Resistance of Multipotent Mesenchymal Stromal Cells to Anoxia In Vitro

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1. Introduction

Oxygen balance is a corner element of tissue physiology. The damaging effects of oxygen deprivation have been under consideration over at least 100 years. Recently, interest to hypoxia and practically complete absence of oxygen referred to as anoxia has been rekindled in context of the great progress made in the studies of stem/progenitor cell function in organism. The hallmark of stem cells is ability to self-renew and maintain multipotency. This ability depends on the balance of complex signals in their microenvironment. One of the most important findings is that oxygen represents a crucial component determining stem cells homeostasis within their native tissue niche. The stem cell niche has come to refer to a defined anatomical compartment that includes cellular and acellular components that integrate both systemic and local cues to regulate the stem cells biology [Jones and Wagers, 2008; Li and Xie, 2005; Scadden, 2006; Yin and Li, 2006; Buravkova & Andreeva, 2010]. The first specialized tissue niche was described for hematopoietic cells in bone marrow [Schofield, 1978]. Cells, blood vessels, matrix glycoproteins, and the three-dimensional space formed the architecture of a highly specialized microenvironment for stem cells [Scadden, 2006]. Oxygen measurements in tissues known to harbor stem cells revealed low level of oxygen, and raised the question of whether such an environment was necessary for the niche to maintain stem cells [Braun et al., 2001; Cipolleschi et al., 1993; Erecinska and Silver, 2001]. Recent evidence has broadened the spectrum of stem cells influenced by limited oxygen supply including cancer stem cells and induced pluripotent stem cells [Brahimi-Horn & Pouysse'gur, 2007]. Low oxygen tension maintains the undifferentiated state of embryonic, hematopoietic, mesenchymal, and neural stem cell phenotypes, and affects proliferation and cell-fate commitment [Mohyeldin et al., 2010].

Multipotent mesenchymal stem/stromal cells (MMSCs) arouse interest of cell biologists because of high proliferating activity and multilineage differentiation capacity. These cells are also shown to be immunoprivilege and to possess immunosuppressive features. The MMSC properties taken together make these cells a very attractive tool for cell therapy and regenerative medicine. By and large, manifestation of the MMSC properties is strictly dependent on oxygen concentration in native milieu. Moreover, the best realization of the regenerative potential is closely associated with low or very low oxygen level in the area of tissue damage.
The chapter highlights the recent progress in evaluation of the pivotal role of low oxygen in MMSC milieu and how it uniquely modulates the MMSC properties.

2. MMSCs and microenvironmental requirements: the role of oxygen

MMSCs (a rare population of non-hematopoietic stem/progenitor cells) are the subject of increasing scientific interest due to the key role they play in physiological renewal and repair. For a long time there was only one special tissue known as a definite source for renewal and substitution of cells in mammals, humans in particular. It was bone marrow capable to produce new mature blood cells from undifferentiated hematopoietic precursors. Marrow stroma contains many cell elements including endothelial cells of vessels, reticular cells, fibroblasts, adipocytes, stromal cells, and macrophages. Among the multiple stromal cells there is a minor population of MMSCs that localizes assumingly in perivascular regions of the bone marrow [Fridenshtein et al., 1976]. According to the modern concept, this population has the capacity to differentiate into cells of mesenchymal lineage (osteoblasts, chondroblasts, adipocytes and some other types of cells) [Kolf et al., 2007; Losito et al., 2009; Mohyeldin et al., 2010].

The direct evidence of stromal progenitor cells entity in vivo was not available due to the lack of a single definitive marker; therefore, demonstration of their existence has relied primarily on retrospective in vitro assays. To date, identification and characterization of bone marrow MMSCs from various animal species and humans have been described in numerous papers. It was recognized that the main phenotypic MMSC features should satisfy the following three basic criteria: adhesion to plastic, extended self-maintenance in culture, and the capacity to differentiate into bone, cartilage, adipose and hematopoiesis-inducing stroma during transplantation in vivo or upon certain inductive stimuli in vitro [Caplan, 2007; Kolf et al., 2007].

As it has been already mentioned, firstly MMSCs were described in bone marrow. Subsequently cells with characteristics similar to MMSCs were isolated from other tissue sources, including trabecular bone, adipose tissue, synovium, skeletal muscle, lung, deciduous teeth, and human umbilical cord perivascular cells derived from the Wharton’s Jelly, peripheral blood, dental pulp, periodontal ligament and etc. (Tabl. 1) [for refs. see also Kolf et al., 2007; Augello et al., 2010].

