The Merit of Histological Evaluation of
Clinically Successful Reconstructive Therapies in
Periodontology and Related Fields

PhD Thesis

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“The more I learn,
the more I learn how little I know.”

*Socrates*

(469-399 BC)
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1. LIST OF ABBREVIATIONS

3D Three-Dimensional
AAP American Academy of Periodontology
ABB Autogenous Bone Block
ABP Autogenous Bone Particulates
ANOVA Analysis of Variance
AR Alveolar Ridge
ARP Alveolar Ridge Preservation
BCP Biphasic Calcium Phosphate
BS Bone Surface Area
β-TCP Beta-Tricalcium Phosphate
BG Bioactive Glass
BIC Bone-to-Implant Contact
BMP Bone Morphogenetic Protein
BoP Bleeding on Probing
cAMP Cyclic Adenosine Monophosphate
CBCT Cone-Beam Computerised Tomography
CEJ Cementoenamel Junction
CENTRAL The Cochrane Central Register of Controlled Trials
CI Confidence Interval
CONSORT Consolidated Standards of Reporting Trials
DBBM Deproteinized Bovine Bone Mineral
DFDBA Demineralized Freeze-Dried Bone Allograft
EDTA Ethylene Diamine Tetraacetic Acid
EMD Enamel Matrix Derivative
e-PTFE Expanded Polytetrafluoroethylene
FMBS Full Mouth Bleeding Score
FMPS Full Mouth Plaque Score
GBR Guided Bone Regeneration
GS Graft Surface Area
GTR Guided Tissue Regeneration
HA Hydroxyapatite

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<th>Acronym</th>
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<td>IL-6</td>
<td>Interleukin-6</td>
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<td>INTRA</td>
<td>Depth of the Intrabony Component of the Periodontal Defect</td>
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<td>IOPA</td>
<td>Intraoral Periapical Radiograph</td>
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<td>ISRCTN</td>
<td>International Standard Randomised Controlled Trial Number Register</td>
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<td>LILACS</td>
<td>Literatura Latino-Americana e do Caribe em Ciências da Saúde</td>
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<tr>
<td>LJE</td>
<td>Long Junctional Epithelium</td>
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<td>LRAP</td>
<td>Leucine-Rich Amelogenin Peptide</td>
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<td>Nano-HA</td>
<td>Unsintered, Nanocrystalline, Phase-Pure Hydroxyapatite</td>
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<td>OFD</td>
<td>Open Flap Debridement</td>
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<td>Periodontal Ligaments</td>
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<td>RCT</td>
<td>Randomised Controlled Clinical Trial</td>
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<td>REC</td>
<td>Gingival Recession</td>
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<td>rhBMP-2</td>
<td>Recombinant Human Bone Morphogenetic Protein-2</td>
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<td>ROI</td>
<td>Region of Interest</td>
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<tr>
<td>SLA</td>
<td>Sandblasted, Large-grit, Acid-etched</td>
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<td>SLActive</td>
<td>Modified (Hydrophilic) Sandblasted, Large-grit, Acid-etched</td>
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<td>Tricalcium Phosphate</td>
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<td>Transforming Growth Factor-Beta</td>
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2. INTRODUCTION

2.1. Tissue regeneration in periodontology. History and principles

Periodontal disease has been considered as one of the major causes of tooth loss, being accountable for 30-35% of extractions [1]. A widely accepted measure to determine the severity of the disease is the periodontal pocket probing depth (PPD). Advanced periodontitis (PPD>5 mm) has been observed in 2-40% of the population in Europe [1] and 7.4% in Hungary, according to a recent survey [2]. The treatment of periodontitis involves non-surgical as well as surgical means, beyond the inevitable enforcement of individual oral hygiene. The main goal of treatment is alleviating inflammation, arresting disease progression and establishing a stable periodontal condition that is maintainable for long run by the patient’s self-performed oral hygiene.

The principles above that were laid down by the great pioneers of periodontology in Scandinavia and North America remain valid today [3]. Nevertheless, the outcome of the conservative (i.e. non-surgical) periodontal therapy or even of the resective/explorative surgical treatment are often associated with gingival recession and very limited regeneration of the periodontium [4]. Despite the clinically stable periodontal conditions of such cases (i.e. PPD≤4 mm, negative bleeding on probing (BoP)), histological results are mostly characterised by reparative tissues. In such type of healing, the genuine attachment apparatus (acellular root cementum – inserting Sharpey fibres – bundle bone) that was lost as a result of periodontitis, has been predominantly replaced by long junctional epithelium (LJE) instead. Such epithelial attachment merely represents a secondary (inferior) connection in terms of quality and tissue adhesion [5]. Optimal treatment outcome would incorporate the reconstitution of the lost periodontal tissues (i.e. cementum, functionally oriented periodontal ligament, alveolar bone and gingiva). In the Seventies, Melcher suggested that periodontal healing might be characterized by the type of tissues first repopulating the periodontal defect following flap surgery [6]. Based on this assumption, it was hypothesized that precluding the undesired cells (epithelium, gingiva, etc.) from the healing hub of the intrabony periodontal pocket by a mechanical barrier device, would enable the cells from the remaining periodontal ligament to repopulate the defect. Consequently, the
principles of Guided Tissue Regeneration (GTR) and subsequently Guided Bone Regeneration (GBR) were conceived and described in Scandinavian research centres by Jan Gottlow, Thorkild Karring, Jan Lindhe, Sture Nyman, Christer Dahlin and many others [7] [8] [9] [10]. Thenceforth, the theory of GTR and GBR was broadly investigated and proved by sound evidence. However, the “mechanical concept” that the clue of regeneration is purely based on the cell-occlusive barrier function of the membrane is being challenged nowadays. Emerging evidence suggests a rather biological way of thinking that the intent of a membrane might be (i) to protect and stabilize the wound by neutralizing and deflecting any wound rupturing forces away from the root surface, (ii) to provide a secluded space to release the native potential for regeneration and therefore, (iii) to maintain the structural integrity of a maturing blood clot [11] [12] [5].

Over the past decades, tremendous efforts have been made by scientists and clinicians to develop novel techniques and materials in order to improve the outcome of regenerative procedures. Treatment of supra-alveolar defects or intrabony defects with missing buccal and/or lingual wall (i.e. non-containing defects) is often associated with least predictable outcome. For such augmentations a combinative approach was suggested and successfully utilized by means of different ‘bone fillers’ in combination with GTR/GBR [13]. Several grafts, bone substitutes and biomaterials were introduced and widely employed to fill such defects in order to exploit their possible osteoconductive, osteoinductive or even osteogenic potentials. However, the additional benefit of the application of these materials has not been unanimously supported by the literature [14] [15] [16]. As described above, the volume of regenerated tissue may rest on the wound stability and the space provided by the barrier membrane beneath the tension-free approximated mucoperiosteal flaps. Therefore, the main mechanism behind the use of bone replacements seems to be blood clot stabilization through their scaffolding architecture, thus space provision, rather than osteoconduction or induction [17].

The use of barrier membranes, with or without bone substitutes, is, however, frequently associated with side effects, such as membrane exposure. It may arise from the compromised re-vascularization of the flaps above an implanted “foreign body”, as well as from the tension of the flaps caused by the increased volume of the augmentation. In the end of the Nineties a biologic device was developed in Sweden by Lars Heijl, Lars
Hammarström and co-workers that promotes regeneration without the side effects and the handling difficulties of the GTR technique [18] [19] [20]. The biological active component of this regenerative gel originates from amelogenin, an enamel-specific protein of porcine tooth buds. Enamel matrix derivative (EMD), as it is predominantly quoted in the literature, refers to amelogenin fraction following a purification process. EMD in combination with a propylene glycol alginate vehicle results in Emdogain®, the brand name of this biological active device. In the past fifteen years a large number of preclinical and clinical trials have proven its ability to regenerate periodontal ligaments (PDL), cementum and alveolar bone in some extent [21].

However, more than a decade following the first publications of EMD, the biologic cascade of the effects of EMD on wound healing at subcellular level has not yet been fully understood. According to data from in vitro studies EMD may modulate wound healing by stimulating the proliferation of preosteoblasts and the differentiation of osteoblasts via upregulating cyclic adenosine monophosphate (cAMP) levels, inducing the synthesis and secretion of transforming growth factor-beta (TGF-β) and interleukin-6 (IL-6) in gingival fibroblasts and PDL [22] [23] [24] [25]. In addition, EMD may retard epithelial cell proliferation and may confound dental plaque homeostasis [26] [27] [28]. Hence, it seems that EMD may stimulate periodontal wound healing by an indirect effect on the release of growth factors and by retarding the downgrowth of junctional epithelium. A very recent in vitro investigation of amelogenin peptides, conducted by the research group of Nikos Donos at the UCL Eastman Dental Institute, elucidated novel possible pathways of the effect of different fractions of EMD on bone repair and regeneration [29]. It was demonstrated that low- and high-molecular-weight fractions of EMD, namely leucine-rich amelogenin peptide (LRAP) upregulated osteogenic differentiation and enhanced terminal differentiation of bone-forming cells, meanwhile tyrosine-rich amelogenin peptide (TRAP) suppressed the formation of bone-like mineralized nodules. This indicates that fractions of EMD might play a therapeutic role not only in periodontal or alveolar bone regeneration, but also in orthopaedic repair (LRAP), or in the opposite, treatment of pathologic or even malignant bone formation (TRAP). A more recent publication of the same group demonstrated that EMD could modulate the differentiation of PDL cells in vitro, such as up-regulating chondrogenic, neovasculogenic and osteogenic genes, but suppressing adipogenesis, gliogenesis and neurogenesis [30]. A very recent review of the effect of EMD at cellular level concluded
that EMD elicited a regenerative response in periodontal tissues through a complex cascade of gene expression, protein production, proliferation and differentiation of various cells, particularly periodontal ligament and osteoblastic cell types [31].

Our research group at the Semmelweis University, led by Péter Windisch and Anton Sculean, has considerably contributed to the understanding of the effect of EMD on human periodontal defect, by illumination of its histological outcomes [32] [33] [34] [35] [36]. Although the regenerative potential of EMD per se was demonstrated both at preclinical and clinical levels [21] [37], when treating large or non-containing defects, clinicians are often facing the challenge of space provision and prevention of flap collapse. For such cases, bone fillers (in combination with EMD) are suggested to overcome these problems. Data from controlled clinical trials have demonstrated greater clinical improvements following treatment of EMD combined with certain types of bone grafts, such as autogenous bone particles (ABP), or deproteinized bovine bone mineral (DBBM) compared to EMD alone [33] [38] [39]. Nevertheless, other substitutes, such as demineralized freeze-dried bone allograft (DFDBA), or bioactive glass (BG) did not demonstrate significant clinical benefit in terms of PPD reduction and CAL gain over EMD alone [40] [41]. Moreover, the histological evidence seems to be even more conflicting, since beside the formation of new cementum, periodontal ligament and alveolar bone up to a various extent, the implanted bone substitute was frequently encapsulated in connective tissue [34] [35] [42].

It has also been investigated over the decades, whether bone grafts/substitutes alone (i.e. without the additional effect of GTR or EMD) might bear the potential to restore the lost periodontal attachment apparatus. The use of certain types of grafting materials such as ABP, DFDBA or DBBM has been shown to result not only in substantial clinical improvements, evidenced by PPD reduction, defect fill and CAL gain, but also to promote, at least to some extent, formation of a new connective tissue attachment (i.e. new cementum with inserting collagen fibres) and of new alveolar bone [43] [44] [45] [46] [47]. However, the use of autografts for regenerative periodontal therapy is limited by its source and increases donor site morbidity. Furthermore, most types of DBBM possess a fairly slow resorption potential, resulting in a mixed hard tissue formation in periodontal osseous defects [48]. In addition, the use of grafts from bovine (e.g. DBBM) or human origin (e.g. DFDBA), still comport the risk (at least theoretical) for
antigenicity and disease transmission [49] [50]. Furthermore, the animal origin of these materials keeps raising concerns for some patients, due to religious purposes or animal right considerations.

To overcome these drawbacks, synthetic materials in conjunction with open flap debridement were tested. The currently available data suggest that in intrabony defects, the implantation of various types of such alloplasts may lead to significant improvements of the investigated clinical parameters [51] [52] [53] [54] [55] [56]. On the other hand, the available histological evidence indicates that in human intrabony defects, the healing following implantation of alloplasts is predominantly characterized by epithelial proliferation, connective tissue encapsulation and limited periodontal regeneration [57] [58] [59] [53] [60] [61] [62] [63] [64] [65].

The above inconsistent findings on the adjunctive benefit of the application of bone replacement biomaterials highlight the current criteria of successful regeneration. In terms of clinical success, beyond arresting inflammation, we aimed at reconstituting the lost periodontal attachment apparatus. Although we are more aware nowadays that the complete reconstruction may remain a daydream, the therapist must possess certain clinical measures that support the evaluation of the treatment outcome. Such measures are PPD reduction, CAL gain, decreased furcation involvement or sometimes reduced gingival recession, discontinued tooth mobility, etc. Albeit the improvement of all of these measurements undoubtedly indicates a level success, none of them is able to demonstrate the presence of true regeneration (i.e. new cementum, inserting Sharpey-fibres and new bundle/alveolar bone). The reduction of PPD for instance may well be a result of reparative attachment (i.e. LJE) between the root surface and the bone substitute that may not bear long-term stability. In order to display the presence of new bone, standardised intraoral periapical radiograph (IOPA) is commonly used. One should bear in mind though, that based on an radiographic image, we will not be able (i) to distinguish the new bone and the implanted, non/slow resorbable radiopaque bone substitute, (ii) to find out the type of connection to the root cementum and (iii) to ascertain, whether the substitute is connected to the newly formed bone, or encapsulated in connective tissue, which is not regarded as functional bonding. Therefore, these measurements are inappropriate for demonstrating the formation of new cementum, new bone and new periodontal ligaments, merely to unveil the presence of the radiopaque...
biomaterial. Thus, from the biological point of view, it seems reasonable that the selection of bone substitutes for the treatment of periodontal defects should be based on histological evidence, indicating positive biological properties of the biomaterial (with or without GTR technique or in combination with biologically active agent) [66].

In order to supply the clinician with generalizable guidelines, the World Workshop in Periodontics of the American Academy of Periodontology (AAP) summarized the criteria for a periodontal treatment to be considered a regenerative procedure [67]:

- Controlled animal histologic studies demonstrating formation of new cementum, periodontal ligament and bone (in absence of human histologic data retrieved from controlled trials)
- Human histologic specimens demonstrating formation of new cementum, periodontal ligament and bone coronal to the former defect base
- Controlled human clinical trials demonstrating improved clinical probing attachment and bone levels.

In addition to these criteria, a further requirement has been proposed, namely, the regenerative procedure should be based on a concept that explains why the treatment resulted in regeneration [68].

Nowadays, the market of regenerative materials offers an overwhelming number of products, making the right choice for the right clinical scenario extremely difficult. A meticulous analysis of the literature reveals that only a few materials meet the above cited criteria. There are, in fact, several data on clinical and radiographic success, but the histological evidence behind is limited. Furthermore, just a few studies have evaluated human histology and ultimately, there are very limited data on controlled human histological trials on periodontal regeneration, such as the one our group published recently [69] [70].

2.2. Current regenerative considerations of the non-contained periodontal defects

As discussed above it has been proven that EMD alone possesses the biological properties to regenerate PDL and cementum in periodontal intrabony defect with supportive anatomy. However, in a wide, non-contained defect EMD gel cannot provide sufficient support for the flap to prevent its collapse, therefore various combinations of
EMD and different types of scaffolds (bone grafts/substitutes), such as ABP, DFDBA, DBBM, BG, or beta-tricalcium phosphate (β-TCP), have been employed [77] [38] [40] [39] [34] [41].

Synthetic bone substitutes bear the advantage of unlimited source, lack of donor site morbidity, lack of disease transmission and lack of religious concerns. Recently a new composite bone substitute was introduced for periodontal and peri-implant reconstructions. This biphasic calcium phosphate (BCP), namely Straumann BoneCeramic®, consists of 60% hydroxyapatite (HA) and 40% β-TCP in a particulate form. Preclinical evidence suggested that this HA/β-TCP ratio may allow for optimised control of bioabsorbability, thus resulting in accelerated new bone formation [60] [78] [50] [79].

Therefore, the use of a combination of EMD and BCP might be of clinical and biologic relevance in the treatment of advanced, non-contained periodontal defects, since the absorption pace of BCP placeholder might be in line with the regenerative process stimulated by EMD. However, as previously underscored, we cannot declare the possible regenerative features of a novel material or combination of materials, until affirmed by human histology.

To the best of our knowledge, no human histologic data have been available so far evaluating the effect of OFD combined with EMD and BCP on intrabony periodontal defects.

Apart from investigating the above treatment combination (Study 1), we wanted to go one step further to examine, whether a synthetic bone substitute per se, without the regenerative potential of EMD would be able to reconstitute periodontium. It would simplify and accelerate the surgical procedure, as well as reduce treatment cost for the sake of the patient.

Therefore, it would be valuable to seek for an osteoconductive scaffold that is able on its own to (i) function as a space-providing buttress for the flap; (ii) foster blood clot stabilization; (iii) induce proliferation, differentiation and maturation/mineralization of periodontal cells; (iv) resorb totally, in harmony with the formation of new bone.
Ideally, this material is synthetic, does not inflict allergic reaction, inexpensive and originates from an unlimited source.

Studies investigating hydroxyapatite (HA) and/or tricalcium phosphates (TCP) in reconstructive periodontal therapy reported promising clinical, but variable histological results [52]. [58] [59] [54] [60] [62] [63] [65]. Incomplete resorption and implant encapsulation in soft tissue seem to be common drawbacks of the sintered HA. In a recent case report where dense hydroxyapatite material (surface area 59 m²/g) was placed in postextraction socket, the implant particles were observable in the histologic sections even 20 years after implantation [80]. This might be the consequence of the sintering process at high temperature (1200°C). Therefore, the surface structure of the biomaterial could not attract osteoclasts, resulting in an undesirable slow resorption pace [81].

Recently, a new, fully synthetic, unsintered, nanocrystalline, phase-pure hydroxyapatite (nano-HA) has been developed (Ostim®; Heraeus Kulzer, Hanau, Germany). The nanometre sized crystalline structure results in a large, 106 m²/g surface, due to the lack of the high temperature sintering phase [82]. It may be a promising candidate for enhancing periodontal and bone regeneration, since its chemical composition and nanocrystalline structure correspond to the calcium phosphate component of natural bone. Ostim® is supplied in a ready-to-use, sterile capsule, containing the injectable, white, aqueous suspension of 35% nano-HA. According to the manufacturer’s instructions, application of biologic barrier above the implanted nano-HA is not required. Hence, the possibility of incomplete primary closure and flap adaptation, thus membrane exposure and infection is reduced to minimum. In comparison with hydroxyapatite porcelains (sintered HA), nano-HA has greater potential for resorption. The nanocrystals can be ingested and broken down by phagocytosis, which did not appear to increase of the serum calcium level [83] [84]. Depending on the site and volume, it is resorbed within few months [85].

Results of in vitro studies indicate that nano-HA bears the potential to stimulate the migration, adhesion, differentiation and proliferation of PDL cells [86] [87] [88] [89]. In preclinical experiments nano-HA demonstrated accelerated angiogenesis, microvascularisation and new bone formation without inducing inflammatory response.
Rapid resorption was observed via phagocytosis by mononuclear macrophages and multinuclear giant cells [85]. Promising clinical and histological results were reported in the field of orthopaedic trauma and reconstructive surgery following nano-HA implantation [94] [95] [96] [97]. In the field of dentistry, case reports have demonstrated substantial clinical improvements following the use of nano-HA, as a filler of jaw cysts, extraction sockets, sinus and ridge augmentation, periodontal and peri-implant defects [98] [99] [100] [101] [102] [103] [104]. Human histology demonstrated the presence of the alloplast surrounded by woven bone six or even 36 months following implantation in lateral ridge augmentation or sinus floor elevation [101] [103]. Furthermore, the use of nano-HA for the surgical treatment of intrabony periodontal defects demonstrated statistically significantly higher clinical improvements compared to OFD alone [105] [106].

In spite of these promising results, to the best of our knowledge, no data are available from human histological studies on the healing of intrabony defects following periodontal surgery with nano-HA. Thus, at the time being, it is virtually unknown to what extent this material may promote periodontal wound healing/regeneration in humans.

2.3. Rationale of peri-implant dehiscence therapy

Comprehensive restorative dentistry that includes the placement of dental implants is considered as a safe and successful therapy aimed at restoring fully or partially edentulous alveolar ridges [126] [127]. However, sufficient quantity and quality of alveolar bone is required for the longevity of a dental implant [128]. Adequate amount of bone is seldom found prior to implantation due to periodontal disease, periapical pathology and trauma before or even during tooth removal. Moreover, the jawbone undergoes atrophy as part of a natural remodelling after tooth extraction [129] [130] [76] [131]. This resorption requires the restoration of the remaining alveolar ridge [132], in order to meet the contemporary demand of the three-dimensional, prosthetically driven implant placement, as a prerequisite of long term success in function and aesthetics [133].
Several surgical methods have been proposed over the past decades to rebuild the alveolar ridge. These procedures comprise the use of autogenous bone block (ABB), ABP, bone substitutes (allografts, xenografts, alloplasts), GBR alone or with grafting, sinus floor augmentation, forced eruption, as well as ridge expansion techniques utilizing “split” osteotomy or distraction osteogenesis [127].

