Introduction

Maintaining the health of teeth and their supporting structures is the goal of modern periodontics. Most periodontal practices focus on prevention of disease, initial therapy and corrective surgical treatment to eliminate deep periodontal pockets. However, restoring supporting tissues to their healthy level is a critical area that offers a much more appealing, and in fact a more desired outcome for the patients. Contemporary periodontal therapy is directed towards controlling the infection and regenerating lost supporting structures. Periodontal regeneration refers to the restoration of supporting tissues of the teeth such as bone, cementum, and periodontal ligament to their original healthy levels before damage from periodontal bacteria has occurred. The key to tissue regeneration is to stimulate a cascade of healing events which, if coordinated, can result in completion of integrated tissue formation. Regeneration of supporting tooth structures is a huge step up in managing advanced periodontal disease and preventing tooth loss. Like other treatment options, it is not a panacea for all patients affected by periodontitis, but research gives us enough evidence to support the use of regenerative therapies in periodontics. Over the last decades different modalities of regenerative treatment have been developed and applied clinically. The positive effects of bone grafts and bone substitutes on the outcome of periodontal regenerative procedures are well documented. Interest in bone replacement grafts has emerged from the desire to fill an intrabony or furcation defect rather than radically resect surrounding intact bone tissue. It is assumed that the application of bone grafts would potentially manipulate the biological response into a regenerative rather than a predominantly reparative pattern of periodontal healing. In addition to bone grafting material, in cases of severe bone loss the use of barrier membranes in regenerative procedures may enhance clinical success by offering better protection and containment of the bone substitute inside the defect and disallowing fast-growing gum tissue to get into the regenerative site and interfere with the process (5). Use of barrier membranes to direct bone regeneration was first described in the context of orthopaedic research by Hurley et al., in 1959 (16). The theoretical principles basic to guided tissue regeneration (GTR) were developed by Melcher in 1976, who outlined the necessity of excluding unwanted cell lines from healing sites to allow growth of desired tissues (19). Based on positive clinical results of regeneration in periodontology research in the 1980’s, research began to focus on the potential for re-building alveolar bone defects using guided bone regeneration. The first application of barrier membranes in the oral cavity was by Nyman, Lindhe, Karring and Gottlow in the context of regeneration of periodontal tissues via GTR, as an alternative to resective surgical procedures to reduce pocket depths (4, 5, 11, 12, 13, 28). Both bone grafts and GTR procedures have become acceptable methods for regenerating lost attachment apparatus, but complete periodontal regeneration is rarely achieved. The recognition and efficacy of bone replacement graft materials used in combination with a barrier membrane in periodontal regeneration piloted to the scripting of this article.

Bone regenerating grafts

The use of bone grafts for reconstructing intra-osseous defects produced by periodontal disease dates back to Hegedus in 1923 (15). It was then revived in 1965 by Nabers and O’Leary (26). Now, with the introduction of advanced bone grafting techniques and the use of sophisticated bone replacement graft materials, it is possible to increase the vo-
lume, width, and height of bone in deficient areas to regen-
erate the tissues supporting affected teeth and also to per-
mit the placement of implants in their ideal positions and angulations.

A bone graft can aid in bone regeneration by three dif-
ferent methods, which include (i) osteogenesis, (ii) osteo-
conduction, and (iii) osteoinduction. Osteogenesis is the for-
mation of new bone by the cells contained within the graft
material. Osteoinduction is a chemical process in which
molecules contained within the graft (bone morphogenetic
proteins) convert the patient’s cells into cells that are capa-
bility of forming bone. Osteoconduction is a physical effect
by which the matrix of the graft forms a scaffold on which
cells in the recipient site are able to form new bone (27).

Classification

Bone replacement grafts can be broadly classified into
human bone and bone substitutes. This can be further clas-
sified into autografts, allografts, xenografts, and alloplasts
(27).

