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1. Introduction

Bone regeneration procedures aim at recapitulating optimal wound healing where tissue components are restored to the form and function required for tissue and organ homeostasis (Zohar & Tenenbaum 2005; Bueno & Glowacki 2009; Dimitriou et al. 2011). Examples of ideal bone regeneration include the healing of a healthy tooth extraction socket or a simple bone fracture. This is not the case in non-union fractures, or extensive damage as a result of tumour removal or bone subjected to chemotherapy, where the overall wound healing ability may be compromised (Dimitriou et al. 2011). Bone is a specialized connective tissue consisting of osteoblasts, osteocytes and osteoclasts embedded in a mineralized matrix capable of remodelling, renewing and load bearing. Optimal bone regenerative therapy will enhance mineralized tissue wound healing through enrichment of the wound/bone defect with a matrix scaffold to support the wound, cells that will give rise to osteoprogenitors and inducer molecules, such as growth factors to amplify activity of cells or events responsible for bone formation. New regenerative approaches may include a combination of these factors in part or as a whole. The temporal, spatial activity and maturation of these three components (i.e. cells, matrix and inducer molecules) during bone regeneration has to be a coordinated and integrative process. Delayed, reduced or lack of activity of any of these components may result in repair and not regeneration of a remodelling functional bone. Cell therapy is compared to the gold standard of autogenous bone marrow grafting, which is considered to be enriched with mesenchymal stem cells, osteoprogenitors and inducer molecules; marrow grafting usually offers predicative regenerative approach. Matrix grafting has to offer mechanical support for the regenerative process to interact with the differentiating osteoprogenitor cells and provide the conditions for the cells to deposit host bone matrix. Grafted inducer molecules need to interact with both the developed matrix and differentiating osteoprogenitors to assure bone matrix deposition and mineralization (Figure 1).

Our earlier studies focused on the isolation and differentiation of bone stem cells, osteoinductive cytokines and matrix development and maturation. The spatial and temporal sequence of matrix molecules expression used to sort stem-like cells population, single application of bone morphogenetic protein-7 (BMP7) induced differentiation of these cells to osteoblasts (Zohar et al. 1997a; Zohar et al. 1998; Zohar et al. 1997b). For bone cells to differentiate or for the bone matrix to mature and mineralize, cross talk between matrix and...
cells is required to activate bone transcription factors associated with signaling pathways and osteogenic protein expression. Communication between matrix and osteoprogenitor cells is crucial to form a mature, weight-bearing bone. This communication is mediated through secreted growth factors, matrix or matrix associated molecules and activated receptors on the differentiating bone cells.

Fig. 1. Wound healing in bone regeneration follows a temporal sequence of ideal healing where a clean wound start healing through bleeding, clot formation and recruitment of mesenchymal stem cells which will differentiate to bone forming cells. The successful differentiation of mesenchymal stem cells to osteoblasts dependent of the temporal and spatial recruitment and expression of cells, matrix and bone related mediators. Matrix would form through adequate blood supply, stable clot formation and deposition of bone matrix that will mineralize. Osteoblasts and osteocytes will differentiate with matrix maturation and will secret mediators and bone specific proteins.

Various animal wound models in number of animal species are used to assess regenerative approaches include rodents, rabbits, sheep, goats, cats, dogs and primates (Gomes & Fernandes 2011; Intini et al. 2007; Kim et al. 2007; Artzi et. al 2003 a,b; Meinig 2002; Lemperle et al. 1998). New experimental approaches attempt to regenerate critical-size defects in the affected bone that won’t heal without therapeutic intervention. Comparing results between animal models is challenging due to different wound models, different bones used, healing rate, unique animal physiology, whether or not the bone is weight bearing and a variety of protocols. Mice are the animals of choice for transgenic analyses for the significance of the permanent present or absence of one or two molecules (Kim et al.
signals between cells and matrix mediate bone regeneration

2007; Masaki & Ide 2007). Large animal models on the other hand are preferred for a slower healing process resembling human physiology; however due to the high cost, control of animals and lower sample number, their use is more limited. The tibia or femur are usually used for the fracture model in a load-bearing area and the calvaria may be used for critical size bony defects in a non-loaded area (Alberius & Gordh 1996; Au et al. 2007; Landry et al. 1996).

Regenerative regimens usually focuses on one of the main components of the missing mineralized tissue: matrix, cells or inducer molecules. While expression patterns were identified for cell differentiation and matrix maturation, ongoing interactions during healing through receptors and signal molecules determines whether the outcome is repair or full regeneration. Thus, evaluating these interactions and the ability of the host wound area to support the process is a major determinant of regeneration. This chapter will focus on the importance of signaling between matrix and bone cells and how growth factors or inducer molecules can mediate this interaction and lead to the regeneration of bone tissue.

2. Bone wound healing

Bone wound healing in primates may involve formation of cartilaginous template, leading to endochondral ossification and/or intramembranous ossification (Dimitriou et al. 2011; Javed et al. 2011). Both processes require the commitment of adult stem cells toward bone-forming or osteoprogenitor cells (Figure 1). It is well recognized that adult bone contains a reservoir of mesenchymal stem cells responsible for physiological remodelling of bone and reconstruction during wound healing (Awad et al. 1999; Pittenger et al. 1999). Notably, mesenchymal stem cells are multipotential and capable of differentiation not only to bone forming cells but also to chondrocytes, adipocytes or fibroblasts, as shown in vitro and in vivo studies (Ghilzon et al. 1999; Owen 1988). Commitment of mesenchymal stem cells is thought to be irreversible, and thus signals during the early stages of the wound healing where mesenchymal stem cells differentiation to osteoprogenitors occurs is crucial for bone regenerative process. The ability to induce mesenchymal stem cells to express osteoblastic markers is dependent on transcription of bone-related genes activated by specific signalling, such as wingless-type MMTV integration site (Wnt) family which control osteoblasts differentiation (Hoepnner et al. 2009; Seroeto et al. 2009). Important mediators in these pathways activated by Wnt will be the Runx2 (Cbfa1) and Osterix transcription factors. These proteins control expression and repression of genes that will direct the commitment of mesenchymal stem cells toward osteoblasts (Liu, W. et al. 2001). Runx2-deficient mice exhibit neonatal lethality due to absence of bone. In the absence of Runx2 there will be no osteoblast differentiation or ossification. Haploinsufficiency of Runx2 in humans results in cleidocranial dysplasia, a disease characterized by abnormal bone development, formation and decreased bone density (Notoya et al. 2004; Post et al. 2008; Xiao et al. 2004). Cytokines derived from the TGFβ superfamily, such as BMP-4, induce the expression of these transcription factors and thus bone-specific proteins such as alkaline phosphatase (AP), collagen I, bone sialoprotein (BSP), osteocalcin(OCN), osteopontin (OPN), integrin and TGFβ receptors. The expression of these markers serves to ascertain osteoblastic differentiation and evaluate the progression of bone formation. Unfortunately, at present, clear markers to identify and isolate mesenchymal stem cells or osteoprogenitors are not available and the lack of hematopoietic stem cells markers, as well as cellular morphological
characteristics, such as undeveloped cytoplasmic structure, are the only reliable criteria for osteoprogenitors (Belmokhtar et al. 2011; Bernardo et al. 2011; Vater et al. 2011).

Following differentiation of osteoprogenitor cells, a stage of the committed cells proliferation and growth cell cycle changes, accompanied by regulation of proliferation-related genes, such as histones, c-myc and c-fos being upregulated; secretion of matrix proteins, such as collagen I, II, III; alkaline phosphatase; fibronectin (Figure 1); as well as cytokines like FGF-2, TGF and BMBs members (Augello & De Bari 2010). Osteoprogenitors mature to secretory osteoblasts with a reduction in mitotic activity and formation of collagenous extracellular matrix (ECM) enriched with bone-specific proteins such as AP, OCN, BSP and OPN. Osteoblasts also secrete osteoprotegerin (OPG) a member of the TNF superfamily to reduce osteoclastic bone resorption by binding with the receptor activator of NF-kappaB ligand (RANKL) (Takahashi et al. 1999). Osteoblasts express receptors to mediate connections between ECM and cells; this connection is primarily mediated through integrins, which will attach to the ECM and intracellular will activate the actin cytoskeleton, initiating cellular signal transduction of proteins such as mitogen-activated protein kinase (MAP kinase) and the SMAD pathway (Blair et al. 2008; Komori 2011). Decrease in matrix formation precedes deposition of hydroxyapatite crystals in the mature collagen organized in a quarter-staggered pattern with 68nm gaps to house hydroxyapatite crystals, which accumulate on the collagen fibers within them and flattening of the active osteoblasts, which may undergo apoptosis or become trapped in the mineralized matrix as osteocytes (Kogianni & Noble 2007).

3. Bone regenerative therapy - Present approaches

There are multiple approaches and various grafting materials available for bone regenerative therapy. The noble regenerative objective is the same for all suggested approaches: living, functional, remodelling bone! Different studies evaluating the success of fracture regeneration or repair estimate the failure rate as 10% or more. Common factors in failure are: lack of vascularity, improper correction, delayed union, non-union and revision surgery (Jones et al. 2000; Lee et al. 2004; Osti et al.; Parker et al. 2011; Smith, T. O. et al. 2009). The tibia is the most common bone to fracture in children and adults. Corruptions that exhibit non-union complications present greater challenge to regenerative therapy (Garrison et al. 2011; Mashru et al. 2005). Other than fixation of fracture, there are also non-invasive approaches used to improve healing, such as electromagnetic field or ultrasound stimulations (Griffin et al. 2011). Distraction osteogenesis is another approach which encourages bone formation through gradual distraction of defect surfaces, requires long treatment, sensitive technique and prolonged healing for the patient; it also serves as a burden to the health system (Heo et al. 2008). Autologous bone marrow grafting is the most treatable and predictable approach to achieve regeneration. Bone can be harvested from the iliac crest of the pelvis, or alternatively, reamers can be used to harvest the intramedullary canal of long bones (Hak & Pittman 2010; Valimaki & Aro 2006). If a larger volume of grafts required, allograft or biomaterials are sometimes used in conjunction with autograft.

