There can be few techniques that have had such a fundamental impact upon restorative dentistry and prosthodontics than osseointegrated implants. It is no great surprise that implants are impacting on the dento-legal front as well.

Not only in terms of problems arising from the provision of implants, but also due to the fact that implants are increasingly being proposed as alternatives to bridgework or dentures as remedial treatment in negligence claims arising from the loss of one or more teeth. This can often drive up the amount of damages claimed by patients, although there is room for doubt that many of the patients receiving these damages ever proceed with the implants that have been proposed for them.

An analysis of the factors that result in negligence claims against dentists relating to implant dentistry (Fig. A) reveals that most of the problems arise from shortfalls in the preliminary stages (ie, patient selection, case assessment, investigations, diagnosis, treatment planning and consent) rather than the treatment itself.

Indeed, many of the problems that result from the procedures themselves can also be traced back to deficiencies in the case assessment and treatment planning stages. Let us now examine some of these issues in more detail.

Preliminary Considerations

Concern that dentists sometimes get involved in implant procedures without having sufficient knowledge, understanding, training and experience to undertake these procedures safely and to an acceptable standard. Furthermore, this same allegation sometimes features in negligence claims as an alleged breach of a dentist’s duty of care. Of particular concern is the short training course that is promoted, organised and conducted by those companies and individuals who have a direct commercial interest in expanding the number of dentists who are carrying out these procedures.

Approaching the Treatment

Patient selection

Not every patient who might seem, at first sight, likely to benefit from implants is going to be a suitable candidate for their provision. A number of risk factors (medical, social and psychological) have been identified in the literature, which have the potential to undermine the prognosis for implant dentistry; all of these need to be carefully considered. The provision of implant-supported restorations may be a last-ditch effort to avert the prospect of becoming edentulous and needing to wear complete dentures. On these occasions it is relevant to look back at the factors that led to the patient being in this situation. These might relate to oral hygiene and patient cooperation, to the patient’s medical history and to a range of other host factors and tissue response generally.

Fig. 1

Fig. A

Fig. B

Problem areas with implants

When do things go wrong?

- Case assessment/investigations/consent
- Unsuitable aesthetics/function
- Implant failure (biological causes or system design)
- Collateral damage
- Other

- Surgical phase
- Joint responsibility
- Restorative phase
- Other issues

28% 9% 14% 27% 22% 34% 36% 3% 27%
Systems selection

The use of an appropriate implant system, with a suitable weight of published research evidence to support its use, is essential. Beware of the “copycat” implant system that adopts some of the principles of various other systems, while having no independent research evidence of its own. The credibility of such a system is easily challenged, and this can raise questions of consent until it has been made entirely clear to the patient, prior to treatment, that the proposed implant system is relatively unproven and/or experimental.

Investigations

This is a critically important stage in the preliminary assessment of any case that involves implants. A detailed assessment of the hard and soft tissue would normally be accompanied by study models, photographs, radiographs and, if appropriate, cephalometric views, CT scans and 3-D reconstructions. It is essential to confirm that any implant fixture can be placed without damage to adjacent structures, and with sufficient bone available. Bone mapping allows a three-dimensional assessment of proposed implant sites to be made, although the quality of the bone in the proposed site may not be fully determined until the time of operation.

Where bone harvesting (or bone grafting) is necessary for a ridge augmentation, or for making the floor of the maxillary sinus, proper consideration must be given to problems that might be encountered at both the donor and recipient sites. These procedures are a frequent source of problems associated with implant dentistry, and they should never be undertaken without extensive training and experience.

Consent

It often becomes clear that a clinician has given little or no consideration to any treatment option except that of placing implants. In some cases this is because the patient has been referred to the clinician in question by the patient’s regular general dental practitioner, specifically for the purpose of assessing the patient’s suitability for implants. In other cases, it appears that the clinician is simply keen to provide implant options rather than to consider any alternative options.

In these cases there is a real risk that patients will be “talked into” implants without going through a detailed consent process. Implants are generally only one of many available treatment options, and each of these options needs to be considered and discussed with the patient in detail. These discussions should include the purpose, nature, risks, benefits, and limitations of each treatment option in turn. Avoid using universal “blanket” information that does not take into account the specific factors that apply to the situation of an individual patient.

Clinicians who are keen to be involved in implant dentistry might be tempted to spend more time explaining the benefits and predictability of implant dentistry—perhaps with the help of persuasive colour brochures provided by implant manufacturers—while spending less time explaining the potential risks and drawbacks. Many negligence claims arise from a failure to understand and manage the patient’s expectations. It is particularly important that the patient should have a clear understanding, at the outset, of the likely appearance and function of the completed restoration, whether this is fixed or removable. By the time the patient is in a position to appreciate the final result at first hand, it is often too late to make fundamental changes to the treatment approach. Given the considerable investment of time, money, inconvenience and discomfort involved in implant procedures, this is likely to result in a very angry and aggrieved patient.