Among these procedures, GBR has found to be one of the most effective according to the current scientific evidence [127] [134] [135] [136]. Successful bone regeneration has been observed when GBR was used alone or in combination with bone grafts, either prior to placement of dental implants (i.e. two stages procedure) [137] [138], or simultaneously with the placement of implants (i.e. one stage procedure) [139] [140] [141] [142]. In addition, the survival rate of implants placed in the augmented alveolar ridge is comparable to that of implants placed in pristine sites [127] [143].

As per definition, GBR requires the placement of an occlusive barrier that prevents the invasion of non-bone-forming cells from the surrounding soft tissues into the defect. At the same time it allows sufficient time and space for bone forming cells to repopulate the defect [144] [145] [146] [147].

One of the most employed and researched non-resorbable barrier, which had proven to be effective in bone regeneration is the expanded polytetrafluoroethylene membrane (e-PTFE) [148] [149] [150] [151] [152]. However, exposure and inflammation, resulting in soft tissue dehiscence, premature membrane removal, thus compromising bone regeneration, were frequently reported [149] [150] [153] [154] [155].

The main disadvantage of non-resorbable materials is the need for a second surgical procedure to remove the device. This led to the development of bioabsorbable barrier membranes, which did not require a re-entry surgery. Resorbable membranes, such as collagen or glycolide and trimethylene carbonate, have shown improved tissue healing, decreased morbidity, complete resorption and in case of exposure, the risk of bacterial contamination is reduced [156] [157] [158] [112] [159]. On the other hand, some of the resorbable materials may elicit tissue reactions, have uncontrolled resorption rate and show poor resistance to collapse [147] [160] [161].
An ideal membrane is (i) highly biocompatible (does not elicit adverse tissue reactions); (ii) totally resorbable in a predictable rate (reliable maturation of the newly formed tissue beneath); (iii) easy to handle (predictable result even in the hand of less experienced surgeons); (iv) inexpensive and synthetic (available for patients with less financial resources and with concern of animal or human origin).

In order to meet the above demand, a novel synthetic bio-degradable polyethylene glycol hydrogel (PEG) membrane (MembraGel®; Straumann; Basel, Switzerland) was developed recently. PEG membrane is composed of two liquid PEG compounds that react upon mixing and form a hydrogel. PEG has been shown to be highly biocompatible and it is presently approved for several pharmaceutical applications or as medical device [162] [163]. Polyethylene glycol hydrogel degrades by hydrolysis and experimental studies have shown that this process is complete within 4–6 months, therefore a second surgery to remove the membrane is not required [164] [165]. This material, applied as a membrane, has been shown to be cell-occlusive and to be able to prevent soft tissue ingrowth and collapse [165] [166]. Recent experimental studies have also demonstrated positive results in bone regeneration with PEG membranes in bone defects and for the treatment of dehiscence defects around implants [167] [168] [169] [170] [166]. Since its biodegradation is significantly slower compared to standard resorbable collagen membranes, the required barrier function may last longer [164]. This novel barrier material is easy to handle, since the two component of the hydrogel is delivered in an automix syringe, thus the amalgamated membrane gel could simply be placed on the top of the defect or bone graft/substitute. The time of application is shown to be significantly reduced, compared to a conventional collagen membrane, hence the length of the surgery is reduced [169].

In order to stabilize the blood clot and provide space-maintenance below the barrier, the use of a recent developed synthetic biphasic bone substitute seemed to be beneficial. The BCP we used (Straumann BoneCeramic®; Straumann; Basel, Switzerland) had been shown to accelerate bone formation in standard, dehiscence bone defects [78] [79].

There is not much evidence in the literature evaluating the response of regenerated bone to functional loading. It has been perceived that once the protection of the secluded space created by a membrane is removed and the newly formed bone is not functionally loaded, then some bone resorption might take place [171] [172]. The influence of
loading on the outcome of GBR in peri-implant dehiscence defects was investigated in a preclinical study [173]. At the loaded sites significant decrease in bone fill was occurred between the three and nine months healing period (loading), whereas no change was observed at non-loaded sites. On the other hand, the one year data of an ongoing five years RCT conducted by our group, failed to demonstrate significant differences between the loaded (immediately provisionalized) and the non-loaded (healing abutments) groups [174].

When looking into the literature for the desired histological evidences of regenerative procedures in other disciplines in the neighbourhood of periodontology, such as ridge augmentation prior to placement of a dental implant, it has to be realised that alike in periodontology, only a limited number of histological studies could be identified. The explanation is presumably the cumbersome patient and case selection, which must be under any circumstances in coherence with the current ethical guidelines. Furthermore, the concomitant cost of such histological analysis either in preclinical, or in clinical setting, prerequisites a wealthy sponsor or department. Finally, a human histological trial usually encompasses a compensative treatment for the patient that extends the overall treatment time and cost.

If we would like to investigate a regenerative method and material with simultaneous dental implant placement, which is quite a frequent clinical scenario in the daily practice, we could face enormous difficulties with a human histological trial design. For such study, an experimental narrow implant should be placed in an experimental defect that would be regenerated with the experimental biomaterial. Then, it is removed for histological evaluation with some surrounding hard and soft tissues resulting in a much larger bony defect that should be eventually restored with a corresponding size compensatory definitive implant. The chance to obtain a positive ethical authorization or to recruit the sufficient number of patients for such experiment is meagre. Thus, designing such a human histological trial could practically be beyond the bounds of possibilities.
2.4. Provision of post-extraction alveolar sockets

Various methods of alveolar bone augmentation could predictably support the prosthetically driven, three-dimensional implant placement that is a contemporary requirement for long-term success in implant therapy [133]. Simultaneous augmentation procedures can be implemented with high predictability, provided that three intact bone walls are present and the implant location is inside the alveolar envelope [139] [140] [141] [142]. However, in case of extensive ridge resorption, simultaneous bone augmentation becomes less predictable [155]. Two-stages implant placement, which is preceded by GBR, block grafting procedure or ridge expansion techniques, could be implemented with success, although the concomitant treatment time is extended [137] [138] [155]. Moreover, this intervention often comprises intraoral bone harvesting as well as extensive soft tissue manipulation aiming at tensionless flap approximation, which considerably increases morbidity and patient’s intra and postoperative discomfort. Therefore, it is desirable to preserve, rather than reconstruct post extraction ridge dimension, thus minimizing morbidity and discomfort.

The reduction of alveolar ridge (AR) dimension may originate in various reasons. Firstly, periodontal disease, periapical pathology and mechanical trauma could result in bone loss prior to tooth removal [131]. Secondly, indelicate extraction has also been associated with additional bone deficit. Finally, alveolar bone undergoes additional atrophy as a result of natural bone remodeling following tooth removal [129] [130] [197] [198] [199]. This process begins subsequently after extraction and continues for years resulting in even 50% reduction of alveolar ridge width [131]. According to a recent review, the horizontal and vertical components of this resorption may amount to 3.87 and 1.67 mm, respectively [200].

Prevention of alveolar bone resorption would apparently maintain acceptable ridge contour for pontics in areas of aesthetic concern. Nowadays, great emphasis is placed on preserving adequate dimensions of alveolar bone in order to facilitate implant placement in prosthetically driven positions [133] [192] [201]. In order to prevent alveolar ridge atrophy, thereby omit extensive augmentative surgeries, alveolar ridge preservation (ARP) procedures have been introduced. This would reduce treatment time, cost and complexity. Again, such treatment should only be regarded as a valid regenerative
therapy, if human histological results corroborated the possible positive clinical benefits. Several methods have already been investigated for ARP in preclinical models [14] [202] [203] [204] and clinical studies, such as socket grafting with autogenous bone [205], DFDBA [205] [206] [207], xenografts, DBBM [208], alloplasts [209]. GBR with or without bone grafts has also been evaluated [210] [211] [212] [213] [214] [192] [201]. In addition, biologically active molecules, like bone morphogenic proteins (BMP) were also tested [215] and for the improvement of early soft tissue closure socket sealing technique was also recommended [216]. Although some of the above procedures were able to limit the resorption of post-extraction alveolar ridge up to a certain extent based on clinical assessments, the quality of the new tissue in the socket varied broadly. The remnants of the graft materials often interfered with the normal healing process according to preclinical results [205] [206] [207] [217].

Due to the increasing interest in ARP technique, an overwhelming number of original articles, as well as some reviews on ARP were published in the last decade [218] [219] [220] [221] [222] [223]. However, a systematic assessment of the nature and quality of the newly formed tissue alongside evaluation of methodological quality and risk of bias of the studies has not been carried out to the best of our knowledge. Furthermore, non-controlled prospective and retrospective studies, as well as case series and solitaire case reports were also included in most of the previous reviews without the comparison to the control group of unassisted socket healing [224] [225] [226] [227]. Conclusions from such articles might not reflect accurately the available highest evidence.

To summarize the background of the present thesis, we can conclude that several ‘regenerative’ methods and bone substitute materials have been claimed as successful in the field of reconstructive periodontology and implantology over the past decades. However, they do not always appear to promote the formation of new bone, periodontal tissue or functional attachment, according to the only reliable histological evidences. In addition, several data sets seem to be contradictory, indicating that some of these materials may even hinder tissue regeneration. Hence, it would be essential to support the clinician’s daily decision-making with evidence-based methods. Within its limits, the present thesis would like to contribute to this decision-making even with a limited extent, with illuminating the true regenerative properties of some of the available materials and methods.
3. OBJECTIVES

The aim of the studies included the present thesis was to investigate the histological appearances and properties of some representative materials and methods in the field of regenerative periodontology and implant dentistry, which had shown favourable clinical outcomes already. In other words, are the treatments and employed materials that were claimed clinically successful in tissue regeneration really effective in the light of histology?

**Regeneration of periodontal defects**

The aims of the human clinical and histological case series were to evaluate the healing of periodontal intrabony defects following surgical treatment with either a combination of EMD and BCP, or with the nano-HA.

**Regeneration of peri-implant dehiscence defects**

The aim of the preclinical study was to evaluate histologically the effect of a (i) novel BCP (ii) with or without the application of the novel PEG barrier and (iii) the effect of functional loading on buccal peri-implant dehiscence defects.

**Regeneration of post-extraction alveolar sockets**

Our objective was to methodically collect, meticulously scrutinise and systematically evaluate the evidence available in the literature on the effect of ARP on the residual alveolar ridge dimensions and on histological characteristics, compared to unassisted socket healing.
4. METHODS AND MATERIALS

4.1. Overview of the applied methods

In order to answer the above questions we aimed at investigating the histological healing characteristics alongside clinical outcomes of selected materials and treatments used for reconstructive periodontal and implant therapy. Therefore, three scenarios were set up.

Firstly, we have examined clinically and histologically the efficiency of two novel bone substitute biomaterials in periodontal settings. It is based on sound grounds in the literature that the highest proof of the possible regenerative properties of a material (i.e. human histology) could be investigated in human intrabony periodontal defects. Furthermore, over the past years, an accepted protocol has been establish and published for such human histological investigations in our Department of Periodontology at the Semmelweis University. [34] [71] [35] [65] [69] [72] [36] [70]

Regeneration of periodontal defects

STUDY 1
To examine the clinical and histological healing characteristics of EMD combined with a new alloplastic defect filler (Straumann BoneCeramic®) for the treatment of human intrabony periodontal pockets. (Clinical case series) [73]

STUDY 2
To examine the clinical and histological healing characteristics of a new synthetic nanocrystalline bone substitute (Ostim®) that may per se promote healing, in absence of any mechanical barrier, for the treatment of human periodontal intrabony pockets. (Clinical case series)[74]

Secondly, we aimed at assess the regenerative potential of the same bone substitute with a novel hydrogel barrier membrane in the field of reconstructive implant dentistry. Practically, it is not feasible to conduct a human histological study to investigate the performance of these materials in case of human peri-implant dehiscence defect, due to obvious ethical considerations. Therefore, a preclinical protocol was established to
assess the regeneration of a critical size, peri-implant dehiscence defect model on Göttingen minipigs.

Regeneration of peri-implant dehiscence defects

STUDY 3
Clinical and histological evaluation of the effect of a novel biodegradable synthetic hydrogel membrane (Straumann MembraGel®) in combination with a novel biphasic synthetic bone substitute (Straumann BoneCeramic®) on critical size porcine peri-implant dehiscence defects. (Preclinical randomised controlled trial) [75]

Finally, we wanted to investigate whether or not the bone loss, associated with tooth extraction, could be prevented with any material or method, hence to limit the need for peri-implant bone augmentation, with its inherent cost, discomfort and morbidity. The pilot search in the literature resulted in overwhelming number of studies in the field of alveolar ridge preservation. However, a meticulous analysis of these data with structured methodology, especially the assessment the risks of bias and the histological healing characteristics in light of clinical results was lacking. Therefore, instead of conducting yet another human clinical and histological trial, we have decided to perform a systematic review in order to extract the available histological and clinical data on ARP, thus to obtain the highest level of evidence in this field.

Regeneration of post-extraction alveolar sockets

STUDY 4
Clinical and histological evaluation of the effect of alveolar ridge preservation on human extraction sockets. (Systematic review of human controlled trials)[76]
4.2. **Regeneration of periodontal defects (Study 1, 2)**

**Experimental design and subject population**

Both trials were designed as prospective, single arm, human histological study in accordance with the latest amendment of Declaration of Helsinki and were approved by the Regional Ethical Committee of Semmelweis University, Budapest, Hungary. Prior to signing the informed consent form each volunteered, enrolled patient received verbal and written explanations of the research protocol, its purpose, risks and the possibility to withdraw at any time without further consequences. All patients were recruited and treated in the Department of Periodontology, Semmelweis University.

Ten (Study 1) and six (Study 2) patients with advanced chronic periodontitis were enrolled in the trials in 2005 (Study 1) and in 2007 (Study 2). Subjects presented with one advanced intrabony defect around the teeth scheduled for extraction due to advanced destruction of the periodontal attachment apparatus and further prosthetic considerations in conjunction with the overall treatment plan (Fig. 1a, 2a). PPD of at least 6 mm and intrabony component of at least 3 mm were present as visualized on the intraoral radiographs (Fig. 1b, 2b). The selected teeth had some potential for regeneration of lost attachment apparatus as diagnosed clinically and radiographically. In every case, the decision of inclusion was based upon agreement of two clinicians, who were entirely independent of the study.

Furthermore, the following inclusion criteria were set: (i) 20-70 years of age; (ii) completed initial phase of periodontal therapy at least six weeks prior to surgery; (iii) good level of oral hygiene as evidenced by a plaque index <1 [107] (Study 1); (iv) FMPS≤20% [108], FMBS≤20% [109] (Study 2); (v) good compliance to follow up visits and to maintain self-performed oral hygiene; (vi) legal ability to sign informed consent form; (vii) absence of untreated endodontic lesion, hypermobility and occlusal overload.

Exclusion criteria were: (i) general medical history that contraindicates elective surgery and may affect treatment outcome (e.g. uncontrolled diabetes, osteoporosis, immunodeficiency); (ii) medication that may affect treatment outcome (e.g. high dose steroid, hormone replacement therapy, bisphosphonate, chemotherapy, immunosuppressant); (iii) systemic antibiotic treatment within three months prior to the
current study; (iv) pregnancy during experimental period; (v) heavy smoking within the past five years (more than 10 cigarettes per day or equivalent); (vi) history of irradiation in head and neck region; (vii) previous periodontal surgery at the selected site.

**Clinical parameters and outcome assessment**

The primary outcome of the studies was the histological evaluation of the healing. Secondary outcomes were the change in PPD, CAL and REC measured at baseline and before biopsy with the same type of periodontal probe (*PCPUNC 15*; Hu-Friedy; Chicago, IL, USA) and by the same calibrated examiner. Furthermore, plaque and bleeding scores and in Study 2 the depth of the intrabony component (*INTRA*) were also calculated. CEJ was used as the reference point. Where CEJ was not visible, the restoration margin was considered instead. Clinical recordings were made at six sites per tooth (mesio-buccal, mid-buccal, disto-buccal, mesio-lingual, mid-lingual and disto-lingual). The studies report the measurements at the same site (deepest at baseline) of the selected defect. Measurements were rounded up to the nearest millimetre. Examiner calibration included CAL and PPD measurements on five periodontal patients with similar disease severity, but other, than the patients enrolled in the study. Data were captured from six sites per tooth from all quadrants by the same way and same type of probe as described above. Measurements were repeated alike, 90 minutes apart. Calibration was accepted, if at least 90% of the collected figures were reproduced within a millimetre difference.

In addition, standardized long cone radiographs were taken at baseline, after three months and before biopsy for the radiological evaluation, utilizing commercial plastic film holder individualized by silicone putty impression material [110].

**Reconstructive periodontal surgery**

All surgeries were performed by the same, experienced periodontist in both trials, respectively. Prior to surgery clinical parameters were recorded as described above. Patients were asked to rinse with 0.2% chlorhexidine (*Curasept ADS 220*; Curaden, Kriens, Switzerland) for two minutes before perioral disinfection. In local anaesthesia (articain 80 mg + epinephrine 0.024 mg; *Ultracain D-S forte*; Aventis Pharma, Frankfurt am Main, Germany) full thickness mucoperiosteal flaps were reflected following intracrevicular incisions at the investigated site with additional one to two teeth apart. No releasing incisions were deemed necessary in Study 2, whereas vertical
releasing incisions were implemented if necessary for improve access in Study 1. Granulation tissue was removed and the roots were meticulously debrided by means of ultrasonic and hand instruments (Fig. 1c, 2c).

The following measurements were then made in Study 1: distance from CEJ to the bottom of the defect (CEJ–BD) and distance from CEJ to the most coronal extension of the alveolar bone crest (CEJ–BC). The intrabony component of the defect was defined as (CEJ–BD) − (CEJ–BC). INTRA was measured in Study 2 thereafter. By using a round diamond bur (1 mm diameter) a notch was placed at the bottom of the defect on the test tooth which was previously scheduled for extraction (Fig. 1d, 2d). Thus, any PDL tissue which later developed coronally to this notch on the root surface would be de novo formed connective tissue and clearly distinguishable in histological sections of biopsies.

After defect debridement, in Study 1, the involved root surfaces were conditioned for 2 minutes with Ethylene Diamine Tetraacetic Acid (EDTA) gel (PrefGel; Straumann; Basel, Switzerland) in order to remove the smear layer, according to the manufacturer’s instruction [111]. The defect, the underlying bone and the surrounding mucoperiosteal flaps were thoroughly rinsed with sterile saline to remove all EDTA residues. Following root conditioning, EMD (Emdogain®; Straumann; Basel, Switzerland) was applied on the root surfaces (Fig. 1e). The defects were then filled with a mixture of EMD + BCP (Straumann BoneCeramic®; Straumann; Basel, Switzerland) (Fig. 1f, g). Where needed, the periosteum, at the base of the mucoperiosteal flaps was incised to allow tension-free flap closure in coronal position.
No root conditioning or any other surface modifications were applied in Study 2. The intrabony defect was subsequently filled with the nano-HA paste (Ostim®, Heraeus Kulzer; Hanau, Germany) and adjusted to the alveolar crest, according to the manufacturer’s instruction (Fig. 2e, f).

The mucoperiosteal flaps were then repositioned and secured with a combination of suspended vertical mattress and single interrupted sutures (non-resorbable, monofilament; Daflon 5/0; Braun Aesculap; Tuttlingen, Germany) in order to achieve tensionless flap closure. Teeth were splinted in case of extreme mobility.

**Postoperative care**

All patients were postoperatively administered antibiotics (amoxicillin 500 mg + clavulanic acid 125 mg; Augmentin 625; GlaxoSmithKline; Brentford, Middlesex, UK)
Fig. 2  Reconstructive periodontal surgery with the use of nano-HA. (a) preoperative clinical measurement demonstrates a PPD of 10 mm; (b) preoperative radiograph demonstrates the presence of a deep intrabony defect; (c) the intraoperative situation revealed a deep one- and two-wall intrabony defect following debridement; (d) placement of the notch by means of a round bur indicating the bottom of the intrabony defect; (e) application of nano-HA; (f) periodontal defect filled with nano-HA.

three times daily for seven days and painkiller (diclofenac 75mg; Diclofenac Duo, Pharmavit, Veresegyház, Hungary) according to individual need. Subjects were advised not to brush the surgical area but rinse with 0.2 % chlorhexidine two times daily for 90 seconds during the postoperative four weeks. Subjects were asked to refrain any mechanical plaque control at the surgical site. Sutures were removed 10 to 14 days after surgery. Then the patients resumed tooth cleaning with the use of a soft brush. Additional appointments including oral hygiene instructions and professional supragingival tooth cleaning were performed fortnightly during the first twelve postoperative weeks. After this period and until biopsy removal, recall appointments were scheduled monthly. Neither subgingival instrumentation nor periodontal probing was performed during the entire experimental period.
Re entry, biopsy and histological procedure

After a healing period of nine (Study 1) or seven months (Study 2) the teeth were removed together with some of their surrounding periodontal tissues following the same type of local anaesthesia and presurgical preparations (Fig. 3a-c, 4a-c).