These findings reveal that MMSCs are diversely distributed in vivo and, as a result, may occupy a ubiquitous stem cell niche. There is a hypothesis that these cells are the common source of multipotent cells in adult organism migrating constantly in various mesenchymal tissues and providing their maintenance, renewal and regeneration.

The stem/progenitor cell microenvironment consists of specific molecular, cellular, and physiological components and is subject to physical and mechanical stimuli. Although stem cells can reside in markedly different locations and have distinctly different developmental paths, low oxygen tension (termed hypoxia or, in case of extremely low $O_2$, anoxia) seems to be a common in vivo feature shared by many types of adult stem cells. Indeed, there is increasing evidence that presence/absence of oxygen is a powerful tool that regulates stem cell proliferation and differentiation [Ma et al., 2009].
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| Tissue source             | Representative References                                                                 |
|---------------------------|------------------------------------------------------------------------------------------|
| Bone marrow               | Bruder et al., 1998; Pittenger et al., 1999; Makino et al., 1999; Majumdar et al., 2000; |
|                           | Bianco et al., 2001; Shake et al., 2002; Shi & Grontos, 2003; Lee et al., 2004; Wagner et al., 2005; Romanov et al., 2006; Fehrer et al., 2007 |
| Adipose tissue            | Zuk et al., 2001, 2002; Lee et al., 2004; Rehman et al., 2004; Wagner et al., 2005; Romanov et al., 2006; Gimble et al., 2007; Schaffler et al., 2007; Buravkova et al., 2009; Madonna et al., 2009 |
| Muscle                    | Bosch et al., 2000; Black, 2001                                                           |
| Umbilical cord blood      | Wagner et al., 2005; Caballero et al., 2010                                              |
| Peripheral blood          | Zvaifler et al., 2000                                                                     |
| Dermis                    | Black, 2001                                                                              |
| Periosteum                | Caballero et al., 2010                                                                   |
| Dental pulp               | Shi & Grontos, 2003                                                                      |
| Synovial membrane         | Kurose et al., 2010                                                                      |

Table 1. MMSC tissue sources in humans.

The role of oxygen in maintaining of both self-renewal and committed status of hematopoietic cells has been described in detail [Ivanovic, 2009; Eliasson & Jonasson, 2010, Valtieri & Sorrenino, 2008]. In bone marrow, the hematopoietic compartments are bound by stromal elements [Kolf et al., 2007], mainly MMSCs, and such way that two cell types form an integral part of each other's niche. Although the importance of understanding the progenitor cell spatial distribution with respect to oxygen supply from blood vessel has long been recognized, a direct noninvasive in vivo measurement of spatial oxygen gradient in bone marrow has been a major technical hurdle. Early direct measurements revealed that bone marrow is generally hypoxic with O$_2$ in some regions as low as ~1–2% [Cipolleschi et al., 1993] and even close to anoxia 0.1% O$_2$ in the osteoblastic niche [Calvi et al., 2003]. Results from the recent in vivo studies provided a direct experimental evidence that long-term repopulating HSCs in mouse reside in hypoxic environment [Parmar et al., 2007] and that hypoxia may in fact be an essential part of microenvironment maintaining cells in the undifferentiated state. On the other hand, hypoxia increases erythropoiesis – one of hematopoietic lineage, - by EPO or, maybe, low O$_2$ [Vlaski et al., 2009].