Record keeping

The clinical records (including any available correspondence and documentation) will often be pivotal in determining the outcome of any complaint or negligence claim relating to implant dentistry. The records should comprehensively demonstrate each of the key stages in the provision of implants.

Consultation and preliminary discussions

- Why are implants being considered, and at whose suggestion?
- Medical history
- Dental history
- Social history (including details of the patient’s employment)
- Assessment of risk factors
- Detailed clinical examination (intra- and extra-oral)
- Investigations (see above)
- Diagnosis
- Provisional treatment plan and costing
- Consent (see above)
- Final treatment plan and costing
- Preliminary treatment (eg, preparation and trial placement of stents or other aids)
- Treatment carried out (including preparatory and preventive treatment and advice)
- Outcome
- Any adverse consequences and their management
- Follow-up and maintenance

Collateral damage

Most such damage relates to the surgical phase, which possibly explains why this phase does seem to be responsible for the larger share of the total problems encountered (Fig. B). Damage to the inferior dental nerve or the mental nerve is the problem most commonly encountered in the mandible, although lingual nerve damage and complications involving the maxillary sinus or adjacent natural teeth, are not uncommon in certain situations.

Contractual issues

Because of the cost involved in implant dentistry, the technical nature of the procedures, and their unfamiliarity to most patients, it is essential to explain the proposed treatment and associated costs in advance and in writing. Try to use language that the patient is likely to understand, and avoid technical jargon.

It should be made clear if the fees quoted are an estimate and/or illustration, or a firm quotation of the treatment that is to be provided and the costs involved. If, as can happen in implant dentistry, the treatment plan subsequently changes for any reason, it is prudent to confirm the revised treatment plan and associated costs in writing once again. Many disputes have arisen from a breakdown in communication where such changes have been explained to the patient verbally, perhaps at a time when they were nervous or distracted, and less able to listen and appreciate the information being provided for them.

Details that have created problems in the past include:
- The number of fixtures to be placed.
- The insertion of reserve fixtures (“sleepers”), which are not subsequently used to support the final restoration.
- The number and type of implant components required.

The materials to be used.

The design of the final restoration.

Patients cannot be expected to appreciate fine distinctions and technical details such as these unless the clinician takes the time to explain them. Similarly, the patient does not see treatment that is different in nature or extent to that agreed with the patient can only botch the lie of the breach of contract, as well as of negligence.

Summary

There has been a steady increase in the provision of implants. It appears that they are being placed in more clinical situations, by more clinicians than ever before. Not all of these clinicians can demonstrate that they have received adequate formal training and supervision, and have sufficient technical knowledge and experience, to carry out these procedures safely and successfully. This factor causes great concern for the future. Furthermore, our patients are living longer and the fast-evolving science and technology of implant dentistry is perhaps leading some clinicians into this field who might otherwise have been prepared to refer their patients to the most experienced and skilled colleagues. While this increased clinical ambition is understandable, it is important for dentists to be aware that this is a potentially high-risk field for the inexperienced. In the longer term, the greatest threat to clinicians may well come not from negligence claims, but from the activities of regulatory bodies around the world. These organisations are becoming increasingly intolerant of dentists (and doctors) who show an apparent disregard for their responsibilities to patients and the public. Because of these increasing demands, more clinicians are undertaking procedures for which they are not sufficiently skilled and trained.

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Bone is formed by osteoblasts derived from uncommitted mesenchymal stem cells. After implant surgery the biomaterial surface is populated by these progenitor cells and eventually bone is formed directly along the surface. Astra Tech embarked on a research program to define a method of modifying the TiOblast™ surface to support even more rapid bone formation by the implant surface adherent cell. The discovery that ionic fluoride modification of the TiOblast™ surface improved the bone-to-implant interface resulted in an intensive research program and development of an improved dental implant surface, namely the OsseoSpeed™ surface.

Clinical challenges to osseointegration

Today, there are indications for dental implants that challenge osseointegration's success, including type IV bone, implant placement in extraction sockets, and immediate loading of dental implants. Further improvement in the rate and the amount of bone formation at implants may overcome these clinical challenges. OsseoSpeed™ has the potential to provide these improvements.

