Cultivation of Rat Marrow-Derived Mesenchymal Stem Cells in Reduced Oxygen Tension: Effects on In Vitro and In Vivo Osteochondrogenesis

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Rat mesenchymal stem cells (rMSCs) represent a small portion of the cells in the stromal compartment of bone marrow and have the potential to differentiate into bone, cartilage, fat, and fibrous tissue. These mesenchymal progenitor cells were maintained as primary isolates and as subcultured cells in separate closed modular incubator chambers purged with either 95% air and 5% CO₂ (20% or control oxygen) or 5% oxygen, 5% CO₂, and 90% nitrogen (5% or low oxygen). At first passage, some cells from each oxygen condition were loaded into porous ceramic vehicles and implanted into syngeneic host animals in an in vivo assay for osteochondrogenesis. The remaining cells were continued in vitro in the same oxygen tension as for primary culture or were switched to the alternate condition. The first passage cells were examined for in vitro osteogenesis with assays involving the quantification of alkaline phosphatase activity and calcium and DNA content as well as by von Kossa staining to detect mineralization. Cultures maintained in low oxygen had a greater number of colonies as primary isolates and proliferated more rapidly throughout their time in vitro, as indicated by hemacytometer counts at the end of primary culture and increased DNA values for first passage cells. Moreover, rMSCs cultivated in 5% oxygen produced more bone than cells cultured in 20% oxygen when harvested and loaded into porous ceramic cubes and implanted into syngeneic host animals. Finally, markers for osteogenesis, including alkaline phosphatase activity, calcium content, and von Kossa staining, were elevated in cultures which had been in low oxygen throughout their cultivation time. Expression of these markers was usually increased above basal levels when cells were switched from control to low oxygen at first passage and decreased for cells switched from low to control oxygen. We conclude that rMSCs in culture function optimally in an atmosphere of reduced oxygen that more closely approximates documented in vivo oxygen tension. J. Cell. Physiol. 187: 345–355, 2001. © 2001 Wiley-Liss, Inc.

The remarkable growth in the use of cell culture as a research tool which followed the pioneering work of Ross Harrison, Alexis Carrel, and others in the early part of the 20th century (Russell, 1969; Harvey, 1975) was accompanied by the equally rapid development of various undefined and defined culture media. Closely linked with some of the changes in culture media were adjustments to the atmosphere in which cells were maintained. Most of these adjustments involved variations in the concentration of carbon dioxide. With a few exceptions, such as Leibovitz’s L-15 (Leibovitz, 1963) and media based on Hanks' balanced salt solution (Freshney, 1987), most media rely on the use of 5% CO₂ in conjunction with sodium bicarbonate or other buffers to maintain proper pH (Freshney, 1987). The slight reduction in oxygen concentration coincident with the use of 5% CO₂ was not found to have any deleterious effects on cell function.

While a small number of investigators (Trowell, 1959; DeRidder and Mareel, 1978) found a requirement for higher than ambient oxygen concentrations in organ (but not cell) culture, others determined that some cells proliferate more rapidly in oxygen concentrations lower than ambient to 5% oxygen, which is sometimes not found to have any deleterious effects on cell function.

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than 20%. Cooper et al. (1958) and Zwartouw and Westwood (1958) were among the first to document these results. An explanation for the phenomenon of increased cellular proliferation in reduced oxygen tension in vitro was articulated by Packer and Fuehr (1977) and expanded by others, including Rich and Kubanek (1982) and Rich (1986), and is briefly summarized as follows. The oxygen tension (or partial pressure) in most in vitro settings (~140 mm Hg) is considerably higher than that found in most mammalian and avian tissues. For example, the partial pressure of oxygen (pO2) in arterial blood has been measured at 60–90 mm Hg by Grant and Smith (1963) and at 75–110 mm Hg by Tendevold et al. (1979). These reports, and that of Kofoid et al. (1985), place the pO2 of bone marrow in the 27–49 mm Hg range. These tensions correspond to an oxygen concentration of approximately 4–7%. While organisms have evolved sophisticated mechanisms, including the enzymes glutathione peroxidase, catalase, and superoxide dismutase, and use the antioxidants ascorbate and vitamin E to defend themselves against the toxic effects of free radicals derived from oxygen (Frank and Massaro, 1980; Halliwell, 1984), it is possible that these mechanisms are inadequate to protect cells when oxygen concentrations are unusually high. Thus, it is conceivable that many cells would function more normally in vitro at oxygen concentrations lower than 20%.