Based on these data, one may assume that MMSC physiology as an integral part of HSCs niche is also governed mainly by hypoxic and even anoxic conditions. Unlike HSCs residing exactly in bone marrow, MMSCs localize in other perivascular tissue depots and, being involved in regenerative and reparative processes are faced with different oxygen conditions; therefore, they should possess a high degree of O$_2$-mediated plasticity. Low/extremely low (hypoxia/anoxia) oxygen partial pressure in extracellular space may be physiologic or damaging in consequence of insufficient blood supply to impaired tissue. Low oxygen can modify drastically morphologic and functional cell properties, such as viability, proliferative status, immunophenotype, and differentiation. On the other hand, it is known that different cells have different tolerance to low oxygen [Csete, 2005; Ivanovic, 2009]. MMSCs, a mixture of stem/progenitor cells that may come to be in different oxygen milieu in vivo, are an attractive experimental model to explore the intrinsic mechanisms of cell adaptation to oxygen limitation.
The major bulk of knowledge concerning mesenchymal stem/progenitors biology came from the *in vitro* studies. The high proliferative activity underlies the MMSC ability to self-renew in culture for an extended period without a dramatic decline of the telomerase activity and change of karyotype [Bruder et al., 1997; Izadpanah et al., 2006], and to form an uniform layer of adhesive spindle-shaped cells with typical fibroblast-like morphology *in vitro* in the normalized conditions, i.e. low glucose content, absence of differentiation stimuli and appropriate seeding density [Pittenger et al., 1999]. The composition of the gaseous phase in cell culture technique is the most conservative parameter and exploring atmospheric O\(_2\) concentration. It should be taken into account that oxygen concentration is significantly lower *in vivo*. Arterial blood contains about 12% oxygen, and the mean tissue level of oxygen is about 3% with considerable local and regional variation. These are values for adult organs and tissues. Mean oxygen tension in embryonic tissue (where stem cells are enriched relative to adult tissues) is considerably less. Although many papers refer to low oxygen levels in embryos as "hypoxic," they are actually "normoxic" for the time and place of development [Csete, 2005]. In the last decade it was demonstrated that low oxygen tension influences greatly biology of both embryonic and adult stem cell *in vitro* [Elfasson and Jonsson, 2010; Panchision, 2009; Silvan et al., 2009] improving the proliferative and migrating abilities and reducing differentiation and proapoptotic reaction of stem cells. These observations fueled a hypothesis that low oxygen tension could be critical, but not damaging to stem cells microenvironment.

Nowadays, *in vitro* studies of the oxygen effect on MMSC functional properties are growing in number. The initial theory that the replicating stem cell microenvironment should provide sufficient oxygen supply to support tissue growth evolved to the understanding of a more complex signaling role of oxygen in regulation of stem cells migration, differentiation, and development [Ma et al., 2009]. Despite the claim that low and extremely low O\(_2\) may be more representative of the physiological conditions for certain cell types than so-called "normoxia", low oxygen is traditionally called hypoxia in consistency with the conventional terminology.

At the moment, there is wealth of data concerning low oxygen effects on the functional properties of MMSCs *in vitro*. It was demonstrated clearly that hypoxia of different severity induces proliferation in cultured MMSCs from various species. Thus, an increased proliferation rate was demonstrated for rat bone marrow MMSCs at 5% O\(_2\) [Lennon et al., 2001; Buravkova & Anokhina, 2007,2008]. Also, proliferation of human bone marrow MMSCs was stimulated by 2% O\(_2\) [Grayson et al., 2006], 3% O\(_2\) [Fehler et al., 2007; D’Ippolito et al., 2006], and 5% O\(_2\) [Zhambalova et al., 2010]. Accelerated cell growth was observed in pig bone marrow MMSC culture at 5% O\(_2\) [Bosch et al, 2006] and murine’s bone marrow MMSCs at 8% O\(_2\) [Ren et al., 2006]. Villarruel S.M. and coauthors [2008] estimated the human bone marrow MMSC colony-forming potential at 1, 5, 10, and 20 % O\(_2\) and found that the number of CFU-F raised most at 5% O\(_2\). The data on MMSCs under low oxygen pressure are also discussed in detail in several review papers [Malda et al., 2007; Ma et al., 2009; Das et al., 2010].

Under reduced oxygen pressure MMSCs also display angiogenic activity. At 1% O\(_2\), murine’s bone marrow MMSCs migrated rapidly, formed a three-dimensional capillary-like structure in Matrigel, and synthesized more vascular endothelial growth factor (VEGF); matrix metalloproteinase (MMP)-2 mRNA expression and protein secretion were down
regulated, while those of membrane-type (MT)MMP-1 were strongly induced by hypoxia [Annabi et al., 2003]. The capillary-like structures were also demonstrated in hypoxic (5%) cultures of human marrow MMSCs [Zhambalova et al., 2010] and adipose tissue MMSCs [Grinakovskaya, personal communication].