Studies confirm greater osteoblast differentiation

One way to examine the role of an implant surface in bone formation is to measure stem cell differentiation to osteoblasts in the cell culture laboratory, as Professor Cooper did at the University of North Carolina. When human mesenchymal stem cells were cultured on TiOblast™ surfaces modified with ionic fluoride, the rate and extent of osteoblast differentiation was greater than on the same surface without fluoride modification. An excellent indicator of osteoblast differentiation is the increased level of Bone Sialoprotein (BSP). Measurement of BSP after 14 days revealed that three times more BSP was made by cells grown on fluoride-modified surfaces than on unmodified ones. This important initial finding was reproduced in three different independent experimental models. These tests were carried out completely 'blind' on the samples sent from Astra Tech in Sweden. "I wanted to know what they were sending me, but I was told that the whole procedure had to be kept completely blind," says Professor Cooper. "They were not going to tell me anything until we had finished."
Positive effects on adherent stem cells

Additional details of the OsseoSpeed™ surface’s effect on the adherent stem cell have recently emerged. For example, human mesenchymal stem cells produced 2.5 times more of the key regulator for osteogenesis (cbfa1) when they were grown for only one day on OsseoSpeed™ compared with TiOblast™. More information from genome-wide microarray analysis of adherent cell behavior indicates that specific signal transduction pathways important to cell proliferation and differentiation are upregulated within the OsseoSpeed™ adherent cell. Microarray analysis comparing cells cultured on the TiOblast™ and fluoride-modified surfaces showed the presence of a number of key ‘enabler’ genes that play an important role in osteogenesis.

‘Human stem cell and molecular research shows there is a bioactive process at work when the OsseoSpeed™ surface is in contact with human bone tissue,’ says Professor Lyndon Cooper, at the Department of Prosthodontics, University of North Carolina, USA.

In vivo tests confirm in vitro findings

Having established in vitro findings, the fluoride-modified TiOblast™ surfaces accelerated the process of osteogenesis, Professor Cooper then conducted in vivo tests to investigate if this would also be reflected in a greater bone-to-implant contact area. The surfaces used in the stem cell research were supplied in implant form and fixed in rat tibiae. The results were consistent with the in vitro findings: after three weeks, it was found that the OsseoSpeed™ surfaces that had stimulated the highest levels of BSP also produced a greater bone-to-implant contact area of contact (55.45% vs 34.21%) at the early 5-week point in time. The parallel cell culture studies suggest this effect is due to signal modification-dependent increases in adherent cell osteogenesis.

Exciting clinical opportunities

Professor Cooper says he is excited about the new opportunities that OsseoSpeed™ can provide for clinicians. The clear conclusion from his work is that the implant surface can be an active component of clinical success. A relatively small, but effective fluoride modification of the TiOblast™ surface is associated with greater osteoblast differentiation of adherent mesenchymal stem cells as well as increased bone-to-implant contact in vivo. The advantages of more rapid and greater bone formation around dental implants may be clinically realized.

Summary

To examine the role of an implant surface in bone formation, Professor Cooper measured stem cell differentiation to osteoblasts in the cell culture laboratory. When human mesenchymal stem cells were cultured on TiOblast™ surfaces modified with ionic fluoride, the rate and extent of osteoblast differentiation was greater than on the same surface without fluoride modification. In fact, measurement of Bone Sialoprotein (BSP) after 14 days revealed that three times more BSP was made by cells grown on fluoride-modified surfaces than on unmodified ones. In addition, microarray analysis comparing cells cultured on the TiOblast™ and fluoride-modified surfaces showed the presence of a number of key ‘enabler’ genes that play an important role in osteogenesis. Results from in vivo studies are consistent with the in vitro findings.

Interviewed:

Professor Lyndon F. Cooper, DDS, PhD, Department of Prosthodontics, University of North Carolina, School of Dentistry, North Carolina, USA

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The truth is out there—looking for the evidence behind bone regeneration

By Anthony Bendkowski oral surgeon and president of the Association of Dental Implantology (UK)

It is widely recognised that one of the key factors contributing to the long term success of dental implants is the quality and quantity of bone around them. When undertaking implant surgery we are frequently faced with defects in the bone that are a consequence of previous underlying periodontal disease, infection or trauma associated with the preceding extractions.

Even a minor bone defect can be a significant barrier where the correct placement of implants is concerned. Without appropriate bone support there is often less than optimal soft tissue profile and consequently compromise in the aesthetic outcome. As clinicians, we are fortunate that bone has a unique potential for regeneration without scarring, provided it is given the correct environment in which to do so.

The vascular and mechanical considerations are important, as is the need for contact with underlying sound living tissue together with passive tension free primary soft tissue closure. The fibrin clot serves as the initial calcifiable matrix containing a concentration of calcium and phosphate ions onto which precursor cells can migrate. This clot needs protection from mechanical stress as any distortion or disruption can profoundly impair the regeneration process.

A further major hindrance to new bone growth is the rapid formation and ingrowth of soft tissue in competition with the slower forming bone regrowth. Fibroblasts have a faster rate of migration than osteoblasts and their proliferation may totally prevent osteogenesis. It is therefore desirable that the soft tissues are excluded from the graft site. This can either be achieved by the use of a barrier membrane, either laid or pinned in place over the underlying particular graft, rather like a tarpaulin covering a mound of sand or by incorporating an additional hard-setting resorbable chemical phase into the graft material, which acts as a barrier to soft tissue downgrowth.

