Preface

In the past 15 years, the use of osseointegrated implants has become the standard of care for the rehabilitation of fully and partially edentulous patients, leading to a rapid expansion of implant therapy in dental offices.

This positive development was supported by several factors and trends. Firstly, implant therapy is meeting with much greater acceptance—not only by patients, but also by dentists. The excellent documentation of osseointegrated implants in prospective clinical studies (with up to ten years of follow-up) and good clinical results have both contributed to this increased acceptance. Secondly, the prosthetic aspect of implant therapy has been simplified by precise prefabricated components, so general practitioners can easily treat patients with implant-supported restorations. Thirdly, there has been significant progress in bone-augmentation procedures (techniques to overcome local bone deficiencies such as guided bone regeneration or sinus grafting). These surgical procedures are routinely used for implant patients today; they have broadened the indications for oral implant therapy, particularly in partially edentulous patients.

As a result, the single-tooth replacement has become the most common indication for implant therapy in recent years. Parallel to this, “novel techniques,” such as immediate implants (with or without flap elevation) and immediate loading, have been promoted to make implant therapy more patient-friendly. Most of these new techniques, however, have not yet been sufficiently documented clinically. Carefully designed, randomized, controlled clinical studies are required to evaluate their value for daily practice.

With this rapid expansion of implant therapy, involving more than 100,000 clinicians worldwide, quality control in implant dentistry has become an increasing challenge. Universities and scientific associations have been asked to make efforts to assure that the implant therapy provided is of high quality in order to maintain the good reputation of dental implants.

The International Team of Implantology (ITI) has responded by establishing the ITI Education Committee. The main objectives of this committee are to discuss and define the
standards of care in the surgical and prosthetic aspects of implant dentistry, to integrate these standards into high-quality continuing-education courses, and to coordinate the worldwide educational efforts. Over the past eight years, the ITI has significantly increased its efforts in the area of implant education, including the establishment of the ITI Scholarship Program, which offers stipends to young clinicians and financial support for Centers of Implant Dentistry in the U.S., Europe, and Japan. In addition, the ITI organized its third ITI Consensus Conference in 2003 to discuss clinical topics of interest to implant dentistry. The proceedings were published in a special supplement of JOMI (Proceedings of the Third ITI Consensus Conference 2004).

The ITI Education Committee has decided to use these consensus proceedings to establish an ITI Treatment Guide. This guide will offer detailed clinical guidelines for specific problems in implant dentistry. The first volumes will discuss the following topics: (i) esthetic implant dentistry; (ii) loading protocols in implant dentistry; and (iii) implant placement in extraction sockets.

These topics will be comprehensively presented with detailed recommendations for step-by-step procedures. Each treatment option will be discussed objectively, taking into account the following parameters:

1. Scientific documentation of the procedure through clinical studies
2. Objective benefits for the patient
3. Risks involved with the procedure
4. Level of treatment complexity according to the SAC (simple—advanced—complex) classification
5. Cost-effectiveness of the procedure

The first volume of the ITI Treatment Guide is devoted to single-tooth replacements in the esthetic zone, a topic of great interest within implant dentistry. It should be of great help to the clinician dealing with esthetic indications in implant patients.

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A 30-year-old female patient presented at the clinic for a consultation to replace a missing tooth (21) lost in a sport accident. Examination of the esthetic risk for her treatment revealed a medium lip line at full smile, exposing three-quarters of the clinical crowns and the tips of the papillae. The gingival biotype was thin, with highly scalloped papillae, tapered clinical crowns, and a thin band of keratinized tissue. The site analysis showed a deficiency in osseous width and height extending apically of the middle third of the adjacent roots (Fig 9).

The periapical radiograph confirmed this bone loss while showing that adequate bone support existed on the interproximal of the adjacent teeth 11 and 22 (Fig 10).

Based upon the clinical findings, the patient was informed of the high risk of esthetic failure with dental implant therapy. A plan was presented to address the hard-tissue and soft-tissue deficiencies prior to implant placement. Six months after the grafting procedure, a clinical examination revealed successful horizontal bone regeneration, while a deficiency in vertical height remained. The patient declined further tissue-enhancement procedures and desired to proceed with implant placement. A radiographic template was fabricated and a periapical radiograph taken (Fig 11).

The dental implant was placed within the orofacial and mesiodistal comfort zones, while entering the coronoapical danger zone (Fig 12).

Six weeks later, the implant was restored with a provisional restoration; the clinical result highlighted the apical position of the soft-tissue margin at site 21 related to the adjacent teeth (Fig 13).
Upon maturation of the transition zone, there was interproximal tissue support, but a mucosal deficiency on the facial aspect was evident (Fig 14).

The implant shoulder was located in the apical danger zone (Fig 15). To compensate for this situation, pink ceramic material was integrated into the implant-supported superstructure. In this way, a pleasing esthetic treatment outcome could be achieved despite the compromised clinical situation (Fig 16).

The esthetic result was acceptable to the patient (Fig 17).

The postoperative periapical radiograph highlights the final position of the supporting structures (Fig 18).

**Case summary:**
This clinical case is an example of an esthetic compromise associated with clinician and patient causes and anatomical limitations. The challenges in this treatment were associated with several factors: patient limitations—time constraints and finances; surgical challenges—extensive grafting procedures with multiple procedure limits; and anatomic limitations—thin biotype coupled with severe localized atrophy. Proper esthetic risk assessment, patient education, treatment planning, and technician expertise allowed a pleasing esthetic treatment outcome, since the necessary compromises were taken into consideration before the treatment began.