Indeed, investigators have reported that reduced oxygen tension enhances proliferation of many cell types, including bovine periosteal cells (Deren et al., 1990), bovine pericytes (Brighton et al., 1992), rat calvarial osteoblasts (Tuncay et al., 1994), chick embryonic chondrocytes (Nevo et al., 1988), fibroblasts from human (Taylor et al., 1974; Balin et al., 1984) and murine (Fisher, 1960) sources, and hematopoietic progenitor cells derived from human (Bradley et al., 1978; Magda et al., 1986; Koller et al., 1992a,b) and murine marrow or umbilical cord blood (Rich and Kubanek, 1982; Rich, 1986; Koller et al., 1992b). Moreover, in vitro cultivation in reduced oxygen affects other aspects of the physiology of some cells, as indicated by increased alkaline phosphatase expression in periosteal cells (Deren et al., 1990; Reilly et al., 1998) and pericytes (Brighton et al., 1992), enhanced responsiveness of erythropoietic cells to erythropoietin (Rich and Kubanek, 1982; Maeda et al., 1986; Rich, 1986), and increased stimulation of macrophage colony formation in response to M-CSF (Broxmeyer et al., 1990). Lastly, this laboratory showed that 5% oxygen is optimal for the cultivation and in vitro differentiation of embryonic chick limb bud mesenchymal cells (Caplan and Koutropas, 1973).

In the current study, we investigated the effect of reduced (5%) oxygen in vitro on rat marrow-derived mesenchymal stem cells (rMSCs); this oxygen tension approximates that of bone marrow in vivo, as described above. These pluripotent cells, which reside in the stromal compartment of bone marrow, have been extensively characterized (Maniatopoulos et al., 1988; Leboy et al., 1991; Dennis et al., 1992; Lennon et al., 1995; Zhang et al., 1995; Cassiede et al., 1996; Hanada et al., 1997) and are capable, given the appropriate environmental cues, of progressing along one of a number of differentiation pathways, including those giving rise to bone (Maniatopoulos et al., 1988; Aubin et al., 1990; Leboy et al., 1991; Dennis et al., 1992), cartilage (Dennis and Caplan, 1993; Lennon et al., 1995; Mann et al., 1996), fat (Wakitani et al., 1995), or muscle (Wakitani et al., 1995).

Two different models have been used in our laboratory to investigate the differentiation of rMSCs along osteochondrogenic pathways. In the first, osteochondral differentiation is induced by the addition of dexamethasone to first passage rMSCs, while untreated control cultures remain in an undifferentiated state (Maniatopoulos et al., 1988; Aubin et al., 1990; Leboy et al., 1991; Cassiede et al., 1996; Hanada et al., 1997). In the second model, rMSCs maintained in vitro as primary cultures are trypsinized and loaded into porous ceramic vehicles. Bone and cartilage develop within these composites when they are implanted subcutaneously in syngeneic host animals (Dennis et al., 1992; Dennis and Caplan, 1993).

Our goal in the present study is to address the following questions. Does cultivating rMSCs in 5% oxygen affect their rate of proliferation? Are dexamethasone-treated rMSCs more likely to express osteoblastic characteristics when cultured in reduced oxygen? Are these results altered when the oxygen concentration is changed at first passage? Finally, are rMSCs maintained in vitro in 5% oxygen affected in their ability to produce bone or cartilage in porous ceramic cubes in vivo.

The data indicate that proliferation of rMSCs is increased in both primary culture and first passage in an atmosphere of reduced oxygen, that markers for osteogenic differentiation are elevated for cells in low oxygen in first passage, and that rMSCs maintained in 5% oxygen in primary culture are likely to produce more bone and cartilage when returned to an in vivo setting than are those cultivated in 20% oxygen.