I) Human bone
  - Autografts or autogenous grafts
    - Extraoral
    - Intraoral
  - Allografts or allogenic grafts
    - Fresh frozen bone
    - Freeze-dried bone allografts (FDBA)
    - Demineralized freeze-dried bone allografts (DFDBA)

II) Bone substitutes
  - Xenografts or xenogenic grafts
    - Bovine-derived hydroxyapatite
    - Coralline calcium carbonate
  - Alloplasts or alloplastic grafts
    - Absorbable
    - Nonabsorbable

Historically, autografts were the first bone replacement
grafts to be reported for periodontal applications. Allogenic
freeze-dried bone was introduced to periodontics in the ear-
ly 1970’s, while demineralized allogenic freeze-dried bone
gained wider application in the late 1980’s. The introduction
of xenografts and alloplasts for periodontal use occurred
during the same time (27).

Autografts

Autogenous grafts are harvested from the patient, from
intraoral sites (such as the maxillary tuberosity of a healing
extraction site) and extraoral sites (such as the iliac crests,
ribs, cranium and tibial metaphyses) (8, 31). The decision
to use autogenous grafts necessitates consideration of the
donor site, procurement technique and handling or pro-
cessing of the harvested material.

Autogenous bone can be harvested intraorally, with or
without processing, to yield graft materials of different
forms, including cortical chips, osseous coagulum and bone
blend. Many investigators have reported on the clinically
successful use of intraoral autogenous grafts in the treat-
mant of intrabony defects (2, 17, 23). Regardless of the in-
traoral donor site, autografts yield regenerative responses
superior to that of surgical debridement alone. Extraoral
autografts such as those obtained from iliac crests have de-
monstrated great potential for supporting new bone growth,
including clinical and histological evidence of crestal bone
apposition and periodontal ligament formation. Schall-
horn, Hiatt and Boyce considered the fill of crestal facial
and furcation defects to be more clinically predictable using
iliac autografts than with intraoral cancellous bone (31).

Autogenous grafts are nonimmunogenic and contain
osteoblasts and osteoprogenitor stem cells, which are capa-
ble of proliferating. These grafts, therefore, are osteoin-
ductive. There are limitations to obtaining autogenous
grafts, however, such as insufficient oral sites, the require-
ment for a second surgical site and morbidity at the donor
site (30).

Allografts

Allografts, bone grafts that are harvested from one per-
son for transplantation in another, are used widely. There
are three main divisions: frozen, freeze-dried and freeze-
dried demineralized. The possibility of disease transfer,
antigenicity and the need for extensive cross-matching has
disallowed the use of fresh frozen bone in modern pe-
riodontics. The evidence that freeze-drying markedly redu-
ces the antigenicity and other health risks associated with
fresh frozen bone, as well as the favorable results obtained
in field trials with freeze-dried bone allografts, have led to
the extensive use of freeze-dried bone allografts in the
treatment of periodontal osseous defects (6, 24). The use
of cortical bone is recommended rather than cancellous
bone allografts since cancellous bone is more antigenic
and there is more bone matrix and consequently more os-
teoinductive components in cortical bone. Freeze-dried
bone allograft is regarded as osteoconductive (10). The
blockade of the effect of bone growth stimulating factors se-
questered in bone matrix, like the bone morphogenic pro-
teins, led to the development of demineralized allografts.
Experimental animal studies have shown that deminera-
lized freeze-dried bone allograft has osteogenic potential
(20, 21). The advantages of using allografts are that the
material is available in large quantities and there is no do-
nor site within the patient. The disadvantages are that the
process for preparing the graft (that is, freeze-drying and ir-
radiation) decreases the material’s integrity and osteogenic
potential, and the immunological response to it may dimi-
nish its incorporation into the recipient bone. A major
concern with allografts in general is the potential for di-
sease transfer, particularly viral transmission and more
particularly HIV (22). Also, there is a need for extensive
cross-matching to decrease the likelihood of both graft re-
jection and disease transmission.
**Xenografts**

