Briefly about Mesenchymal Stem Cells - One of the Main Players in Bone Tissue Engineering

Abstract
Mesenchymal stem cells (MSCs) are multipotent stem cells with high self-renewal capacity, ability to differentiate into osteoblasts, adipocytes and chondroblasts, immunomodulating properties and lack of teratogenic potential. These cells have been recognized to be suitable for the needs of regenerative medicine and bone tissue engineering. The presented mini-review summarizes information about biological characteristics (advantages and challenges) of MSCs of various tissue origin.

Keywords: Mesenchymal stem cells; Bone diseases; Bone tissue engineering; Regenerative medicine

Bone Tissue Engineering
Bone tissue engineering (BTE) is a strategy combining the knowledge and principles of orthopedics, bioengineering, molecular/cellular biology, cell transplantation, and materials' science to construct substitutes that can restore and maintain normal functions of injured and diseased bone. This innovative approach involves the use of stem cells that are seeded into 3D biocompatible scaffolds and induced by appropriate stimulation (growth factors, cytokines, etc) to generate genuine new bone.

Stem cells
Stem cells (SCs) are unspecialized types of cells with the ability to proliferate, self-renew and differentiate under certain physiologic conditions and signaling. The initial concept of stem cells appeared more than one century ago. Although the major advances in clarifying the biology / behavior of SCs that occurred over the last three decades, our knowledge about their nature and ability to explore therapeutic potential are still insufficient [6,7]. As compared to embryonic stem cells (ESCs) the adult (or somatic) stem cells are rare, quiescent cells with a more limited self-renewal and differentiation. At the same time, adult stem cells have two very important advantages: they circumvent ethical issues and show less tumorigenicity. Thousands of clinical trials using stem cells are currently in progress [8].

Mesenchymal stem cells
Mesenchymal stem cells (MSCs - multipotent stem cells with fibroblast-like morphology) are of mesodermal origin, except for the facial bones, which arise from neural crest [9]. MSCs can be isolated from several adult tissues (bone marrow, adipose tissue, peripheral blood, etc) as well as from birth associated
neonatal tissues (BA/NTs) (placenta, amnion, umbilical cord and cord blood) and are found in both embryonic and adult tissues in humans. The minimal set of criteria for identification/characterization of MSCs have been proposed by the International Society for Cellular Therapy (ISCT): First, MSCs must be plastic-adherent when maintained in standard culture conditions. Second, MSCs have to be positive for the presence of CD105, CD73 and CD90 surface molecules (these markers should be expressed by more than 95% of the anticipated MSCs population) with no expression (less than 2% of the cells) of CD45, CD34, CD14 or CD11b, CD79a/CD19 and HLA-DR. Third, MSC must differentiate (at least) to osteoblasts, adipocytes and chondroblasts under standard in vitro inductive conditions [10]. The yield and quality (proliferation activity, differentiation potential) of MSCs depend on tissue source, donor age (decrease with increasing donor age) and disease stage / health status. The decreased quantity and quality of MSCs due to the advance age of the patient may limit the application of autologous MSCs in clinical practice. MSCs have been recognized to be suitable for BTE because of their relatively easy access and high availability, capacity for extensive self-renewal or expansion to generate sufficient amount, the ability to differentiate into osteoblasts and chondroblasts, immunomodulating properties, lack of teratogenic potential [2,11,12].

Bone marrow mesenchymal stem cells

The first multipotent mesenchymal stem cells identified were the bone marrow stromal/stem cells (BM-MSCs), described as colony-forming unit-fibroblasts in vitro. They are still the most frequently investigated cell type and are often designated as the gold standard [13]. BM-MSCs are very attractive for the needs of regenerative medicine because of their high differentiation potential. The main obstacles associated with their application in this field are: i) MSCs are a rare population in bone marrow (0.001 – 0.01% of the total nucleated cells); ii) harvesting bone marrow is an invasive procedure carrying a potential risk for infection. BM-MSCs are limited to a growth potential of 30-50 population doubling following ex vivo expansion [11,14,15].

Adipose stem cells

Adipose stem cells (ASCs) possess some specific features that makes them particularly suitable for the needs of BTE. For example, as compared to BMSCs, ASCs have the following advantages: i) they can be harvested in large amounts and readily expanded diminishing in this way the complications that accompany cultivation in laboratory conditions; ii) more pronounced proangiogenic activity; iii) higher genetic stability in long-term cultures [16,17]. ASCs produce biologically active molecules that play important roles in bone regeneration / remodelling (bone morphogenetic proteins-2 and -4; receptor activator of nuclear factor-kappaB ligand – RANKL; macrophage colony-stimulating factor; fibronectin; type I collagen); wound healing (fibroblast growth factor-2; keratinocyte growth factor; insulin-like growth factor-1) and angiogenesis (vascular endothelial growth factor – VEGF; insulin-like growth factor-1; matrix metalloproteinase enzymes MMP-3 and MMP-9; interleukin-8). The plasma membrane-derived vesicles (MV) secreted by ASCs allow them to influence the behaviour of targeted cells situated not only in the neighborhood (paracrine effects) but also at distant locations throughout the body stimulating their proliferation and differentiation as well as the regenerative and reparative processes. These MVs contain and deliver various growth factors, cytokines, RNAs and micro RNAs. Last, but not the least, ASCs can survive under conditions of hypoxia, making them extremely valuable for bone tissue engineering applications, where the limited blood flow to the implant can be a challenge [18,19].