**MATERIALS AND METHODS**

**Cell culture**

* Rat mesenchymal stem cells: isolation and primary culture. Rat MSCs were isolated as previously described (Dennis et al., 1992; Lennon et al., 1995). Briefly, male F344 rats, 6–12 weeks of age, were sacrificed by CO2 asphyxiation according to Institutional Animal Care and Use Committee (IACUC) guidelines. The tibiae and femora were aseptically removed and adherent soft tissue was thoroughly debrided. After the epiphyses were removed, an 18-gauge needle attached to a 10-ml syringe was used to bore a small opening through the growth plate on the distal end of the femora and the proximal end of the tibiae. A 10-ml syringe fitted with a fresh 18-gauge needle was filled with Dulbecco’s Modified Eagle’s Medium-Low Glucose (DMEM-LG; Sigma Chemical, St. Louis, MO) supplemented with 10% fetal bovine serum (FBS) (Hyclone Laboratories, Logan UT) from a lot previously tested and found to support the preferential attachment, proliferation, and differentiation of rMSCs (Lennon et al., 1996). The needle was inserted into the opening previously prepared, and a small volume of medium was ejected to expel the bone marrow from the medullary canal. Marrow samples were collected and mechanically disrupted by passing them successively through 16-, 18-,
and 20-gauge needles. Disaggregated marrow was centrifuged at 500g in a Beckman TJ-6 centrifuge and resuspended in serum-supplemented medium; an aliquot of the cell suspension was combined with an equal volume of 4% (v/v) acetic acid to lyse red blood cells, and nucleated cell numbers were determined with a hemacytometer. Cells were seeded at 5 x 10^3/100-mm tissue culture dish.

Equal numbers of culture dishes were placed inside two air-tight modular incubator chambers (Billups-Rothenburg, Del Mar, CA). Humidity was supplemented by placing an open tissue culture dish containing 10–20 ml of sterile distilled water in the bottom of each chamber. Gas mixtures, consisting of 5% oxygen, 5% CO₂, and 90% nitrogen (referred to as low or 5% oxygen) for one chamber and 5% CO₂ and 95% air for the other (referred to as control or 20% oxygen), were allowed to flow into the chambers through one of the two ports, thus flushing ambient air from the chamber. The airflow was stopped after 15 min and both ports were immediately closed. The chambers were then placed inside a standard tissue culture incubator.

In order to ensure that pressure inside the chamber was comparable to that inside the incubator, gas was periodically vented from the chamber until the temperature inside the chamber equaled that inside the main incubator. The modular chambers were flushed with fresh gas only, and excess gas was vented in the same manner. Cells were continued in control or low oxygen conditions for the duration of their time in primary culture. However, at first passage, half of the cultures generated for each of the two oxygen conditions were continued in the same atmosphere, while the other half were switched to the alternate condition. Culture medium was removed and replaced with fresh serum-supplemented medium after 3 days and every 3–4 days thereafter. In one experiment, the pH of medium collected from culture dishes at each medium change was determined with a pH meter (Accumet model 10, Fisher Scientific, Pittsburgh, PA).

**Cell subculture**

When colonies of rMSCs increased in size, but prior to the time they became multilayered and the cells came in contact with one another (usually on day 12 of primary culture), the cells were subcultured by treatment with 0.25% trypsin in 1 mM EDTA (Life Technologies, Grand Island, NY) for 5 min at 37°C. Trypsinization was arrested with the addition of ½ volume of calf serum (Hyclone Laboratories), and the resulting cell suspension was centrifuged at 500g for 5 min, resuspended in serum-free DMEM-LG, and counted with a hemacytometer. A portion of the cell suspension was reserved for an in vivo osteogenic assay, while the remainder was resuspended in serum-containing medium and seeded into 35-mm culture dishes at a density of 10^5 cells per dish for use in an assay for in vitro osteogenesis; both assays are described in detail below. A flow chart outlining the experimental design is illustrated in Figure 1.

**In vitro osteogenesis**

In order to assess the effect of oxygen concentration on in vitro osteogenesis, cultures of first passage rMSCs maintained in primary culture in 5% or 20% oxygen were each divided into two equal groups immediately after being subcultured (day 0). One group was continued in the same oxygen condition in which it was maintained in the previous passage, while the other was switched to the alternate oxygen condition. Thus, four groups of cultures were generated: those which were in either control or low oxygen throughout their time in culture (referred to as C-C and L-L respectively) and those switched from control to low (C-L) or from low to control (L-C) oxygen after being seeded into 35-mm dishes.