MMSCs are considered as a perspective tool for regenerative medicine and approaches to improve MMSCs quality are being developed rapidly. The preconditioning in low oxygen medium is one of the attractive ways. It was demonstrated that preconditioning of human marrow-derived MMSCs in 1%-3% oxygen activated the Akt-signaling pathway while maintaining cell viability and cell cycle rates, induced expression of cMet, the major receptor for hepatocyte growth factor (HGF), and enhanced cMet signaling. MMSCs cultured in hypoxic conditions increased migration rate. Preconditioned normoxic and hypoxic MMSCs equally improved revascularization after surgical hind limb ischemia; however, restoration of blood flow was observed significantly earlier in mice that had been injected with hypoxic preconditioned MMSCs [Rosova et al., 2008]. According to Hu et al. [2008], subletally hypoxic close to anoxia (0.5%) preconditioning of murine bone marrow MMSCs increased expression of prosurvival and proangiogenic factors including hypoxia-inducible factor 1, angiopoietin-1, vascular endothelial growth factor and its receptor, Flk-1, erythropoietin, Bcl-2, and Bcl-xL. Caspase-3 activation in hypoxic MMSCs and population of apoptotic cells were significantly lower compared with normoxic cells in vitro. Transplantation of hypoxic vs normoxic MMSCs after myocardial infarction resulted in an increase in angiogenesis, as well as enhanced morphologic and functional benefits of stem cell therapy [Hu et al., 2008].

The adipose tissue-derived MMSCs under low oxygen pressure are of special interest because of the considerable promise for regenerative medicine and cell therapy. Most of the data on MMSCs at low O2 were gathered using bone marrow MMSCs. Much less investigations have been concerned with the hypoxia effects on stromal cells derived from adipose tissue.

In a few papers hypoxia has been shown to affect the differentiation potential of the adipose tissue-derived MMSCs. Wang et al. [2005] demonstrated that human adipose MMSCs in alginate beads did not display proliferative activity at 5% O2 in normal expansion medium; however in chondrogenic medium its growth rates was lower at 5% in comparison with 20% O2. Still, under these conditions they exhibited enhanced chondrogenic differentiation markers including collagen II, glucosaminoglycan, and chondroitin-4-sulfate production [Wang et al., 2005]. The hypoxia effect on adipose tissue MMSCs is strongly dependent on the cultivation conditions. For this reason, there are conflicting data regarding chondrogenic gene expression under induction in hypoxic conditions [Khan et al., 2007; Betre et al., 2006]. Adipose tissue MMSCs expanded in 20% O2 and transferred into a 2% O2 environment failed to differentiate robustly to either adipogenic or osteogenic lineages as compared with adipose MMSCs differentiated in normal atmospheric conditions [Lee & Kemp, 2006].

We have developed an experimental approach utilizing permanent expansion of adipose tissue derived MMSCs at a reduced oxygen tension [Buravkova et al., 2009]. In hypoxia (5% O2) MMSCs demonstrated enhanced growth exceeding that in normoxia (20% O2) in 2.9±0.2 folds (p<0.05) [Buravkova et al., 2009]. The osteogenic differentiation capacity of MMSCs was significantly reduced in hypoxia vs normoxia [Grinakovskaya et al., 2009].
After expansion at low oxygen (2%) adipose tissue derived MMSCs were able to enhance the wound-healing function. Conditioning medium of hypoxic MMSCs promoted significantly collagen synthesis and migration of human dermal fibroblasts in vitro, and reduced the wound area in animal studies. These effects were based on up-regulation of growth factors such as the vascular endothelial growth factor (VEGF) and basic fibroblast growth factor (bFGF) [Lee et al., 2009].

The data above demonstrated clearly that low oxygen concentration in MMSCs microenvironment in vitro leads to modulation of MMSCs functions rather than impairment. The question arises whether extremely low oxygen (~ 0% O2) brings damage to MMSCs?

Papers dedicated to MMSCs and anoxia are few and far between. Nevertheless, the data on the “true” anoxia effects on MMSCs in vitro do exist. For example, studies of Annexin V-positive cells in rat bone marrow MMSC culture during 12 h anoxia exposure revealed a time-dependent increase in apoptotic cells from 3% to 15%. When following a 3 h anoxic exposure these MMSCs (apoptotic rate ~ 4%) were cocultured with rat’s cardiomyocytes or injected into infarcted zone in the heart, cardiomyocyte death reduced significantly owing to the treatment with both normoxic MMSCs and anoxic MMSCs, the Bcl-2/Bax protein ratio increased and cleaved cysteine-aspartic acid protease-3 decreased; anoxic MMSCs were superior to MMSCs in the normoxic condition. Consequently, MMSCs exert the antiapoptotic effect on cardiomyocytes, partially by paracrine action. The authors assume that anoxic preconditioning may be an effective and convenient way to enhance the cardioprotective effect of MMSCs [He et al., 2009].