Fig. 3  Re-entry and biopsy after nine months of healing following treatment of EMD + BCP. (a) stable-looking soft tissue conditions; (b) radiograph indicates radiopaque tissue in the former defect; (c) decoronated tooth with some of the surrounding tissues

Fig. 4  Re-entry and biopsy after seven months of healing following treatment of nano-HA. (a) at 7 months following surgery, a substantial reduction of PPD was measured; (b) radiograph indicates mineralisation in the former defect; (c) removed tooth with some of the surrounding tissues

Specimens were subsequently placed in 10% buffered formalin for fixation. The extraction sites were then augmented with the use of various types of bone substitutes and barrier membranes (Study 1) and with DBBM (TutoDent Microchips; Tutogen; Neunkirchen, Germany) in conjunction with a resorbable collagen membrane of bovine pericardium origin (TutoDent Membrane; Tutogen; Neunkirchen, Germany) in Study 2.
According to local need, connective tissue graft was transplanted in order to increase the volume of keratinised gingiva [112]. After the healing of the extraction sites, subjects received either implant-based crowns or fixed partial dentures as part of the prosthetic rehabilitation.

The block biopsies were fixed in 10 % buffered formalin, decalcified in EDTA for a period of 4–6 weeks (depending on tooth/root volume) and dehydrated in graded series of ethanol. Immediately prior to embedding in paraffin, the roots/teeth were split in two along their long axis either in bucco-oral or mesiodistal direction (depending on the location of the deepest site of the defect) exactly at the notch indicating the site of interest. Thus, each biopsy provided two specimen blocks without the need for extensive cutting. Twenty sections from each of the two blocks per specimen were obtained with the microtome, set at 5 to 8 μm, and subsequently stained with hematoxylin–eosin and further with the oxone-aldehyde-fuchsin-Halmi staining method in Study 2. One experienced examiner measured the parameters by means of a computer-assisted toolbox. The field of view in the light microscope (Olympus DH 50; Olympus Denmark AS; Ballerup, Denmark) was examined on an LCD flat screen monitor through live streaming, which were also captured by a digital camera (Olympus DP 71; Olympus Denmark AS; Ballerup, Denmark). The following parameters were measured: (i) height of cementum regeneration: distance between the apical extension of the defect (bottommost point of the notch) and the coronal extension of a continuous layer of new cementum or cementum-like deposit on the planed root (millimetres); (ii) bone regeneration height: distance between the bottommost point of the notch and the coronal extension of regenerated alveolar bone along the planed root (millimetres) (The coronal extension of regenerated bone was defined as the most coronal level where the PDL space had an almost normal width); (iii) root resorption: combined linear heights of distinct resorption lacunae on the planed root (millimetres); (iv) ankylosis: combined linear heights of ankylosic union between the regenerated alveolar bone and the planed root (millimetres). Digital images were evaluated using a software program (SIS AnalySIS Auto Software 3.2; Soft Imaging System; Münster, Germany). The histomorphometric evaluation was carried out under ×25 to ×100 magnification. In the absence of a visible notch, the apical extension of instrumentation was used as landmark for the histomorphometric measurements.
Sources of support
Study 1 was supported by Institute Straumann, Basel, Switzerland. Study 2 was self-funded and supported by a research grant from Heraeus Kulzer, Hanau, Germany.

Candidate’s contribution to the studies
Fine-tuning of the protocol, study co-ordination, patient recruitment, assessment, data acquisition, patient management, compensatory treatment provision, revision and approval of the manuscript (Study 1); study design and protocol, study co-ordination, patient recruitment, surgical treatment (both stages), patient management, compensatory treatment provision, analysis and interpretation of data, preparation of the manuscript were carried out by the candidate of the present thesis. (Study 2)

4.3. Regeneration of peri-implant dehiscence defects (Study 3)

Experimental model and management
In the present experiment twelve female Göttingen minipigs (20 months of age and average body weight of 35 kg) were used. Before each surgical procedure, the animals were fasted overnight, weighed and pre-medicated. Each subjects received an injection of analgesia (Temgesic®; Schering-Plough; Brussels, Belgium) on the day of the surgery and on the next 3 days. StreptocillinVet® 250 + 200 mg/ml (Boehringer Ingelheim; Copenhagen, Denmark) was started one day before the procedure and was continued for seven days. The animals were anaesthetized according to a standard procedure using ketamine (Ketalar® 50 gm/ml; Pfizer; Sollentuna, Sweeden) and midazolam (Hameln 5 mg/ml; Pharmaceuticals GmbH; Hameln, Germany). After disinfection of the surgical site with 0.2% chlorohexidine solution (Corsodyl®; GlaxoSmithKline; Brentford, Middlesex, UK), local infiltration anaesthesia (Lidocaine 2% with 1:80,000 of epinefrine; HenrySchein Inc; Port Washington, NY, USA) was given. During the operation, additional 10 ml of ketamine and 1.5 ml of midazolam were administered when needed. Soft diet was provided daily during the whole course of the study. For documentation purposes digital photography and radiographs of the inserted implants were taken. The experimental protocol was approved by the Ethical Committee of the University of Lund (Sweden).
Study sequence

All surgical procedures were performed by two experienced periodontists. The treatment sequences in the study were:

1. Extractions and creation of “chronic” defect. Day 0/Baseline
2. Randomised implantation with/without creation of acute defect, with/without the use of test materials (4 groups: T1, T2, N, P). 3 Months
3. Uncovering of the implants and connection of abutments in different length (2 groups: loaded/non-loaded). 6 Months
4. Termination, histological analysis. 8 Months

Extraction and creation of chronic defect – Day 0/Baseline

Following intracrevicular incisions and careful elevation of a full thickness mucoperiosteal flap, the premolars and the first molar were extracted in both hemi-mandibles in all animals (Fig. 9a). Particular care was taken to ensure that no root remnants were left in the extraction sockets. Following tooth extraction, bone defect was created on both sides of the mandible by removal of the buccal plate with a chisel, on a length of 40 mm and a height of 6 mm (Fig. 9b). The lingual cortical bone wall was left intact. The surgical area was then rinsed with saline. Flap closure was achieved with absorbable sutures (Vicryl® 4–0; Ethicon GmbH; Norderstedt, Germany) (Fig. 9c) and the defect was allowed to heal for three months (Fig. 9d).
Implantation with/without creation of acute defect, with/without the use of test materials (first stage surgery) – 3 months

After a healing period of three months, following midcrestal incision and full thickness flap reflection, novel, titanium, bone level, 4.1x8 mm dental implants with a modified hydrophilic surface (Straumann®, Bone Level, SLActive implants; Straumann, Basel, Switzerland) were inserted either into the reduced thickness alveolar ridge (chronic defect and pristine site), or into a combined chronic and a surgically created standardised acute buccal dehiscence-type defect. Triangular shaped acute defects (height: 6 mm, apical mesio-distal width: 12 mm, bucco-lingual width: 2 mm) were prepared with a rotating surgical drill with ample saline irrigation. Defects were created in order to mimic real clinical circumstances of a partially edentulous ridge. Consequently, the coronal 6 mm of the implants were exposed on the buccal aspect. Only the apical 2 mm of the implants was in the alveolar bone around the whole circumference (Fig. 10). In total, 48 bone level implants with a diameter of 4.1 mm and a length of 8 mm were placed in 12 minipigs (four implants/animal; two in each hemi-mandible). All implants were inserted according to the guidelines of the manufacturer. Profile drilling (without tapping) was performed and the implants were inserted in such a way that the implant shoulder was aimed to match the bone crest level. Insertion torque was always higher than 35 Ncm.

According to a computer-generated randomization scheme, four groups were created: 

P (12 implants) – In the positive control group, twelve implants were inserted into the
chronic defects resulting from the 3-month healing following buccal bone plate removal at the time of teeth extraction (pristine sites) (Fig. 11).

**Fig. 11 Three months after baseline: Implant placement in group P (pristine site). Minor buccal and/or lingual dehiscence defects were occasionally present.**

_N (12 implants) – In the negative control group the chronic and acute dehiscence defects were left untreated as such the buccal surface of the implant remained exposed (negative control) (Fig. 12)._

**Fig. 12 Three months after baseline: Creation of standardized acute dehiscence-type defect. (a) Standardized defects were created around the osteotomy by removing part of the buccal bone. (b) The resulting dehiscence defects presented triangular-shaped base and the following dimensions: 6 mm apico-coronally, 12 mm mesio-distally, 2 mm bucco-lingually.**

*T1 (12 implants)* – The exposed implant surface in the dehiscence defects was treated with BCP (*Straumann BoneCeramic<sup>®</sup>*; particle size 500–1000 µm; Straumann; Basel, Switzerland). The BCP particles were mixed with autologous blood collected in the surgical site. The graft material was then applied in the dehiscence defect in contact with the exposed implant threads. Care was taken to avoid overfilling or coverage of the implant head (Fig. 13).
Fig. 13 Three months after baseline: Group T1: (a) BCP was mixed with autologous blood from the surgical site; (b) The exposed implant surface in the dehiscence defects was covered with the BCP.

T2 (12 implants) – BCP and the novel PEG membrane (Straumann MembraGel®, Straumann; Basel, Switzerland) was used for the treatment of the dehiscence defects. The synthetic bioresorbable polyethylene glycol hydrogel membrane was activated by mixing its components according to the manufacturer’s guidelines. The membrane was applied as a gel with a continuous flow over the bone substitute and the adjacent bone. (Fig. 14a-c). After the cross-linking reaction of the membrane components was completed and the membrane set, the margins were smoothed with a sharp scalpel blade. The membrane extended on the adjacent bone surface approximately 3 mm further than the grafted area and covered completely the implant head (Fig. 14d).

Fig. 14 Three months after baseline: Group T2: (a) BCP was placed onto the defect; (b) The two component of the hydrogel was mixed prior to application; (c) The polyethylene glycol hydrogel membrane (PEG) was applied on and around the BCP and the implant; (d) The set, polimerised membrane in place.
All implants (48) of all groups received closure screws and were fully covered by the mucoperiosteal flaps for submerged healing. The surgical area was carefully sutured with absorbable material using single interrupted and vertical mattress sutures.

**Implant uncover and abutment connection according to two different loading protocols – 3 months after implant placement**

At 3 months after implant placement (6 months after extraction), a second stage surgery was performed to uncover all implants. Midcrestal incisions were performed on the sites of implant placement. Minor buccal and lingual flap elevation was performed to limit periosteal exposure. The distance from the implant shoulder to the first bone-to-implant contact was measured on the buccal side with a UNC 15 periodontal probe. According to a computer generated randomization scheme, two different abutments were connected to the fixture in a split-mouth design (Fig. 15).

![Split mouth design](image)

**Fig. 15 Loading model (randomised):**
- one side loaded (two implants), contralateral hemimandible non-loaded (two implants).

On one side, the closure screw was replaced with a new, transmucosal closure screw, for all treatment groups. These implants were not functional loaded (non-loaded implants) (Fig. 16a). On the contralateral side, the implants in all treatment groups were connected to long (8 mm), custom-made, abutments so that the implant would be subjected to occlusal functional forces with their antagonists (loaded implants) (Fig. 16b). The flaps were then sutured using single interrupted sutures.
Fig. 16 Three months after implant placement: (a) Non-loaded implants. Transmucosal closure screws in one hemimandible; (b) Loaded implants. In the contralateral hemimandible, the implants were connected to long abutments (8 mm) so that the implant would be subjected to occlusal functional forces with their antagonists.

Termination – 5 months after implant placement
At five months after implant placement (eight months following extraction and after two months of functional loading), the study was terminated by an intravenous injection of a 20% solution of pentobarbital (Pentobarbital Veterinary; Apoteksbolaget, Sweden).

Histological processing
The jaws with the surrounding soft tissues were dissected and embedded in neutral buffered formalin solution (5%) for at least three weeks. After fixation, the specimens were thoroughly rinsed in running water and dehydrated in ascending concentrations of ethanol (50%, 70% and 100%). Following dehydration, the biopsies were embedded in polymethylmetacrylate (PMMA; Merck AG; Darmstadt, Germany). The samples were orientated parallel to the longitudinal axis of the implant and one central section was cut in a bucco-lingual direction. The Exact Cutting–Grinding technique was used for the preparation of the histological sections with an approximate thickness of 70 μm [175]. The sections were stained with toluidine blue.

Histomorphometric measurements
Histological analysis was carried out using a light microscope at ×10 and ×40 magnification. Histometric measurements were performed on digitally captured images. All measurements were performed by the same experienced investigator with the help of an image analysis system (Image I/Metacolor®; Universal Imaging Corporation, West Chester, PA, USA). For intraexaminer calibration, measurements were repeated on 10% of the histological sections (with 1 h interval) and resulted in high agreement (kappa-score 0.83).

The following reference points were identified on each section:

- $S$: implant shoulder
- $D$: apical limit of the previous acute dehiscence defect in pristine bone
- $NBC$: bone crest height on the buccal side
- $BIC$: most coronal bone-to-implant contact on the buccal side.

Linear measurements between these points were calculated in mm as follows (Fig. 17):
The defect height (S-D) was measured on the implant surface from the implant shoulder to the apical border of the former acute dehiscence defect.

The residual dehiscence defect was calculated from the implant shoulder to: (i) the bone crest on the buccal side (S-NBC) and (ii) the most coronal bone-to-implant contact on the buccal side (S-BIC).

**Fig. 17** Histomorphometry. Reference points and linear measurements. Defect height from the implant shoulder to the apical border of the acute dehiscence defect in native bone measured on the implant surface (S-D); Residual dehiscence defect from the implant shoulder to the bone crest on the buccal side (S-NBC); the most coronal bone-to-implant contact on the buccal side (S-BIC); bone-to-implant contact in the region of interest (S-D).

The bone-to-implant contact percentage (BIC%) was defined as the percentage of the defect height (S-D) occupied by newly formed bone in direct contact with the implant surface without the presence of fibrous membrane inside the defect area.

Surface measurements were performed by the use of the same image analysis software where, the following surfaces were calculated in mm²:

- The area of regenerated bone (bone surface - BS) was calculated as the area of newly formed bone in the previous defect area.
- The graft area (graft surface - GS) was calculated as the area occupied by the BCP particles in the previous defect area.

All measurements were reported for each implant and categorized according to the treatment groups (T1, T2, P, N) and loading protocols.

The primary variable was the BIC%. Secondary variables were S-BIC, S-NBC, BS and GS (where applicable).

**Statistical analysis**

The statistical analysis was performed using a software program (SPSS 14.0; SPSS Inc; Chicago, IL, USA). Mean values and standard deviations were calculated for each variable and each group. Differences between means for the groups (T1, T2, P, N) and the loading protocols (loaded and non-loaded) were assessed by performing a 2 way
repeated measures analysis of variance (ANOVA) for each variable, with repeated measures over loading sites and treatments. The assumptions of normality and constant variance underlying the ANOVA were checked by a study of the residuals. If the assumptions were not fulfilled, a square root transformation was applied to the data. When a statistically significant difference between the group means was indicated in the ANOVA, a post-hoc Bonferroni test was used to determine which group means differed significantly. Result was considered statistically significant if $P < 0.05$.

**Source of support**

The present study was supported by Institute Straumann, Basel, Switzerland.

**Candidate’s contributions to the study**

Fine-tuning of the design and protocol, study co-ordination, surgical treatment (all the three stages), revision and approval of the manuscript were carried out by the candidate of the present thesis.

**4.4. Regeneration of post-extraction alveolar sockets (Study 4)**

Prior to commencement of the study a detailed protocol was developed and agreed upon by the authors based on the Cochrane Collaboration guidelines and previous reviews published by our research group [127] [228] [229] [230] [231].

**Focused question**

Following tooth/root extraction in humans, what is the effect of ridge preservation on the residual alveolar ridge dimension and on histological characteristics, compared to unassisted socket healing?

**Definition**

Whilst ‘socket preservation’ has widely been employed to depict a certain procedure, we believe that the objective of these interventions is to preserve the dimension of the AR. Therefore, we have used the term ‘Alveolar Ridge Preservation’ to define such procedures.
Types of studies
Longitudinal prospective studies were included i.e. RCTs, CCTs and cohort studies with control group.

Populations of studies
Healthy individuals, without any age limit, who underwent any type of ridge preservation following permanent tooth extraction, were included. Smokers and patients with history of periodontal disease were not excluded. The minimum number of subjects per group was five. However, no limit was set for study follow-up period.

Types of Interventions
Test groups
Studies reporting on any of the following types of interventions were included: socket grafting (autograft, allograft, xenograft, alloplastic materials); socket sealing (soft tissue grafts); GBR (resorbable/non-resorbable barriers); biological active materials (growth factors) and combinations of the above techniques/materials.

Control groups
The control groups of the included studies comprised empty sockets, i.e. unassisted socket healing.

Outcome variables
The primary outcome was the change in oro-facial (horizontal) and apico-coronal (vertical) AR dimensions. Secondary outcomes were the followings: (i) change in buccal plate thickness; (ii) bone volume alteration following extraction; (iii) complications; (iv) histological healing characteristics; (v) site eligibility for placement of an adequate size dental implant with or without further augmentation; (vi) patient-reported outcomes, such as quality of life and (vii) health economics.

Risk of bias and methodological quality assessment
In order to evaluate the methodological quality and risk of bias of individual studies, we used a combination of parameters from the Cochrane Collaboration and Consolidated Standards of Reporting Trials (CONSORT) statement. The following parameters were assessed and taken into consideration in the final analysis: sample size calculation, statement of eligibility criteria, ethics approval, informed consent, baseline
homogeneity, randomisation method, allocation concealment, masking, calibration, follow up, protocol violation, method of statistics, unit of analysis, CONSORT implementation, International Standard Randomised Controlled Trial Number Register (ISRCTN) and funding disclosure.

Randomisation was accepted as adequate, in case the allocation sequence was correctly generated either by computer, toss of a coin, throwing dice, etc. Quasi randomisation e.g. birth dates, hospital numbers were not accepted. Adequacy of allocation concealment was accepted if the sequence was concealed, until intervention was assigned (e.g. in sequentially numbered and sealed opaque envelopes, remote computer or central telephone). Statistical analysis was judged as adequate if appropriate statistical method was selected to accommodate to the characteristic of the each individual data (e.g. number of groups and investigated categories, size of samples, normally distributed or skewed data, parametric or non-parametric, paired or unpaired, numerical or categorical variables). Statistical significance was accepted in case of confidence interval (CI) >95% (p<0.05), while ‘statistically highly significant’ referred to CI>99.9% (p<0.001).

Based on the above, we attempted to categorize the possible risk of bias as low, moderate or high. Low risk referred to studies with adequate randomisation method, sequence concealment and masking of examiner. Studies were classified as moderate, if one of the above key categories were missing, or high risk of bias, if more than one were lacking.

Inclusion criteria

1. All prospective longitudinal studies (i.e. RCTs, CCTs and cohort studies) were included, where one of the above mentioned types of interventions were carried out in the test group, whereas unassisted socket healing served as control.
2. Studies on healthy individuals, without any age limit, who underwent ARP following tooth extraction, were included.
3. Studies had to report on minimum of five patients per group.
4. Studies, performing clinical or three-dimensional (3D) radiographic evaluation of hard tissue or histological assessment, were included.
Exclusion criteria

1. Case reports, case series, retrospective analyses were excluded.
2. Studies without a control group comprising unassisted socket healing were excluded.
3. Studies on medically compromised patients e.g. uncontrolled diabetes mellitus or cancer were excluded.
4. Studies reporting on immediate placement of dental implant were excluded.
5. Studies describing extraction of third molars were excluded.

Search strategy

A sensitive search strategy was designed as we anticipated that relevant studies might be difficult to locate. The search strategy incorporated both electronic and hand searches. The following electronic databases were utilized in Apr 2010: (i) MEDLINE In-Process & Other Non-Indexed Citations and MEDLINE 1950 to present via Ovid interface; (ii) EMBASE Classic + EMBASE 1947 to present via Ovid interface; (iii) The Cochrane Central Register of Controlled Trials (CENTRAL); (iv) Literatura Latino-Americana e do Caribe em Ciências da Saúde (LILACS).

The electronic search strategy used the following combination of key words and MeSH terms:

(“tooth extraction” OR “tooth removal” OR “socket” OR “alveol$” OR “ridge” OR “crest” OR “tooth socket” OR “alveolar bone loss” OR “bone resorption” OR “bone remodeling”) AND (“preserv$” OR “reconstruct$” OR “augment$” OR “fill$” OR “seal$” OR “graft$” OR “repair$” OR “alveolar ridge augmentation” OR “bone regeneration” OR “bone substitutes” OR “transplantation”).

Cochrane search filters for RCTs and CCTs were implemented. In addition, cohort trials were also searched. The results were limited to humans only.

Periodontology, Oral Surgery, Oral Medicine, Oral Radiology, Oral Pathology and Endodontics, Periodontology 2000. No language restrictions were applied. Translations were carried out as necessary by two reviewers.

The extracted data were copied into Reference Manager 10 software (Thomson Reuters; New York, NY, USA). Thus, the further steps of screening were performed on this interface. A three-stage selection of the resulted hits was performed independently and in duplicate by two reviewers. In order to reduce errors and bias, a calibration exercise was performed with the first 500 titles, resulting in 96.4% agreement. In case of disagreement at the title selection stage, the trial was included in the abstract stage. At the abstract and full text selection any disagreements between the above reviewers were resolved by discussion. If unresolved, a third reviewer was involved for arbitration. The reasons for exclusion were recorded either in the Reference Manager (abstract stage) or in a specific data extraction form (full text stage). The level of agreement was determined by Kappa score calculation.