Medium was changed for all cultures on day 1 of first passage. Serum-supplemented DMEM-LG, further augmented with osteogenic supplements (OS), consisting of 10 nM dexamethasone (Sigma Chemical) and 80 μM ascorbic acid-2-phosphate (Wako Chemicals, Richmond, VA), was added to one half of the cultures in each of the four groups, while control medium (DMEM-LG supplemented with 10% FBS and 80 μM ascorbic acid-2-phosphate) was added to the remainder. On day 10, all culture media were further augmented with 10 mM β-glycerophosphate (BGP; Sigma Chemical). Cells were continued in control or OS medium for the remainder of their time in vitro. The goal of this part of the study was to determine whether low oxygen concentration in primary culture affects the commitment of cells to the osteogenic lineage, or whether the observed effects are on osteogenic expression and are limited primarily to the time of exposure to the respective oxygen tensions. The frequency of medium changes and of gas replenishment was the same as for primary culture. In vitro osteogenesis was evaluated quantitatively through assays for calcium content and alkaline phosphatase activity as described below.

**Alkaline phosphatase assay**

Alkaline phosphatase activity was determined for first passage rMSCs in control or OS medium on days 3, 6, 9, 12, and 16 of culture. Triplicate cultures from each experimental group were rinsed twice with Tyrode's balanced salt solution (Sigma Chemical), and 1 ml of a 1 mg/ml solution of alkaline phosphatase substrate (p-nitrophenyl phosphate; Sigma Chemical) in a buffer consisting of 50 mM glycine and 1 mM MgCl₂·6H₂O (pH 10.5) was added per 35-mm dish. After 10 min the solution was removed and transferred to a tube containing an equal volume of 1 M NaOH. Appropriately diluted samples of the resulting solutions were transferred to a 96-well culture dish, and the absorbance was read at 405 nm on a model 3550 microplate reader (Bio-Rad Laboratories, Hercules, CA). A standard curve generated from a series of dilutions of p-nitrophenol (Sigma Chemical) was used to determine the concentration of the enzyme reaction product. After the alkaline phosphatase assay solution was removed, the cultures were rinsed twice with Tyrode's salt solution and fixed with 100% ethanol. After 15 min, the ethanol was removed and the plates were dried and stored at 4°C until used for quantification of DNA.

**Calcium**

On day 21, triplicate rMSC cultures were rinsed twice with Tyrode's salt solution and fixed with 1% (v/v)
glutaraldehyde in Tyrode’s for 30 min. Following fixation, cultures were rinsed twice with distilled water, allowed to dry, and stored at 4°C. Calcium was extracted with 1 ml of 0.6 M HCl per dish. Dishes containing this solution were placed on a rotary shaker and maintained at 50 rpm overnight. Aliquots of the extract, diluted as needed, were added to wells of a 96-well culture dish (4 wells/sample); reagents from a commercial calcium assay kit (Sigma Chemical) were added to the wells, and the absorbance was read at 575 nm with a model 2550 microplate reader (Bio-Rad Laboratories). Calcium concentration was determined with a standard curve generated from a series of dilutions of CaCl₂.

DNA

DNA content was quantified by a modification (Hanada et al., 1997) of a technique described by Gillery et al. (1993). Triplicate cultures were rinsed twice with Tyrode’s salt solution and fixed with 100% ethanol for 15 min, either after quantification of alkaline phosphatase activity (day 3 through 16) or independent of other assays (day 21). Dehydrated cultures were stored at 4°C until all were collected. All samples from each preparation were analyzed for DNA content at the same time. One ml of 40% (w/v) 3,5-diaminobenzoic acid (DABA, Sigma Chemical), twice clarified by adsorption to granular charcoal and then filtered through a 0.22-μm Nalgene filter unit (Nalgene Corp.), was added per 35-mm tissue culture dish. The dishes were incubated at 60°C for 45 min and the DABA-DNA reaction was stabilized by the addition of 2.5 ml of 3N HCl. The optical density of the resulting solution was determined with a spectrophotofluorometer at an absorbance of 420 nm and an emission of 490 nm. A standard curve generated from a series of dilutions of calf thymus DNA was used to determine the DNA concentration of the samples. Differences in DNA content are assumed to reflect differences in cell number.