3. Resistance of rat’s marrow MMSCs to extremely low oxygen

Cell morphology and immunophenotype. We have examined the direct effects of 96 h anoxia (~0%O2) on rat bone marrow MMSCs. Anoxia did not affect cell morphology. The percentage of MMSCs bearing CD90, CD54, CD44, CD29 (more than 95%), CD45, CD11b (less than 0.6%) molecules, and pattern of molecules expression were identical in normoxic and anoxic cells. The slight reduction in percentage of positive cells at anoxia was observed only for the CD73 marker (72% in normoxia vs 67% in anoxia).

Cell growth. Assessment of bone marrow MMSC proliferation in 1-4th passages did not reveal inhibition of the MMSC proliferative activity [Fig. 1; Anokhina et al., 2009]. It should be mentioned that during MMSC exposure in hypoxia proliferative rate displayed a more pronounced excess over normoxic MMSCs [Buravkova et al., 2007] as compared to anoxic MMSCs. Nevertheless, the fact that MMSCs can really proliferate in anoxia seems amazing and deserves special attention.

Absence of evidence for cell proliferation inhibition/deceleration in anoxia disagrees with the results of experiments demonstrating the cell cycle arrest in the conditions of anoxia [Amellem et al., 1994; Gardner et al., 2001; Goda et al., 2003].

The mechanism of inactivation of enzymes involved in nucleic acids production and subsequent inhibition of DNA replication was proposed as an explanation of the phenomenon of the cell growth arrest in murine embryonic fibroblasts and splenic B lymphocytes in low oxygen environment. Moreover, this held true for total oxygen deprivation (0.01%) or anoxia only, but not for 0.1-1% O2 [Goda et al., 2003]. The drop in
bromodeoxyuridine incorporation into murine embryonic fibroblast DNA also confirmed growth arrest due to blockade of replication in anoxia [Gardner et al., 2001]. At very low O\(_2\) (0.01-0.13%) NHIK 3025 cells were able to enter into S-phase of cell cycle but failed to complete DNA synthesis [Amellem et al., 1994].

However, in anoxic conditions MMSCs showed a normal process of cell division. It is known that different cell types are characterized by varying resistance to O\(_2\) deprivation in microenvironment [Ivanovic, 2009; Mohyeldin et al., 2010]. If a trace amount of O\(_2\) remaining in pericellular space after gas force-out is sufficient for DNA replication, cell division can be completed successfully, otherwise cells will undergo apoptosis. On the other hand, it was shown that fibroblasts are able to recover after the cell cycle arrest in anoxic conditions [Gardner et al., 2001]. To sum up, cellular mechanisms underlying MMSCs proliferation in anoxia are not clear.

**Cell viability.** The data on MMSCs viability in normoxic vs anoxic conditions are presented on Fig. 2. Though the share of damaged (apoptotic+necrotic) cells in anoxia did not differ from that in normoxic conditions significantly, the rate of apoptotic cells was doubled in anoxia (p<0.05) (Fig. 2).

Therefore, 96 h anoxia did not lead to a considerable increase in the number of damaged cells. At the same time, anoxia appeared to induce apoptosis in MMSCs. It is believed that apoptosis may be associated with anoxia, since a significant O\(_2\) reduction increases frequency of point mutations accumulation of which can be prevented by cell death through the apoptotic path [Greijer & van der Wall, 2004]. Apoptosis due to O\(_2\) deprivation can be triggered by different mechanisms. The key role is played by ROS, JNK kinase, and cytochrome C release from mitochondria mediated, in its turn, by various factors including
p53. p53 induction may result from stabilization of HIF-1α protein, the key transcription factor involved in hypoxia and capable to start other, p53-independent mechanisms of apoptosis [Greijer & van der Wall, 2004]. Besides apoptosis, O₂ depletion can mobilize some other cell death pathways based on indirect effects of O₂ reduction. Intracellular acidosis triggered by O₂ deprivation is the most important one [Schmaltz et al., 1998].

Fig. 2. Rat bone marrow MMSC viability after 96 h of exposure in 20% O₂ - □, and ~0% O₂ - ■. AnnV+ cells - MMSCs, stained with Annexin V-FITC. PI+ cells - MMSCs stained with Propidium Iodide. Total damaged cells - (AnnV+ cells) + (PI+ cells). The averaged data of 12 independent experiments are presented as M+m.

Earlier we described the antiapoptotic effects of hypoxia on rat bone marrow MMSCs [Buravkova et al., 2007]. The antiapoptotic effect of hypoxia and proapoptotic effect of anoxia may be explained using the data on the impact of HIF phosphorylation level on cell viability. The dephosphorylated HIF-1α subunit may indicate the proapoptotic HIF effect through p53 binding, whereas phosphorylated HIF-1α does not [Suzukiet al., 2001]. It may also come in line with the data suggesting that the proapoptotic effect of low oxygen depends on HIF stabilization occurring mostly at less than 5% oxygen, while the antiapoptotic hypoxia effect takes place regardless of HIF [Jiang, Semenza, Bauer 1996; Greijer & van der Wall, 2004].