Xenografts are made of naturally derived deproteinized cancellous bone from another species (such as bovine or porcine bone). The risk of transmission of diseases such as bovine spongiform encephalopathy is negligible because the bone’s organic component is extracted. After the extraction of the organic components, the remaining inorganic structure provides a natural architectural matrix as well as an excellent source of calcium. The inorganic material also maintains the physical dimension of the augmentation during the remodeling phases. Bovine-derived hydroxyapatite bone replacement grafts increase the available surface area that can act as an osteoconductive scaffold due to their porosity and have a mineral content comparable to that of human bone, allowing then to integrate with the host bone. These grafts are prepared by chemical or low-heat extraction of the organic component from the bovine bone. Examples of commercially available bovine-derived bone replacement grafts are Bio-Oss® (Osteohealth Co., Shirley, NY) and Osteografin® (CeraMed Dental, LLC, Lakewood, CO). Coraline calcium carbonate graft is obtained from a natural coral, genus *Porites*. It is hugely porous similar to that of spongy bone and so provides a large surface area for resorption and replacement by bone (14, 34). An example for such type of grafts is Biocoral® (Inotech, Saint Gonnery, France). Biocoral has a high osteoconductive potential because no fibrous encapsulation has been reported.

The main advantages of xenografts are that they are osteoconductive and readily available. A major disadvantage of bovine-derived grafts is due to the fact that it can cause disease transmission, which was evident in the case of bovine spongiform encephalopathy reported in Great Britain (27).

**Alloplasts**

The alloplastic grafts or synthetic bone graft substitutes as yet offer only a part solution to the management of localized bone loss. They possess some of the desired mechanical qualities of bone as well as osteoconductive properties but are largely reliant on viable periosteum/bone for their success. They primarily function as defect fillers. Ideally synthetic bone graft substitutes should be biocompatible, show minimal fibrotic reaction, undergo remodeling, and should have a similar strength and elasticity to that of the bone being replaced, thereby supporting the new bone formation. They do not induce adverse local tissue reactions, immunogenicity or systemic toxicity. They can be classified, by their ability to be bioabsorbed, into absorbable and non-absorbable.

The absorbable materials include alpha and beta tricalcium phosphate, non-sintered hydroxyapatite, and calcium sulfate. The non-absorbable materials include sintered hydroxyapatite, bioglass and HTR™ polymer. Bioceramic alloplasts are comprised mainly of calcium phosphate, with the proportion of calcium and phosphate similar to bone. The two most widely used forms are tricalcium phosphate and hydroxyapatite.

Tricalcium phosphate is a porous form of calcium phosphate. Alpha and beta tricalcium phosphate are produced similarly, although they display different resorption properties. The crystal structure of alpha tricalcium phosphate is monoclinic and consists of columns of cations while the beta tricalcium phosphate has a rhombohedral structure. The former is formed by heating the latter above 1180 °C and quenching in air to retain its structure (9). Alpha form is less stable than beta and forms the stiffer material calcium-deficient hydroxyapatite when mixed with water (32). The most commonly used form is beta tricalcium phosphate. It was one of the earliest calcium compounds to be used as a bone graft substitute. Structurally porous beta tricalcium phosphate has a compressive strength and tensile strength similar to that of cancellous bone. It undergoes resorption over a 6–18 month period. Unfortunately, the replacement of beta tricalcium phosphate by bone does not occur in an equitable way. That is, there is always less bone volume produced than the volume of the graft material resorbed. For this reason, the clinical use of beta tricalcium phosphate has been rather as an adjunctive or other less resorbable bone graft substitutes or as an expander for autogenous bone graft. Examples of commercially available beta tricalcium phosphate graft material are Synthograft™ (Bicon, Boston MA, USA) and Cerasorb® (Curasan Pharma GmbH, Kleinostheim, Germany).