von Kossa staining

The presence of mineralized deposits in first passage rMSC cultures was demonstrated with von Kossa staining.
staining. On day 21, triplicate cultures of each experimental group were rinsed twice with Tyrode’s salt solution, fixed with 1% glutaraldehyde (v/v) for 15 min, and rinsed three times with distilled water. One ml of 2% (w/v) silver nitrate (Sigma Chemical) was added per dish, and the cultures were placed in a dark environment for 10 min. Cultures were then rinsed three times with distilled water and exposed to bright light (while covered with water) for 15 min. Culture dishes were rinsed again with water and then dehydrated with 100% ethanol.

Colony counts

For some rMSC preparations, colony numbers were determined between days 7 and 12 of primary culture (when colonies were large but not overlapping). Cultures maintained in either control or low oxygen were rinsed twice with Tyrode’s salt solution, fixed for 15 min with 1% (v/v) glutaraldehyde, rinsed twice with distilled water, and air dried. Fixed cultures were stained with 4 ml of 0.1% (w/v) crystal violet in distilled water for 30 min, rinsed three times with distilled water, and air dried. Inverted culture dishes were examined on a dissecting microscope at low magnification (7X), and the number of colonies (of more than 50 cells) was determined. All cultures were evaluated by the same individual, who was unaware of the identity of the cultures.

In vivo osteogenic assay

Implantation of MSC-loaded ceramic cubes into syngeneic or immunocompromised host animals as an in vivo assay for the osteochondrogenic potential of these cells has been described previously (Dennis et al., 1992, Dennis and Caplan, 1993). Briefly, blocks of porous ceramic (mean pore size of 200 μm), consisting of 60% tricalcium phosphate and 40% hydroxyapatite, were cut into cubes measuring 3 mm per side. The ceramic cubes were washed with water to remove ceramic dust, dried under a heating lamp, and sterilized in an autoclave. The cubes were placed in a 30-ml syringe with 1% (v/v) glutaraldehyde and air was withdrawn with a 20-ml syringe as for fibronectin coating. Cell-loaded cubes were incubated at 37°C for 2 h and then implanted subcutaneously into pockets created by blunt dissection on the dorsal surface of syngeneic rats anesthetized with sodium pentobarbital (90 mg/kg; Veterinary Laboratories, Inc., Lenexa, KS). Host animals were sacrificed after 3 or 6 weeks by CO₂ asphyxiation. Ceramic cubes were removed, fixed with 10% buffered formalin phosphate (Fisher Scientific), decalcified with RDO (Apex Engineering, Plainfield, IL), embedded in Tissue Prep 2 (Fisher Scientific), and cut into sections 5 μm in thickness. Each 10th and 11th section of a cube was collected and placed onto alternate glass slides; paired slides were stained with either toluidine blue or Mallory-Heidenhain. Stained sections were examined for the presence of bone or cartilage by brightfield microscopy with an Olympus IMT-2 inverted microscope. The total number of pores for each ceramic cube and the number of those pores which contained bone or cartilage were determined for each section, and the percentage of bone- or cartilage-positive pores was calculated. Histologic sections were evaluated on a double blind basis, and all were graded by the same individual (DPL).

Statistical analysis

Statistical analysis was conducted with Sigma Stat software (SPSS Inc., Chicago, IL). Student’s paired t-test was used to analyze cell yields at the end of primary culture and ceramic cube data from the in vivo osteogenic assay. Colony counts for cells in low and control oxygen were compared with Student’s t-test. DNA, alkaline phosphatase, and calcium data for first passage cultures were evaluated with one way analysis of variance (ANOVA) after the data passed normality and equal variance tests. All pairwise multiple comparisons were conducted with the Student-Newman-Keuls method. Comparisons between cells in control and OS medium for these assays were done with the t-test.