Thus, path of cell death may be determined by severity of O₂ deprivation which is illustrated by the opposing apoptotic trends in hypoxia and anoxia.

Despite the relative MMSC resistance to 96 h anoxia, a prolonged exposure (up to 3 weeks) to anoxic conditions resulted in a significant reduction in cell viability (Tabl. 2). During 3 weeks in culture the percent of damaged MMSCs in normoxia and hypoxia varied insignificantly (Tabl. 2). Prolonged MMSCs exposure in anoxia led to a drastic decrease in viability up to 22% after one week, 50% after 2 weeks and to practically total cell death after
3 weeks. Necrosis was the predominant path of cell death in anoxic MMSCs, though the percentage of apoptotic cells increased also significantly (Tab.2).

|                  | AnnV+ (%)  | PI + (%)  | Damaged cells, total (%) |
|------------------|------------|-----------|--------------------------|
| **After 1st week** |            |           |                          |
| Normoxia (20% O₂) | 5,7 ± 0,7  | 5,0 ± 0,2 | 10,8 ± 0,9               |
| Hypoxia (5% O₂)  | 4,9 ± 0,8  | 10,0 ± 0,7| 14,9 ± 0,1               |
| Anoxia (~0% O₂)  | 8,1 ± 0,7  | 13,9 ± 0,6| 22,0                     |
| **After 2 weeks** |            |           |                          |
| Normoxia (20% O₂) | 1,2 ± 0,2  | 4,9 ± 0,04| 6,2 ± 0,2                |
| Hypoxia (5% O₂)  | 3,0 ± 0,01 | 10,3 ± 0,5| 13,3 ± 0,5               |
| Anoxia (~0% O₂)  | 16,8 ± 0,4 | 42,2 ± 2,0| 58,9 ± 1,6               |
| **After 3 weeks** |            |           |                          |
| Normoxia (20% O₂) | 1,4 ± 0,4  | 8,2 ± 0,1 | 9,7 ± 0,5                |
| Hypoxia (5% O₂)  | 0,4 ± 0,1  | 7,4 ± 0,4 | 7,8 ± 0,3                |
| Anoxia (~0% O₂)  | 8,2 ± 0,4  | 89,0 ± 0,7| 97,2 ± 1,1               |

The averaged data of 5 independent experiments are presented as M±m.

Table 2. Rat bone marrow MMSC viability after prolonged exposure in normoxia, hypoxia and anoxia.

The predominance of necrosis over apoptosis in MMSCs in the course of prolonged exposure to anoxia seems quite natural. In contrast to necrosis, apoptosis is a programmed energy-dependent cell death and, therefore, cell death path is determined by energy state of cells [Leist et al., 1997].

Consequently, MMSC death due to short-term anoxia is minimal and realized primary through the apoptotic pathway. Prolonged anoxic exposure induced massive cell death associated mainly with necrosis.

Cell differentiation. Evaluation of the MMSC differentiation capacity was carried out after 8 days in anoxia because of quite rapid decline of MMSC viability with exposure extension, as was described above. Spontaneous and induced osteogenic differentiation was revealed in anoxic MMSCs with alkaline phosphatase staining; the intensity of the process was found significantly less pronounced than in normoxic MMSCs (Fig.3). The mechanism of O₂-mediated suppression of osteogenic capacity of human bone marrow MMSCs and murine osteoblasts in the condition of O₂ deprivation (hypoxia (2% O₂) and anoxia (0.02% O₂)) was demonstrated earlier. The authors made a supposition that anoxia rather than hypoxia provoked inhibition of Runx2 protein expression, the key transcription factor in osteogenesis. Runx2 suppression resulted in inhibition of nodule formation and a significant reduction in mineralization of the extracellular matrix [Salim et al., 2004].
Accumulation of lipid droplets in anoxic MMSCs indicated of differentiation in the adipogenic direction was both spontaneous and induced (Fig. 4). Further extension of exposure in anoxia caused death of differentiating MMSCs. It appears that alteration of MMSCs viability rather than of differentiating capacity was the cause of differentiation suppression in anoxia.