Present descriptors of grafting materials other than their source (i.e. allograft, autogenous, alloplast, cancellous, cortical), refer to grafts as being capable of osteoconduction, osteoinduction, mechanical support, cell exclusion, cement and filler. Regeneration of bone is a very clear outcome, and unless osteoblast differentiation taking place, new bone matrix
deposition and interaction between the two during de novo bone formation and remodelling, no real regeneration could occur. The assumption that placing an allograft that may contain BMP’s, collagen matrix or an even high number of mesenchymal stem cells will result in regeneration in every case cannot be true. Without receptive wound environment where osteoprogenitors have signals for differentiation and deposition of new bone matrix, healing by fibrous or cartilage or adipose tissues may occur. Thus, using terminology like osteoconduct and osteoinduction would only suggest of the potential of regenerative approach or material, but it is not necessarily predictive of the desired outcome in specific host, specific wound, and specific surgical approach. The clinical results suggest variability of wound healing (Garrison et al. 2011).

Since most new bone graft or regenerative product is first tested for its biological activity, rather than focusing on osteoinduction and osteoconduction, this chapter classifies present grafts by their contribution to one of the major missing components of the missing bone: cells, matrix, and mediators (Figure 1). To evaluate the present state of bone regenerative therapy, it is worthwhile to see how each approach can contribute to the restoration of one of these three components.

4. Matrix grafting

Matrix serves as an organized framework for bone as a tissue and organ, offering mechanical support, and facilitate preservation of form and adaptive protection to internal organs through ongoing remodelling (Grabowski 2009; Scott et al. 2008). Osteogenic cells, like most other matrix-associated cells, cannot survive or differentiate without adhesion to their matrix (Popov et al. 2011). Thus, the importance of bone matrix in addition to acting as mechanical scaffold, is to mediate the biological activities of bone cells and signals that maintain homeostasis, remodelling and ability for wound healing. The mature mineralized bone matrix is composed of ~20% organic components, primarily collagens I, III and V and less than 5% noncollagenous proteins. The latter consists of proteoglycans, such as versican, decorin, and hyaluronan, adhesions molecules such as fibronectin and vitronectin, and specialized proteins like OCN, BSP, OPN and cytokines (Nagata et al. 1991). The collagen fibrils structure house the hydroxyapatite crystals which tend to be oriented in the same direction as the collagen fibrils. The collagen network also mediates adhesion to cells primarily through integrin receptors connected to collagen or the associated non-collagenous proteins. A mineralized bone matrix not only increases the mechanical strength of the bone but also act as reservoir for specialized proteins and cytokines, such as BMP’s.

Matrix proteins mediate not only maturation and mineralization of bone matrix, but also bone cell differentiation and signalling. Bone cell differentiation is detected through the differential expression of matrix molecules such as collagen OPN, BSP, AP and OCN. Expression of AP, collagen I and OPN are considered an early markers, while BSP, OCN and a second peak of OPN are considered a late mineralization associated marker (Aronow et al. 1990; Binderman et al. 2011; Lynch, M. P. et al. 1995). Our studies of OPN expression, which is not restricted to bone, but can be used as a useful marker for early and late differentiation of osteogenic cells. We isolated a population of small cells that do not express OPN, AP, collagen I and that are enriched with stromal stem cells capable of generating bone, fat and cartilage (Zohar et al. 1998; Zohar et al. 1997b). We have isolated BMP-responsive cells, which will undergo chondrogenic differentiation with continuous
stimulation of BMP-7 or osteogenic differentiation with single dose (Zohar et al. 1998). Thus, evaluating the expression of matrix proteins can help determine the status of mesenchymal stem cells differentiation.

4.1 Matrix-based grafts can be autologous, allogeneic or biomaterials

Autologous bone grafts, such as the marrow graft, marrow aspirates, will contain cancellous and/or cortical or blocks such as vascularised Graft and will carry cells, matrix and potentially inductive molecules (Friedrich et al. 2009; Sotereanos et al. 1997). A vascularised graft will carry blood vessels to enrich the wound with nutrients and soluble mediators, which may support or inhibit bone formation and carry periosteum enriched with osteoprogenitor cells. There is less necrosis of grafted material during healing and vascularised grafts are thought to be a very reliable option for reconstructing non-union or osteonecrosis defects (Friedrich et al. 2009; Gaskill et al. 2009; Sotereanos et al. 1997). The difficulty with all autogenous grafts is the quantity and morbidity, such as non-stress fracture for donor sites (Friedrich et al. 2009). The cancellous or cortical block graft may carry cells and cytokines, and their quantity and effectiveness is related to the age and state of the donor area. Cortical block graft will contain the least amount of cells and mediators and considered to function primarily as scaffolding which is more susceptible to infection and necrosis.

In allogeneic bone matrix grafts, cadaver bone is a common source of allograft. To generate a safe allograft, the bone is subjected to irradiation or freeze-drying and is thus devoid of any cellular components (Nguyen et al. 2007). Allografts are prepared as particulate, morselized or block, with mineralized or demineralized bone particles that are easy to shape and mold. Demineralized bone matrix serves as a natural matrix as well as decellularized matrices that could derive from dura or intestine of various animals (Costain & Crawford 2009; Kligman et al. 2003; Mroz et al. 2006). Allografts have very limited, if any, biological activity and serve primarily as osteoconductive and mechanical support. The main advantage is ample supply (Hamer et al. 1996). Reports of infection transfer, matrix alteration during processing and limited remodelling of the grafted bone reduce the likelihood of full regeneration (Nguyen et al. 2007) unless combined with autologous bone (Matejovsky et al. 2006) to add osteoprogenitors and mediators that can append biological activity to the dead bone particles.

Matrix proteins-based polymers are very popular, as are collagen, fibrin, hyaluronic acid, fibronectin and BSP. These proteins are delivered as membranes, sponges, gels, demineralized bone particles, small intestinal submucosa, dura or even urinary bladder (Chajra et al. 2008; Smith, I. O. et al. 2009; von der Mark et al. 2010). The problem with generating these polymers is fairly low solubility; the organic purified polymers is costly and hard to extract, purify and stabilize; risk of immunogenicity; and variations based on the batch.

Biomaterials and synthetic bone substitutes are currently used as fillers and/or scaffolds for the missing bone structure (Gosain et al. 2009; Healy et al. 1999; Shekaran & Garcia 2010; Wojtowicz et al. 2010). The design and fabrication of matrix-based regenerative materials is aimed at restoring the natural bone matrix properties as a whole or in part. Reconstruction of missing bone using matrices involves the planning of macrostructures as well as
Signals Between Cells and Matrix Mediate Bone Regeneration

Microstructure of the engineered matrix (Cholewa-Kowalska et al. 2009; Huang & Miao 2007; Vater et al. 2009). Macrostructures to fill and adapt to the space to assure sufficient quantity and/or provide mechanical support for the surrounding tissue or cells carried. Microstructures of micron or nanotechnology designs of particles or pores are used to encourage cell adhesion, colonization and absorption of proteins or required molecules. An ideal scaffold will have highly interconnected macroporosity to allow host bone tissue and blood vessels to grow into the scaffold (Healy et al. 1999). Popular building blocks are hydroxyapatite (HA), calcium phosphates (CP), tricalcium phosphate (TCP) and bioactive glasses (Behnamghader et al. 2008; Muschler et al. 1996; Valimaki & Aro 2006). They form a carbonated apatite layer when grafted, which is very similar to bone mineral; this will attract attachment of collagen fibres and eventually should be replaced by host tissue, mineralized matrix and cells. Other scaffolds consist of combinations of poly (lactic-co-glycolic acid) (PLGA), alginate and chitosan (Huang & Miao 2007; Jose et al. 2009; Liu, X. et al. 2009; Mishra et al. 2009; Renghini et al. 2009). These polymers can also be used to carry cytokines for controlled release at the wound and/or to carry mesenchymal stem cells. Different studies use different mixtures of these materials or different preparation protocols. The requirement for most preparations is to offer bioactivity and mechanical support. Bioactivity of the scaffold is measured by the number of host cells attached to its surface and interaction with the material to transform them into functional osteoblasts. The mineralized bone matrix will appose directly onto the surface of the material which ideally will have the ability to degrade over time (Holy et al. 2000). It is important that the material will degrade at a rate that allows the newly formed tissue to gradually replace the scaffold, both as a mechanical structure and in terms of space occupied. Finally, and this is where most current materials fail, the material needs mechanical properties that allow the device to be implanted without losing mechanical properties, still allowing sufficient loading of the newly formed tissue (Au et al. 2007; Smit et al. 2010). As of yet, no one has reported a material that fulfills all these requirements. The new scaffolds, usually termed composite scaffolds, maybe coated with proteins to increase cell adhesion, carry cells or cytokines with sustained release (Ameer et al. 2002; Bueno & Glowacki 2009; Gupta et al. 2011; Nie et al. 2008).

Bioactivity of biomaterial can be modified through chemical and physical alterations. Nanotechnology approaches try to mimic cell surface properties through approaches such as controlling space between ligands connected to biomaterials (Smith, I. O. et al. 2009). Using the proper spacing will enable, for example, integrin receptor clustering to enable propagation of signals through ligation. Another line of research focuses on molecules that work in synergy with receptors to promote cell adhesion and differentiation; for example, fibronectin, laminin and BSP contain heparan sulphate binding domains that interact with molecules on the cell surface in conjunction to integrin binding. Thus, using cell membrane molecule, such as syndecan which has three sites of heparan sulphate, would augment ligation of fibronectin or RGD sequence by integrin receptors (Whiteford et al. 2007; Yamada et al. 2010).