Birth associated /Neonatal tissues

BA/NTs have some specific advantages as a source for MSCs isolation such as: i) as medical waste tissues some of them (placenta, umbilical cord) are easily accessible thus avoiding invasive procedures and ethical problems; ii) the presence of various ESCs, MSCs, endothelial progenitor cells and hematopoietic stem cells (CD34+; CD133+); iii) There are data suggesting that MSCs derived from BA/NTs exhibit improved proliferative capacity, life span and differentiation potential as compared to BM-MSCs [20-23]. Placental tissue can be of fetal or maternal origin which has to be taken into account during isolation and characterization of MSCs. Maternal MSCs are generally derived from the placental tissue, whereas fetal MSCs are usually obtained from the umbilical cord, amniotic membrane, and amniotic fluid [24]. Cord blood (CB) has been reported to be the most reliable and attractive source for fetal MSCs because of at least two reasons: i) no ethical and regulatory limitations associated with ESCs; ii) CB-MSCs have been suggested to be safer for clinical application as compared to ESCs (no teratoma produced by CB-MSCs have been reported) [2,25]. The availability of private and public CB banks facilitates the proper storage and possible therapeutic application of CB-MSCs. Human endometrium is a highly dynamic tissue that undergoes more than 400 menstrual cycles of proliferation, differentiation and shedding during a woman’s lifetime. It has recently been identified as an accessible source of MSCs that can be easily harvested via endometrial biopsy tissue or from menstrual blood. The biological characteristics of endometrial MSCs have been reported to be similar to those of BM-MSCs and ASCs [24,26].

Dental / Oral stem cells

Mesenchymal stem cells can be isolated from several dental locations (pulp, exfoliated primary deciduous teeth, apical papilla, dental follicle, periodontal ligament) as well as from some other oral tissues (gingiva, oral mucosa, craniofacial bone) [27]. Dental stem cells (DSCs) are derived from neural crest post natal stem cell populations with MSC-like characteristics. Compared to BM-MSCs, the DSCs appear to be more related to odontogenic than osteogenic development. At the same time, it has been reported that human dental pulp stem cells (DPSCs) can co-differentiate into osteoblasts and endothelial cells and have been suggested to be beneficial for solving one of the major problems in tissue engineering regarding angiogenesis [28]. Although widely studied for their promising clinical potential, there are some obstacles in using DSCs for cell therapy, such as their limited tissue sources and the requirement for tooth extraction. Fox example, a very low number of DPSCs can be obtained because of the small size.
of the pulp (especially in the case of exfoliated deciduous teeth). Obtaining a large amount of cells sufficient for clinical application requires long term cultivation in laboratory conditions that might reduce the differentiation potency of the cells and could result in undesired genetic and/or epigenetic changes [29]. There are data revealing that DPSCs can be cultured for 6 months without (even after cryopreservation) alterations in their morphology, expression of stem cell markers, chondrogenic and myogenic differentiation potential [30]. Gingival mesenchymal stem cells (GMSCs, isolated from gingival lamina propria) are a promising alternative to BMSCs because they are abundant, readily accessible and easily obtainable via minimal discomfort for the patient. The challenge is that GMSCs are usually highly heterogeneous cell population that can result in impaired self-renewal ability and multipotent differentiation capacity. The isolation of high purity GMSCs without fibroblast contamination is crucial for their successful application. Achieving this goal is not an easy task, because specific GMSC markers are not defined yet. On the other hand, in the course of cultivation, markers currently used in determination of the heterogeneous populations gradually decrease and disappear [31,32].

Concluding Remarks

Successful introduction of BTE in clinical practice requires overcoming of several challenges, including better clarification of biology and behavior of MSCs – one of the main players in this innovative therapeutic approach. We need to increase our knowledge about mechanisms of self-renewal and regulation of stem cell differentiation; to understand better the interactions between immune system and MSCs, to improve isolation, identification and cultivation methods, to learn more about specific characteristics of MSCs of various tissue origin and about the influence of individual patient/donor features on the quality of the cells. Current and future joint efforts of biologists, chemists, engineers and clinicians focused on the design of optimized 3D scaffolds; better study of the interactions between the scaffold, the cells and the environment, and searching the ways of influencing them in the desired direction; establishing strategies to provide vascular support to the large osseous construct - all this will contribute to the advancement of regenerative medicine and its transformation into part of the routine orthopedic practice.

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Conflict of Interest

None.

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