tissue remodelling. We observed in a 3-year time span nearly full papillary re-growth and a low risk for advanced midfacial recession with 19/24 (79%) cases showing acceptable to excellent aesthetics. In this context, we wish to emphasize the favourable starting point in all cases excluding high-risk patients with a thin-scalloped gingival biotype and/or buccal bone defect. Besides careful case selection, appropriate surgical and restorative procedures and clinical experience are also considered of pivotal importance. More prospective studies monitoring soft tissue dynamics over longer time periods are needed. Since it is currently unclear to what extent treatment modalities influence aesthetics, comparative studies specifically focusing on this aspect of treatment outcome would be valuable.

References


### Table 1. Marginal bone loss in relation to baseline (implant installation) at 1 and 3 years

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<td>Mesial bone loss (mm)</td>
<td>0.98 (0.50) [0.00; 2.10]</td>
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<td>Distal bone loss (mm)</td>
<td>0.78 (0.55) [0.10; 2.10]</td>
<td>0.86 (0.54) [0.30; 2.30]</td>
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n = number of patients

Mean (SD) [range]
Table 2. Clinical conditions at 1 and 3 years

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<tr>
<th>Location</th>
<th>1 year (n = 28)</th>
<th>3 years (n = 25)</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Plaque score (%)</td>
<td>17 (18) [0; 50]</td>
<td>18 (17) [0; 50]</td>
<td>0.668</td>
</tr>
<tr>
<td>Probing depth (mm)</td>
<td>3.46 (0.69) [2.00; 4.70]</td>
<td>3.17 (0.63) [2.00; 4.30]</td>
<td>0.015</td>
</tr>
<tr>
<td>Bleeding on probing (%)</td>
<td>41 (16) [0; 75]</td>
<td>24 (19) [0; 50]</td>
<td>0.002</td>
</tr>
</tbody>
</table>

n = number of patients
Mean (SD) [range]
Table 3. Changes in soft tissue dimensions in relation to the pre-operative status at 1 and 3 years

<table>
<thead>
<tr>
<th>Location</th>
<th>1 year (n = 28)</th>
<th>3 years (n = 25)</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mesial papilla (mm)</td>
<td>-0.41 (0.71) [-2.0; 0.5]</td>
<td>-0.05 (0.83) [-1.5; 1.5]</td>
<td>0.015</td>
</tr>
<tr>
<td>Distal papilla (mm)</td>
<td>-0.31 (0.83) [-2.0; 1.0]</td>
<td>-0.08 (1.24) [-2.5; 2.5]</td>
<td>0.117</td>
</tr>
<tr>
<td>Midfacial mucosa level (mm)</td>
<td>-0.53 (0.76) [-2.0; 0.5]</td>
<td>-0.34 (0.80) [-2.0; 1.0]</td>
<td>0.135</td>
</tr>
</tbody>
</table>

n  = number of patients
Mean (SD) [range]; negative value indicates recession in relation to the pre-operative status
Table 4. Aesthetic outcome at 3 years

<table>
<thead>
<tr>
<th>Parameter</th>
<th>0</th>
<th>1</th>
<th>2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mesial papilla</td>
<td>2</td>
<td>10</td>
<td>13</td>
</tr>
<tr>
<td>Distal papilla</td>
<td>4</td>
<td>8</td>
<td>13</td>
</tr>
<tr>
<td>Midfacial level</td>
<td>2</td>
<td>6</td>
<td>17</td>
</tr>
<tr>
<td>Midfacial contour</td>
<td>2</td>
<td>11</td>
<td>12</td>
</tr>
<tr>
<td>Alveolar process deficiency</td>
<td>5</td>
<td>4</td>
<td>16</td>
</tr>
<tr>
<td>Soft tissue colour</td>
<td>1</td>
<td>7</td>
<td>17</td>
</tr>
<tr>
<td>Soft tissue texture</td>
<td>1</td>
<td>8</td>
<td>16</td>
</tr>
</tbody>
</table>

Pink Esthetic Score (n = 25)  
Mean (SD) [range]  
10.48 (2.47) [5-14]

| Tooth form                      | 0  | 10 | 14 |
| Tooth volume                    | 0  | 5  | 19 |
| Tooth colour                    | 5  | 10 | 9  |
| Tooth texture                   | 0  | 3  | 21 |
| Translucency                    | 1  | 4  | 19 |

White Esthetic Score (n = 24)  
Mean (SD) [range]  
8.17 (1.52) [5-10]
LEGEND FOR FIGURE 1

Fig. 1. One-year and 3-year result of a single-implant case.
Fig. 2. Cumulative percent of the Pink Esthetic Score.
Fig. 3. Cumulative percent of the White Esthetic Score.