5. Cellular grafting

Cellular grafting for bone regeneration is a rapidly developing area. This approach had been used for many years through autologous bone grafting, which contains high numbers of
Bone-committed cells in marrow aspirates, or in bone particles or blocks containing cells embedded in their own matrix (Hak & Pittman 2010; Papakostidis et al. 2008; Tiedeman et al. 1995). The objective of new approaches is to obtain an unlimited amount of adult stem cells, comparing new cellular sources to the gold standard of autogenous bone marrow stromal cell, which are considered to be enriched with osteoprogenitors. Notably, the frequency of osteoprogenitors in young rodent marrow is about 0.0005% (Falla et al. 1993) and up to 0.3% in fetal periosteal tissues. Adult marrow shows a reduction of these precursor cells in number and quality (Stolzing et al. 2008). We have used single cell flow cytometric sorting to isolate osteoprogenitors from fetal rodent periosteal tissues. These cells when plated and stimulated exhibit high proliferative capacity and enhanced osteogenic potential. Notably, these cells consisted of only a very small fraction of the fetal bone tissues. Thus, even in young fetal tissues osteoprogenitors consist of only a very small fraction of bone tissue and usually reside in a well-protected niche. Moreover, during seeding, grafting and transfer of cells to the wounded area there is loss of cells through apoptosis or cytotoxic effects of mediators in the wound area (Giannoni et al. 2009). Regeneration efforts focus on the ability to deliver mesenchymal stem cells to the wound, which will differentiate to the osteoblastic lineage. Differentiation requires the commitment of mesenchymal stem cells to osteoblasts, exhibiting bone-specific gene expression. Osteoblast-specific gene expression is a fairly clear analysis of proteins like AP, OPN, BSP, OCN that are selectively expressed in bone. For the mesenchymal stem cells to form new bone and regenerate the wound, cells need to attach, proliferate, differentiate and survive. Mesenchymal stem cells from marrow seem to be the most predictable source for osteoprogenitor cells and a safe autologous grafting. Unfortunately, bone marrow stromal cell consists of heterogenous population that are subject to age changes; not only does their number deplete, but also their quality and ability to generate new bone is reduced (Benayahu 2000; Stolzing et al. 2008; Zhou et al. 2008). Thus, in the aging population where bone wound healing is compromised, harvesting autologous sufficient number of mesenchymal stem cells from marrow may not be that predictable.

Other sources for bone forming cells could be the umbilical cord, peripheral blood, adipose tissue, dental pulp or periodontium (Goodwin et al. 2001; Honda et al. 2011; Rhee et al. 2010; Yamamoto et al. 2007). Human embryonic stem (hES) cells also being considered as an option due to their fast growth and the fact that these cells, if kept as undifferentiated cell lines, are pluripotential and capable of differentiating to many tissue types under the right conditions (Bahadur et al. 2011; Lerou & Daley 2005). The hES has the advantage of unlimited supply, minimal immune response and no need for a second surgical site (Watt & Hogan 2000). Ethical dilemmas, as well as work needed to control their growth in the targeted tissue, seem to be the main concerns limiting their use. Animal experimentations results are inconsistent and complexed by grafted cell death, formation of teratomas and tumours have been observed (Blum & Benvenisty 2008; Brederlau et al. 2006).

Autologous mesenchymal stem cells derived from bone marrow is still the preferred cellular source and iliac crest harvesting is the most common source. The simple approach could be through bone marrow aspirates or the harvest of cancellous bone enriched in osteoprogenitors. These cells can sometimes go through in vitro expansion before being loaded onto a scaffold or other carrier (Bernardo et al. 2011; Caplan & Correa 2011; Kuo et al.
2011). Gene therapy for insertion or activation of selected genes through transfection or electroporation is often attempted on mesenchymal stem cells (Stender et al. 2007). Due to the morbidity associated with marrow mesenchymal stem cells harvesting, need for a second surgical site, limited amounts of grafting material and lack of mechanical stability in extensive defects composites of mesenchymal stem cells with non-autologous grafting materials are frequently used (Caplan et al. 1997; Dimitriou et al. 2011). The question is if delivery of bone marrow stromal cell containing stem cells to different wounds will assure a predictable and consistent outcome. Mesenchymal stem cells differentiation, proliferation and survival is dependent on their surrounding matrix, signals to express receptors and secrete signaling molecules. Large size defects with a potentially compromised host may offer a local environment that is not supportive or even inhibitory for bone formation. For example, it has been shown that disruption of integrin activity in mesenchymal stem cells will result in cell death and lack of differentiation (Popov et al. 2011). Various combinations have been prepared in an attempt to find a predictable and consistent graft (Schofer et al. 2011). Notably, at present, even if the number of mesenchymal stem cells is high, without the right matrix and cytokine’s support bone differentiation and maturation may not occur.

6. Inducer molecules

The ability of demineralized bone matrix to induce bone formation in the subcutaneous sites of rodents, as reported by Dr. Urist, revolutionized our approach to bone therapy and studies of bone regeneration (Urist 1965; Urist et al. 1967). These studies demonstrated that the non-mineralized fraction of the bone stores molecules that can derive osteogenic differentiation and initiate bone formation in ectopic sites. Factors such as BMP’s consist of only a very small fraction of the bone matrix and cannot be purified from bone for scientific or clinical use; however, these factors were cloned and prepared as recombinant molecules or peptides with very potent biological activity (Reddi &Cunningham 1993; Sampath et al. 1992). Inducer molecules can be delivered in a carrier or integrated into expression vehicles through ex vivo transfer to grafted cells, or infected through viruses that will target the tissues; these approaches fall under the category of gene therapy (Table 1)(Franceschi et al. 2000; Mason et al. 1998). Transient transfection and conditional expression approaches achieved in mice and other animals, thorough adeno and lentiviral, as well as non-viral, approaches such as electroporation (Franceschi et al. 2000; Holstein et al. 2009; Kawai et al. 2006). Gene delivery approaches being used in an ex vivo and in vivo gene delivery can also be utilized in humans to deliver genes to marrow stromal cells (Belmokhtar et al. 2011; Chen et al. 2011). Expression control modifications at embryo through transgenic animals or conditional modifications which, dependent on the initiator or temporary gene alteration in adult animals, assist in determining the relative importance of cell, matrix or inducer molecules to mineralized tissue healing. Gene therapy is still not available for regular clinical use, due to inability to assure target of specific cells only and adequate control over the gene transfer transcription, translation and expression in a temporal and spatial manner that will support bone regeneration. Other issues limiting clinical use are concern of viral vectors, control on the expression, immune response and potential for other non-controlled mutations. Moreover, The applications of gene transfer and control in human is not always
as efficient or predictable as shown in rodents or primates in animal experiment models (Gomes & Fernandes 2011; Sharma et al. 2011).

| Matrix grafting | Cellular grafting | Inducer molecules | Techniques |
|-----------------|-------------------|-------------------|------------|
| Vascularised Graft | mesenchymal stem cells - bone marrow aspirate | (bone morphogenetic proteins (BMPs) | Gene therapy - transfection, Transduction |
| Matrix molecules - Collagen, fibrin, hyaluronic acid, BSP, OPN | Cancellous graft - iliac, distal femur, proximal or distal tibia | platelet-derived growth factor-PDGF, Fibroblast growth factor, Vascular Endothelial growth factor, | Recombinant proteins |
| Mineral - Hydroxyapatite, β- Tricalcium phosphate (TCP), | Other sources of adult stem cells - peripheral blood adipose | Transforming growth factor’s | Peptides |
| Polymers - poly (lactic-coglycolic acid) (PLGA), alginate and chitosan | ES-Embryonic stem cells | insulin-like growth factor-I,II | Nanotechnology |
| Calcium phosphate or sulphate, glass ceramics | Umbilical cord | endothelial growth factor | Cellular in-vitro expansion, differentiation induction, |
| DBM- Demineralised bone matrix | Dental – follicle, pulp, periodontal | Hormones - parathyroid hormone, Growth Hormone | Scaffolds - Three-dimensional porous scaffolds, coated, biodegradable |
| cancellous bone allograft | | Pepetides- FHRRIKA, FNIII 7-10, P15, DGEA (Asp-Gly-Glu-Ala), RGD, PTH 1-34, and PTH 1-84 | Morcellized bone grafting, freeze-drying |
| Cortical | | Denosumab-antibody to RANKL | Purified proteins Membranes, Mesh |
| Block graft | | agonists of the prostaglandin receptors EP2 and EP4 | Distraction osteogenesis |

Table 1. Classification of Grafting
At present, about 20 BMPs have been identified, with about eight having osteogenic effects: BMP-2, 3, 4, 6, 7, 8, 12, 14. BMP-7 or OP-1 is the subject of many studies approved for clinical use and exhibits a very potent osteoinductive effect in vivo and in vitro. BMP-7 effects on mesenchymal stem cells include increased migration, differentiation and induction of bone formation through endochondral as well as intramembranous ossification (Giannoudis et al. 2009). Many other cytokines are the subject of ongoing investigations and use such as platelet-derived growth factor (PDGF), transforming growth factor-β (TGF β), insulin-like growth factor-I and II (IGF), vascular endothelial growth factor (VEGF), endothelial growth factor (EGF), parathyroid hormone (PTH), growth hormone (GH) and fibroblast growth factor (FGF). Some are prepared as synthetic peptides where only the active sequence is synthesized; often the peptide will be more potent that the whole molecule. Examples of these peptides include PTH [PTH(1-34); Forteo (or teriparitide) and PTH 1-84, P24 is a 24-amino acid peptide derived from BMP2 capable of induction of ectopic bone (Lin et al. 2010; Wu et al. 2008). The growth factors that are approved for clinical use in human and received the Food and Drug Administration (FDA) approval for bone regeneration are BMP-2, 7 and PDGF-BB (Caplan & Correa 2011; Kanakaris et al. 2008; Lynch, S. E. et al. 2006; Mulconrey et al. 2008). These growth factors will predictably stimulate bone formation, and when compared to the gold standard of autologous bone grafting, these growth factors meet the expectations of inducing bone regeneration in a high percentage of the clinical cases (Garrison et al. 2011). Advantages include ample supply, convenient grafting carriers, osteoinduction, no need for a second surgical site and no significant immune responses. The reported concerns are no cellular component, no osteoconduction support, lower mechanical strength of the newly formed bone, expensive and variability in induction. These growth factors are carried or released by various materials that may alter their effects and potency (Nauth et al. 2011). It is beyond the scope of this chapter to describe the molecular mechanism known for each of these growth factors or the expression of their receptors and associated signaling pathways. Each of these growth factors is a subject of numerous clinical trials and reports and suggestions on its most potent use for bone regeneration. Their effects are dependent on the availability of cells, the expression of the appropriate receptors and biological half-life at the bone defect.