The general term Guided Bone Regeneration (GBR) has been applied to procedures that attempt to regenerate bone, either prior to or at the same time as the placement of dental implants. GBR is accomplished using bone graft materials such as autogenous bone, xenografts, human donor grafts, as well as various synthetic ceramic materials, usually also in association with a biocompatible membrane.

As already mentioned, the use of a membrane ensures that competitive cells do not invade the area where we want new bone to regenerate. The mechanical barrier provided by the membrane offers a means of excluding mucosal tissue and epithelial cells that would interfere with bone healing from the clot. This permits the slower bone producing osteoblasts to facilitate clot organization and produce unimpeded osseous healing.

GBR membrane materials must maintain the integrity of their barrier function long enough to allow osteoblasts to migrate into the wound. Both resorbable and non-resorbable membranes have been used as a GBR barrier. However, non-resorbable membranes such as e-PTFE, although effective, must be surgically removed after the healing period. A resorbable membrane that can transmit tissue fluid, but excludes undesired cells from the clot, has the advantage of not requiring surgical removal. Examples of resorbable membranes include bovine and porcine collagen, PLLA-PGA polymers and calcium phosphate.

The science, application and clinical effectiveness of the various bone graft materials, whether autograft, allograft, xenograft or alloplast, is currently one of the most controversial in implantology and periodontology. Autografts may still be considered by many surgeons to be the graft of choice for specific indications, but there is no current consensus regarding the most appropriate materials for each clinical situation.

There is still much work to be done in this field and we still await the production of good quality randomised controlled trials. Recent years have seen more development of synthetic based bone augmentation materials and a decrease in the prescription of human derived bone substitutes.

The ideal graft material should preferably be gradually resorbed and fully replaced by vital bone that is subsequently remodelled into a natural bone structure that is capable of supporting implants re-softed into the occlusion. If the implanted material does not fully resorb, as is the case with some hydroxyapatites, the mass incorporation is restricted to bone apposition to the material surface, but no substitution occurs during the remodeling phase. This may be desirable for cases where simple ridge preservation or augmentation is required, to help stabilise conventional removable prostheses or improve soft tissue outcomes around conventional pontics, but is not so desirable in implantology.

As clinicians, we have a professional duty to discuss treatment options with our patients and only undertake those procedures which have a reasonable likelihood of a favourable outcome. This is particularly true of elective surgery such as dental implantology and associated bone augmentation procedures.

The gold standard for evidence supporting clinical practice is the randomised controlled trial. Unfortunately in dental implantology, as in many other branches of clinical practice, we are faced with a relative lack of high quality trials of this type. Around the world, researchers are working hard to achieve this gold standard, but much of the important evidence is still to emerge.

Before prescribing materials and treatments for bone augmentation, clinicians need to arrive at their own conclusions from the available information regarding the suitability of these materials and their limitations.

In recent years I have come to the conclusion that it is no small challenge to assimilate this information in an impartial way and free from commercial influence. Discussion with colleagues and reading of the literature only really extends to anecdotal information and small groups of case studies or presentations. For some time I have nurtured a desire to put together a specialist meeting dedicated to reviewing the choice of materials available as bone substitutes.

One of my first initiatives, therefore, on recently taking over the presidency of the Association of Dental Implantology was to organise the forthcoming meeting - Focus on Bone Substitutes - to be held at the Manchester Conference Centre on Monday 28th April 2008.

This full day symposium will examine both the biological basis, as well as the evidence supporting the clinical use of the currently available bone augmentation materials. During the day, seven renowned experts will provide the evidence for both xenografts and synthetic materials. Specifically, the performance of these materials in both alveolar ridge reconstruction and sinus augmentation will be highlighted.

Following this meeting, clinicians should be in a better position to critically evaluate the evidence supporting bone augmentation materials currently available and bone regeneration in day to day clinical practice. The truth is certainly out there to be discovered.
Oral hygiene and dental implants maintenance

By Gregori M. Kurtzman, DDS, MAGD, DICOI and Lee H. Silverstein, DDS, MS

Dentistry has become so exciting and challenging since predictability has been recognized for long-term dental implant and restoration success. As the number of patients selecting dental implants as a treatment option continues to grow, the dental team must accept the challenges of maintaining these sometimes complex restorations.

Proper monitoring and maintenance is essential to ensure the longevity of the dental implant and its associated restoration through a combination of appropriate professional care and effective patient oral hygiene. The value of using conventional periodontal parameters to determine peri-implant health is not clearly evident in the literature. Therefore, it is paramount that the dental implant team understands the similarities and distinctions between the dental implant and the natural tooth. Subsequently, by examining the similarities and differences between a natural tooth and a dental implant, basic guidelines can be provided for maintaining the long-term health of the dental implant.