The next calcium phosphate preparation to become available was synthetic hydroxyapatite in the 1970’s. It is available in resorbable and non-resorbable forms. Whether synthetic hydroxyapatite is resorbable or non-resorbable depends on the temperature at which it is prepared. High-temperature preparation (sintering) of hydroxyapatite results in a nonresorbable, nonporous, dense material (18). Dense non-resorbable hydroxyapatite grafts are osteophilic, osteoconductive and act primarily as inert biocompatible bone defect fillers. Histologically, new attachment is not achieved but a more stable clinical improvement is attained than with open flap debridement alone in the treatment of periodontal osseous defects (7, 35). The resorbable form is processed at a low temperature. As it resorbs, a readily available source of calcium becomes available in sites that have osteogenic potential (29). Its reported advantage is the slow resorption rate, allowing it to act as a mineral reservoir and at the same time acting as a scaffold for bone replacement (33). It is marketed in different trade names like Osteogen® (Impladent, NY, USA).

Calcium sulfate or plaster of Paris was first documented as being used for fracture treatment by the Arabs in the 10th century, who would surround the affected limb in a tub of plaster. In 1852 a Dutch army surgeon named Mathysen incorporated plaster into the bandageable form which we are familiar with today (25). Calcium sulfate is thought to act as an osteoconductive matrix for the

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growth of blood vessels and associated fibrogenic and osteogenic cells. For this to occur, it is critically important that the implanted calcium sulfate is adjacent to viable periosteum or endosteum (3). Over a period of 5–7 weeks, calcium sulfate is reabsorbed by a process of dissolution (1). Currently, medical grade calcium sulfate impregnated with tobramycin is commercially available (Osteoset®; Wright Medical Technology, Arlington, TN, USA). Calcium sulfate in its set form has a compressive strength greater than cancellous bone and a tensile strength slightly less than cancellous bone. Calcium sulfate, however, requires a dry environment to set and if it is re-exposed to moisture it tends to soften and fragment. For this reason it has no reliable mechanical properties in vivo and its application should be limited to a contained area. Hence the primary use of calcium sulfoates should be as a bone void filler.

Bioactive glass is a silicate-based, osteoconductive material that binds to bone through the formation of carbonated hydroxyapatite. When exposed to tissue fluids, bioactive glasses are covered by a double layer composed of silica gel and a calcium-phosphorous rich (apatite) layer. The later promotes adsorption and concentration of proteins utilized by osteoblasts to form a mineralized extracellular matrix. It is believed that these bioactive properties guide and promote osteogenesis, allowing rapid formation of bone. Examples of bioactive glasses commercially available are Perioglass® (Block Drug Co., NJ, USA) and Biogran® (Orthovita, PA, USA).

HTR™ synthetic bone (Bioplanet, CT, USA) is a bio-compatible microporous composite of methylmethacrylate and hydroxymethylmethacrylate polymers and calcium hydroxide. HTR stands for hard tissue replacement. Its hydrophilicetchness enhances clotting, and its negative particle surface charge allows adherence to bone. It appears to serve as a scaffold for bone formation when in close contact with alveolar bone. Histological evidence of new bone formation on HTR™ particles has been reported.

Alloplasts can be mixed with autogenous grafts or allografts in the management of large structural defects. Some alloplastic materials are mixed together to achieve superior results. Fortoss® Vital (Biocomposites, Staffordshire, UK) is such a mixture of beta tricalcium phosphate and calcium sulfate. This can be used for guided tissue regeneration without an additional membrane as calcium sulfate serves the purpose of a membrane.

Conclusion

Bone grafting is now a well-recognized choice in the treatment of periodontal osseous defects, especially when used along with barrier membranes. Various types of bone grafts and also their combinations are used with varying degrees of success. Rapid developments in this particular field are leading us towards achieving the ultimate goal in periodontal therapy, which is the regeneration of lost periodontal tissues. Although complete regeneration is now a distant dream, the use of bone grafts enabled us to make slow but sure progress. Autografts are still considered the ideal grafts except for the difficulty in obtaining them. So with the source limitations of autologous bone, the role of bone substitutes will likely increase. The future of bone grafts is likely to lie in the industrially manufactured biomaterials in combination with laboratory-grown cells developed by tissue-engineering.

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