Research synthesis
Studies were grouped by research design and their chief characteristics. Outcomes were recorded in evidence tables. In view of the marked heterogeneity, no meta-analysis was conducted. Instead, a narrative synthesis was undertaken.

Sources of support
The present study was self-funded and supported by the Research Discretionary Account of the Unit of Periodontology, UCL Eastman Dental Institute, which received a proportion of funding from the UK Department of Health’s NIHR Biomedical Research Centres funding scheme.

Candidate’s contributions to the study
Study design and protocol, study co-ordination, selection and review of all included and excluded trials, data collection, analysis and interpretation of data, manuscript preparation were carried out by the candidate of the present thesis.
5. RESULTS

5.1. Periodontal regeneration
5.1.1. Treatment with EMD and SBC

All ten subjects (six women and four men), aged between 27 and 62 years (mean age, 48.6 years) completed the study. Postoperative healing period was uneventful in all cases. No complications, such as allergic reactions, abscesses or infections were observed throughout the duration of the study.

In one of the 10 subjects, the clinical evaluation and radiographs demonstrated excellent clinical healing. Therefore, putting patient’s benefit first, a decision was made not to remove the tooth and to perform only a re-entry. In this particular case, the re-entry revealed that the intrabony defect, which was located at the mesial aspect of tooth #24, presented with almost complete healing with a newly formed hard tissue resistant to probing.

The clinical measurements demonstrated a reduction in mean PPD from 8.6 ± 1.9 mm to 5.3 ± 2.0 mm at baseline and at nine months, respectively. The mean CAL changed from 10.8 ± 2.0 mm to 7.8 ± 1.7 mm resulting in a mean CAL gain of 3.0 ± 1.6 mm (Table 1).

Table 1 Clinical and histologic results following treatment of human intrabony periodontal defects with EMD + BCP (mm)

In six out of nine biopsies, the histological findings indicated the formation of new cementum with inserting collagen fibres to a varying extent (Fig 5a, b). The newly formed cementum was a mixed acellular and cellular type in all specimens. Collagen
fibres were inserting into the newly formed cementum in all specimens showing new attachment. In three out of the nine specimens, the healing resulted in a long junctional epithelium extending to the bottom of the defect (subjects 1, 5, and 7) (Table 1). Mean new connective tissue attachment (i.e. new cementum with inserting collagen fibres) varied from 0.0 to 2.1 mm. The amount of newly formed bone was limited and varied from 0.0 to 0.7 mm. In most specimens, the remaining BCP particles were encapsulated in connective tissue, whereas formation of a bone-like tissue around the graft particles was observed only occasionally. Direct contact between the graft particles and the root surface (cementum or dentin) was not observed in any of the analysed specimens.

Fig. 5 The histologic analysis demonstrated (a) formation of new cementum (NC), new periodontal ligament (NPL), and new bone (NB) along the debrided root surface coronally to the notch (N). The presence of BCP particles is evident. Bone formation is visible only around some of the grafted BCP particles (hematoxylin and eosin; original magnification, ×12.5). (b) Higher magnification of the photomicrograph. The BCP particles are encapsulated in connective tissue and do not show signs of resorption (hematoxylin and eosin; original magnification, ×100).
5.1.2. Treatment with nano-HA

Baseline Characteristics
All six subjects completed the study. Mean age was 44.7±14.2 years and the male/female ratio was 1:1. All patients displayed clear medical history apart from one (#03) with treated hypertension. Two patients (#01, #06) were light smokers (≤10 cigarettes/day). All six defects displayed a combined one- and two-wall configuration. Clinical measurements at baseline revealed the following mean values: FMPS: 17.8%, FMBS: 14.3%, PPD: 8.7±2.0 mm, CAL: 12.2±1.6 mm, REC: 3.5±1.8 mm. The mean values of CEJ-BD and INTRA were measured as 12.9±1.9 mm and 3.4±0.7 mm respectively during the first surgery (Table 2).

Table 2 Clinical and histologic results following treatment of intrabony defects with OFD and nano-HA (mm)

<table>
<thead>
<tr>
<th>Patient No.</th>
<th>Tooth type</th>
<th>PPD before 7 months</th>
<th>PPD after 7 months</th>
<th>Diff.</th>
<th>CAL before 7 months</th>
<th>CAL after 7 months</th>
<th>Diff.</th>
<th>CAL gain</th>
<th>INTRA</th>
<th>New cementum</th>
<th>New bone</th>
</tr>
</thead>
<tbody>
<tr>
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<td>11</td>
<td>11</td>
<td>6</td>
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<td>1</td>
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<td>3</td>
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<td>4.5</td>
<td>3.5</td>
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<td>6.0</td>
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<td>3.5</td>
<td>2.6</td>
<td>3.5</td>
<td>0.4</td>
</tr>
<tr>
<td>SD (mm)</td>
<td>1.8</td>
<td>1.7</td>
<td>0.0</td>
<td>1.6</td>
<td>1.4</td>
<td>0.7</td>
<td>1.5</td>
<td>0.7</td>
<td>0.6</td>
<td>0.4</td>
<td>0.6</td>
</tr>
</tbody>
</table>

Clinical outcomes
Postoperative healing was uneventful in all cases. Neither adverse event nor complication such as allergic reaction, infection, ulceration or abscess was detected. On the contrary, signs of accelerated early wound healing were observed clinically. At 7 months following surgery, mean PPD reduction and mean CAL gain were measured 4.0 ± 0.8 and 2.5 ± 0.8 mm, respectively (Table 2).

Radiological observations
Mineralization of the newly formed tissue in the intrabony defect was observable on the IOPAs. The phenomenon of increased radiopacity was already visible at three months (Fig. 6a-d).
Fig. 6 Increasing density of the former defect is observable on the radiographs (a) prior to surgery; (b) right after surgery; (c) after three months of healing; (d) after seven months of healing.

Histological outcomes

The histological analysis revealed that in three out of the six defects, the healing occurred through formation of a long junctional epithelium along the debrided root surfaces extending until the most apical part of the defects (Fig. 7a, b). In the remaining three specimens limited formation of new cementum with inserting connective tissue fibres and new bone were observed with a magnitude varying from 0.53 to 0.86 mm and from 0.86 to 1.33 mm, respectively (Table 2). Neither ankylosis, nor root resorption was observed in any of the biopsies. In two out of the six biopsies, some remnants of the grafting material were visible (Fig. 8a, b). The nano-HA particles were predominantly surrounded by connective tissue, without signs indicating a potential to promote periodontal or bone regeneration.

Fig. 7 The histological evaluation revealed (a) healing predominantly characterized by a long junctional epithelium and limited regeneration of cementum and bone. Red arrowhead: coronal extension of new cementum, blue arrowhead: coronal extension of new bone, red dotted line: apical extension of the notch. Original magnification × 5; (b) Higher magnification of the defect. Formation of new cementum (NC) and new periodontal ligament (PDL) was confined to the area of the notch. White dotted line: margin between the newly formed bone and old bone, red asterisk: artifact. Original magnification × 25.
Fig. 8 (a) Representative photomicrograph revealing limited periodontal regeneration (case 5 from Table 1). Red dotted line: apical extension of the notch, red arrowhead: coronal extension of new cementum, blue arrowhead: coronal extension of new bone. Original magnification × 25. (b) Higher magnification of the defect. Formation of new cementum (NC), new periodontal ligament (PDL) and new bone (B) was confined to the area of the notch. White dotted line: margin between the newly formed bone and old bone, red asterisk: artefact, white asterisk: remnant nano-HA particle encapsulated in connective tissue. Original magnification × 50

5.2. Provision of peri-implant dehiscence defects with SBC and PEG

Clinical observations

Extractions and creation of chronic defect – Day 0/Baseline
Following extractions and removal of the buccal plate, all sites were closed by primary intention and healed without complications.

Creation of acute dehiscence defects (T1, T2, N) and implantation (T1, T2, P, N) – First stage surgery 3 months after baseline
As a result of the previous buccal plate removal, a remarkable decrease in the buccolingual ridge width, which resulted in thin (knife-edge) ridge, was observed in all these cases (Fig. 18). Consequently, in group P, the implant placement was occasionally
associated with the presence of buccal and/or lingual dehiscence around the 4.1 mm implants. Dense (type 1) bone was observed in all sites during implantation.

Fig. 18 Removal of the buccal plate after extraction resulted in knife–edge ridge.

Good primary implant stability was achieved at all sites of all experimental groups, despite the extensive dehiscence defects that were created. The final insertion torque was always higher than 35 Ncm for all implants.

In group T2, the membrane was applied successfully in all cases. However, in 4/12 sites, the membrane had to be reconstructed prior to flap closure, because of membrane rupture (three cases) or instability on the underlying graft/bone (one case) (Fig. 19). Primary closure of the flaps was achieved in all sites of all groups.

Fig. 19 Three months after baseline. Rupture of the PEG membrane above the implant head, before flap closure is visible.

Implant uncover and abutment connection – 3 months after implant placement
Exposure of the cover screw of the implant was observed in 17/48 implants. This occurred in all groups with similar frequency (four exposures in groups N, T1, T2 and
five exposures in group P). One implant was found to be mobile and was removed (Group T2). All other sites healed uneventfully.

The distance from the implant shoulder to the first clinical bone-to-implant contact was measured on the buccal side in all groups.

In group T1, after flap reflection, graft granules were found to be encapsulated in the soft tissue flap in 3/12 sites. No attempt was made to remove the mobile graft particles. The mean clinical distance from the implant shoulder to the first clinical bone-to-implant contact was 2.55 mm (range: 0-6 mm).

In group T2, no remnant of the PEG membrane was visible in any of the sites during second stage surgery. In this group, no loose graft particles were observed. The mean clinical distance from the implant shoulder to the first clinical bone-to-implant contact was 2.1 mm (range: 2-6 mm).

In group P, the mean clinical distance was 1.25 mm (range: 1-3 mm).

In group N, the mean clinical distance was 3.35 mm (range: 1-7 mm).

**Histological analysis**

Histological measurements and the relevant statistical analysis were performed in a total of 47 of the 48 implants initially placed.

These assumptions of normality and constant variance were satisfied for all the ANOVAs apart from that for GS when the constant variance assumption was violated. In this case, a square root transformation of GS was taken and the ANOVA was repeated on the transformed data.

**Descriptive histology**

**Group P**

One implant (loaded) showed complete lack of osseointegration. In this implant, a layer of loose connective tissue was in contact with the implant surface around its
circumference. In the 11 remaining implant sites, the new bone crest (S-NBC) was always located within 3 mm apical to the implant shoulder (in 7 sites, the bone crest was located 0–2 mm from the implant shoulder). Similarly, the most coronal bone-to-implant contact was located within 4 mm apical to the implant shoulder in 10/11 cases. Implant osseointegration was achieved in both groups (Fig. 20).

**Fig. 20** Representative histological views of wound healing at 5 months after implant placement. Group P: Non-loaded (left) and loaded implants (right). Implant osseointegration was observed in 11 specimens irrespective of the loading (Toluidine blue, original magnification ×10).

**Group N**
This group exhibited significant bone growth in the defect area. None of the specimens showed complete resolution of the defect with newly formed bone, indicating that the created defect met the criteria of ‘critical size’.

**Non-loaded implant sites**
New bone formation was observed in connection with significant bone-to-implant contact alongside SLActive implant surface in most of the specimens, varying from 6% to 67% (mean 40.5%). The newly formed bone crest always corresponded (within 1 mm) to the most coronal bone-to-implant contact. The bone formation was continuous with the floor of the defect and it resembled trabecular bone (Fig. 21).

**Loaded implant sites**
The histological observations at the loaded implants were not significantly different
from those at the unloaded sites (Fig. 21). The newly formed bone crest often corresponded to the most coronal bone-to-implant contact and only in 1/6 specimens, the bone crest was located coronally to the first bone-to-implant contact. New bone formation proceeded in connection with significant bone-to-implant contact along the SLActive implant surface (BIC%) in most of the specimens, varying from 47% to 79% (mean 64.1%).

**Group T1**

*Non-loaded implant sites*

In this group, moderate to marked bone formation at the area of the defect was observed (Fig 22). Bone regeneration proceeded both along the implant surface and in contact with the BCP particles. However, while bone formation around the graft particles could be always observed in the apical portion of the defect, a significant amount of particles were also encapsulated by soft tissue in the middle and coronal portion of the defect. BCP particles were surrounded by soft tissue in the marginal portion of the defect in 3/6 sites. In 2/6 sites, bone formation was only observed in the apical half of the defect, while in the remaining half, the BCP particles were embedded in soft tissue (Fig 22).
Loaded implant sites

Similar histological results were observed in the loaded sites in terms of bone regeneration and osseointegration. Bone formation could be always observed in the middle and apical portion of the defect and along the implant surface. However, a significant amount of BCP particles were encapsulated by soft tissue in the coronal portion of the defect at 3/6 sites (Fig. 22).

![Representative histological views of wound healing at five months after implant placement. Group T1: Non-loaded (left) and loaded implants (right): apically, bone regeneration was obvious in the vicinity of the implant and the BCP. Soft tissue encapsulation of the bone substitute was observed in the middle and coronal portion of the defect (*) (Toluidine blue, original magnification ×10)](image)

Group T2

Non-loaded implant sites

In 3/5 of the unloaded specimens, new bone formation of trabecular appearance was observed occupying all the defect area up to the level of the implant shoulder (Fig. 23). The PEG membrane seemed to be utterly resorbed. The bone substitute was in the majority of the case in contact with the newly formed bone and were smaller in size and shape than in the T1 group. In 4/5 specimens, a layer of connective tissue interposed between the newly formed bone and the implant surface indicating lack of bone-to-implant contact in these areas (Fig. 23).
Loaded implant sites

Similar histological results were observed in the loaded sites compared to the non-loaded sites. New bone formation of trabecular appearance was observed occupying the defect area in 4/6 specimens. An extensive soft tissue downgrowth interposed between the newly formed bone and the SLActive implant surface was observed in 5/6 specimens (Fig. 23).

Histometric results

Linear measurements

Defect height

S-D was calculated in groups T1, T2, and N. The mean height was measured 5.95 mm ranging from 4.3 to 7.4 mm. No statistically significant differences were observed among these three group means ($P = 0.82$). No statistically significant difference was observed between the means of the loaded and non-loaded groups ($P = 0.47$) (Table 3).
Bone-to-implant contact percentage

BIC% in the defect area was calculated in all the groups (Table 4). The statistical analysis showed no statistically significant differences between the means of the treatment groups T1, T2 an N ($P = 0.43$) or between those of the loaded and non-loaded implants within groups ($P = 0.43$). A trend for higher values was observed in the loaded sites compared with non-loaded sites in all treatment groups with exception of group T2, where lower BIC% values were observed at loading sites.

Implant shoulder to bone crest

The mean distance from the implant shoulder to the bone crest on the buccal side (S-NBC) was calculated in all groups (Table 5). No statistically significant differences were observed between the group means ($P = 0.91$) or between those of the loaded and non-loaded sites ($P = 0.25$). However, a trend for higher values was observed in the non-loaded sites compared with loaded sites in all groups with exception of group T2, where values were higher in loaded sites compared with non-loaded sites.
Implant shoulder to the most coronal bone-to-implant contact

Analysis of the values of S-BIC showed no statistically significant differences between the means of groups N, T1, T2 ($P = 0.63$). No statistically significant differences were observed between loaded and non-loaded group means ($P = 0.53$). Similar to S-NBC measurements, a tendency for higher values was observed in the non-loaded sites compared with loaded sites in all groups with exception of group T2, where values were higher in loaded sites compared with non-loaded sites (Table 6). All the linear measurements (S-BIC, S-NBC, BIC%) seem to indicate a possible positive influence of functional loading on new bone formation and bone-to-implant contact. However, when the dehiscence site was treated with GBR using a PEG membrane (group T2), loading seemed to negatively influence new bone formation and osseointegration.

<table>
<thead>
<tr>
<th>S-BIC (mm)</th>
<th>Loading ±</th>
<th>Non-loading</th>
<th>Loading</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>non-loading</td>
<td></td>
<td></td>
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<tr>
<td>T1</td>
<td>2.43 ± 1.74</td>
<td>2.85 ± 1.60</td>
<td>2.01 ± 1.92</td>
</tr>
<tr>
<td>T2</td>
<td>3.71 ± 1.83</td>
<td>2.98 ± 1.91</td>
<td>4.33 ± 1.66</td>
</tr>
<tr>
<td>N</td>
<td>2.67 ± 1.40</td>
<td>3.41 ± 1.58</td>
<td>1.93 ± 0.69</td>
</tr>
<tr>
<td>P</td>
<td>2.50 ± 2.17</td>
<td>2.66 ± 1.45</td>
<td>2.35 ± 2.87</td>
</tr>
</tbody>
</table>

**Table 6** Mean ± standard deviation for: Distance from the implant shoulder to the most coronal bone-to-implant contact on the buccal side (S-BIC) for groups P, N, T1, T2

Surface measurements

Graft surface

The BCP surface was calculated for groups T1 and T2 (Table 7). A limited number and size of BCP particles could be identified in specimens of the group T2. In contrast with this, approximately 1/3 of the new hard tissue area in group T1 was occupied by graft particles. This was evident in both loaded and non-loaded subgroups. The differences between the means of the two groups were statistically significant ($P = 0.031$).

<table>
<thead>
<tr>
<th>GS (mm²)</th>
<th>Loading ±</th>
<th>Non-loading</th>
<th>Loading</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>non-loading</td>
<td></td>
<td></td>
</tr>
<tr>
<td>T1</td>
<td>4.94 ± 2.81*</td>
<td>5.30 ± 3.32</td>
<td>4.58 ± 2.47</td>
</tr>
<tr>
<td>T2</td>
<td>0.71 ± 0.80*</td>
<td>0.77 ± 1.06</td>
<td>0.67 ± 0.60</td>
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</tbody>
</table>

*statistically significant difference between the groups T1 and T2, $P = 0.031$.

**Table 7** Mean ± standard deviation for: Graft surface (GS) for groups T1, T2
Bone surface

Bone surface area was calculated in groups T1, T2 and N in both loaded and non-loaded subgroups (Table 8), with group T2 exhibiting the largest amount of bone formation among the three groups (P < 0.05). There was a trend for higher BS values, in groups T1 and N when functional loading was applied. However, in group T2, the newly formed bone surface was higher in the non-loading group. This was in accordance with what was observed with the linear measurements.

<table>
<thead>
<tr>
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<th>Loading + non-loading</th>
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<th>Loading</th>
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<tbody>
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<td>9.32 ± 5.28</td>
<td>10.05 ± 3.00</td>
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<tr>
<td>T2</td>
<td>13.18 ± 5.85</td>
<td>15.28 ± 4.98</td>
<td>11.43 ± 6.37</td>
</tr>
<tr>
<td>N</td>
<td>8.81 ± 4.48</td>
<td>7.92 ± 5.14</td>
<td>9.17 ± 3.97</td>
</tr>
</tbody>
</table>

**Table 8** Mean ± standard deviation for: Bone surface (BS) for groups N, T1, T2

Summated surfaces

The total surface occupied by bone and substitute (BS + GS, hard tissue surface) was also calculated for group T1, T2 and N (Table 9). Groups T1 and T2 showed similar amounts of hard tissue formation (14.6 and 13.9 mm² respectively). Limited hard tissue formation was observed in group N too.

<table>
<thead>
<tr>
<th></th>
<th>Loading + non-loading</th>
<th>Non-loading</th>
<th>Loading</th>
</tr>
</thead>
<tbody>
<tr>
<td>GS + BS (mm²) T1</td>
<td>14.63 ± 5.48</td>
<td>14.62 ± 6.98</td>
<td>14.64 ± 4.15</td>
</tr>
<tr>
<td>T2</td>
<td>13.90 ± 5.58</td>
<td>16.05 ± 4.41</td>
<td>12.11 ± 6.17</td>
</tr>
<tr>
<td>N</td>
<td>8.81 ± 4.48</td>
<td>7.92 ± 5.14</td>
<td>9.17 ± 3.97</td>
</tr>
</tbody>
</table>

**Table 9** Mean ± standard deviation for: Hard tissue surface (BS + GS) for groups N, T1, T2

5.3. The effect of ARP on extraction sockets

Search sequence

The electronic search yielded 6,216 relevant hits after removal of duplicates (Fig. 24). Subsequently, 157 titles were selected for the abstract stage. Following investigation of the abstracts, 42 articles qualified for full text evaluation. Four extra papers were then added as a result of the hand search. Assessment of these articles resulted in the
following 14 publications eligible for the review [232] [233] [210] [234] [215] [207] [235] [212] [213] [214] [236] [237] [209] [238].

![Flow chart of the screening process](image)

**Fig. 24** Flow chart of the screening process

The excluded full text papers along with the reasons for exclusion are listed in table 10. The most typical reasons for exclusion were lack of control group with unassisted socket healing; use of retrospective design; assessment of dimensional changes of the AR only on periapical two dimensional radiographs or on casts taken from soft tissue level; and surgical removal of third molars.