Direct anchorage of alveolar bone to a dental implant body provides a foundation to support a prosthesis and transmits occlusal forces to the alveolar bone. This is the definition of osseointegration. With the increased acceptance of dental implants as a viable treatment option for the restoration of a partially edentulous or edentulous mouth, the dental team is faced with maintaining and educating those patients.

Recently, the focus of implant dentistry has changed from obtaining osseointegration, which is highly predictable, to the long-term maintenance health of the peri-implant hard and soft tissues. This can be achieved through appropriate professional care, patient cooperation, and effective home care. Patients must accept the responsibility for being co-therapists in maintenance therapy, as the dental team essentially must screen the potential implant patient. Diagnosis and treatment planning based on a risk-benefit analysis should be performed subsequent to a thorough medical, dental, head-and-neck, psychological, temporomandibular disorder and radiographic examination.

There is convincing evidence that bacterial plaque not only...
leads to gingivitis and periodontitis, but also can induce the development of peri-implantitis. Thus, personal oral hygiene must begin at the time of dental implant placement and should be modified using various adjunctive aids for oral hygiene to effectively clean the altered morphology of the peri-implant region before, during, and after implant placement. For instance, interproximal brushes can penetrate up to 5 mm into a gingival sulcus or pocket and may effectively clean the peri-implant sulcus. In addition to mechanical plaque control, daily rinses using 0.1% chlorhexidine gluconate or Listerine provide a welcome adjunct.

Hygiene with dental implants is so tedious and critical to their long-term success that the patient and dental professional must exercise considerable effort. During the maintenance visit, the dental professional should concentrate on the peri-implant tissue margin, implant body, prosthetic abutment to implant collar connection, and the prosthetic crown.

Clinical inspection for signs of inflammation, ie, bleeding on probing, exudate, mobility, probe-able pockets, and a radiographic evaluation of the peri-implant housing still remains the standard mode for evaluating the long-term status of endosseous dental implants. For instance, successful and stable endosseous dental implants exhibit no mobility. But, if there is clinically perceptible mobility, then subsequent to radiographic evaluation of the implant and its surrounding bony housing, the abutment retaining screws, and/or prosthetic abutment collar interface should be examined for looseness or breakage. All these modes of clinical assessment are used routinely, except for periodontal probing around peri-implant tissues that appear to be in a state of good health. The baseline data and data from subsequent recare visits should be recorded in the daily progress notes to properly assess the peri-implant status longitudinally.

Subsequent to a thorough intrasural examination, unless there is visual evidence of soft tissue changes, ie, inflammation of peri-implant tissue with even slight attachment loss or mucositis, routine probing of the peri-implant tissue should not be performed. Usually during the first year subsequent to restoring dental implants, a 3-month recare schedule should be implemented, especially if the patient lost teeth because of periodontal disease. But if after 12 months, the patient’s implants are stable and peri-implant tissues are healthy, then a 4-6 month recare regimen can be implemented. However, cognizant of each patient’s level of home care effectiveness, systemic health, and periodontal status of the peri-implant tissue when determining these recare intervals.

With dental implant patients, the dental professional must evaluate the prosthetic components for plaque, calculus, and the stability of the implant abutment. Radiographs of dental implants should be taken every 12 to 18 months for maintenance visits. For dental implant restorations that are screw retained, the dental professional needs to re-probe the tissue at least once on every 12-monthly interval to more easily assess the status of the peri-implant’s hard and soft tissues, the presence of any symptoms of inflammation, ie, bleeding on probing, inflammation, or radiographic evidence of peri-implant alveolar bone loss. Lastly, routine periodontal probing around dental implants should not be performed, because this procedure could damage the weak epithelial attachment around dental implants, possibly creating a pathway for the ingress of periodontal pathogens. Commercially available plastic probes should be used when investigating the crevicular depth around dental implants. The probing depth around dental implants may be related closely to the thickness and type of bone surrounding the implant. A healthy peri-implant sulcus has been reported to range from 1.5 to 3.5 mm, which is greater than those depths reported for natural teeth. In essence, the best indicator for evaluating an unhealthy site would be probing data gathered longitudinally.

For all of these reasons, personal home care and consistent professional monitoring and treatment are proven to be critical to the success and longevity of endosseous dental implants. This is especially true in an environment with adjacent natural teeth which if affected by periodontal disease, could act as a reservoir for pathogenic bacteria, as gram-negative anaerobic rods, and seed the peri-implant sulcus.

The physical characteristics of the peri-implant soft tissue are the focus of all oral hygiene instruction. The presence or absence of keratinized tissue in this critical area has not been unequivocally documented to state in which probing, inflammation, or probing are more vulnerable to the ingress of pathogenic bacteria with or without radiographic evidence of alveolar bone loss. However, the ability of the patient to maintain good home care around dental implants is facilitated by the presence of keratinized tissue around dental implants. Thus, if a patient has no keratinized tissue around the implant, and a pull from a frenum or a chronic probing peri-implant mucositis exists, then placement of a soft tissue autogenous or alloplastic connective tissue graft is recommended to facilitate the establishment of keratinized gingival tissue.