Thus, 96 h anoxia didn’t lead to changes in MMSC morphology, proliferation rate and immunophenotype, which may obviously indicate MMSC functional stability under reduced oxygen tension. Also, anoxia didn’t increase significantly the percentage of damaged cells despite some activation of apoptosis. Further MMSC propagation in anoxia led to progressive damage of cells mainly by necrosis in contrast to apoptosis as a main death pathway in short-term anoxia and the antiapoptotic effect of hypoxia. Short-term anoxia did not inhibit the initial stages of stimulated adipo- and osteo-differentiation [Tuncay et al., 1994; Matsuda et al., 1998]. Probably, trace oxygen is enough for some MMSCs to start differentiation and the only limiting factor is viability in anoxia rather than termination of the differentiation signaling pathways.

Fig. 3. Spontaneous (a,c,e) and induced (b,d,f) osteodifferentiation in rat bone marrow MMSCs after 8 days exposure in normoxic (20% O\textsubscript{2}) (a,b), hypoxic (5% O\textsubscript{2}) (c,d) and anoxic (~0% O\textsubscript{2}) (e,f) conditions. Alkaline phosphatase, representative images of MMSCs on 3\textsuperscript{rd} passage, 100x.
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4. Susceptibility of human adipose-tissue derived MMSCs to anoxia

In order to expand our understanding of stromal precursor adaptation to extremely low oxygen we continued studies of anoxia effects on human MMSCs from adipose tissue. MMSCs of 2-4 passages were subjected to anoxic condition over 240 hours. We did not change culture medium in the course of exposure in anoxia, which induced an additional stress from nutrient “starvation”. Cells of the same passage were placed in normoxia (20% O₂) as reference “starvation” cells, and also reference“standard” cells with regular medium replacement every third day were used.

Cell growth. To characterize cell growth, we evaluated increase in MMSC population in each experimental condition. In the “standard” condition at 20% O₂ MMSC population grew 7.1 folds (Fig. 5). After 240 h w/o medium change, i.e. under nutrients deprivation, growth of normoxic MMSCs made up only 4.4 folds. In anoxic cultures cell population increased 3.9 folds (Fig. 5). Against expectations, anoxia did not suppress MMSCs growth. Increase in

Fig. 4. Spontaneous (a,c,e) and induced (b,d,f) adipodifferentiation in rat bone marrow MMSCs after 4 days exposure in normoxic (20% O₂) (a,b), hypoxic (5% O₂) (c,d) and anoxic (~0% O₂) (e,f) conditions. Lipid droplets were evaluated with Oil Red O staining, representative images of MMSCs on 3rd passage, 400x.
MMSC number was slightly less pronounced than in normoxic MMSCs in the condition of starvation. It appears, that starvation decreased oxygen demand but did not stop cell proliferation. The mechanism regulating MMSC proliferation under increasing oxygen limitation is still unclear.

Fig. 5. MMSC increment during expansion in the following conditions: ■ - 20% O₂, with regular medium changes (control), □ -20% O₂, w/o medium replacement and ▼ ~0% O₂, w/o medium replacement. Results are representative of three independent experiments. Data are shown as M+m.

**Cell viability.** We compared MMSC viability in the experimental conditions described in the context of cell death path (Tabl. 4). After 240 h in the standard normoxic condition, the share of necrotic and apoptotic MMSCs was similar and fairly low. Growth of MMSCs in normoxia and anoxia w/o medium replacement shifted the ratio of cell death path toward necrotic. Comparison of these data with MMSC proliferation makes it evident that cell population increase is higher in anoxia and that percent of necrotic cells is same as in normoxia, which means MMSCs growth prevails in anoxia.

| Condition | AnnV+ cells (%) | PI+ cells (%) | Damaged cells, total (%) |
|-----------|-----------------|---------------|-------------------------|
| Standard culture condition | 2,25 ± 0,12 | 2,20 ± 0,11 | 4,45 ± 0,22 |
| Growing cells, 240 hrs without medium replacement | 20% O₂ | 0,70 ± 0,09 | 9,17 ± 0,42 | 9,87 ± 0,45 |
| | 20% → 0% O₂ | 2,51 ± 0,03 | 9,46 ± 0,08 | 11,97 ± 0,05 |

The averaged data of 3 independent experiments are presented as M+m.