<p>| Table 10 | List of excluded full text papers and reasons for exclusion (on subsequent page) |</p>
<table>
<thead>
<tr>
<th>First author (year of publication)</th>
<th>Journal</th>
<th>Reasons for Exclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bolouri (2001)</td>
<td>Comp Cont Educ Dent</td>
<td>Reported on optical density on two-dimensional radiographs</td>
</tr>
<tr>
<td>Brawn (2007)</td>
<td>Impl Dent</td>
<td>Case report</td>
</tr>
<tr>
<td>Carmagnola (2003)</td>
<td>Clin Oral Impl Res</td>
<td>Lack of real control group, resembles to a retrospective analysis (extreme difference in follow-up period between tests and controls. T1: 4 months; T2: 7 months; C: 1-15 years, mean: 7.8 years)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Retrospective study as stated by the authors in the discussion</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Healing period varied between 1.5 months and 1.5 years</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Neither histomorphometry nor clinical or radiographic measurements reported in the results</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Extraction of fully impacted third molars</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Linear measurements on OPG and scintigraphy</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Primary outcome was to analyze bone metabolism</td>
</tr>
<tr>
<td>Hoad-Reddick (1994)</td>
<td>Eur J Prosthrest Dent</td>
<td>Two-dimensional linear measurements obtained from OPG and cephalometry</td>
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<tr>
<td></td>
<td></td>
<td>Lack of defined landmarks</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Surgical procedure was not described</td>
</tr>
<tr>
<td>Howell (1997)</td>
<td>Int J Periodont Rest Dent</td>
<td>Neither socket preservation procedure nor the results were described. Soft tissue punch technique only</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Case series without control group</td>
</tr>
<tr>
<td>Kangvonkit (1986)</td>
<td>Int J Oral Maxillofac Surg</td>
<td>Primary outcome was soft tissue healing</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Based on OPG and lateral cephalogram only</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Evaluation method remains unclear</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Primary outcome was the biocompatibility of HA cones</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Primary outcome was to assess the periodontal status of second molars after extraction of third molars</td>
</tr>
<tr>
<td>Kerr (2008)</td>
<td>J Periodontol</td>
<td>No biomaterials were used to preserve the ridge dimensions, therefore did not address the focused question</td>
</tr>
<tr>
<td>Kwon (1986)</td>
<td>J Oral Maxillofac Surg</td>
<td>Based on OPG and lateral cephalogram only</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Evaluation method remains unclear</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Lack of description of the measurement methods</td>
</tr>
<tr>
<td>Molly (2008)</td>
<td>J Periodontol</td>
<td>Control group was covered by an e-PTFE membrane, thus lack of unassisted control sockets</td>
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<td></td>
<td></td>
<td>Case series without control group</td>
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<tr>
<td></td>
<td></td>
<td>Resembles to a retrospective analysis (healing period ranged from 3 to 11 months)</td>
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<tr>
<td>Pape (1988)</td>
<td>Deutsche Zahnarztliche Zeitschrift</td>
<td>Augmentation of a resorbed ridge</td>
</tr>
<tr>
<td>Quinn (1985)</td>
<td>J Am Dent Assoc</td>
<td>Immunohistochemical analysis</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Did not address the focused question</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Resembles to a retrospective analysis</td>
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<tr>
<td>Schepers (1993)</td>
<td>Impl Dent</td>
<td>Retrospective case series without control group</td>
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<tr>
<td></td>
<td></td>
<td>Evaluated soft tissue healing and radiographic analysis based on the two-dimensional periapical radiographs</td>
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<tr>
<td></td>
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<td>Titanium implants placed simultaneously</td>
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<td></td>
<td></td>
<td>No control group</td>
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<tr>
<td></td>
<td></td>
<td>Primary outcome was microbiological analysis</td>
</tr>
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<td></td>
<td></td>
<td>No compatibility of the follow-up periods of the different groups</td>
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<tr>
<td></td>
<td></td>
<td>Measurements based on two-dimensional periapical radiographs</td>
</tr>
</tbody>
</table>
The Kappa score for agreement between the reviewers at the abstract and full text selection level, was 0.96 and 0.90, respectively, indicating a high level of agreement [239].

**Study characteristics**

In the 14 included articles (eight RCTs and six CCTs) the efficacy of ARP techniques was evaluated clinically by means of direct measurements of the residual alveolar ridge dimensions during re-entry procedures, radiographically by means of computer tomography or histologically from trephine biopsies taken at re-entry during osteotomies for implant placement (Tables 11-13). Limited data were reported on confounding factors, such as periodontitis, smoking, systemic disease and medication.

The extraction site distribution was fairly heterogeneous. In some studies ARP was performed only in maxillary anterior sockets [232] [236] [237], whereas such restriction was not employed in other studies. The residual bone volume around the investigated sockets, e.g. the presence/absence and width of the buccal bone plate varied from severely compromised [215] [236], to completely intact, buccal bone (Table 12) [207] [210] [232].
<table>
<thead>
<tr>
<th>Study</th>
<th>First author</th>
<th>Year of publication</th>
<th>Type</th>
<th>Reference number</th>
<th>Randomisation</th>
<th>Masking</th>
<th>Calibration</th>
<th>Eligibility Criteria</th>
<th>Follow up</th>
<th>Ethical considerations</th>
<th>Funding</th>
<th>Statistical analysis</th>
<th>Miscellaneous</th>
<th>Estimated Risk of Bias</th>
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<tbody>
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</tbody>
</table>
Table 11  Quality assessment of the included studies (see above)

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</thead>
<tbody>
<tr>
<td>Lekovic 1998 RCT #214</td>
<td>Yes</td>
<td>Yes</td>
<td>N/R</td>
<td>N/A</td>
<td>No</td>
<td>No</td>
<td>Yes</td>
<td>Yes</td>
<td>N/R</td>
<td>1. N/R</td>
<td>2. Patient</td>
<td>3. Insufficient data to determine</td>
<td>Yes</td>
<td>N/R</td>
<td>N/R</td>
<td></td>
</tr>
</tbody>
</table>

N/A not applicable; N/R not reported, T test; C control; RCT randomised controlled trial; CCT controlled clinical trial; PRGF platelet-rich growth factor; CONSORT Consolidated Standards of Reporting Trials; ISRCTN International Standard Randomised Controlled Trial Number Register
<table>
<thead>
<tr>
<th>First author</th>
<th>Year of publication</th>
<th>Type</th>
<th>Methodology</th>
<th>Reference number</th>
<th>Trial characteristics</th>
<th>Population characteristics</th>
<th>Confounding factors</th>
<th>Defect characteristics</th>
<th>Test material (number of sockets/subjects)</th>
<th>Control (number of sockets/subjects)</th>
<th>Surgical management</th>
<th>Follow-up</th>
<th>Alveolar ridge dimension changes in horizontal width Mean/median mm (reference point)</th>
<th>Alveolar ridge dimension changes in vertical height Mean/median mm (reference point)</th>
<th>Implant</th>
<th>Implant complications</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aimetti 2009</td>
<td>RCT Parallel Clin + Histo #232</td>
<td>1 Italy</td>
<td>2 University</td>
<td>3 Private practice</td>
<td>1. 36-68 (51.27 ±8.4) 2. 40 (40)</td>
<td>1. No 2. N/R</td>
<td>1. Maxillary anterior 2. 4-wall configuration</td>
<td>Calcium sulphate Hemitetrhydrate (22/22)</td>
<td>Empty (18/18)</td>
<td>1. Flapless 2. No primary closure 3. Amoxicillin 2g/day for 5 days, Chlorhexidine 0.12% for 2 weeks</td>
<td>1. 3 months 2. N/R 3. Unventful healing</td>
<td>1. T: -2.0 ± 1.1**, C: -3.2 ± 1.8**, *** 2. N/R</td>
<td>1. T: -0.3±1.1*, C: -1.2±0.6*, ** 2. T: -0.2±0.6, C: -0.5±1.2 3. T: -0.4±0.9, C: -0.5±1.1 4. T: 11.3±2.8**, C: 10.0±2.3** (Acrylic stent)</td>
<td>1. Implants were inserted 2. N/R</td>
<td></td>
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</tr>
<tr>
<td>Anitua 1999</td>
<td>CCT Parallel + Split-mouth Histo #233</td>
<td>1 Spain</td>
<td>2 Private practice</td>
<td>3 Hospital</td>
<td>1. T: ±55 (41) C: 38-54 (42) 2. 23 (26)</td>
<td>1. Yes 2. Yes</td>
<td>1. Any 2. Variable</td>
<td>T1: PRGF (5+3/5+3) T2: PRGF-Autologous bone (5/5)</td>
<td>Empty (10+3/10+ 3)</td>
<td>1. Full-thickness 2. Primary closure 3. Amoxicillin 1.5g/day for 5 days</td>
<td>1. 2.5 – 4 months 2. 0 3. N/R</td>
<td>N/A</td>
<td>N/R</td>
<td>1. N/R 2. N/R</td>
<td>1. 'Implants were inserted in both groups’ 2. Some GBR were needed due to buccal dehiscence in the control group</td>
<td></td>
</tr>
<tr>
<td>Barone 2008</td>
<td>RCT Parallel Clin + Histo #210</td>
<td>1 Italy</td>
<td>2 Hospital</td>
<td>3 Private practice</td>
<td>1. 26-69 2. 40 (40)</td>
<td>1. &lt;10/day 2. Yes (treated)</td>
<td>1. Non-molars 2. 4-wall configuration</td>
<td>Corticocancellous porcine bone + collagen membrane (20/20)</td>
<td>Empty (20/20)</td>
<td>1. Full-thickness 2. Primary closure 3. Amoxicillin 2g/day for 4 days + Chlorhexidine 0.12% for 3 weeks</td>
<td>1. 7-9 months 2. 0 3. Unventful healing (pain, swelling)</td>
<td>1. T: -2.5 ± 1.2*, C: -4.5 ± 0.8*, *** 2. N/R</td>
<td>1. T: -0.7±1.4*, C: -3.6±1.5*, ** 2. T: -0.2±0.8, C: -0.4±1.2 3. T: -0.4±0.8, C: -0.5±1.0 4. N/R (Acrylic stent)</td>
<td>1. 'Implants were inserted in both groups’ 2. Some GBR were needed due to buccal dehiscence in the control group</td>
<td></td>
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</tr>
<tr>
<td>Camargo 2000</td>
<td>CCT Split-mouth Clin #234</td>
<td>1 USA, Yugoslavia</td>
<td>2 Private practice</td>
<td>3 University</td>
<td>1. 28-60 (44±15.9) 2. 16 (32)</td>
<td>1. N/R 2. N/R</td>
<td>1. Maxillary anterior, premolars 2. N/R</td>
<td>Bioactive glass + covered by calcium sulphate layer (16/8)</td>
<td>Empty (16/8)</td>
<td>1. Full-thickness with 4 vertical releasing incisions 2. No primary closure 3. Penicillin 1.5g/day for 7 days + Chlorhexidine 0.12% for 2 weeks</td>
<td>1. 6 months 2. N/R 3. Unventful healing</td>
<td>1. T: -3.48±2.68**, C: -3.08±2.41** 2. N/R</td>
<td>1. T: -0.38±0.18, C: -1.00±2.25 (titanium tack) 2. N/R 3. N/R 4. T: 6.43±2.78**, C: 4.00±2.33**, *** (to buccal bone crest)</td>
<td>1. ‘Reintry only’ 2. N/A</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Study</td>
<td>Design</td>
<td>Study Country</td>
<td>Included Centres</td>
<td>Study Sample Size</td>
<td>Study Population</td>
<td>Data Collection</td>
<td>Study Outcome</td>
<td>Study Conclusion</td>
<td></td>
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<tr>
<td>Fiorelli 2005</td>
<td>RCT</td>
<td>USA</td>
<td>28 centres</td>
<td>474</td>
<td>Maxillary anterior, premolars 2. ≥50% buccal bone loss</td>
<td>T1: 1.5mg/ml rhBMP-2 (7/217) T2: 0.75mg/ml rhBMP-2 (7/227) T3: Collagen sponge (7/177)</td>
<td>Empty (7/207)</td>
<td>1. Full-thickness with vertical incisions 2. Primary closure 3. Penicillin (7mg) for 7-10 days + Chlorexidine 0.12%</td>
<td>1. 4 months 2. No drop-outs reported. (3 patients incorrectly randomized, 1 patient received different graft) 3. 250 (T&gt;2)</td>
<td>1. Coronal: T1: +3.27±2.53*, T2: +1.76±1.67*, T3: +0.82±1.40, C: +0.57±2.56, *** (T1 vs T/C) 2. N/R</td>
<td>1. F1: -0.02 ± 1.2, T2: -0.02±1.39*, T3: -1.00±1.40, C: -1.17±1.23*, *** (T1 vs C) 2. N/R 3. N/R 4. N/R</td>
<td></td>
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<tr>
<td>Froum 2002</td>
<td>RCT</td>
<td>USA</td>
<td>Single centre</td>
<td>35-77 (54±11.9)</td>
<td>Any 2. 4-wall, configuration, ≥2mm buccal plate loss</td>
<td>T1: Bioactive glass (10/8) T2: DFDBA (10/8)</td>
<td>Empty (10/10)</td>
<td>1. Full-thickness without vertical incisions 2. Primary closure 3. Doxycycline 100mg/day for 13 days + Chlorexidine 0.12% for 30 days</td>
<td>1. 6-8 months 2. 0 3. Uneventful healing</td>
<td>N/A</td>
<td>N/A</td>
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<tr>
<td>Guarneri 2004</td>
<td>RCT</td>
<td>Italy</td>
<td>35-58</td>
<td>10 (25)</td>
<td>Maxillary, mandibular anterior, premolars 2. socket with ridge resorption ≥50% were excluded</td>
<td>Calcium sulphate Hemihydrate (10/10)</td>
<td>Empty (5/5)</td>
<td>1. Full-thickness without vertical incisions 2. Primary closure 3. Amoxicillin (7mg) for 1 week + Chlorexidine 0.2% for 2 weeks</td>
<td>1. 3 months 2. N/R 3. N/R</td>
<td>N/A</td>
<td>N/A</td>
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<tr>
<td>Lasella 2003</td>
<td>RCT</td>
<td>USA</td>
<td>28-76 (51±13.6)</td>
<td>10 (24)</td>
<td>Maxillary, mandibular anterior, premolars and mandibular premolars 2. N/R</td>
<td>Tetraacycline hydrated DFDBA + collagen membrane (12/12)</td>
<td>Empty (12/12)</td>
<td>1. Full-thickness without vertical incisions 2. No primary closure 3. Doxycycline 200mg/day for 1 week + Chlorexidine 0.12% for 2 weeks</td>
<td>1. 4 or 6 months (combined) 2. 0 3. N/R</td>
<td>1. T: -1.2 ± 0.9*, C: -2.6 ± 2.3*, 2. N/R</td>
<td>1. T: +1.3±2.0, C: -0.9±1.6*** 2. T: -0.1±0.7, C: -1.0±0.8*** 3. T: -0.1±0.7, C: -0.8±0.8*** 4. N/R (Acrylic stent) 1. Implants successfully placed at all sites 2. Some sites had slight dehiscence and required further augmentation</td>
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<tr>
<td>Lekovic 2007</td>
<td>RCT</td>
<td>Yugoslavia</td>
<td>49.8</td>
<td>10 (20)</td>
<td>Maxillary and mandibular incisions, premolars 2. N/R</td>
<td>e-PTFE membrane (10/10)</td>
<td>Empty (10/10)</td>
<td>1. Full-thickness with vertical releasing incisions 2. Primary closure 3. Penicillin 1g/day for 7 days + Chlorexidine 0.2%</td>
<td>1. 6 months 2. 3/10 drop-outs due to premature membrane exposure 3. 3/10 exposed, 7/10 no infection</td>
<td>1. 10/10: T: -1.8±0.51, C: -2.6±2.0**, 2.0±1.3*** 7/10: T: -1.7±1.75, C: -4.43±2.42*, *** 3/10: T: -2.00±0.00, C: -4.33±0.88* 2. N/R</td>
<td>1. 10/10: T: -0.5±0.22, C: -1.2±0.13**, *** 7/10: T: -0.28±0.18, C: -1.0±0.04**, *** 3/10: T: -1.0±0.58, C: -1.6±0.33 (Titanium lack) 2. N/R 3. N/R 4. N/R</td>
<td>1. 10/10: T: -2.0±0.86*, C: -3.0±0.63, *** 7/10: T: -5.4±2.1*, C: -2.9±1.6, *** 3/10: T: -3.6±1.20, C: 4.3±1.45 (to buccal bone crest) 1. Reentry only 2. N/A</td>
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<tr>
<td>Study</td>
<td>Country</td>
<td>Sample Size</td>
<td>Treatment</td>
<td>Control</td>
<td>Outcomes</td>
<td>Note</td>
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<tr>
<td>Lekovic 2006 RCT</td>
<td>Yugoslavia</td>
<td>28-70 (47.5±10.3)</td>
<td>N/R</td>
<td>N/R</td>
<td>Empty (15/6)</td>
<td>N/A</td>
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<tr>
<td>Nevins 2006 RCT</td>
<td>USA / Italy</td>
<td>2.9 (35)</td>
<td>N/R</td>
<td>N/R</td>
<td>Empty (16/16)</td>
<td>N/A</td>
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<tr>
<td>Serino 2003 CCT</td>
<td>Italy</td>
<td>35-64</td>
<td>N/R</td>
<td>N/R</td>
<td>Empty (13/12)</td>
<td>N/A</td>
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<tr>
<td>Serino 2008 CCT</td>
<td>Italy</td>
<td>32-64</td>
<td>N/R</td>
<td>N/R</td>
<td>Empty (9/9)</td>
<td>N/A</td>
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* p<0.05; statistically significant intra-group difference, baseline to final; ** p<0.01 statistically highly significant intra-group difference, baseline to final; *** p<0.005 statistically significant inter-group difference, between test and control (bold); N/A not applicable; N/R not reported; T test; C control; M = month(s); Clin clinical analysis; Histo histological analysis; Radiogr radiographic analysis; RCT randomised controlled trial; CCT controlled clinical trial; PRGP plasma rich in growth factors; DFDBA demineralised freeze-dried bone allograft; FDBA freeze-dried bone allograft; ePTFE expanded-polytetrafluorethylene; PG/PL polyglycolide/polylactide; DBBM demineralised bovine-bone mineral

Table 12 Clinical and radiographic assessment of the included studies (see above)
<table>
<thead>
<tr>
<th>First author</th>
<th>Year of publication</th>
<th>Type</th>
<th>Reference number</th>
<th>Number of biopsies (test material)</th>
<th>Histomorphology</th>
<th>Histomorphometry (mean or median %)</th>
<th>Statistical difference between test and control</th>
</tr>
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<tbody>
<tr>
<td>Aimetti 2009</td>
<td>RCT</td>
<td>3 M</td>
<td>#232</td>
<td>T: N/R 22? (MGCSH) C: N/R 18?</td>
<td>No residual graft material. No inflammatory infiltrate. New bone formation in all specimens, 100% living trabecular bone with woven and lamellar structure. 100% living bone (mostly woven) in all biopsies. Lamellar bone remodeling was starting.</td>
<td>Trabecular bone: T: 58.8±3.5 C: 47.2±7.7 Residual substitute material: T: 0.0 C: N/A Woven bone: Coronal: T: 83.6±6.6 C: 89.9±7.6 Middle: T: 59.6±13.2 C: 81.1±7.6 Apical: T: 56.4±10.9 C: 77.8±8.1</td>
<td>Lamellar bone: Coronal: T: 16.4±6.6 C: 11.1±7.6 Middle: T: 40.4±13.2 C: 18.9±7.6 Apical: T: 43.6±10.9 C: 22.2±8.1</td>
</tr>
<tr>
<td>Anitua 1999</td>
<td>CCT</td>
<td>2.5 – 4 M</td>
<td>#233</td>
<td>T: N/R (PRGF± autogen bone) C: N/R</td>
<td>Compact mature bone with well-organized trabeculae and morphology in 8/10 patients. Connective tissue with non-organized trabeculae in 2/10 patients. Significant intra-group differences 10 vs. 16 weeks! Connective tissue fills the main part of the defect. No mature bone.</td>
<td>N/R</td>
<td></td>
</tr>
<tr>
<td>Barone 2008</td>
<td>RCT</td>
<td>7 – 9 M</td>
<td>#210</td>
<td>T: 20 (Corticocancellous porcine bone+ collagen membrane) C: 20</td>
<td>Residual graft material embedded in newly formed bone in all specimens. Complete bone fill. Typically trabecular bone pattern. Large marrow spaces filled with adipocytes. Lamellar bone was also present within the bone marrow.</td>
<td>Total bone volume: T: 35.5±10.4 C: 25.7±9.5 Connective tissue: T: 36.6±12.6 C: 59.1±10.4 Residual graft material: T: 29.2±10.1 C: N/A</td>
<td>Bone: T&gt;C* Connective tissue: T&lt;C*</td>
</tr>
<tr>
<td>Fiorellini 2005</td>
<td>RCT</td>
<td>4 M</td>
<td>#215</td>
<td>T1: 16 (hBMP-2 1.5mg/ml) T2: 15 (0.75mg/ml) T3: 11 (Collagen sponge) C: 14</td>
<td>No evidence of inflammation or residual graft. Trabecular bone formation in 2/3 of the samples. Mineralized tissue formation presented with different level of remodeling. Minor osteoclastic activity. No comparison reported between T and C!</td>
<td>N/R</td>
<td></td>
</tr>
<tr>
<td>Study</td>
<td>Type</td>
<td>Duration</td>
<td>#</td>
<td>Treatment</td>
<td>Control</td>
<td>Findings</td>
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<tr>
<td>Froum 2002</td>
<td>RCT</td>
<td>6 – 8 M</td>
<td>#207</td>
<td>T1: 10 (Bioactive glass) T2: 10 (DFDBA) C: 10</td>
<td></td>
<td>T1: New bone formation. Osteoid surrounding and penetrating the bioactive glass particles. T2: Varying degrees of reossification around DFDBA. N/R Vital bone: T1: 59.5 T2: 34.7 C: 32.4 Connective tissue: T1: 35.3 T2: 51.6 C: 67.0 Residual bone substitute: T1: 5.5 T2: 13.5 C: N/A Connective tissue: T1&lt; T2 or C*</td>
<td></td>
</tr>
<tr>
<td>Guarnieri 2004</td>
<td>CCT</td>
<td>3 M</td>
<td>#235</td>
<td>T: 10 (MGCSH) C: 5</td>
<td></td>
<td>Almost complete absence of MGCSH. Absence of connective tissue and inflammatory cells. In all sections trabecular bone formation with no differences between the apical, middle and coronal levels. Less bone formation compared to test sites. Trabecular bone area: T: Coronal: 58.6±9.2 Middle: 58.1±6.2 Apical: 58.3±7.8 C: ≤ 46 No statistical significance could be drawn due to small number of control specimens.</td>
<td></td>
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<tr>
<td>Lasella 2003</td>
<td>RCT</td>
<td>4 – 6 M</td>
<td>#212</td>
<td>T: 4M: 5, 6M:7 (Tetracycline hydrated FDBA + Collagen membrane) C: 4M: 5, 6M: 5</td>
<td></td>
<td>Residual graft particles surrounded by woven bone or by connective tissue. Similar amount of total bone and trabecular spaces as in test. (No biopsy from 2 C sites due to minimal bone fill) Vital bone: 4M T: 31±9 C: 58±11 6M T: 25±17 C: 50±14 Combined T: 28±14 C: 54±12 No statistical significance could be drawn due to small number of control specimens.</td>
<td></td>
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<tr>
<td>Nevins 2005</td>
<td>RCT</td>
<td>6 M</td>
<td>#236</td>
<td>T: 5 (DBBM) C: 5</td>
<td></td>
<td>DBBM granules present. Apically integrated in cancellous bone but coronally in soft tissue. No signs of inflammation or foreign body reaction. New bone formation No comparison made.</td>
<td></td>
</tr>
<tr>
<td>Pelegrine 2010</td>
<td>RCT</td>
<td>6 M</td>
<td>#237</td>
<td>T: 7 (Autologous bone marrow) C: 6</td>
<td></td>
<td>Mineralized bone: T: 45.0 C: 43.75 No significant difference.</td>
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</table>