Specific criteria for obtaining clinical data around dental implants that would allow proper monitoring and detect early possible failure of osseointegrated dental implants has not been clearly defined. Presently, the presence of mobility is the best indicator for diagnosis of implant failure. As opposed to natural teeth, dental implants exhibit minimal clinically undetectable movement because of the absence of a periodontal ligament. Therefore, healthy implants should appear nonmobile, even in the presence of peri-implant bone loss, if an adequate amount of supporting alveolar bone still exists.

When monitoring the health of the peri-implant soft tissues, the practitioner should be cognizant of changes in soft tissue color, contour, and consistency. The presence of a fistulous tract could indicate the presence of a pathologic process or implant fracture.

Bleeding

There is controversy in the literature regarding accuracy and significance of bleeding upon probing around dental implants. Presently, the literature advocates the use of bleeding on probing as an indicator of peri-implant disease. However, use of this indicator to histologic signs of inflammation or concurrently with other signs of implant failure, ie, bone loss. However, in an environment with adjacent natural teeth, routine probing is not recommended.

Radiographic evaluation

Radiographic interpretation is one of the most useful clinical parameters for evaluating the status of an endosseous dental implant. Invasion of biologic width, predictable remodeling, or so-called sauceration, is an average marginal bone loss of 1.5 mm, therefore, progressive bone loss around a dental implant that exceeds these averages may be indicative of an ailing or failing implant. Lastly, during radiographic evaluation, no evidence of a peri-implant radiolucency should be found, because such a radiolucency usually indicates infection or failure to osseointegrate.

Professional cleaning instrumentation

Instruments made of metal, such as stainless steel, should be limited to natural teeth and not to be used to probe or scale dental implants. The rationale for this well-documented and spoken conclusion is that this metal is so hard it can scratch, contaminate,
or cause a galvanic reaction at the implant-abutment interface.

Ideally, hand periodontal scalers for cleaning dental implants can be plastic, Teflon, gold-plated, or made of wood (Fig. 3 and 4). When using gold-plated curettes, the manufacturer recommends not sharpening these hygiene instruments, as the gold surface could be chipped exposing the hard metal underneath this coating. Stainless steel scaling instruments may abrade the implant surface, stripping off any surface treatment such as hydroxyapatite (HA) as the instrument hardness is greater than the titanium alloy the implant is fabricated from. (Fig. 7)

Other cleaning armamentarium contraindicated for use with dental implants are air powder abrasive units, flour or pumice for polishing, and sonic and ultrasonic scaling units. Ultrasonic, piezo or sonic scaler tips may mar the implants surface leading to microroughness and plaque accumulation. The stainless steel tip may also lead to gouging of the implants polished collar. (Fig. 9) However, some clinicians advocate using a sonic instrument with a plastic sleeve over the tip for scaling dental implants. Air powder polishing units may also damage the implant surface and should be avoided during hygiene appointments. (Fig. 9) Even the use of baking soda powder in these units may strip off any surface coating on the implant. Additionally, the air pressure may detach the soft tissue connection with the coronal of the implant leading to emphysema.

Titanium or titanium alloy surfaces of dental implants can be polished using a rubber cup along with a nonabrasive polishing paste or a gauze strip with tin oxide. Not only is the hygiene armamentarium important, but so are the home care techniques used to maintain endosseous dental implants. Patients should be taught the modified bass technique of brushing using a medium-sized head, soft-bristled toothbrush. The use of intradental brushes should be used by implant patients after being shown their proper use. The plastic-coated wire brush is the only type to be used with dental implants to clean and not scratch the implant surface (Fig. 10).

Recently, automated mechanical toothbrushes have been advocated as a daily mode of tooth cleansing. These devices may be a rotary, circular, or sonic type. With these home care instruments, the key to their effective-ness is proper instruction on their use and then diligent daily use by the implant patient.

As with natural dentition, adjunctive cleaning aids such as flossing are still valuable. As with dentated patients, an implant patient’s home care requirements should be individually tailored according to each patient’s needs. Individual needs are based on the location and angulation of the dental implants, the position and length of transmucosal abutments, the type of prosthesis, and the dexterity of each patient.

The other popularized type of cleansing device is the use of oral irrigators with or without the addition of antimicrobial solutions. Also, oral rinses with antimicrobial properties such as Listerine or chlorhexidine have been widely advocated throughout the literature.

Summary

During the infancy years of dental implantology, the emphasis for long-term success of osseointegrated implants was the surgical phase of dental implantology. In the years that followed, the emphasis for success had switched from a purely surgical influence to focusing more on the proper fixture placement which would be dictated by the prosthetic and aesthetic needs of each particular case.