RESULTS

Morphology

Rat marrow-derived MSCs maintained in either 20% or 5% oxygen in primary culture were essentially identical as viewed by phase contrast microscopy. Small colonies of fibroblast-like cells were seen after several days of primary culture (Fig. 2A). These colonies increased in size and were subcultured after 12–14 days. Colonies appeared to be larger and were significantly more numerous in cultures maintained in low oxygen (Fig. 3). Individual cells, however, appeared to be identical in either oxygen condition in these primary colonies. The pH of culture medium collected when medium was changed (days 3, 7, 11, and 13) was statistically identical for rMSC cultures maintained in the two oxygen environments (data not shown). Individual subcultured cells in control medium were also similar to one another in morphology regardless of the oxygen condition (not shown). First passage cells were more uniformly distributed on the culture substrate and were somewhat more spread than in primary culture.

When cultured in OS medium in first passage, rMSCs assumed a less elongated, polygonal appearance. By approximately day 7, small foci of small round cells, presumed to be osteoblasts, could be seen at multiple sites among confluent fibroblast-like cells throughout the cultures (Fig. 2B). These foci, referred to as nodules, increased in size and number through time in culture, becoming multilayered (Fig. 2C) and then mineralizing as confirmed by brown to black von Kossa staining.
Internodular cells maintained their polygonal morphology, became multilayered, and were negative for von Kossa staining; these morphologies are as previously reported by us and others (Herbertson and Aubin, 1995; Cassiede et al., 1996; Hanada et al., 1997). First passage cells in control medium (which contained ascorbic acid-2-phosphate and BGP but not dexamethasone) were more elongated than the non-nodular cells in OS medium. The cells in control medium became multilayered, but did not form nodules and were always von Kossa-negative.

**Cell proliferation**

For each of five separate rMSC preparations, the number of cells harvested at first passage, as determined by hemacytometer count, was higher for cultures maintained in low oxygen than for those in control oxygen (Table 1). When these numbers were analyzed with Student's paired *t*-test, the difference was found to be significant (*P* < 0.01). The effect of oxygen concentration was also evaluated by determining the percent change in cell yield for each preparation (Table 1). Thus, a positive change (increase) in cell number for rMSCs cultured in low oxygen was calculated. When the data from all preparations were considered, the mean increase in cell number for rMSCs cultured in low oxygen was determined to be 41.2% (±18.8%).

**Table 1. Effect of oxygen concentration on rMSC yield at first passage**

<table>
<thead>
<tr>
<th>Preparation</th>
<th>Control oxygen cell yield (×10⁶)</th>
<th>Low oxygen cell yield (×10⁶)*</th>
<th>% Increase in cell yield for rMSCs in low oxygen**</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1.24</td>
<td>1.93</td>
<td>55.7</td>
</tr>
<tr>
<td>2</td>
<td>2.18</td>
<td>3.17</td>
<td>45.4</td>
</tr>
<tr>
<td>3</td>
<td>1.59</td>
<td>2.51</td>
<td>57.9</td>
</tr>
<tr>
<td>4</td>
<td>1.46</td>
<td>1.97</td>
<td>34.9</td>
</tr>
<tr>
<td>5</td>
<td>2.01</td>
<td>2.25</td>
<td>11.9</td>
</tr>
</tbody>
</table>

Cell yield per 100-mm culture dish at first passage is illustrated for rMSCs cultured in control (20%) and low (5%) oxygen. Cell numbers were determined by hemacytometer count.

*Significantly higher (*P* < 0.01) than cell yield for rMSCs maintained in control oxygen when data are analyzed with the paired *t*-test.

**Data indicate the difference in yield for rMSCs maintained in the two conditions according to the following formula (after Pennathur-Das and Levitt, 1987)

\[
\text{Cells/platein5%/oxygen - Cells/platein20%/oxygen} \times 100
\]

Fig. 3. The number of rMSC colonies on day 7 of primary culture in control oxygen (solid column) or low oxygen (open column) is represented on the Y axis. Values indicate the mean and standard deviation for nine cultures for each condition; all cultures were from the same cell preparation. *Significantly higher (*P* < 0.0001).
in control or OS medium in the four oxygen conditions (Figs. 4A, B). The data depicted in Figure 4 are derived from a single preparation of rMSCs, and are representative of the data from five separate rMSC preparations. As illustrated in Figure 4A, the amount of DNA per culture for cells in control medium increased from day 3 to 21 except for a slight decline at day 12. On most days, DNA values were higher for cells which had been in low oxygen in primary culture (L-L and L-C) than for those in control oxygen in primary culture (C-C and C-L). On
days 6 and 21, DNA values for those in low oxygen in both passages were significantly higher ($P < .05$) than those for any of the other groups.