*p<0.05; statistically significant difference between test and control (bold)

T test; C control; M month(s); N/R not reported; N/A not applicable; vs. versus; TBV total bone volume; MGCSH medical grade calcium sulphate hemihydrate; DFDBA demineralised freeze-dried bone allograft; FDBA mineralised freeze-dried bone allograft; DBBM demineralised bovine-bone mineral; PG/PL polyglycolide/polylactide

Table 13  Histological assessment of the included studies (see above)
Intervention characteristics

With regard to the techniques or materials used for ARP, the included studies were grouped into three categories (Table 12);

1. Bone grafts/substitutes
2. GBR
3. Biological active materials.

In the majority of the included studies various bone grafts were utilised, such as autologous bone marrow [237], plasma rich in growth factor (PRGF) with or without autologous bone [233], DFDBA [207], DBBM [236], calcium sulphate hemihydrates [232] [235] and bioactive glass [207]. Alloplastic polyglycolide/polylactide (PGPL) sponge was also employed [209] [238]. GBR technique was applied using non-resorbable e-PTFE [213] or resorbable (PGPL) barrier [214]. Resorbable collagen membrane was also employed in combination with FDBA [233] or corticocancellous porcine bone [210]. Biological active material, namely recombinant human bone morphogenetic protein-2 (rhBMP-2) was used on a collagen sponge carrier in one study [215]. Neither the ‘soft tissue punch’/socket seal, nor the ‘ice-cream cone’/bio-col techniques were identified among the included studies [216] [240]. Primary flap closure was achieved in nine out of 14 studies, while the sockets left uncovered in the rests. Various types and amounts of antibiotics and antiseptic rinses were administered for different duration in studies reporting on postoperative care. Finally, average healing period ranged from one to nine months.

Outcome characteristics

Clinical outcomes

Eight out of the 14 included studies investigated the efficacy of various ARP techniques to preserve the pre-extraction ridge dimensions using intra-surgical hard tissue measurements taken during re-entry procedure [209] [215] [212] [213] [214] [232] [234] [237]. In these studies ARP was performed in 137 sockets of 119 patients and compared to 120 sockets that left to heal without any treatment in a total of 92 patients (Table 12, Fig. 25, 26).
Bone ‘graft’

Four studies evaluated changes in AR dimensions following grafting of the socket. Two studies were RCTs [232] [237] and two were CCTs [209] [234]. Healing time varied from three to six months [209] [234] [232] [236]. The horizontal (bucco-lingual) changes of the alveolar ridge were assessed in three studies [234] [232] [237]. The AR reduced in width from baseline to re-entry between -1.0 mm and -3.5±2.7 mm following ARP (p<0.05) and between -2.5 mm and -3.2±1.8 mm in the control groups (p<0.05). In two out of the three studies the width reduction was statistically significantly smaller in the test groups compared to the controls [232] [237].

Four studies investigated the mean change in ridge height at the mid-buccal aspect [237] [209] [234] [232]. The AR height changed from baseline to re-entry between +1.3±1.9 and -0.5±1.1 mm following ARP, and between -0.8±1.6 and -1.2±0.6 mm in the control groups. The height reduction between baseline and re-entry was not statistically significant in one study in both test and control groups [234], while one study reported an increase in height instead of loss following ARP with a PGPL sponge (p<0.05 [209]). In two out of the four studies the height reduction was statistically significantly smaller in the test groups compared to the controls [232] [237].
The vertical dimension changes at the mesial and distal aspects of the socket were measured in two studies [232] [209] and did not present any statistically significant difference for both groups.

Three studies captured data on socket fill and reported statistically significant differences between baseline and re-entry in both groups [232] [234] [237], but only one reported statistically significantly higher socket fill, where bioactive glass was covered by calcium sulphate, compared to the unassisted healing [234].

**GBR**

Four studies evaluated changes in AR dimensions following ARP with GBR alone [213] [214], or in combination with bone graft [210] [212]. Three studies were RCTs [210] [212] [214] and one was CCT [213]. Healing time varied between four and nine months.

Horizontal (bucco-lingual) changes of the AR were assessed in all four studies. AR width reduction from baseline to re-entry varied between -1.2±0.9 and -2.5±1.2 mm in the GBR treated sockets and between -2.6±2.3 and -4.6±0.3 mm in the control groups. With the exception of one study [212], a statistically significantly smaller reduction of the alveolar ridge width was observed when e-PTFE [213], PGPL [214], or collagen membranes in combination with xenograft [210] were used.

All the four studies investigated the mean change in AR height at the mid-buccal aspect. The AR height changed from baseline to re-entry between +1.3±2.0 and -0.7±1.4 mm in the ARP groups and between -0.9±1.6 and -3.6±1.5 mm in the control groups. The resorption in the ARP group was not statistically significant in three out of four studies [212] [213] [214]. All studies reported statistically significantly less post extraction reduction in AR height when the socket was treated by GBR compared to unassisted healing.

Vertical dimension changes at mesial and distal aspects of the socket were measured in two studies [210] [212]. The observed differences between baseline and re-entry were not statistically significant in both groups. In one out of the two studies the height
reduction was statistically significantly smaller in the test group compared to the control [212].

Two studies captured data on the socket fill [213] [214] and reported statistically significant socket fill in both groups between baseline and re-entry, as well as between tests and controls.

No data were found on either initial buccal plate thickness or alteration of bone volume. However, one study measured the buccal bone thickness loss and reported statistically significantly less reduction in the ARP group [237].

**Radiographic measurements**

Two RCTs, reporting on 3D radiographic assessment, met the inclusion criteria [215] [236]. The healing time varied from one to four months. In one study, where the post extraction socket was grafted with a radiopaque material (DBBM), treatment resulted in significantly less reduction in radiographic AR height compared to unassisted socket healing [236]. The test group in the other study, where the higher concentration (1.5mg/ml) of rhBMP-2 was utilised [215], resulted in a mean increase of the radiographic AR width by 3.27±2.53 mm at the most coronal part, compared to the 0.57±2.56 mm increase in the group of unassisted healing. AR height was reduced by 0.02±1.2 mm in the same test group and by 1.17±1.23 mm in the control group (Table 12). The differences between test and control were statistically significant.

**Histological results**

Eleven studies carried out a histological analysis based on trephine biopsies retrieved at re-entry [232] [233] [210] [215] [207] [235] [212] [236] [237] [209] [238]. Seven studies were RCTs [232] [207] [215] [210] [212] [236] [237] and four were CCTs [209] [238] [233] [235]. In these studies, ARP was performed in 181 sockets of 158 patients and compared to 149 sockets that left to heal without any treatment in 131 patients (Table 13). Only two studies reported statistically significantly higher trabecular bone volume following ARP in comparison to unassisted socket healing [232] [210] and two studies reported statistically significantly more connective tissue in the post extraction socket when no ARP was performed [210] [207]. On the contrary, one study reported more vital bone in the unassisted socket healing group compared to
the ARP group [212]. None of the differences of the investigated histomorphometric parameters reached statistical significance in other studies (Fig. 27).

![Fig. 27 Histological differences between test and control.](image)

**Bone 'graft'**

Eight studies evaluated histologically the healing of post-extraction sockets following the application of some type of bone grafts/substitutes [232] [233] [207] [235] [236] [237] [209] [238]. Four studies were RCTs [232] [207] [236] [237] and four were CCTs [233] [235] [209] [238]. New mineralised bone was observed at various levels in all studies in both ARP and control groups in a healing period from 2.5 to eight months. Connective tissue occupied a portion of the socket in both groups. When DBBM, DFDBA or BG were used, the graft particles were embedded either in new bone or in connective tissue. In most studies there was no significant difference in the type of healing or amount of bone formation between test and controls.

**GBR in combination with graft**

GBR in combination with graft was utilized in two RCTs. ARP with a collagen membrane and deproteinized porcine bone resulted in statistically significantly higher new bone and lower connective tissue formation after seven to nine months of healing in comparison to unassisted socket healing [210]. However, residual graft materials were present in the ARP biopsies. FDBA and collagen membrane resulted in similar amounts of new bone formation to untreated sockets, although more vital bone was observed in the untreated sockets at four to six months of healing (p>0.05) [212].

DOI:10.14753/SE.2015.1708
Biological active material

RhBMP-2 in a collagen sponge carrier was completely resorbed at four months following ARP regardless of the concentration of the growth factor [215]. Mineralised tissue was found and trabecular bone formation was noticed in two third of both the test and control biopsies in the RCT.

Adverse events, complications

Adverse events were reported in six RCTs [215] [207] [210] [214] [232] [237] and four CCTs [209] [213] [234] [238] including oedema, pain, erythema and membrane exposure/infection. In two studies more adverse events i.e. oedema, erythema [215] or membrane exposure [213] were observed in the ARP group compared to the natural socket healing. No comparison between tests and controls were reported in the other studies (Table 12).

Feasibility of implant placement

Seven studies [207] [209] [212] [232] [235] [236] [238] reported that implant placement in the previous sockets were successful, but no differences between the ARP and untreated sites were revealed. The outcome of implant placement remained unclear in one article [233] and only re-entry without implantation was performed in three trials [213] [214] [234]. Three studies reported that at the stage of implant placement less need of further augmentation was observed in the ARP group compared to the controls (i.e. less [215] or no sites [210] [237] in the ARP group presented with residual dehiscence or fenestration defects around the inserted implants) (Table 12, Fig. 28).
Patient-reported outcome and health economics

No data were found for patient-reported outcome measures or health economic evaluation.

Quality assessment

Considerable heterogeneity was found among the studies in terms of methodological quality. Detailed description of the quality assessment of the included studies is presented in Table 11. Among the 14 included controlled studies, eight were randomised [232] [210] [215] [207] [212] [214] [236] [237], although in four of them the randomisation technique was not reported [215] [232] [236] [237]. None of the RCTs reported the method of allocation concealment. Masking of the examiner was reported at the clinical level in two out of eight [212] [214], at radiological level in one out of two [215] and at histological level in four out of 11 studies [232] [233] [210] [207]. Examiner calibration was declared in three articles [232] [215] [212], whilst inclusion and exclusion criteria were defined in seven publications [207] [210] [212] [232] [233] [236] [237]. Apart from three studies [233] [210] [236], all the other reported the approval of the ethical committee. Three studies were funded by industry [234] [207] [215], two studies by academic institution [235] [238] and the remaining nine did not report the source of funding.

Nine trials implemented patient-based analysis [232] [234] [215] [210] [212] [213] [214] [237] [238], whilst the extraction site served as unit of analysis in the remaining five investigations [207] [209] [233] [235] [236]. Sample size calculation was reported only in three studies [232] [215] [212], although with insufficient data to evaluate the validity of the calculations. Statistical analysis was appropriately carried out and described in one study only [237]. Appropriate statistics were either not carried out [207] [209] [215] [210] [233] [236] [235], or the reported data were insufficient to determine the validity [232] [212] [213] [214] [233] [238]. In addition, no RCTs were either registered with ISRCTN or reported using the CONSORT guidelines (Table 11).

Risk of bias

Four studies were classified as moderate risk of bias [210] [207] [212] [214] and the rest were categorised as high risk of bias (Table 11, Fig. 29).
Fig. 29 Quality assessment of the included trials
6. DISCUSSION

6.1. Limits of periodontal regeneration

The main focus of the present investigations were to provide histological insight on the regenerative potential of EMD combined with BCP (Study 1) or nano-HA (Study 2) for the treatment of human periodontal intrabony defects.

6.1.1. Treatment with EMD and SBC

This study demonstrated that treatment of intrabony defects with a combination of EMD + BCP may lead to improvement of clinical parameters, such as PPD reduction and CAL gain (Table 1). The lack of adverse reactions observed in all treated subjects is in accordance with findings from previous clinical and histological studies that showed that EMD and BCP are well tolerated and do not elicit allergic or foreign body reactions [113] [77] [20] [78] [114] [60] [79] [115] [21] [116] [117] [36]. The mean CAL gain of 3.0 mm obtained in this study is in line with previous findings from human histologic studies evaluating the treatment of advanced intrabony defects with EMD alone or in combination with DBBM or BG [33] [34] [41] [117] [36].

Considerations of the clinical outcomes

In contrast to our findings, the increased recession is an often observed phenomena following surgical periodontal therapy. It suggests that in very advanced deep and wide intrabony defects the additional use of graft material may prevent flap collapse and consequently soft tissue recession. This finding corroborates the results from controlled, clinical studies comparing the use of EMD in conjunction with ABP, DFDBA or DBBM in the treatment of large intrabony defects [38] [40] [39]. The combined treatment in these studies resulted in significantly less recession and consequently greater CAL gain compared to treatment with EMD alone that points to the potential clinical indication of a combination approach. BCP was employed to fill the intrabony periodontal defect in a recent case-control study [118]. The authors reported a mean CAL gain of 4.6 mm (3.0 mm in our trial) following the treatment with a combination
of BCP and guided tissue regeneration, showing the biocompatibility and promising clinical properties of BCP.

To the best of our knowledge only two recent clinical trials investigated the treatment of periodontal defects with a combination of EMD+BCP. In a randomized controlled multicenter trial EMD+BCP (test; n=38) was compared to EMD alone (control; n=35) [119]. At six months after surgery all investigated clinical parameters represented significant improvements in both groups (PPD: 1.93±1.8 vs. 2.55±1.8; CAL: 1.31±1.8 vs. 1.83±1.6; REC: 0.62±1.1 vs. 0.72±1.1; bone sounding: 2.01±2.1 vs. 2.07±1.2; when compared test to control, in millimeters, respectively). These results are slightly inferior, but still comparable to the clinical outcomes of our investigation. The intergroup differences failed to show statistical significance. Their results did not support the theory that the additional bone substitute scaffold might improve the clinical outcome, especially the hard tissue fill of the intrabony defect, compared to EMD alone. Nevertheless, the investigators concluded that the null hypothesis that the applied combined treatment of EMD+BCP is inferior to EMD alone, was refuted.

Another, more recent, randomised controlled clinical trial (RCT) that was conducted in collaboration with the University of Szeged, evaluated the long-term effect of EMD+BCP [120]. The authors reported similar outcomes in the test (EMD+BCP; n=12), compared to the control group (EMD; n=12) at 12 months. Both groups lead to significant improvement in terms of PPD reduction (from 8.8±1.5 to 4.3±0.9 mm vs. from 8.8±1.0 to 4.1±0.5 mm) and CAL gain (from 10.8±1.6 to 7.4±1.6 mm vs. from 10.4±1.3 to 6.9±1.0 mm) in the test compared to control, respectively. No significant differences were reported for any of the variables between the two groups indicating that the added BCP did not additionally improve the outcomes obtained with EMD alone. Slight, but not statistically significant deterioration of the investigated clinical parameters between 1 and 4 years was reported, concluding that the results, irrespective of the treatment modalities, were sustainable for four years.

**Histological considerations**

Human histological studies evaluating the regenerative potential of BCP alone or in combination with EMD or other regenerative materials (e.g. barrier membranes), in surgical treatment of periodontal intrabony defects are currently not available; therefore, direct comparison is not possible. The histological evaluation of our experiment showed
that new cementum with associated PDL was found in six out of nine biopsies, whereas
the formation of new bone or mineralized bone-like tissue around the graft particles was
observed only occasionally (Table 1). These observations seem to be in contrast with
the findings from a human histologic study by Windisch et al [36]. They compared the
effect of EMD and GTR on human periodontal defects by clinical, radiographical and
histological means. The use of EMD alone resulted in 5.00±0.63 mm PPD reduction and
2.67±1.03 mm CAL gain, which are comparable to our clinical results. However, the
histological results demonstrated 1.81±1.71 mm new cementum and 0.78±0.97 mm new
bone formation that looks superior to our data. Regarding this indirect comparison, the
added BCP does not seem to improve the regenerative effect of EMD alone.

Furthermore, our human histological observations are in contrast with the findings from
a histologic study in dogs evaluating the tissue response of intrabony periodontal defects
treated with BCP [60]. At 6 months following surgery, histological analysis indicated
formation of new connective tissue attachment and accelerated bone formation around
the BCP particles.

The probable osteoconductive capacity of BCP in non-periodontal defects is indeed
scrutinized in histological studies [78] [121] [122] [123] [79]. Comparable findings
were reported in a histologic and histomorphometric study in minipigs evaluating the
healing of standardized mandibular bone defects filled with BCP [78]. At 24 weeks, the
graft material was surrounded by newly formed bone, but showed only limited
remodeling. These findings seem to corroborate the observations from the present study
where, in most specimens, the graft particles were still present at 9 months following
surgery.

BCP covered by a collagen membrane was investigated in dehiscence-type defects
around dental implants in dogs [79]. Histomorphometric and immunohistochemical
analysis, performed at 9 weeks following reconstructive surgery, revealed that BCP
particles were completely integrated into a secondarily formed network of spongiosa.
However, the results failed to show any signs of graft resorption as evidenced by a lack
of osteoclastic activity at the surface of the particles. In a very recent experiment of the
same group led by Frank Schwarz, BCP or a collagen-coated DBBM were placed into
the dehiscence defects around dental implants of dogs and covered by a resorbable
collagen barrier [123]. At nine weeks, both types of bone substitutes were integrated in
spongiosa, in addition, dissolution of some of the BCP granules were observed. Bone remodeling was more pronounced in the BCP group compared to the DBBM. The above positive histological results of animal studies highlight the importance of the ultimate test of the new material in a human histological setting. It also corroborates the frequently cited principle of Thorkild Karring that the necessary positive preclinical histological outcome should not be considered more just a chance to step forward and see, whether the material or method may or may not work in a human (histological) environment.

Taken together, the above findings suggest that in human intrabony defects, the bone substitute itself does not seem to significantly influence the rate and type of healing. The combined treatment of BCP and EMD failed to promote additional bone formation over EMD alone in our study that was later corroborated by two clinical RCTs [119] [120]. In addition, the small amount of newly formed tissue, observed histologically may query the necessity of the adjunctive use of the present alloplast and in the same time highlights the significance of human histological investigations.

6.1.2 Treatment with nano-HA

No adverse reactions, such as allergies or abscesses occurred in any of the patients, indicating that the used grafting material is biocompatible and well tolerated. This corroborates the findings from previous reports [94] [95] [96] [97] [98] [99] [100] [101] [102] [103] [104] [106] [105].

Surgically, minuscule mechanical stability of the implanted material was observed. However, its creamy consistence enabled to establish an intimate interface between the bony wall of the pocket. This direct contact might enable a close “bridging” to the osteoprogenitor cells.