Table 3. Human adipose-tissue derived MMSC viability after exposure in normoxia and anoxia.
Cellular organelle status. To clarify the mechanisms underlining MMSC resistance to extremely low oxygen in microenvironment, we characterized alterations in some vital parameters of cellular homeostasis (Fig. 6). After 240 h of anoxia, MMSCs demonstrated an increase in percent of MMSCs containing intracellular ROS (4% vs 1.2% in normoxia) w/o increase in mean fluorescence intensity (MFI) per a cell (the parameter describes the relative amount of fluorochrome in the cell). The number of MMSCs with active lysosomes and unchanged MFI decreased drastically in anoxia. It is interesting, that long-term anoxia neither affected the mitochondrial compartment (all cells imposed active mitochondria) nor
influenced the mitochondrial transmembrane potential. Thus, in comparison with normoxia, anoxia increased slightly the number of cells with intracellular ROS and decreased significantly the number of MMSCs with active lysosomes.

It is necessary to underline that cell growth potential and viability of human adipose-derived MMSCs depend more on nutrient’s supply than on the concentration of oxygen. The drastic increase in the number of cells with active lysosomes under combination of nutrients starvation and anoxia should be clarified further.

Taking together, presented data confirm that MMSCs from different source are extraordinary resistant to externally low oxygen in the microenvironment. In anoxic conditions MMSCs retain their properties to proliferate and differentiate in mesenchymal-specific lineages and also possess fairly high viability. Nevertheless, long-term anoxia provoked cell death in MMSCs mainly through necrotic pathway.

5. Concluding remarks

The data reviewed above and our own results demonstrate clearly that low oxygen tension is undoubtedly an important regulator of MMSCs maintenance and plays a pivotal role in architecturing the MMSCs microenvironment. With oxygen partial pressure sublethal or lethal for other cell types, MMSCs not only survive but enhance proliferative activity and slow down differentiate capacity supporting “stemness”. Low oxygen conditions accentuate the paracrine role of MMSCs by altering the soluble factor release which is also plays an important role in mobilizing MMSCs and recruiting them to site of injury. MMSC ability to outlive low/extremely low oxygen (anoxia) probably, is based on their capacity to upregulate survival pathways and increase glycolytic metabolism. MMSCs are very tolerant of oxygen starvation keeping their morphology, immunophenotype and proliferation rate, demonstrating slightly affected metabolism and potency to mesenchymal lineage differentiation. Short-term anoxia gives start to apoptosis in contrast to hypoxia which exerts the antiapoptotic effects on MMSCs. Long-term anoxia provokes progressive MMSC damage mainly through the necrotic pathway. The effects of hypoxia and anoxia may be diverse and accounted to the fact, that low oxygen simulates the in vivo conditions and can be regarded as an approximation of the physiological MMSC milieu rather than the hypoxic impact. On the contrary, anoxia may represent a real hypoxic microenvironment for these cells.

These results are very encouraging both for understanding particular mechanisms of MMSC existence in different microenvironments and cell therapy as an instrument of MMSC ex vivo modification. In view of the outstanding properties, MMSCs are considered as a perspective tool for cell therapy and regenerative medicine. These cells have already shown a great regenerative potential in preclinical studies and clinical trials. The quality of cell product for these purposes is very important. Up to now, in a few studies MMSCs ex vivo expanded at low oxygen and anoxia demonstrated regenerative properties superior to MMSCs propagated by the standard cultivation. The possibility to modify MMSC properties ex vivo opens great opportunities for implication of ‘hypoxic” or even “anoxic” protocol for MMSC expansion to meet the needs of cell therapy. Nevertheless, the question about the true hypoxic environment for mesenchymal stromal progenitor cells still has not got the final answer and invites further investigations.
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Zvaifler NJ, Marinova-Mutafchieva L, Adams G, Edwards CJ, Moss J, Burger JA, Maini RN: Mesenchymal precursor cells in the blood of normal individuals. Arthritis Res. 2000; 2:477-488.
This book reviews how severe oxygen deprivation affects biological systems - from the molecular to the ecological level. The contributing authors come from diverse regions of the world, which proves the interest in the academic analysis of oxygen deprivation. The diversity in the experimental approach scientists take, in order to understand the influence oxygen deprivation has on living systems, is apparent throughout this book. One of the presented ideas deals with the exploration and examination of the physiological, cellular and genetic characteristics of killifish embryos and nematodes exposed to anoxia. Furthermore, the book includes material on the mechanisms regulating hypoxia and anoxia tolerance and their implications of on human health issues. Finally, new methodologies to examine oxygen deprivation and the impact of human-related activities on oxygen level, within important ecological systems such as Lake Victoria, are presented. There is no doubt that the oxygen molecule is central to every stratum of biological systems.

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