Bone formation can also be induced by non-growth factor molecules, such as matrix components or proteins that will encourage mesenchymal stem cells cell adhesion, migration, proliferation, differentiation and survival (Popov et al. 2011). Matrix components like collagen will not only induce bone cells directly but also their ability to bind other potent molecules, such as growth factors, thrombospondin, decorin, biglycan, OPN, OCN, BSP, fibronectin, vitronectin and hydroxyapatite (Bentley & Tralka 1983; Ber et al. 1991; Bergmann et al. 1990). Control of expression of receptors to mediate bone matrix adhesion would be another approach, through antibodies or fragments that will induce their expression; for example, the Denosumab human monoclonal antibody that inhibits osteoclastic activity through binding to RANKL and safe even for systemic use (Miller 2009).

Matrix proteins can be used as purified proteins or synthetic peptides. Purified collagen is one source of a primary matrix molecule derived from human, bovine or porcine sources as purified fibrillar collagen or composite with other minerals that can be use to fill
defects or mixed with other grafts (Gleeson et al. 2011; Muschler et al. 1996; Thula et al. 2011). Proteoglycans, such as hyaluronan, can be purified from the human umbilical cord, cultures of cells or bacteria. The non-collagenous proteins of the bone, such as OPN, BSP and OCN, can also be purified and used to coat biomaterials, or mixed with grafting materials. The use of synthetic peptides as a whole molecule or just active sequence is a more accurate approach, as it may be missing post-translational modifications found on the native purified protein. It would be a cleaner and safer product as far as immune reactions or carrying impurities for clinical use. Recombinant molecules and synthetic peptide technologies are becoming more popular as well as more accurate, pure and have reduced variability in mediating osteogenic cell adhesion and bone formation. RGD (arginine - glycine - aspartate) is a well-characterized sequence in number of matrix proteins including fibronectin, OPN, BSP and vitronectin that mediate attachment of osteogenic cells to integrin receptors (Hsiong et al. 2009; Pallu et al. 2009). RGD will usually ligate αVβ3-integrin, but also αvβ1, α8β1, αvβ8, αvβ6, αvβ5, and αIIbβ3. RGD being synthesised as linear as well as cyclic peptide as some studies also suggest that the cyclic form may offer better presentation that is more potent in inducing osteoblastic differentiation (Hsiong et al. 2009). Collagen I adheres to bone cells via α2β1 integrin receptor (Mizuno et al. 2000) through DGEA (Asp-Gly-Glu-Ala) motif. Its recognition sequences and competition for this association with DGEA peptide could inhibit osteoblastic differentiation (Takeuchi et al. 1996). Fibronectin fragments FNIII 7-10, α5β1 integrin specific enhanced osteoblastic differentiation in bone marrow stromal cells and can upregulate adherence to titanium implants (Petrie et al. 2008). P15 is a 15-amino acid sequence derived from Collagen I, α1 chain and in clinical use (Gomar et al. 2007; Pettinicchio et al.). P15 enhances osteoblastic cell adhesion and differentiation to osteoblasts. Other peptides will be FHRRIK, derived from the heparin binding site of BSP, human vitronectin peptide HVP (351-359) and osteopontin-derived peptides (Healy et al. 1999; von der Mark et al. 2010).

Most of these peptides and growth factors show great promise in in vitro studies and great potential in human trials and therapy (Bosetti et al. 2007; Nauth et al. 2011; Rose et al. 2004). Unfortunately, the animal and human analyses seem to exhibit wide variability (Faour et al. 2011; Giannoudis & Dinopoulos 2010; Papakostidis et al. 2008; Shekaran & Garcia 2011). An important factor in the application of these peptides and growth factors is the delivery system, as are the biochemical properties of the surrounding matrix and accessibility of the cells and the relevant receptors for their signaling.

The nature of the biomaterial, the surface to be coated or the carrying polymer, scaffold or gel will have an impact on the availability of the inducer or the ligand used to attach the differentiating bone cells. A common problem will be the hydrophobic surfaces of biomaterials, which will be covered by plasma and absorb abundant proteins such as albumin. This will make any ligand attached to the biomaterial less accessible, while more hydrophilic surfaces, such as culture dishes coated with ECM proteins, will encourage cell adhesions. Nanotechnology used to space ligands, such as RGD, affects cells adhesion, clustering and increases affinity between ligand and the receptors through both chemical and physical modifications. These approaches will enable osteoprogenitors to differentiate and migrate in the desired direction (Hirschfeld-Warneken et al. 2008). Designs aimed at
creating the right topography of the biomaterial, as well as chemical alteration of serine residues or energy molecules such as purines that will change the availability of the inducer, will have impact on the ability of osteoprogenitors to differentiate (Costa et al. 2011; Mager et al. 2011; Vater et al. 2009).

7. Concluding remarks

The number of bone regeneration tools is growing every day, some but not all of which are listed above (Table 1). Unfortunately, there is no single tool available that can predictably match the gold standard of autologous marrow bone grafting. To restore the missing bone matrix, cells and inducer molecules need to act in a synergistic manner. Indeed, the new regenerative approaches are based on composite grafting, including matrix replacement, mesenchymal stem cells and inducer molecules. Most composites grafts focus on merging osteoconductive scaffolds with osteoinductive agents, such as BMP, or with cells (Bueno & Glowacki 2009; Lin et al. 2010). Nanotechnology improves matrix characteristics for cell adherence, survival and differentiation, delivery vehicle for cell, proteins or gene carriers also improve macro mechanical properties (Shekaran & Garcia 2010; Smith, I. O. et al. 2009; Zhang et al. 2007). The research of forming a scaffold with organic and non-organic parts, which is mechanically strong, bioreosorbable, carries inducer molecules and cells, and will adhere to the newly forming bone and still be affordable, is challenging. These are hard objectives to achieve. At present, a composite graft that can match the success of autologous marrow bone grafting does not exist.

The question is whether our quest for an ideal composite graft that will fit and regenerate most, if not all, bone wounds in every host is a realistic one. This chapter classified the three main components needed to restore missing bone tissue and outlined some of the tools and techniques (Figure 1). It is unlikely that composite grafts will be successful as autogenous grafting without having individual “custom made composite graft”. We can mix autogenous marrow aspirates with the scaffold, but still most of the grafted components will not derive from the host. Host factor variables should dictate our regenerative approach for supplementing either matrix, cellular and inductive molecules at the right composition to increase our success. Bony defects are rarely uniform and healing patterns may vary, especially in human subjects. Other than local factors, host factors such as age, medications and chronic conditions may impact wound healing in general. Our future ability to design and adapt our regenerative tools may aid in boosting critical wound healing factors required in a compromised site or individual.

A different approach is suggested, in which the clinical team will be able to identify the difficulties associated with particular wounds, such as size, mechanics, blood supply and whether or not the bone is load bearing. Host factors to be considered include age, medications and other systemic conditions that may compromise wound healing. Based on these analyses of the available tools (Table 1), a list will be presented to the lab with physical, chemical and inductive requirements. An individual composite graft will be constructed for the wound that will meet and boost the particular requirements of the specific wound. With advancement of clinical diagnosis and scientific and biotechnological tools, this approach may be more predictable in achieving bone regeneration.
8. References

Alberius, P. and M. Gordh (1996). Failure of onlay bone grafts to integrate over the calvarial suture: observations in adult isogeneic rats. *J Craniomaxillofac Surg* 24(4): 251-5

Ameer, G. A., T. A. Mahmood and R. Langer (2002). A biodegradable composite scaffold for cell transplantation. *J Orthop Res* 20(1): 16-9.

Aronow, M. A., L. C. Gerstenfeld, T. A. Owen, M. S. Tassinari, G. S. Stein and J. B. Lian (1990). Factors that promote progressive development of the osteoblast phenotype in cultured fetal rat calvaria cells. *J Cell Physiol* 143(2): 213-21

Artzi, Z., Givol, N., Rohrer, M.D., Nemcovsky, C.E., Prasad, H.S. & Tal, H. (2003) Qualitative and quantitative expression of bovine bone mineral in experimental bone defects. Part 1: Description of a dog model and histological observations. Journal of Periodontology 74, 1143-1152.