Radiographical observations

The above feature may elucidate the radiographic observation at three months following implantation of nano-HA, namely the rapid dissolution of the dense shade of the cortical pocket wall, and the increased density inside the former intrabony periodontal defect, compared to the immediate postoperative IOPA (Fig. 6a-d). Since nano-HA does not
bear high radiopacity, such as sintered HA ceramics, β-TCP, ABP or DBBM, the observable increased radiopacity could be explained by the rapid mineralization of the newly formed tissue in the former pocket.

**Clinical results**

The clinical findings correspond to those reported in the randomized controlled studies [106] [105]. In the first RCT intrabony defects were treated either with OFD alone or combined with nano-HA [105]. The additional application of the alloplast yielded a mean PPD reduction of 3.9 mm and a mean CAL gain of 3.6 mm compared to 2.6 mm and 1.8 mm following OFD alone. The intergroup comparison demonstrated significantly more PPD reduction (P = 0.012) and CAL gain (P = 0.005) in the nano-HA group compared to the OFD group. A subsequent study has evaluated the treatment of intrabony defects by papilla preservation flap surgery with or without the application of nano-HA [106]. At 6 months after surgery, both treatments yielded significant improvements compared to baseline. Intergroup comparison demonstrated statistically significantly higher mean PPD reduction (4.3 ± 1.6 mm) and mean probing bone level gain (4.3 ± 1.4 mm) in the nano-HA group compared to the controls (2.9 ± 1.1 and 2.6 ± 1.4 mm, respectively).

In a recent randomised controlled split mouth clinical study, nano-HA was compared to ABP in periodontal intrabony defects as adjunct to OFD [124]. Six months after surgery both therapies resulted in statistically significant PPD reduction (3.21 vs. 3.37 mm) and CAL gain (2.62 mm vs. 2.38 mm) in the nano-HA vs. ABP group, respectively. The PPD changes reported in the aforementioned studies well concord to those obtained in our investigation (i.e. 4.0 mm versus 3.9, 4.3 and 3.21 mm) (Table 2). The detected slight discrepancies in CAL gain (i.e. 2.5 mm versus 3.6, 4.3 and 2.6 mm) may be related to differences in defect configuration. In the present study the majority of the defects displayed a predominantly one- and two-wall configuration, while the defects in the clinical RCTs presented with either two or three-wall configuration. It is well known that three-walled defects have a significantly higher healing potential compared to the one-walled ones [72].

**Histological considerations**

The histological evaluation has, however, indicated that in all six biopsies the healing was predominantly characterized by formation of a long junctional epithelium along the
debrided root surfaces, while some limited periodontal regeneration was only observed in the other three defects (Table 2). In those three cases, formation of cementum, periodontal ligament and bone was confined to the apical portion of the defects. Remnants of the implanted material were encapsulated in connective tissue without apparent signs of circumferential bone formation. This was detected in two out of the six biopsies. The present outcome indicates that the material is resorbable, although it has no visible effect on enhancing periodontal regeneration. It should be kept in mind that two subjects (#01, #06) among the three with no regeneration, were smokers, which is considered as a detrimental factor in periodontal regenerative therapy [34] [255]. The decision to include light smokers was taken in order to amplify the external validity of the trial [256].

The outcome of the present study emphasizes that neither clinical nor radiographical evaluation are adequate means to demonstrate periodontal regeneration [125] [67]. It has been extensively demonstrated that positive clinical outcomes such as PPD reduction, CAL gain or defect fill may not necessarily reflect a regenerative type of periodontal healing [57] [58] [59] [60] [61] [62] [63] [64] [65]. Our results may well be in harmony with the evidence in the literature suggesting that the surgical treatment of human intrabony defects with alloplastic materials predominantly results in LJE with limited or no periodontal regeneration. The alloplast is frequently found to be embedded in connective tissue [58] [61] [62] [63] [64] [65].

The present findings corroborate the results of the above human histological studies and provide further evidence that the use of alloplastic materials alone, for the treatment of periodontal intrabony defect, possess limited biological value.

Finally, a crucial aspect needs to be addressed when interpreting the histologic findings of both investigations (Study 1 and 2), namely that human histological studies are naturally associated with inherent weaknesses. Merely the cases, promising minute success, could be considered as biopsy specimens for ethical reasons. Thus, the sometimes inconsistent finding of periodontal regeneration may be due to the histologic evaluation of hopeless or irrational-to-treat cases. All teeth selected for the present investigation displayed very advanced destruction of the periodontal supporting apparatus, which may have limited their regenerative potential. These intrabony defects
probably would not be considered as ideal candidates for reconstructive therapy in a controlled clinical trial or in the daily periodontal practice. Nevertheless, previous similar human histological case series have indeed provided evidence for periodontal regeneration following the use of EMD, GTR or rhGDF-5 with and without the use of bone substitutes [35] [32] [34] [72] [73] [69].

In both of our studies, only a small number of defects were evaluated and a control group was lacking. Undoubtedly, it would have been better to include more defects and a control group, treated with OFD alone, allowing for adequate statistical analysis. One shall bear in mind though that it is extremely cumbersome to recruit, manage and compensate a large number of patients in a powered human histological parallel armed RCT. Furthermore, it requires considerable amount of human and financial resources that was not achievable in this case.

6.2. Peri-implant dehiscence defect therapy with SBC and PEG

The present investigation demonstrated that osseointegration and bone regeneration in dehiscence defects at implants with hydrophilic surface, treated with or without PEG membrane and BCP, can be achieved and is not negatively affected by functional loading. However, the amount and quality of the newly formed mineralized tissue along with their contact to the implant is not predictable.

In this study, the particular defect configuration (chronic + acute) was aimed to resemble clinically challenging dehiscence defects, which occurs during implant placement in thin resorbed alveolar ridges. The standardized acute dehiscence defects were created by gradually thinning the buccal bone plate, hence the hydrophilic SLActive implants were inserted in the ‘knife-edge’ ridges. This type of complex defects that we encounter in clinical practice are usually more demanding in terms of bone regeneration. However, by using solely chronic type of osseous defects, the standardization of the defect in terms of dimensions and configuration could not have been assured [176] [177]. Such standardization of dehiscence type defects allowed the accurate comparison between the different groups [178].
Hydrophilic implant surface

This recently developed chemically modified hydrophilic implant surface (SLActive), which represents high free surface energy, has been shown to influence pro-osteogenic and pro-angiogenic gene expression, to enhance production of local growth factors and osteoblast activity and to accelerate osteogenic differentiation and mineralization compared to smooth surface, conventional SLA surface or dual acid-etched surface with a calcium phosphate modification [179] [180] [181] [182] [183] [184] [170] [185] [186].

In the present investigation, histological observation of the SLActive implant inserted in pristine sites showed mature lamellar bone in contact with the implant surface. Furthermore, this study demonstrates the stability of the established osseointegration after short-term functional loading at SLActive implants. The successful osseointegration of SLActive titanium surfaces is in line with previous findings from histomorphometric studies reporting promoted healing and enhanced bone apposition even in circular peri-implant defects and dehiscence-type defects in comparison with non-hydrophilic implant surface [180] [187] [188] [189] [184].

Irrespective of loading, osseointegration in the dehiscence defect seemed to be supported by the presence of the SLActive titanium surface in situations where no graft/GBR was applied. A thin layer of bone was observed in direct contact with the implant surface in the defect area and new bone formation proceeded coronally, along the SLActive implant surface. Furthermore, bone regeneration had taken place to a significant extent within the defect. However, the defect was in no case completely filled and a number of threads were always exposed at the most coronal point of the implant, demonstrating that the chosen model was indeed of critical size.

It has to be highlighted that the loaded control group of dehiscence defect, where no material was placed onto the exposed buccal side of the implants (N), presented the highest percentage of bone-to-implant contact (BIC%: 64.1 ± 12.8) and the smallest remaining dehiscence i.e. the lowest distance between the most coronal BIC to the implant shoulder (S-BIC: 1.93 ± 0.69 mm) among all other groups. This data indicates that the modified hydrophilic SLActive implant surface per se may have an osteoconductive effect that improves osteogenic response.
Biphasic calcium phosphate bone substitute
In the present investigation, when a synthetic bone graft was used alone, or combined with GBR osseointegration was observed along the implant surface. However, while bone formation around the graft particles could be observed in the apical portion, in a number of specimens, a variable amount of particles were encapsulated by soft tissue in the coronal portion of the defect. This is in agreement with previous studies where xenografts have been used in combination with GBR [190] [147] [172] or when synthetic bone substitute was used [191] [192]. Thus, again, the positive effect in bone-to-implant contact may be attributed to the SLActive surface rather than the present alloplastic bone replacement material.

In addition, it should be noted that the seemingly inferior bone promoting property of the present BCP for peri-implant dehiscence defects, demonstrated histologically, could not be detected by merely clinical and radiographic observation [193]. This finding is in harmony with the result of the experiment of the treatment of periodontal defects, presented in this thesis, namely that the positive clinical outcome following use of the same BCP material does not necessarily supported by histological evidence [73].

Polyethylene Glycol-Based Hydrogel Membrane
One of the main aims of the present study was to test the novel polyethylene glycol hydrogel membrane. It is activated by mixing the two liquid components and is applied in a gel state. Thus, the membrane has limited space-maintaining ability by itself and was designed by the manufacturer to be placed in combination with a bone substitute. Previous experimental studies have demonstrated that this type of membrane is cell-occlusive [165] and has positive results on bone regeneration around implants inserted in jaw bone as well in calvaria defects [167] [168] [169] [170] [166].

In the present investigation group T2 (PEG membrane and BCP) exhibited the largest bone surface area (BS) in terms of new bone formation, followed by T1 and N (T2: 13.18 ± 5.85, T1: 9.69 ± 4.11, N: 8.81 ± 4.48 mm²). These differences were not statistically significant. However, in group T2, soft tissue was frequently interposed between the newly formed bone and the implant surface, irrespective of the functional loading. This was reflected in a lower mean bone-to-implant contact (BIC%) compared with groups N and T1, although differences were not statistically significant (T2: 33.9 ± 26.5, T1: 54.3 ± 27.1, N: 52.3 ± 20.9). Furthermore, the greatest distance
between the implant shoulder to the first BIC (S-BIC), i.e. the greatest residual vertical defect alongside the buccal surface of the implant was also evident in this group (T2: 3.71 ± 1.83, T1: 2.43 ± 1.74, N: 2.67 ± 1.40, P: 2.50 ± 2.17 mm). According to these unexpected results, not only the other test group, but both controls performed superior compared to the PEG group. A possible explanation of the limitation of this barrier may reside in its poor flexibility and elasticity after the membrane set following polymerization. In the present study, the PEG membrane was extended 2–3 mm around the bone defect and covered completely the coronal part of the implant. In a number of implants mesio-distal rupture of the membrane above the occlusal part of the implant was observed after polymerization was completed. When this occurred, the membrane was removed and re-applied. Even though all animals were fed by soft diet, this may have not prevented the rupture of the membrane above the implant after suturing of the flap above the membrane or during the early healing phase. It can be speculated that this has allowed soft tissue proliferation between the bone substitute and the implant surface, preventing the functional attachment formation between the implant and the newly formed hard tissue. These results might have an impact on possible clinical application of this PEG membrane indicating that the application should be limited on buccal surface avoiding complete coverage (i.e. occlusal surface) of the implant to avoid rupture.

Significant differences were observed in the mean residual bone substitute surface between groups T1 and T2. This value was several times higher in T1 (4.94 ± 2.81 mm²) compared with that in T2 (0.71 ± 0.80 mm²). This increased resorption/dispersion of the BCP particles when combined with GBR was also reported in previous studies. In an experimental study in dogs, Jung et al. observed the absence of BCP particles in 2/5 defects at six months [168], and Schwarz et al. showed a decrease in the mean graft surface between two and eight weeks [170]. It has been speculated that this could be correlated with the hydrolytic disruption of the PEG membrane, causing dispersion of the BCP [170]. Polyethylene glycol hydrogels degrade by hydrolysis and this process may take several weeks and is accompanied by an increase in the PEG volume of 30–50% [165]. It is not clear, if the increase in volume could exert pressure on the graft causing dislocation of its particles. However, in the present study, after five months of healing, the graft particles were reduced in number and size when the PEG membrane was used. In this treatment group, the small BCP particles were surrounded by
mineralized tissue. These observations may indicate an increased resorption rate of the
graft when the PEG membrane was used, rather than the dispersion of the particles.

**Loading**

It has been demonstrated that GBR could lead predictably in significant amounts of
bone formation when used prior to implant placement [136], or for treatment of
dehiscence/fenestration defects around implants [194] [127]. It has been suggested that
if the newly formed bone is not functionally loaded, then some bone resorption in height
might take place, once the protection of the secluded space created by a membrane is
removed [171] [172]. In the present study, where the regenerated bone was subjected to
functional loading for a short period, no significant differences were observed between
loaded and non-loaded implants in any of the examined parameters. However, in groups
P, N and T1, there was a clear tendency for increased bone regeneration (BIC%, S-BIC)
under functional loading which, however, never reached statistical significance. An
opposite trend was observed in group T2 (GBR). This group showed lower BIC% and
higher S-BIC and S-NBC at loaded sites. It could be suggested that these observations
are in agreement with previous studies in the literature where marginal bone loss around
implants after GBR had been observed [173] [195] [196].

This could also imply that in this study, the regenerated bone did not result in the
expected BIC in the PEG group and this may lead to bone resorption after a period.
These results emphasise not only the use of barrier membranes with appropriate
occlusive properties but also the importance on the surgical approach (membrane
application/ location) in order to avoid interference with the healing.

**Clinical versus histological data**

The present experiment provided some clinical as well as histological data on bone
regeneration. The distance between the newly formed tissue and the implant shoulder
was measured clinically prior to abutment connection and histologically after study
termination. According to clinical measurements, T2 performed most favorably with the
smallest remaining dehiscence along the implant surface (ROI) compared to T1 and N
(T2: 2.10, T1: 2.55, N: 3.35 mm). However, histological evidence failed to support this
clinical observation with T2 demonstrating the least favorable outcome (T2:
3.71 ± 1.83, T1: 2.43 ± 1.74, N: 2.67 ± 1.40 mm). This result corroborates that merely
clinical assessment of novel materials (i.e. without histological evaluation) used for GBR procedures may result in delusive conclusions.

6.3. Current options for the treatment of extraction sockets by ARP

Key findings
This systematic review has demonstrated that different ARP techniques cannot totally eliminate post extraction alveolar ridge resorption nor predictably promote new bone formation. However, the reduction in ridge width and height following ARP may be less than that which occurs following natural socket healing. The clinical data suggest that the horizontal ridge contraction was most successfully limited in the two studies applying GBR without additional bone grafts [213] [214], whereas the vertical shrinkage was most efficiently limited by employing GBR with additional bone graft [210] [212].

Strengths of the review
The present systematic review was limited to randomised controlled trials, controlled clinical trials and prospective cohort studies with a control group of empty untreated sockets. Furthermore, the inclusion criteria of our systematic review were based on the fact that the clinical merit of applying the different ARP techniques could only be validated, if the clinical as well as histological outcomes, following the application of a technique, are superior to that of unassisted socket healing.

In comparison to the previous systematic reviews [219] [223] the present review has evaluated the histological characteristics of the alveolar socket healing with or without ARP. The amount and the quality of the newly formed osseous tissues in the socket area are essential, especially when the justification of ARP is to facilitate the placement of a dental implant in the position of a previously extracted tooth. It is doubtful, whether an ARP technique should be claimed successful if it only preserves the external contour of the AR, but the newly formed tissue is of not superior or even inferior quality and quantity (percentage of matured trabecular bone) to what is normally achieved following extraction.
Finally, the quality of the included studies has also been meticulously assessed in this review. Such a quality evaluation of the retrieved data is essential to estimate the source and magnitude of potential bias that may lead to delusive conclusions.

**Strength of evidence – risk of bias**

The quality assessment of the included studies in this systematic review revealed that none of the trials have qualified for a low risk of bias category. Ten out of the 14 studies presented with high risk of bias, thus their results must be evaluated with caution. The lack of clear reporting of research methodology elements, such as adequate randomization and concealment and/or masking of the therapist and the examiner were among the primary reasons for the high risk of bias [241]. We did not contact authors for clarification of unclear methodology. Therefore, it is possible that actual study conduct was better than that reported in the publication. Statistical considerations played important role as well, since appropriate analytical statistics was completed and reported merely in one study [237]. Power calculation was conducted in three trials only [210] [212] [232], nevertheless the reported data were insufficient to determine the validity of the calculation.

**Dimensional changes and histological characteristics**

Sufficient ridge width and height have been considered as one of the key requirements for successful implant therapy and for the establishment of an aesthetically pleasing emergence profile at fixed partial dentures [128] [133] [242]. Therefore, the alteration in oro-facial (horizontal) and apico-coronal (vertical) AR dimensions was selected as the primary outcome of the present review. Direct intrasurgical measurements on the AR at re-entry are considered as the most precise method to evaluate the bone volume changes following ARP. It is desirable though to establish and validate a surrogate measure that avoids the need for re-entry surgery, while providing the clinician with a reliable measure. Two dimensional radiographs, such as IOPA or orthopantomogram, are not ideal to estimate the three dimensional changes of the AR [243]. Also, measurements of the alveolar mucosa level or study casts incorporate not only the alveolar bone, but also the overlaying soft tissue. For these reasons only studies performing clinical or three dimensional radiographic evaluation of hard tissue were included in this review. Cone-beam computerised tomography (CBCT) appears to offer a valid technique to assess alveolar ridge changes, with newer models greatly reducing radiation exposure [244].
However, a prerequisite of this technique would be some type of standardisation, so that the captured image is always being taken from exactly identical positions [245]. None of the two included radiographic studies reported on such standardisation [215] [236].

For the interpretation of the results we attempted to cluster the studies in respect to the type of intervention.

**Unassisted sockets**

In the present review the mean reduction of the AR width of the untreated sites varied between 2.6±2.3 and 4.6±0.3 mm and the mean reduction of the AR height was between 0.8±1.6 and 3.6±1.5 mm after one to nine months of healing. This corroborates the result of a previous clinical study which indicated that 95% of AR reduction should be expected after three months of extraction [131]. Furthermore, it is in agreement with a recent systematic review, which reported that the average reduction of the AR width seemed to be higher (3.87 mm), than the reduction in AR height (1.67 mm) [200].

Even though both AR width and height present resorption, histologically, new bone formation up to a variable extent was also observed in most studies as result of unassisted socket healing [209] [215] [210] [212] [232] [235] [236] [238]. In addition, a large area was occupied by bone marrow [210] [209] [238], such as reported in preclinical studies [14] [203] [45]. Only a single study reported on connective tissue fill and lack of mature bone in the control sites [233].

**Bone grafts and substitutes**

Effective grafting procedures for bone augmentation have been associated with the osteoconductive, osteoinductive or osteogenetic properties of the graft [45] [246] [247]. This led to the assumption that the placement of these materials in the extraction socket may accelerate new bone formation by the above biological properties and may also reduce AR resorption by stabilising the blood clot, providing a scaffold and external source of minerals and/or collagen [248] [14] [202] [249]. The placement of DBBM with collagen in fresh extraction sockets resulted in limited reduction of the AR dimensions, although delayed initial socket healing in terms of new bone formation was also observed [14] [202]. Human studies reported similar unfavourable histological observations when DFDBA was employed for ARP [205] [206].
In the present review of human experiments, two out of three studies reported that socket grafting with autologous bone marrow [237] or alloplastic material [232] have significantly limited the reduction of the AR width compared to the unassisted socket healing. Three out of five studies reported that reduction of the resorption in AR height was significant [232] [236] [237], while the ridge height was even increased in one study, where sockets were grafted with polymer sponge [209]. We should emphasise though that since the graft material (DBBM) in a radiographic study possessed radiopaque characteristic [236], the alteration of the AR contour on the CBCT image should be interpreted with caution.

Based on the histological evaluation of these studies, the above AR dimensional changes were not necessarily accompanied by higher amount of new bone formation in the socket, since the quality of newly formed tissue in the ARP sites was comparable to that in the control sites. Furthermore, the sockets were occupied by a mixture of new bone and connective tissue which in many occasions was surrounding the graft particles [207] [210] [236].

**GBR (membrane alone or in combination with bone substitute)**

The conception of guided bone and tissue regeneration [136] was translated to ARP procedures in order to exclude epithelial cells from the extraction socket by the use of barrier membrane in four studies of the present review [210] [212] [213] [214].

*a) GBR with membrane alone*

ARP with GBR resulted in statistically significantly less resorption in ridge width and height compared to unassisted socket healing, regardless of the type of membrane [213] [214]. It should be noted that in one study [213], in three out of 10 cases, the exposed non-resorbable e-PTFE barrier had to be removed prematurely, highlighting the importance of sufficient soft tissue closure and timing of removal of the barrier. The outcomes in these three cases were similar to that in the control sites. Where healing was uncompromised, a statistically significant difference was found after six months in width and height changes in favour of the ARP group.

*b) GBR with membrane and bone substitute*

ARP resulted in statistically significantly less resorption in width [210] [212] and height [212] in comparison to unassisted socket healing. The histological evaluation of the
GBR procedures in the included studies demonstrated new bone formation, but the presence of graft particles was also evident in both studies, embedded either in newly formed bone [210] or in connective tissue [212]. This is in agreement with a recent trial, where a collagen membrane in combination with DBBM or a biphasic bone substitute was used for ARP [192] [188].