In more recent years, the dental professional has recognized professional implant maintenance and diligent patient home care as two critical factors for the long-term success of dental implants. The microbiota and clinical presentation of peri-implantitis is the same as periodontitis around a natural tooth.

A complete list of references is available from the publisher.
Prevention of peri-implant soft tissue recessions

By Dr. André P. Saadoun, D.D.S., M.S.

Introduction

A beautiful aesthetic result is difficult to obtain with im-
plants in the anterior areas. Both the alignment of the gingi-
val margin and the presence of papillae are essential elements in
resolving aesthetic implant problems to achieve an harmo-
nious smile. These two soft tissue entities, however, are
closely related to the patient’s biotype and to the quality/
quantity of underlying structural alveolar bone.

The peri-implant gingiva, particularly if it is narrow, with a
thin-scalloped biotype, inevitably retracts six months after
the abutment connection and restoration, owing to the refor-
mation of the biologic space (Small and Tarnow, 2000).

The process of soft and hard tissue healing must be under-
stood and incorporated into a carefully coordinated se-
quence of therapy. It is also im-
portant to identify complica-
tions and clinical mistakes and
their implications on the final
esthetic outcome (Saadoun et
al., 1999).

How, then, should soft tissue recession (bone and gingiva)
around an implant be pre-
vented or treated?

Prevention of peri-implant recession

Marginal bone loss of 1 mm in the first year following the
abutment connection, fol-
lowed by loss of 0.2 mm per
year, were among the criteria
defined for implant suc-
cess (Albrektsson et al., 1986). Sav-
ing a few tenths of a millimetre
of bone around an implant does
not increase the longevity of the
implant, and should be done only for aesthetic rea-
s. To prevent or to decrease peri-implant bone resorption
and consequent gingival re-
cession following implant restora-
tions in the anterior zone, several strategies have
been suggested, which are ex-
plained in detail in the follow-
ing points.

1) Implant design and diameter

The design of the collar of the
implant should stabilize the
crestal bone by bringing the roughened surface right up
to the platform, and the
threads/microgrooves as close
as possible to the platform,
with no divergence of the col-
lar walls.

The thread position of the
implant determines the effec-
tive level of remodeling after
loading, and this is perhaps
even more important than the
position of the implant abut-
ment microgap. (Rompen et al.,
2005).

Placement of the implant platform 1.5-2 mm above the
bone, helps to minimize bone
loss as the biological space
around the implants is estab-
lished on the collar (Lezly
Miler, 2003).

2) Implant placement and

extration timing

To make the best choice be-
tween different alternatives of
implant placement, a precise
pre-surgical diagnosis is nec-
essary in order to evaluate the
gingivopalatal parameters, to
determine the optimal mo-
tem to extract the tooth and
place the implant, and to de-
cide whether implant place-
ment and loading should be
immediate, early or delayed
(Saadoun and Landsberg,
1997).

Orthodontic treatment is
the best solution for patients
who wish to limit the surgery
required for the placement of
implants to a single session, and
to enhance the hard and
soft tissue profile prior to ex-
tration and implant place-
ment (Salama et al, 1993).

3) Flap design

On healed site the limited
flap design minimizes inter
proximal bone and papilla-
loss. Many flap design have
been described for healed sites, some raising the total in-
 ter proximal papillae with su-
cular incision around adjacent
 teeth, others using mid-crest/
palatal crest incision with sul-
cular envelope flap and, fi-
nally, tissue punch flap recom-

mended in large amount of
keratinized gingiva.

Flapless approach using tis-
ssue punch procedure has many
advantages: less trauma to the
bone and disturbances to the
soft tissue stability, reduction
of pain and oedema, and less
post surgical information.

Immediate implant place-
ment after extration is usually
a flapless surgical procedure,
the extraction being done us-
ing a periostome to minimize
traumatic damage to the hard
and soft tissues.

4) Tridimensional implant

placement

Satisfactory morphology of
the papilla and of the gingival
margin after anterior implant
restoration depends ultimately
on two factors: implant place-
ment (Esposito et al.,1995,
(Saadoun et al, 1994, Ja-
vanovic, 1999, Grunler et al,
2005) and implant restoration.

The tridimensional criteria
for implant placement in the
esthetic zone are:

• Mesio-Distal: 1.5-2 mm
between implant and adjacent
tooth 3.5-4 mm between im-
plant and adjacent implant
• Bucco-Lingual: 2.5-3 mm
from the cervical height of
corona-alepal to the buccal surface
of the implant platform.
• Corono-Apalical: 2.5-3 mm
apical to the buccal gingival
margin depending on the bio-
type

Therefore, if immediately
post extration implant place-
ment is indicated, the ex-
tration must be performed
against the palatal wall to pre-
vent any damage to the re-
main (and usually thin)
buccal cortical bone (Testori,
2003).