Artzi, Z., Givol, N., Rohrer, M.D., Nemcovsky, C.E., Prasad, H.S. & Tal, H. (2003) Qualitative and quantitative expression of bovine bone mineral in experimental bone defects. Part 2. Morphometric analysis. Journal of Periodontology 74, 1159-1160

Au, A. G., V. James Raso, A. B. Liggins and A. Amirfazli (2003). Contribution of loading conditions and material properties to stress shielding near the tibial component of total knee replacements. *J Biomech* 40(6): 1410-6.0021-9290 (Print) 0021-9290 (Linking)

Augello, A. and C. De Bari (2010). The regulation of differentiation in mesenchymal stem cells. *Hum Gene Ther* 21(10): 1226-38.1557-7422 (Electronic) 1043-0342 (Linking)

Awad, H. A., D. L. Butler, G. P. Boivin, F. N. Smith, P. Malaviya, B. Huibregtse and A. I. Caplan (1999). Autologous mesenchymal stem cell-mediated repair of tendon. *Tissue Eng* 5(3): 267-77

Bahadur, G., M. Morrison and L. Machin (2011). Beyond the ‘embryo question’ : human embryonic stem cell ethics in the context of biomaterial donation in the UK. *Reprod Biomed Online* 21(7): 868-74.1472-6491 (Electronic)1472-6483 (Linking)

Behnamghader, A., N. Bagheri, B. Raissi and F. Moztarzadeh (2008). Phase development and sintering behaviour of biphasic HA-TCP calcium phosphate materials prepared from hydroxyapatite and bioactive glass. *J Mater Sci Mater Med* 19(1): 197-201.0957-4530 (Print) 0957-4530 (Linking)

Belmokhtar, K., T. Bourguignon, M. E. Worou, G. Khamis, P. Bonnet, J. Domenech and V. Eder (2011). Regeneration of Three Layers Vascular Wall by using BMP2-Treated MSC Involving HIF-1alpha and Id1 Expressions Through JAK/STAT Pathways. *Stem Cell Res*.1558-6804 (Electronic)1550-8943 (Linking)

Benayahu, D. (2000). The Hematopoietic Microenvironment: The Osteogenic Component of Bone Marrow: Cell Biology and Clinical Application. *Hematology* 4(5): 427-435,1607-8454 (Electronic)1024-5332 (Linking)

Bentley, S. A. and T. S. Tralka (1983). Fibronectin-mediated attachment of hematopoietic cells to stromal elements in continuous bone marrow culture. *Exp Hematol* 11(2): 129-38.0301-472X (Print)0301-472X (Linking)

Ber, R., T. Kubota, J. Sodek and J. E. Aubin (1991). Effects of transforming growth factor-beta on normal clonal bone cell populations. *Biochem Cell Biol* 69(2-3): 132-40

Bergmann, P., N. Nijs-De Wolf, T. Pepersack and J. Corvilain (1990). Release of parathyroid hormonelike peptides by fetal rat long bones in culture. *J Bone Miner Res* 5(7): 741-53
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Bernardo, M. E., A. M. Cometa, D. Pagliara, L. Vinti, F. Rossi, R. Cristantielli, G. Palumbo and F. Locatelli (2011). Ex vivo expansion of mesenchymal stromal cells. Best Pract Res Clin Haematol 24(1): 73-81.1532-1924 (Electronic) 1521-6926 (Linking)

Binderman, I., A. Yaffe, R. Zohar, D. Benayahu and H. Bahar (2011). Tissue engineering of bone: an ectopic rat model. Front Biosci (Schol Ed) 3: 61-81.1495-0524 (Electronic) 1945-0516 (Linking)

Blair, H. C., M. Zaidi, C. L. Huang and L. Sun (2008). The developmental basis of skeletal cell differentiation and the molecular basis of major skeletal defects. Biol Rev Camb Philos Soc 83(4): 401-15.1469-185X (Electronic)0006-3231 (Linking)

Blum, B. and N. Benvenisty (2008). The tumorigenicity of human embryonic stem cells. Adv Cancer Res 100: 133-58.0065-230X (Print) 0065-230X (Linking)

Bosetti, M., F. Boccafosci, M. Leighheb and M. F. Cannas (2007). Effect of different growth factors on human osteoblasts activities: a possible application in bone regeneration for tissue engineering. Biomol Eng 24(6): 613-8.1389-0344 (Print)1389-0344 (Linking)

Brederlau, A., A. S. Correia, S. V. Anisimov, M. Elmi, G. Paul, L. Roybon, A. Morizane, F. Bergquist, I. Riebe, U. Nannmark, M. Carta, E. Hanse, J. Takahashi, Y. Sasai, K. Funa, P. Brundin, P. S. Eriksson and J. Y. Li (2006). Transplantation of human embryonic stem cell-derived cells to a rat model of Parkinson's disease: effect of in vitro differentiation on graft survival and teratoma formation. Stem Cells 24(6): 1433-40.1066-5099 (Print)1066-5099 (Linking)

Bueno, E. M. and J. Glowacki (2009). Cell-free and cell-based approaches for bone regeneration. Nat Rev Rheumatol 5(12): 685-97.1759-4804 (Electronic)1759-4790 (Linking)

Caplan, A. I. and D. Correa (2011). PDGF in bone formation and regeneration: New insights into a novel mechanism involving MSCs. J Orthop Res.1554-527X (Electronic)0736-0266 (Linking)

Caplan, A. I., M. Elyaderani, Y. Mochizuki, S. Wakitani and V. M. Goldberg (1997). Principles of cartilage repair and regeneration. Clin Orthop(342): 254-69

Chajra, H., C. F. Rousseau, D. Cortial, M. C. Ronziere, D. Herbage, F. Mallein-Gerin and A. M. Freyria (2008). Collagen-based biomaterials and cartilage engineering. Application to osteochondral defects. Biomater Eng 18(1 Suppl): S33-45.0959-2989 (Print)0959-2989 (Linking)

Chen, J., H. Chen, P. Li, H. Diao, S. Zhu, L. Dong, R. Wang, T. Guo, J. Zhao and J. Zhang (2011). Simultaneous regeneration of articular cartilage and subchondral bone in vivo using MSCs induced by a spatially controlled gene delivery system in bilayered integrated scaffolds. Biomaterials 32(21): 4793-805.1878-5905 (Electronic)0142-9612 (Linking)

Cholewa-Kowalska, K., J. Kokoszka, M. Laczka, L. Niedzwiedzi, W. Madej and A. M. Osyczka (2009). Gel-derived bioglass as a compound of hydroxyapatite composites. Biomater 4(5): 055007.1748-605X (Electronic)1748-6041 (Linking)

Costa, M. A., A. Barbosa, E. Neto, A. Sa-e-Sousa, R. Freitas, J. M. Neves, T. Magalhaes-Cardoso, F. Ferreirinha and P. Correia-de-Sa (2011). On the role of subtype selective adenosine receptor agonists during proliferation and osteogenic differentiation of human primary bone marrow stromal cells. J Cell Physiol 226(5): 1353-66.1097-4652 (Electronic)0021-9541 (Linking)

www.intechopen.com
Costain, D. J. and R. W. Crawford (2009). Fresh-frozen vs. irradiated allograft bone in orthopaedic reconstructive surgery. *Injury* 40(12): 1260-4.1879-0267 (Electronic)0020-1383 (Linking)

Derwin, K. A., A. R. Baker, J. P. Iannotti and J. A. McCarron (2009). Preclinical models for translating regenerative medicine therapies for rotator cuff repair. *Tissue Eng Part B Rev* 16(1): 21-30.1937-3376 (Electronic)1937-3368 (Linking)

Dimitriou, R., E. Jones, D. McGonagle and P. V. Giannoudis (2011). Bone regeneration: current concepts and future directions. *BMC Med* 9: 66.1741-7015 (Electronic)1741-7015 (Linking)

Falla, N., V. Van, J. Bierkens, B. Borremans, G. Schoeters and U. Van Gorp (1993). Characterization of a 5-fluorouracil-enriched osteoprogenitor population of the murine bone marrow. *Blood* 82(12): 3580-91

Faour, O., R. Dimitriou, C. A. Cousins and P. V. Giannoudis (2011). The use of bone graft substitutes in large cancellous voids: Any specific needs? *Injury*.1879-0267 (Electronic)0020-1383 (Linking)

Franceschi, R. T., D. Wang, P. H. Krebsbach and R. B. Rutherford (2000). Gene therapy for bone formation: in vitro and in vivo osteogenic activity of an adenovirus expressing BMP7. *J Cell Biochem* 78(3): 476-86.0730-2312 (Print)0730-2312 (Linking)

Friedrich, J. B., S. L. Moran, A. T. Bishop and A. Y. Shin (2009). Free vascularized fibula grafts for salvage of failed oncologic long bone reconstruction and pathologic fractures. *Microsurgery* 29(5): 385-92.1098-2752 (Electronic)0738-1085 (Linking)

Garrison, K. R., I. Shemilt, S. Donell, J. J. Ryder, M. Mugford, I. Harvey, F. Song and V. Alt (2011). Bone morphogenetic protein (BMP) for fracture healing in adults. *Cochrane Database Syst Rev*(6): CD006950.1469-493X (Electronic)1361-6137 (Linking)

Gaskill, T. R., J. R. Urbaniak and J. M. Aldridge, 3rd (2009). Free vascularized fibular transfer for femoral head osteonecrosis: donor and graft site morbidity. *J Bone Joint Surg Am* 91(8): 1861-7.1558-1373

Ghilzon, R., C. A. McCulloch and R. Zohar (1999). Stromal mesenchymal progenitor cells. *Leuk Lymphoma* 32(3-4): 211-21 the above report in

Giannoni, P., S. Scaglione, A. Daga, C. Ilengo, M. Cilli and R. Quarto (2009). Short-time survival and engraftment of bone marrow stromal cells in an ectopic model of bone regeneration. *Tissue Eng Part A* 16(2): 489-99.1937-335X (Electronic)1937-3341 (Linking)

Giannoudis, P. V. and H. T. Dinopoulos (2010). Autologous bone graft: when shall we add growth factors? *Orthop Clin North Am* 41(1): 85-94; table of contents.1558-1373 (Electronic)0030-5898 (Linking)

Giannoudis, P. V., N. K. Kanakaris, R. Dimitriou, I. Gill, V. Kolimara and R. J. Montgomery (2009). The synergistic effect of autograft and BMP-7 in the treatment of atrophic nonunions. *Clin Orthop Relat Res* 467(12): 3239-48.1528-1132 (Electronic)0009-921X (Linking)

Gleeson, J. F., N. A. Plunkett and F. J. O’Brien (2011). Addition of hydroxyapatite improves stiffness, interconnectivity and osteogenic potential of a highly porous collagen-based scaffold for bone tissue regeneration. *Eur Cell Mater* 20: 218-30.1473-2262 (Electronic)1473-2262 (Linking)
Gomar, F., R. Orozco, J. L. Villar and F. Arrizabalaga (2007). P-15 small peptide bone graft substitute in the treatment of non-unions and delayed union. A pilot clinical trial. *Int Orthop* 31(1): 93-9.0341-2695 (Print)0341-2695 (Linking)

Gomes, P. S. and M. H. Fernandes (2011). Rodent models in bone-related research: the relevance of calvarial defects in the assessment of bone regeneration strategies. *Lab Anim* 45(1): 14-24.1758-1117 (Electronic)0023-6772 (Linking)

Goodwin, H. S., A. R. Bicknese, S. N. Chien, B. D. Bogucki, C. O. Quinn and D. A. Wall (2001). Multilineage differentiation activity by cells isolated from umbilical cord blood: expression of bone, fat, and neural markers. *Biol Blood Marrow Transplant* 7(11): 581-8.