Biological active materials
The potential benefit of biological active molecules were investigated in periodontal and bone regeneration through fostering the proliferation and differentiation of different mesenchymal cells in various preclinical models [250] [251]. The safety and feasibility of rhBMP-2 on human ARP or ridge augmentation was evaluated and shown to be safe in a bi-center clinical study [226]. Dimensional changes of the alveolar ridge were measured on CBCT scans in an RCT [215]. Treatment with rhBMP-2 resulted in an increase in ridge width, which was statistically significantly greater than in the controls. However, this observation needs to be interpreted in light of the surprise finding of an increase in ridge width of the untreated controls. This was a unique observation amongst the studies that we reviewed. Histologically, no comparison between ARP and controls sites was reported.

The human histological findings of the included papers of the present review were generally found to be comparable to preclinical studies [248] [14] [202] [252] [203]. There are a number of aspects to be considered in the interpretation of the results. Firstly, it has to be kept in mind that whilst the biopsies of the animal model incorporate the cross section of the whole AR, the biopsy retrieval at human studies is limited to a trephine core sample of part of the former socket. This location may not necessarily coincide with the exact position of the previous extraction, thus making interpretation of the results challenging. Furthermore, the differentiation between apical, mid and coronal, as well as the central and lateral aspects of the biopsies was not always apparent.

Another important parameter when considering a histological overview of the studies was the variation in healing time. Due to the nature of post-extraction healing, the direct comparison of the new tissue formation in studies between one and nine months of healing could be misleading. This was highlighted in three studies which did not make a
distinction between the variable healing times within the groups, ranging from 2.5 months to nine months [207] [210] [233]. It must also be noted that the single study, which completed and reported appropriate statistical methodology [237], did not observe statistically significant difference between the test and control biopsies. Furthermore, small sample sizes in the majority of the studies may also limit the generalisability of the histological findings.

Merely two studies found statistically significant histological differences in new bone formation favouring the test group [210] [232]. It is hard to draw generalizable conclusions from these two experiments, since the test groups differed in many respects compared with each other, including different technique (bone substitute only [232] vs. GBR+graft [210]), different material (MGCSH [232] vs. porcine bone with collagen membrane [210]), different flap management (flapless, no primary closure [232] vs. mucoperiosteal flap, primary closure [210]), different healing time (3 months [232] vs. 7-9 months [210]). One common feature was that both groups limited their intervention to sockets with four intact walls. It is noteworthy that all three studies that included intact socket walls only, reported statistically significant differences both on AR width and height in favour of ARP [232] [210] [237], while only one [215] out of two studies [215] [209] with initial buccal bone loss reported similar significant difference between test and control. Hence, socket morphology could be an important predictor of improved ARP. The need for ARP in such sockets, in terms of future clinical success/implant placement needs further investigation.

**Other factors affecting interpretation of the findings**

**Flap management**

All studies reporting statistically significant inter-group differences in both horizontal and vertical clinical measurements achieved either primary flap closure [210] [213] [214] [237], or did not detach the periosteum in a flapless procedure [232]. Furthermore, none of the studies without primary closure demonstrated statistically significant differences between test and control in terms of both horizontal and vertical clinical measurements [209] [212] [234]. Therefore, both achieving and maintaining the epithelial seal above the socket may be essential to improve ARP. Further corroboration of this concept was suggested where e-PTFE barriers were prematurely
exposed. The healing of these three exposed cases demonstrated no statistically significant differences compared to the control sites [213].

**Healing time**

The optimal timing of re-entry following ARP is determined by the implant insertion. The volume of the AR is gradually decreasing, while the quality of the newly formed tissue is gradually increasing during the post extraction remodelling [131] [198]. Therefore, the implant placement should be considered as early as possible, but as late as necessary, in order to maintain AR volume, but also to achieve complete epithelial seal with some extent of osseous fill. The healing periods of the trials in the present review varied considerably (one to nine months). Hence, the interpretation of the results is difficult due to the heterogeneity present in the included studies.

**Antimicrobials**

Improvement of clinical parameters was demonstrated as a result of regular rinsing with chlorhexidine following tooth extraction [253]. Subjects of the included trials in the present review were prescribed various types of antibiotics and instructed to rinse with chlorhexidine for two to three weeks. Thus, no conclusion could be drawn on the necessity or benefit of employment of antibiotics/antimicrobials following ARP.

**Smoking**

Smoking is associated with delayed wound and socket healing and increased reduction in post extraction alveolar width [254] [255]. Three trials in this review included smokers [210] [212] [233] and the half of the studies did not report on smoking as an exclusion factor, thus any conclusions about the impact of this well-recognised risk factor for impaired healing are difficult to draw.

**Periodontal treatment/health**

Four studies included patients whose periodontal treatment was carried out prior to the ARP [210] [209] [214] [238]. ARP resulted in statistically significant difference between tests and controls in clinical [210] [214] and in histological parameters [210]. In addition, in the studies where periodontitis was present, but periodontal treatment was not reported, no statistically significant histological differences were demonstrated.
This suggests that treated periodontitis may not hinder the success of ARP.

Hard and soft tissue morphology
No data were reported on factors, such as gingival biotype, width of the keratinised gingiva, thickness of buccal plate or total volume of AR that may modify the outcome of ARP. Therefore, the possible impact of these factors on ARP cannot be determined.

Clinical relevance
The clinical rationale for ARP is to minimise the necessity for one or two stage alveolar ridge reconstruction to allow successful implant placement. If the ARP procedure fails to meet this requirement, it may be considered as an unnecessary or even unsuccessful procedure. Therefore, a statistical significance favouring ARP does not necessarily lead to a clinical benefit, unless the whole treatment is simplified or made more successful [192]. In the present systematic review seven out of 10 studies did not report differences in feasibility of implant insertion at re-entry [207] [209] [212] [232] [235] [236] [238]. Merely two studies reported that there was no need for further reconstruction in the ARP group, whilst GBR or ridge expansion were carried out in some of the control sites alongside implant insertion [210] [237]. One study reported that statistically significantly less augmentation had to be performed in the ARP group, compared to the control [215]. In relation to illuminating the understanding of possible long term benefits of ARP, the success rate of the inserted dental implants in the former test, versus control sites should be examined [193]. No studies have yet reported this.

Patient-reported outcome and health economics
It would be helpful to understand patient experiences such as concomitant discomfort at/following ARP in order to avoid a further, extensive reconstructive surgery. On the other hand, the additional costs of ARP at the time of extraction might not be desirable, if the outcome and benefit of such extra treatment were not predictable. There are no data yet to inform on these questions.
6.4. Overall considerations

Why did we select these materials for our experiments?

Stabilization of blood clot and fibrin matrix in the wound healing process, scaffolding for osteoblasts during new mineralized tissue formation as well as a ‘buttress’ of the space maintaining materials, as well as an external source of collagen and/or minerals believed to support tissue regeneration [66]. For these purpose ABP as adjunct to GTR/GBR is still considered as the gold standard in GBR procedures. However, autogenous tissues are always associated with their drawbacks, i.e. their harvest usually requires a second surgical site (increased surgical trauma and donor site morbidity), presents with limited volume, resorbs rapidly that may result in decrease in volume. In order to overcome these shortcomings, the emphasis was placed on alloplastic bone substitute materials. Various alloplasts have then been available and have desperately been promoted by their manufacturers. In many cases though, a sound clinical and histological evidence seemed (seems) to be lacking behind these novel materials. In other words, the products of some (small) companies could earlier be discovered on the shelves of their suppliers, than on to the pages of scientific journals.

The recently developed materials that we aimed to investigate histologically, appeared to bear with very promising properties, in terms of new tissue formation and resorption, according to previous in vitro or in vivo investigations. In addition, their manufacturers are known for prioritizing the scientific evidence before launching a new material to the market. Therefore, both BCP, nano-HA and PEG seemed to be worth to examine in these different, but representative settings.

Why these study designs?

We aimed at scrutinizing the performance of these novel materials in reconstructive periodontology and related fields. The ultimate evidence of a material or method being ‘regenerative’ is based on human histology, or where inappropriate or less feasible, histological proof of randomized controlled preclinical trial [67] [68]. Therefore, the effect of the novel BCP combined with EMD or the nano-HA alone on periodontal intrabony pockets were investigated in human, single arm, clinical and histological case series. The same BCP with the novel, innovative PEG membrane was designed to
improve bone regeneration in case of dehiscence defects around dental implants. Due to the extremely challenging nature of performing human histology in such cases, randomized, controlled preclinical trial was designed. To the best of our knowledge, these were the first time to test the above materials in such settings in periodontology and implant dentistry. In contrast to this, broad variety of materials has been indeed tested for ARP procedures. Therefore, instead of conducting yet another human clinical and histological trial, a systematic review of these procedures and materials was carried out. To the best of our knowledge, no such review that systematically investigates not only the clinical, but also the histological healing characteristic of the result of ARP procedures compared to unassisted socket healing, alongside quality assessment of the included trials, have yet been published.

Clinical ‘success’ vs. histological evidence

All the selected and investigated novel materials have been used successfully in previous clinical trials and they all have shown improvements in clinical parameters.

The use of BCP + EMD in advanced periodontal defects resulted in significant PPD reduction, CAL gain and defect fill at six month in a multicenter study [119]. These results were comparable to that of their controls (EMD). In addition, these favourable clinical results were corroborated by a recent long-term RCT [120]. In accordance with these findings, the clinical results of our experiment (Study 1) were similarly beneficial, with mean PPD reduction of 3.3 ± 1.4 mm and mean CAL gain of 3.0 ± 1.6 mm. In spite of these promising clinical data, histology has revealed none to limited periodontal regeneration i.e. 0.69 ± 0.67 mm mean new cementum and 0.16 ± 0.25 mm mean bone formation. One third of the biopsies displayed LJE instead. One may consider that this result looks inferior to that achieved by EMD only in a similar human histological investigation i.e. formation of mean 1.87 ± 1.17 mm new cementum and 0.78 ± 0.97 mm new bone [36]. In addition, in our study the additional use of BCP resulted in soft tissue encapsulation.

Study 2 has actually led to similar conclusions. Application of nano-HA for the surgical treatment of intrabony periodontal defect in randomized controlled clinical trials resulted in significant PPD reduction and CAL gain [124] [106] [105]. Furthermore, the result in the test group (nano-HA + OFD) appeared to be significantly better than that in
the control counterparts (OFD) [106] [105]. The clinical results of our experiment (Study 2) resembled to the above i.e. mean PPD reduction of 4.0 ± 0.8 mm and mean CAL gain of 2.5 ± 0.8 mm. However, the histological findings appeared to be again in contrast (or at least inferior) to that of the clinical ones, since LJE was observed in 3/6 cases, resulting in the other half of the cases merely 0.36 ± 0.41 mm and 0.54 ± 0.61 mm of mean new cementum and new bone formation, respectively. When compared this histological data to that of a control group (OFD) of a recent similar study by our group [69], it may be noted that the present histological findings of the nano-HA appears to be even inferior to that of the aforementioned OFD group, i.e. 1.2 ± 1.1 mm and 0.8 ± 1.0 mm of mean newly formed cementum and bone, respectively. Furthermore, histomorphology revealed that the remaining implant particles (2/6 biopsies) were embraced by connective tissue.

The clinical measurement between the implant shoulder to the first BIC at the remaining buccal defect during re-entry surgery in Study 3 revealed T2: 2.10, T1: 2.55, N: 3.35 mm. This suggests that the treatment of BCP + PEG led to the most favourable defect fill. In contrast to this clinical measurement, histometry revealed quite the opposite, resulting in the least favourable S-BIC in BCP + PEG group (T2: 3.71 ± 1.83, T1: 2.43 ± 1.74, N: 2.67 ± 1.40 mm). Furthermore, histomorphology displayed bone formation around the graft particles in T2 group and in the apical portion of the defect in T1 group, however, a significant amount of particles were embedded in soft tissue in the middle and coronal portion of the defect. This histological result seems to be similar to that of previous investigations [191] [192], as well as to Study 1, where the same bone substitute was applied in a periodontal setting [73]. Nevertheless, it may be noted that a recent investigation comparing the clinical performance of the new PEG to a standard collagen membrane for the treatment of peri-implant dehiscence bone defects concluded that the PEG was as successful as the collagen membrane, with simplified clinical handling [169].

The clinical data of Study 4 indicated that ARP procedure may effectively minimize ridge resorption both in horizontal and in vertical dimensions. This is in agreement with other reviews [218] [219] [220] [221] [222] [223]. Histologically, however, ARP does not clearly appear to promote new bone formation in the extraction socket. Instead, some bone substitute may delay the natural healing process [207] [212] [236].
Moreover, our results reflected various degree of new bone formation in both test and control sockets. Statistically significantly more trabecular bone formation in the test groups was reported merely in 2 of 11 studies [232] [210]. In addition, in one study, more vital bone was detected in the control sockets [212].

The results of the investigations in the present thesis indicate that clinical success may not necessarily be routinely reflected by histology.

**Recommendations for further research**

- Randomised controlled trials on adequately powered sample sizes are needed where unassisted socket healing serves as the negative control.
- Appropriate follow-up periods are required. Ideally, this should reflect implant insertion protocols, such as six weeks (Type 2), three to four months (Type 3) or >6 months (Type 4) placement following extraction.
- Clinical studies should be designed to perform not only clinical (quantitative), but also histological (qualitative) assessment.
- The role of additional factors like smoking, reason for extraction, tooth location, initial buccal plate thickness, flap reflection and closure, antimicrobial regime should also be investigated.
- Comparative studies should also be designed in order to identify the most successful treatment options.
- It may be beneficial to seek for a cell occlusive barrier membrane that does not require extensive soft tissue and muco-gingival junction mobilization for flap approximation.
- Necessity of re-augmentation at implant placement should be investigated.
- Survival and success rates of implants, placed in former ARP sites should be evaluated.
- Outcome evaluation should ideally incorporate patient’s preference, quality of life, as well as treatment economy.
7. CONCLUSIONS

Within the limitations of each study the following conclusions could be drawn:

1. The combination of EMD and BCP for the surgical treatment of human periodontal intrabony defects is safe and well tolerated; could lead to improvement of clinical periodontal parameters; could result in meagre formation of new cementum with associated PDL and in none or minimal new bone formation. Therefore, this combination does not seem to possess additional benefit for the regenerative periodontal treatment. (Study 1)

2. The use of nano-HA for the surgical treatment of human periodontal intrabony defects is safe and well tolerated; could lead to improvement of clinical periodontal parameters; could result in the formation of minuscule amount of new cementum with associated PDL and little amount of new bone. Therefore, the use of nano-HA seems to have limited potential to promote periodontal regeneration. (Study 2)

3. For simultaneous augmentation of critical size, porcine, peri-implant dehiscence defects, PEG membrane is safe and well tolerated; shows appropriate occlusive property, hence may be effective in bone formation. However, the fragile property of the polymerised material could lead to early rupture of the membrane, which could have a negative impact on the whole healing process, particularly on the bone-to-implant contact. (Study 3)

4. BCP alone does not result in predictable new bone formation in such defects. (Study 3)

5. The hydrophilic implant surface (SLActive) per se could support peri-implant new bone formation in these defects. (Study 3)

6. The short-term loading of SLActive implants inserted in augmented dehiscence defects does not have a significant negative influence on osseointegration and new bone formation. (Study 3)

7. The postextracion resorption of the AR cannot be totally prevented by ARP, but some of the ARP techniques can limit dimensional changes of the AR. However, ARP could be associated with increased incidence of adverse events. (Study 4)
8. Conflicting evidence exists on the benefit of ARP at the histological level, since ARP does not appear to promote de novo hard tissue formation routinely, in addition, some graft materials could interfere with the healing. (Study 4)

9. The strength of evidence of the included trials ranges from weak to moderate, therefore the results should be interpreted with caution. (Study 4)

10. Due to the broad variety of employed materials, techniques, defect morphologies, healing periods and small sample sizes, meta-analysis or comparative assessment of ARP could not be made. Consequently, no material or method can be claimed to serve superior to another, however, in certain cases, GBR appears to be most effective. (Study 4)

11. Only limited evidence supports the ultimate clinical benefit of ARP, namely the reduction of necessity of further augmentation in conjunction with implant placement. (Study 4)

Generally, the results of the present thesis strongly indicate, that meticulous (human) histological assessment should also be carried out prior to addressing ‘regenerative’ properties to a new material or method.
8. SUMMARY

Several ‘regenerative’ methods and materials have been claimed as successful in the field of reconstructive periodontology, implantology and related fields. However, they do not always appear to promote the formation of new bone, periodontal tissue or functional attachment, according to the only reliable histological evidences. The aim of the studies included the present thesis was to investigate histologically some representative materials and methods, which had shown favourable clinical outcomes already. In the first scenario we have examined two novel materials used in periodontology. The investigations demonstrated that the combination of a biphasic calcium phosphate (Straumann BoneCeramic®; BCP) and enamel matrix derivative (Emdogain®; EMD) (Study 1) or an unsintered, nanocrystalline, phase-pure hydroxyapatite (Ostim®, nano-HA) alone (Study 2) for surgical treatment of human intrabony periodontal defect is safe and well tolerated. Both treatments improved the investigated clinical parameters, however, they resulted in none to meagre new cementum and bone formation histologically. In the second scenario we examined the regenerative potential of BCP with a novel polyethylene glycol-based hydrogel barrier membrane (Membragel®; PEG) in the field of reconstructive implant dentistry (Study 3). Since it is not feasible to conduct a human histological study due to obvious ethical considerations, we assessed the regeneration of a critical size, peri-implant dehiscence defects on Göttingen minipigs. The investigation demonstrated that osseointegration around hydrophilic dental implants (SLActive®), treated with or without BCP and PEG, can be achieved and is not negatively affected by functional loading. However, the amount and quality of the newly formed hard tissue, as well as their attachment to the implant were not predictably achieved by the applied materials. In the third scenario we investigated whether or not the bone loss, associated with tooth extraction, could be prevented, hence to limit the need for peri-implant bone augmentation. Therefore, a systematic review was conducted in order to obtain the highest level of histological and clinical evidence (Study 4). Our results demonstrated that different alveolar ridge preservation (ARP) techniques could limit post extraction shrinkage in width and height, however the natural resorption cannot be totally eliminated. In addition, ARP does not predictably promote new bone formation histologically. The results of the thesis strongly indicate that meticulous (human) histological assessment should also be carried out prior to addressing ‘regenerative’ properties to a new material or method.
ÖSSZEFOGLALÁS

Többféle anyagokat és módszereket neveztek már sikeresnek a parodontológiában, implantológiában és határtételeiken. Ugyanakkor, a szövettani vizsgálatok mint egyedül megbízhatók tükrében, ezek nem minden esetben segítik elő az új csont és parodontális szövet képződését, sem az új tapadás létrejöttét. A jelen tézisben bemutatott vizsgálatok célja az volt, hogy szövettani megvizsgáljunk olyan anyagokat és módszereket, amelyek már klinikailag jó eredményeket mutattattak fel. A disszertáció első részében a parodontológiában használt két új anyagot vizsgáltunk. Kutatásunk bemutatta, hogy a kétfázisú kalcium foszfát csontpótló (Straumann BoneCeramic®; BCP) és a zománc mátrix protein (Emdogain®; EMD) együttes alkalmazása (1. vizsgálat), illetve a szinterezetlen, nanokristályos hidroxilapatit (Ostim®; nano-HA) önmagában (2. vizsgálat), humán parodontális intraosszer defektusok sebészi kezelésére biztonságos és jól tolerált. Mindkét kezelés javította a vizsgált klinikai paramétereket, ugyanakkor szövettaniag nulla, vagy minimális új cement- és csontképződést eredményeztek. A második részben a BCP és egy új polietilén-glikol alapú hidrogél membrán (Membragel®; PEG) regeneratív potenciálját tanulmányoztuk a rekonstruktív implantológia területén (3. vizsgálat). Mivel itt humán szövettani kutatás nyilvánvaló etikai okokból kivitelezhetetlen, ezért a kritikus méretű, implantátum körüli dehiszcencia defektusok gyógyulását Göttingen miniserteseken vizsgáltuk. Kutatásunk bemutatta, hogy létrejön osszeointegráció BCP és/vagy PEG membránnal kezelt hidrofil (SLActive®) implantátumok körül és mindezt a funkcionális terhelés nem befolyásolja. Mindazonáltal az újonnan képződött keményszövet mennyisége és minősége, valamint mindezek tapadása az implantátum felszínéhez előre nem kiszámítható. Harmadikban azt vizsgáltuk, hogy a fogelvölgyitást követő csontpusztulás megelőzhető-e, következésképp az implantátum körüli csontaugmentáció szükségessége csökkenthető-e. Ezért egy ún. systematic review-t végeztünk annak érdekében, hogy a lehető legmagasabb klinikai és szövettani bizonyítékot demonstráljunk (4. vizsgálat). Eredményeink bebizonyították, hogy a különféle állcsongerinc-megtartó technikák (ARP) csökkenthetik a csontpusztulás vertikális és horizontális dimenzióját, ugyanakkor a csontpusztulás nem küszöbölhető ki teljesen. Továbbá az ARP nem képes előre megjósolhatóan új csontot képezni szövettanilag. A tézis eredményei kifejezetten megerősítik, hogy csak akkor nevezhetünk egy új anyagot, vagy módszert regeneratívnak, ha ezt megfelelő (humán) szövettani vizsgálatok is alátámasztják.
10. REFERENCES


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11. LIST OF PUBLICATIONS (IF: 20.193)

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