5) Connective osseous grafts

An autogenous bone and
xenograft with a membrane is
used to gain buccal thickness
knowing that bone resorption/
gingival recession always oc-
curs after extraction/implant
placement.

Gingival biotype plays an
important role in determining
tissue levels achieved around
implants. A thin biotype is
generally more susceptible to peri-
implant recession, induced by
the resorption of a thin labial
cortical plate. The use of os-
seous and connective grafts
converts a thin gingival biotype
into a thick gingiva (Mathews,
2000), which can enhance gin-
gival marginal stability and
simply tissue management
during the restorative treat-
ment phase.

6) Abutment and restoration

Optimal aesthetics will be
promoted if the final abutment
is installed at the time of im-
plant placement, and left in
place undisturbed, throughout
the final restoration phase,
avoiding disturbance of bone
and soft tissue architecture.

Fig. 1: Deformed ridge following traumatic extration (right view)

Fig. 2: Deformed ridge following traumatic extration (central view)

Fig. 3: Deformed ridge following traumatic extration (left view)

Fig. 4: Implant insertion after flap elevation

Fig. 5: Bio-Oss graft combined with PRF particu-
lates

Fig. 6: Implant covered with PRF mem-
brane

Fig. 7: Coronally advanced flap (frontal view)

Fig. 8: Coronally advanced flap (left view)
Disconnection and reconnection of the abutment disrupts the biological zone, inducing the junctional epithelium to migrate apically beyond the implant-abutment junction until it can adhere again. This often results in marginal bone loss, particularly in cases of thin gingival biotype.

It is important to minimize the bacterial contamination in and around the implant-abutment junction. The seal provided by an abutment of locking-tapered design has been demonstrated to be optimal in this respect, in vitro (Dibart et al., 2005).

Implant abutments of gold or glazed ceramic should be avoided. Only titanium or zirconium abutments are recommended because hemidesmosomes have been shown to attach to them (Touati and Guez, 2002).

In order to retain soft and hard tissue around the implant-abutment connection, the transmucosal aspect of the implant abutment should not be oversized and divergent, but rather narrow and concave in order to induce thickening and immobilization of the peri-implant tissues, thus increasing the interface between the implant and the soft tissue, and creating an “O-ring connective tissue”. This will ensure the long-term stability of the biological width (Rompen et al., 2007).

Beneath the restoration, the concave abutment should provide maximum space to the soft tissue and clearly avoid a flared geometry. Its submerged profile should be negative to avoid compression of and to allow maximum thickness and stability of the soft tissue, as well as more room for the biological width (Touati 2004). On the buccal aspect, the emergence profile of both the provisional and the final restorations should be flat or concave (under-contoured), to minimise pressure-induced apical migration of the gingival margin.

Design of final crowns to comply with the following “norms” will go a long way toward optimising papillary form (Salama et al., 1999; Elian et al., 2002):  
- Distance from interdental bony crest to contact point between natural crown and implant-borne crown: 4.5 mm
- Distance from inter-implant bony crest to contact point between two implant-borne crowns: 5.5 mm
- Distance between bony crest and connection point between an implant-borne crown and a pontic: 5.5 mm

7) Occlusal trauma
It has been proven that an excessive occlusal load during

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function can cause the loss of peri-implant bone (Misch et al., 2003). The control of horizontal, trans-axial forces on an implant during the first months of function is a determining factor in reducing stress in the crestal zone, in enabling bone adaptation, and in minimizing crestal bone loss (Legall et Saadoun, 2002).

Conclusion

The essential prerequisites for an optimally aesthetic implant restoration should always remain a careful, precise, comprehensive, biologically- and prosthetically-based diagnosis, as well as the choice of the most appropriate implant materials, most conservative, and least traumatic treatment techniques, aimed at conserving, and where necessary augmenting gingival and bone to achieve a successful outcome.

References


About the author

Dr. Saadoun has received his De- gree in Dental Surgery from the Faculty of Paris and completed his Post-Graduate Certificate in Periodontology at the University of Pennsylvania and Post-Graduate Certificate in Implantology at the University of California in Los Angeles.

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He is also on the Editorial Board of Scientific Journals, Practical Procedures and Aesthetic Den- tistry, Implant Dentistry, Dental Implantology Update.

Dr. Saadoun maintains a private practice in Paris, limited to Aes- thetic Periodontics and Implant Surgery.

Fig. 11: Zirconium abutment in place using the guide

Fig. 12: Zirconium abutment with its guide and screw driver

Fig. 13: Zirconium abutment in place using the guide

Fig. 14: Temporary crown 6 weeks later

Fig. 15: Soft tissue aspect after removal of the temporary crown

Fig. 16: Soft tissue aspect around the abutment 6 months later

Fig. 17: Soft tissue aspect around the abutment 6 months later

Fig. 18: Proximal restoration on the day of cemen- tation

Fig. 19: Restoration 3 months later