Gosain, A. K., H. Chim and J. S. Arneja (2009). Application-specific selection of biomaterials for pediatric craniofacial reconstruction: developing a rational approach to guide clinical use. *Plast Reconstr Surg* 123(1): 319-30.1529-4242 (Electronic)

Grabowski, P. (2009). Physiology of bone. *Endocr Dev* 16: 32-48.1662-2979 (Electronic)1421-7082 (Linking)

Griffin, X. L., M. L. Costa, N. Parsons and N. Smith Electromagnetic field stimulation for treating delayed union or non-union of long bone fractures in adults. *Cochrane Database Syst Rev*(4): CD008471.1469-493X (Electronic)1361-6137 (Linking)

Gupta, V., G. H. Mun, B. Choi, A. Aseh, L. Mildred, A. Patel, Q. Zhang, J. E. Price, D. Chang, G. Robb and A. B. Mathur (2011). Repair and Reconstruction of a Resected Tumor Defect Using a Composite of Tissue Flap-Nanotherapeutic-Silk Fibroin and Chitosan Scaffold. *Ann Biomed Eng.*1521-6047 (Electronic)0090-6964 (Linking)

Hak, D. J. and J. L. Pittman (2010). Biological rationale for the intramedullary canal as a source of autograft material. *Orthop Clin North Am* 41(1): 57-61; table of contents.1558-1373 (Electronic)0030-5898 (Linking)

Hamer, A. J., J. R. Strachan, M. M. Black, C. J. Ibbotson, I. Stockley and R. A. Elson (1996). Biochemical properties of cortical allograft bone using a new method of bone strength measurement. A comparison of fresh, fresh-frozen and irradiated bone. *J Bone Joint Surg Br* 78(3): 363-8.0301-620X (Print)

Healy, K. E., A. Rezania and R. A. Stile (1999). Designing biomaterials to direct biological responses. *Ann N Y Acad Sci* 875: 24-35.0077-8923 (Print)0077-8923 (Linking)

Heo, C. Y., S. Kwon, G. H. Back and M. S. Chung (2008). Complications of distraction lengthening in the hand. *J Hand Surg Eur Vol* 33(5): 609-15.0171-9335 (Print)0171-9335 (Linking)

Hirschfield-Warneken, V. C., M. Arnold, A. Cavalcanti-Adam, M. Lopez-Garcia, H. Kessler and J. P. Spatz (2008). Cell adhesion and polarisation on molecularly defined spacing gradient surfaces of cyclic RGDFK peptide patches. *Eur J Cell Biol* 87(8-9): 743-50.0171-9335 (Print)0171-9335 (Linking)

Hoeppner, L. H., F. J. Secreto and J. J. Westendorf (2009). Wnt signaling as a therapeutic target for bone diseases. *Expert Opin Ther Targets* 13(4): 485-96.1744-7631 (Electronic)1472-8222 (Linking)

Holstein, J. H., P. Garcia, T. Histing, A. Kristen, C. Scheuer, M. D. Menger and T. Pohlemann (2009). Advances in the establishment of defined mouse models for the study of fracture healing and bone regeneration. *J Orthop Trauma* 23(5 Suppl): S31-8.1531-2291 (Electronic)0890-5339 (Linking)
Holy, C. E., M. S. Shoichet and J. E. Davies (2000). Engineering three-dimensional bone tissue in vitro using biodegradable scaffolds: investigating initial cell-seeding density and culture period. *J Biomed Mater Res* 51(3): 376-82.

Honda, M. J., M. Imaizumi, S. Tsuchiya and C. Morsczeck (2011). Dental follicle stem cells and tissue engineering. *J Oral Sci* 52(4): 541-52.1880-4926 (Electronic)1343-4934 (Linking)

Hsiong, S. X., T. Boontheekul, N. Huebsch and D. J. Mooney (2009). Cyclic arginine-glycine-aspartate peptides enhance three-dimensional stem cell osteogenic differentiation. *Tissue Eng Part A* 15(2): 263-72.1937-3341 (Print)1937-3341 (Linking)

Huang, X. and X. Miao (2007). Novel porous hydroxyapatite prepared by combining H2O2 foaming with PU sponge and modified with PLGA and bioactive glass. *J Biomater Appl* 21(4): 351-74.0885-3282 (Print)0885-3282 (Linking)

Intini, G., S. Andreana, F. E. Intini, R. J. Buhite and L. A. Bobek (2007). Calcium sulfate and platelet-rich plasma make a novel osteoinductive biomaterial for bone regeneration. *J Transl Med* 5: 13.1479-5876 (Electronic)1479-5876 (Linking)

Javed, A., H. Chen and F. Y. Ghori Genetic and transcriptional control of bone formation. *Oral Maxillofac Surg Clin North Am* 22(3): 283-93, v.1558-1365 (Electronic)1042-3699 (Linking)

Jones, G. L., G. M. McCluskey, 3rd and D. T. Curd (2000). Nonunion of the fractured clavicle: evaluation, etiology, and treatment. *J South Orthop Assoc* 9(1): 43-54.1059-1052 (Print)1059-1052 (Linking)

Jose, M. V., V. Thomas, K. T. Johnson, D. R. Dean and E. Nyairo (2009). Aligned PLGA/HA nanofibrous nanocomposite scaffolds for bone tissue engineering. *Acta Biomater* 5(1): 305-15.1878-7568 (Electronic)1742-7061 (Linking)

Kanakaris, N. K., G. M. Calori, R. Verdonk, P. Burssens, P. De Biase, R. Capanna, L. B. Vangosa, P. Cherubino, F. Baldo, J. Ristiniemi, G. Kontakis and P. V. Giannoudis (2008). Application of BMP-7 to tibial non-unions: a 3-year multicenter experience. *Injury* 39 Suppl 2: S83-90.1879-0267 (Electronic)0020-1383 (Linking)

Kawai, M., K. Bessho, H. Maruyama, J. Miyazaki and T. Yamamoto (2006). Simultaneous gene transfer of bone morphogenetic protein (BMP) -2 and BMP-7 by in vivo electroporation induces rapid bone formation and BMP-4 expression. *BMC Musculoskelet Disord* 7: 62.1471-2474 (Electronic)1471-2474 (Linking)

Kim, J. B., P. Leucht, K. Lam, C. Luppen, D. Ten Berge, R. Nusse and J. A. Helms (2007). Bone regeneration is regulated by wnt signaling. *J Bone Miner Res* 22(12): 1913-23.0884-0431 (Print)0884-0431 (Linking)

Kligman, M., A. Rotem and M. Roffman (2003). Cancellous and cortical morselized allograft in revision total hip replacement: A biomechanical study of implant stability. *J Biomech* 36(6): 797-802.0021-9290 (Print)0021-9290 (Linking)

Kogianni, G. and B. S. Noble (2007). The biology of osteocytes. *Curr Osteoporos Rep* 5(2): 81-8.1544-1873 (Print)1544-1873 (Linking)

Komori, T. (2011) Signaling networks in RUNX2-dependent bone development. *J Cell Biochem* 112(3): 750-5.1097-4644 (Electronic)0730-2312 (Linking)

Kuo, H. C., C. C. Chiu, W. C. Chang, J. M. Sheen, C. Y. Ou, R. F. Chen, T. Y. Hsu, J. C. Chang, C. C. Hsiao, F. S. Wang, C. C. Huang, H. Y. Huang and K. D. Yang (2011). Use of proteomic differential displays to assess functional discrepancies and
adjustments of human bone marrow- and Wharton jelly-derived mesenchymal stem cells. J Proteome Res 10(3): 1305-15.1535-3907 (Electronic)1535-3893 (Linking)

Landry, P. S., A. A. Marino, K. K. Sadasivan and J. A. Albright (1996). Bone injury response. An animal model for testing theories of regulation. Clin Orthop Relat Res(332): 260-73.0009-921X (Print)0009-921X (Linking)

Lee, C., J. Dorcil and T. E. Radomisli (2004). Nonunion of the spine: a review. Clin Orthop Relat Res(419): 71-5.0009-921X (Print)0009-921X (Linking)

Lemperle, S. M., C. J. Calhoun, R. W. Curran and R. E. Holmes (1998). Bony healing of large cranial and mandibular defects protected from soft-tissue interposition: A comparative study of spontaneous bone regeneration, osteoconduction, and cancellous autografting in dogs [In Process Citation]. Plast Reconstr Surg 101(3): 660-72

Lerou, P. H. and G. Q. Daley (2005). Therapeutic potential of embryonic stem cells. Blood Rev 19(6): 321-31.0268-960X (Print)0268-960X (Linking)

Lin, Z. Y., Z. X. Duan, X. D. Guo, J. F. Li, H. W. Lu, Q. X. Zheng, D. P. Quan and S. H. Yang (2010). Bone induction by biomimetic PLGA-(PEG-ASP)n copolymer loaded with a novel synthetic BMP-2-related peptide in vitro and in vivo. J Control Release 144(2): 190-5.1873-4995 (Electronic)0168-3659 (Linking)

Liu, W., S. Toyosawa, T. Furuiuchi, N. Kanatani, C. Yoshida, Y. Liu, M. Himeno, S. Narai, A. Yamaguchi and T. Komori (2001). Overexpression of Cbfa1 in osteoblasts inhibits osteoblast maturation and causes osteopenia with multiple fractures. J Cell Biol 155(1): 157-66.

Liu, X., W. Huang, H. Fu, A. Yao, D. Wang, H. Pan, W. W. Lu, X. Jiang and X. Zhang (2009). Bioactive borosilicate glass scaffolds: in vitro degradation and bioactivity behaviors. J Mater Sci Mater Med 20(6): 1237-43.1573-4838 (Electronic)0957-4530 (Linking)

Lynch, M. P., J. L. Stein, G. S. Stein and J. B. Lian (1995). The influence of type I collagen on the development and maintenance of the osteoblast phenotype in primary and passaged rat calvarial osteoblasts: modification of expression of genes supporting cell growth, adhesion, and extracellular matrix mineralization. Exp Cell Res 216(1): 35-45

Lynch, S. E., L. Wisner-Lynch, M. Nevins and M. L. Nevins (2006). A new era in periodontal and periimplant regeneration: use of growth-factor enhanced matrices incorporating rhPDGF. Compend Contin Educ Dent 27(12): 672-8; quiz 679-80.1548-8578 (Print)

Mager, M. D., V. Lapointe and M. M. Stevens (2011). Exploring and exploiting chemistry at the cell surface. Nat Chem 3(8): 582-9.1755-4349 (Electronic)1755-4330 (Linking)

Masaki, H. and H. Ide (2007). Regeneration potency of mouse limbs. Dev Growth Differ 49(2): 89-98.0012-1592 (Print)0012-1592 (Linking)

Mashru, R. P., M. J. Herman and P. D. Pizzutillo (2005). Tibial shaft fractures in children and adolescents. J Am Acad Orthop Surg 13(5): 345-52.1067-151X (Print)1067-151X (Linking)

Mason, J. M., D. A. Grande, M. Barcia, R. Grant, R. G. Pergolizzi and A. S. Breitbart (1998). Expression of human bone morphogenic protein 7 in primary rabbit periosteal cells: potential utility in gene therapy for osteochondral repair. Gene Ther 5(8): 1098-104 the above report in

www.intechopen.com
Matejovsky, Z., Jr., Z. Matejovsky and I. Kofranek (2006). Massive allografts in tumour surgery. *Int Orthop* 30(6): 478-83.0341-2695 (Print)0341-2695 (Linking)

Meinig, R. P. (2002). Polylactide membranes in the treatment of segmental diaphyseal defects: animal model experiments in the rabbit radius, sheep tibia, Yucatan minipig radius, and goat tibia. *Injury* 33 Suppl 2: B58-65.0020-1383 (Print)0020-1383 (Linking)

Miller, P. D. (2009). Denosumab: anti-RANKL antibody. *Curr Osteoporos Rep* 7(1): 18-22.1544-2241 (Electronic)1544-2241 (Linking)

Mishra, R., B. Basu and A. Kumar (2009). Physical and cytocompatibility properties of bioactive glass-polyvinyl alcohol-sodium alginate biocomposite foams prepared via sol-gel processing for trabecular bone regeneration. *J Mater Sci Mater Med* 20(12): 2493-500.1573-4838 (Electronic)0957-4530 (Linking)

Mizuno, M., R. Fujisawa and Y. Kuboki (2000). Type I collagen-induced osteoblastic differentiation of bone-marrow cells mediated by collagen-alpha2beta1 integrin interaction. *J Cell Physiol* 184(2): 207-13.0021-9541 (Print)0021-9541 (Linking)

Mroz, T. E., E. L. Lin, M. C. Summit, J. R. Bianchi, J. E. Keeling, Jr., M. Roberts, C. T. Vangsness, Jr. and J. C. Wang (2006). Biomechanical analysis of allograft bone treated with a novel tissue sterilization process. *Spine* J 6(1): 34-9.1529-9430 (Print)1529-9430 (Linking)

Mulconrey, D. S., K. H. Bridwell, J. Flynn, G. A. Cronen and P. S. Rose (2008). Bone morphogenetic protein (RhBMP-2) as a substitute for iliac crest bone graft in multilevel adult spinal deformity surgery: minimum two-year evaluation of fusion. *Spine (Phila Pa 1976)* 33(20): 2153-9.1528-1159 (Electronic)0362-2436 (Linking)

Muschler, G. F., S. Negami, A. Hyodo, D. Gaiser, K. Easley and H. Kambic (1996). Evaluation of collagen ceramic composite graft materials in a spinal fusion model. *Clin Orthop Relat Res* (328): 250-60.0009-921X (Print)0009-921X (Linking)

Nagata, T., H. A. Goldberg, Q. Zhang, C. Domenicucci and J. Sodek (1991). Biosynthesis of bone proteins by fetal porcine calvariae in vitro. Rapid association of sulfated sialoproteins (secreted phosphoprotein-1 and bone sialoprotein) and chondroitin sulfate proteoglycan (CS-PGIII) with bone mineral. *Matrix* 11(2): 86-100

Nauth, A., B. Ristevski, R. Li and E. H. Schemitsch (2011). Growth factors and bone regeneration: how much bone can we expect? *Injury* 42(6): 574-9.1879-0267 (Electronic)0020-1383 (Linking)

Nguyen, H., D. A. Morgan and M. R. Forwood (2007). Sterilization of allograft bone: effects of gamma irradiation on allograft biology and biomechanics. *Cell Tissue Bank* 8(2): 93-105.1389-9333 (Print)1389-9333 (Linking)

Nie, H., B. W. Soh, Y. C. Fu and C. H. Wang (2008). Three-dimensional fibrous PLGA/HAp composite scaffold for BMP-2 delivery. *Biotechnol Bioeng* 99(1): 223-34.1097-0290 (Electronic)0006-3592 (Linking)

Notoya, M., E. Otsuka, A. Yamaguchi and H. Hagiwara (2004). Runx-2 is not essential for the vitamin D-regulated expression of RANKL and osteoprotegerin in osteoblastic cells. *Biochem Biophys Res Commun* 324(2): 655-60.0006-291X (Print)0006-291X (Linking)

Osti, M., H. Philipp, B. Meusburger and K. P. Benedetto Analysis of failure following anterior screw fixation of Type II odontoid fractures in geriatric patients. *Eur Spine J* 14(32):0940-6719 (Linking)

www.intechopen.com
Owen, M. (1988). Marrow stromal stem cells. *J Cell Sci Suppl* 10: 63-76

Pallu, S., J. C. Fricain, R. Bareille, C. Bourget, M. Dard, A. Sewing and J. Amedee (2009). Cyclo-DfKRG peptide modulates in vitro and in vivo behavior of human osteoprogenitor cells on titanium alloys. *Acta Biomater* 5(9): 3581-92.1878-7568 (Electronic)1742-7061 (Linking)

Papakostidis, C., G. Kontakis, M. Bhandari and P. V. Giannoudis (2008). Efficacy of autologous iliac crest bone graft and bone morphogenetic proteins for posterolateral fusion of lumbar spine: a meta-analysis of the results. *Spine (Phila Pa 1976)* 33(19): E680-92.1528-1159 (Electronic)0362-2436 (Linking)

Parker, M. J., J. Kendrew and K. Gurusamy (2011). Radiological predictive factors in the healing of displaced intracapsular hip fractures. A clinical study of 404 cases. *Hip Int* 21(4): 393-8.1724-6067 (Electronic)1120-7000 (Linking)

Petrie, T. A., J. E. Raynor, C. D. Reyes, K. L. Burns, D. M. Collard and A. J. Garcia (2008). The effect of integrin-specific bioactive coatings on tissue healing and implant osseointegration. *Biomaterials* 29(19): 2849-57.0142-9612 (Print)0142-9612 (Linking)

Pettinicchio, M., T. Traini, G. Murmura, S. Caputi, M. Degidi, C. Mangano and A. Piattelli. Histologic and histomorphometric results of three bone graft substitutes after sinus augmentation in humans. *Clin Oral Investig.* 1436-3771 (Electronic)1432-6981 (Linking)

Pittenger, M. F., A. M. Mackay, S. C. Beck, R. K. Jaiswal, R. Douglas, J. D. Mosca, M. A. Moorman, D. W. Simonetti, S. Craig and D. R. Marshak (1999). Multilineage potential of adult human mesenchymal stem cells. *Science* 284(5411): 143-7 the above report in

Popov, C., T. Radic, F. Haasten, W. C. Prall, A. Aszodi, D. Gullberg, M. Schieker and D. Docheva (2011). Integrins alpha2beta1 and alpha11beta1 regulate the survival of mesenchymal stem cells on collagen I. *Cell Death Dis* 2: e186.2041-4889 (Electronic)

Post, S., B. M. Abdallah, J. F. Bentzon and M. Kassem (2008). Demonstration of the presence of independent pre-osteoblastic and pre-adipocytic cell populations in bone marrow-derived mesenchymal stem cells. *Bone* 43(1): 32-9.8756-3282 (Print)1873-2763 (Linking)

Reddi, A. H. and N. S. Cunningham (1993). Initiation and promotion of bone differentiation by bone morphogenetic proteins. *J Bone Miner Res* 8(Suppl 2): S499-502

Renghini, C., V. Komlev, F. Fiori, E. Verne, F. Baino and C. Vitale-Brovarone (2009). Micro-CT studies on 3-D bioactive glass-ceramic scaffolds for bone regeneration. *Acta Biomater* 5(4): 1328-37.1878-7568 (Electronic)1742-7061 (Linking)

Rhee, S. C., Y. H. Ji, N. A. Gharijian, E. S. Dhong, S. H. Park and E. S. Yoon (2010). In vivo evaluation of mixtures of uncultured freshly isolated adipose-derived stem cells and demineralized bone matrix for bone regeneration in a rat critically sized calvarial defect model. *Stem Cells Dev* 20(2): 233-42.1557-8534 (Electronic)1547-3287 (Linking)

Rose, F. R., Q. Hou and R. O. Oreffo (2004). Delivery systems for bone growth factors - the new players in skeletal regeneration. *J Pharm Pharmacol* 56(4): 415-27.0022-3573 (Print)0022-3573 (Linking)

Samapth, T. K., J. Maliakal, P. V. Hauschka, W. K. Jones, H. Sasaki, R. F. Tucker, K. H. White, J. E. Coughlin, M. M. Tucker, R. H. Pang, C. Corbett, E. Ozkaynak, H. Oppermann and D. C. Rueger (1992). Recombinant human osteogenic protein-1
(hOP-1) induces new bone formation in vivo with a specific activity comparable with natural bovine osteogenic protein and stimulates osteoblast proliferation and differentiation in vitro. J Biol Chem 267(28): 20352-62

Schofer, M. D., A. Veltum, C. Theisen, F. Chen, S. Agarwal, S. Fuchs-Winkelmann and J. R. Paletta (2011). Functionalisation of PLLA nanofiber scaffolds using a possible cooperative effect between collagen type I and BMP-2: impact on growth and osteogenic differentiation of human mesenchymal stem cells. J Mater Sci Med 22(7): 1753-62.1573-4838 (Electronic)0957-4530 (Linking)

Scott, A., K. M. Khan, V. Duronio and D. A. Hart (2008). Mechanotransduction in human bone: in vitro cellular physiology that underpins bone changes with exercise. Sports Med 38(2): 139-60.0112-1642 (Print)0112-1642 (Linking)

Secreto, F. J., L. H. Hoepchner and J. J. Westendorf (2009). Wnt signaling during fracture repair. Curr Osteoporos Rep 7(2): 64-9.1544-1873 (Linking)

Sharma, A. K., M. I. Bury, A. J. Marks, N. J. Fuller, J. W. Meisner, N. Tapaskar, L. C. Halliday, D. J. Matoka and E. Y. Cheng (2011). A nonhuman primate model for urinary bladder regeneration using autologous sources of bone marrow-derived mesenchymal stem cells. Stem Cells 29(2): 241-50.1549-4918 (Electronic)1066-5099 (Linking)

Shekaran, A. and A. J. Garcia (2010). Nanoscale engineering of extracellular matrix-mimetic bioadhesive surfaces and implants for tissue engineering. Biochim Biophys Acta 1810(3): 350-60.0006-3002 (Print)0006-3002 (Linking)

Shekaran, A. and A. J. Garcia (2011). Extracellular matrix-mimetic adhesive biomaterials for bone repair. J Biomed Mater Res A 96(1): 261-72.1549-4965 (Electronic)1549-3296 (Linking)

Smit, T. H., T. A. Engels, S. H. Sontjens and L. E. Goveraat (2010). Time-dependent failure in load-bearing polymers: a potential hazard in structural applications of polylactides. J Mater Sci Med 21(3): 871-8.1573-4838 (Electronic)0957-4530 (Linking)

Smith, I. O., X. H. Liu, L. A. Smith and P. X. Ma (2009). Nanostructured polymer scaffolds for tissue engineering and regenerative medicine. Wiley Interdiscip Rev Nanomed Nanobiotechnol 1(2): 226-36.1939-0041 (Electronic)1939-0041 (Linking)

Smith, T. O., C. Hedges, R. MacNair, K. Schankat and J. A. Wimhurst (2009). The clinical and radiological outcomes of the LISS plate for distal femoral fractures: a systematic review. Injury 40(10): 1049-63.1879-0267 (Electronic)0020-1383 (Linking)

Sotereanos, D. G., A. Y. Plakseychuk and H. E. Rubash (1997). Free vascularized fibula grafting for the treatment of osteonecrosis of the femoral head. Clin Orthop Relat Res(344): 243-56.0009-921X (Print)0009-921X (Linking)

Stender, S., M. Murphy, T. O’Brien, C. Stengaard, M. Ulrich-Vinther, K. Soballe and F. Barry (2007). Adeno-associated viral vector transduction of human mesenchymal stem cells. Eur Cell Mater 13: 93-9; discussion 99.1473-2262 (Electronic)1473-2262 (Linking)

Stolzing, A., E. Jones, D. McGonagle and A. Scutt (2008). Age-related changes in human bone marrow-derived mesenchymal stem cells: consequences for cell therapies. Mech Ageing Dev 129(3): 163-73.0047-6374 (Print)0047-6374 (Linking)

Takahashi, N., N. Udagawa and T. Suda (1999). A new member of tumor necrosis factor ligand family, ODF/OPGL/TRANCE/RANKL, regulates osteoclast differentiation and function. Biochem Biophys Res Commun 256(3): 449-55

www.intechopen.com
Takeuchi, Y., K. Nakayama and T. Matsumoto (1996). Differentiation and cell surface expression of transforming growth factor-beta receptors are regulated by interaction with matrix collagen in murine osteoblastic cells. J Biol Chem 271(7): 3938-44.0021-9258 (Print)0021-9258 (Linking)

Thula, T. T., D. E. Rodriguez, M. H. Lee, L. Pendi, J. Podschun and L. B. Gower (2011). In vitro mineralization of dense collagen substrates: a biomimetic approach toward the development of bone-graft materials. Acta Biomater 7(8): 3158-69.1878-7568 (Electronic)1742-7061 (Linking)

Tiedeman, J. J., K. L. Garvin, T. A. Kilé and J. F. Connolly (1995). The role of a composite, demineralized bone matrix and bone marrow in the treatment of osseous defects. Orthopedics 18(12): 1153-8

Urist, M. R. (1965). Bone formation by autoinduction. Science 150(698): 893-9.0036-8075 (Print)0036-8075 (Linking)

Urist, M. R., B. F. Silverman, K. Buring, F. L. Dubuc and J. M. Rosenberg (1967). The bone induction principle. Clin Orthop Relat Res 53: 243-83.0009-921X (Print)0009-921X (Linking)

Valimaki, V. V. and H. T. Aro (2006). Molecular basis for action of bioactive glasses as bone graft substitute. Scand J Surg 95(2): 95-102.1457-4969 (Print)1457-4969 (Linking)

Vater, C., A. Lode, A. Bernhardt, A. Reinstorf, C. Heinemann and M. Gelinsky (2009). Influence of different modifications of a calcium phosphate bone cement on adhesion, proliferation, and osteogenic differentiation of human bone marrow stromal cells. J Biomed Mater Res A 92(4): 1452-60.1552-4965 (Electronic)1549-3296 (Linking)

von der Mark, K., J. Park, S. Bauer and P. Schmuki (2010). Nanoscale engineering of biomimetic surfaces: cues from the extracellular matrix. Cell Tissue Res 339(1): 131-53.1432-0878 (Electronic)0302-766X (Linking)

Watt, F. M. and B. L. Hogan (2000). Out of Eden: stem cells and their niches. Science 287(5457): 1427-30

Whiteford, J. R., V. Behrends, H. Kirby, M. Kusche-Gullberg, T. Muramatsu and J. R. Couchman (2007). Syndecans promote integrin-mediated adhesion of mesenchymal cells in two distinct pathways. Exp Cell Res 313(18): 3902-13.0014-4827 (Print)0014-4827 (Linking)

Wojtowicz, A. M., A. Shekaran, M. E. Oest, K. M. Dupont, K. L. Templeman, D. W. Hutmacher, R. E. Guldberg and A. J. Garcia (2010). Coating of biomaterial scaffolds with the collagen-mimetic peptide GFOGER for bone defect repair. Biomaterials 31(9): 2574-82.1878-5905 (Electronic)0142-9612 (Linking)

Wu, B., Q. Zheng, X. Guo, Y. Wu, Y. Wang and F. Cui (2008). Preparation and ectopic osteogenesis in vivo of scaffold based on mineralized recombinant human-like collagen loaded with synthetic BMP-2-derived peptide. Biomater 3(4): 044111.1748-605X (Electronic)1748-6041 (Linking)

Xiao, Z. S., A. B. Hjelmeland and L. D. Quarles (2004). Selective deficiency of the "bone-related" Runx2-II unexpectedly preserves osteoblast-mediated skeletogenesis. J Biol Chem 279(19): 20307-13.0021-9258 (Print)0021-9258 (Linking)

Yamada, Y., K. Hozumi, F. Katagiri, Y. Kikkawa and M. Nomizu (2010). Biological activity of laminin peptide-conjugated alginate and chitosan matrices. Biopolymers 94(6): 711-20.0006-3525 (Print)0006-3525 (Linking)
Yamamoto, N., H. Akamatsu, S. Hasegawa, T. Yamada, S. Nakata, M. Ohkuma, E. Miyachi, T. Marunouchi and K. Matsunaga (2007). Isolation of multipotent stem cells from mouse adipose tissue. *J Dermatol Sci* 48(1): 43-52.0923-1811 (Print)0923-1811 (Linking)

Zhang, Y., J. Song, B. Shi, Y. Wang, X. Chen, C. Huang, X. Yang, D. Xu and X. Cheng (2007). Combination of scaffold and adenovirus vectors expressing bone morphogenetic protein-7 for alveolar bone regeneration at dental implant defects. *Biomaterials* 28(31): 4635-42.0142-9612 (Print)0142-9612 (Linking)

Zhou, S., J. S. Greenberger, M. W. Epperly, J. P. Goff, C. Adler, M. S. Leboff and J. Glowacki (2008). Age-related intrinsic changes in human bone-marrow-derived mesenchymal stem cells and their differentiation to osteoblasts. *Aging Cell* 7(3): 335-43.1474-9726 (Electronic)1474-9718 (Linking)

Zohar, R., W. Lee, P. Arora, S. Cheifetz, C. McCulloch and J. Sodek (1997a). Single cell analysis of intracellular osteopontin in osteogenic cultures of fetal rat calvarial cells. *J Cell Physiol* 170(1): 88-100

Zohar, R., C. A. McCulloch, K. Sampath and J. Sodek (1998). Flow cytometric analysis of recombinant human osteogenic protein-1 (BMP-7) responsive subpopulations from fetal rat calvaria based on intracellular osteopontin content. *Matrix Biol* 16(6): 295-306.

Zohar, R., J. Sodek and C. McCulloch (1997b). Characterization of Stromal Progenitor cells enriched by flow cytometry. *Blood* 90(9): 3471-81

Zohar, R. and H. C. Tenenbaum (2005). How predictable are periodontal regenerative procedures? *J Can Dent Assoc* 71(9): 675-80
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