Rat MSCs in osteoinductive (OS) medium had lower quantities of DNA than their sister cultures in control medium except on day 21 (Fig. 4B). As in the case of rMSCs in control medium, DNA for cells in OS medium increased through time in culture. There was, however, no decline between day 9 and 12, and the increase in DNA after day 12 was greater than for control medium. Cells in L-L and L-C conditions in OS medium were not significantly different from one another except on day 3, but both had higher DNA values than for those in C-C and C-L conditions on all assay days except day 6; in most cases the differences were significant. C-L cultures in OS medium had higher DNA values than those in C-C conditions.

Thus, oxygen tension affected cell proliferation both in primary culture and in first passage. Cell numbers were higher at the end of primary culture for cells in low oxygen than for those in control oxygen. Rat MSCs maintained in low oxygen in first passage (L-L) had higher DNA values than those continued in control oxygen (C-C). When data from all preparations are considered (not all data shown), the number of cells was slightly lower, on most days on which they were evaluated, for rMSCs in L-C conditions than for those in L-L conditions. Conversely, cell numbers were somewhat higher for C-L cultures than for those in C-C conditions. That is, switching cultures from low to control oxygen at first passage resulted in lower cell numbers, while switching from control to low oxygen resulted in higher cell numbers.

**Alkaline phosphatase activity**

While the data depicted in Figure 4C are representative of the effect of oxygen concentration on alkaline phosphatase (AP) activity (normalized per µg of DNA) for rMSCs, there was a greater degree of variability from one preparation to another than in the case of DNA. In most instances, as shown in Figure 4C, AP activity was higher in cells maintained in control medium than in OS medium, through day 6, regardless of the oxygen environment. By day 9, however, the levels of AP activity in rMSCs in OS medium usually exceeded those for cells in control medium. AP activity for rMSCs in control medium rose slightly from day 9 to 12, and then declined by day 16, except for cells in the L-L and L-C groups. On the other hand, AP activity of rMSCs in OS medium increased approximately two-fold from day 9 to 12 and then doubled again from day 12 to 16.

With respect to first passage rMSCs in OS medium, L-L and L-C cultures were approximately equivalent in AP activity on day 9 and thereafter, and C-C and C-L cultures were comparable to one another in the same interval. AP activity was significantly higher ($P = 0.05$) for the former groups. Although L-C cultures had higher AP/DNA values than L-L cultures, the differences, except for day 12, were not significant.

**Calcium content**

For each of five separate preparations of rMSCs, day 21 calcium content, normalized per µg of DNA, was greater for cultures in OS medium than for those in control medium, regardless of the oxygen concentration in which the cells were maintained. There was a clear correlation between calcium content and oxygen concentration for rMSCs maintained in OS medium. Calcium content per µg of DNA on day 21 was highest in cells in L-L conditions and lowest in cultures in C-C conditions (Fig. 4D). Moreover, the influence of oxygen tension was also evident in the calcium values of cells switched from one oxygen condition to the other at first passage. Cultures transferred from low to control oxygen had lower calcium values than those continued in low oxygen, while those transferred from control to low oxygen had higher calcium content than those maintained continuously in control oxygen.

**von Kossa staining**

Staining of rMSCs by the von Kossa procedure visually demonstrated that oxygen tension affected the extent of mineralized bone nodule formation in first passage cultures in OS medium in the same manner that it influenced calcium content. The number of mineralized nodules was much greater in cultures in L-L and C-L conditions than for those in C-C conditions (Fig. 5).

**In vivo osteogenic assay**

Histologic evaluation of porous ceramic cubes harvested from syngeneic host animals 3 and 6 weeks after implantation revealed that greater amounts of bone and cartilage are present in cubes loaded with rMSCs from primary cultures cultivated in low oxygen than in those containing cells cultured in control oxygen in primary culture. The results for all ceramic cubes harvested at 6
weeks are presented in Table 2. For each of the ten individual host animals (two rats for each of five separate cell preparations), the mean score (expressed as the percentage of pores which contained bone or cartilage) for cubes loaded with cells cultured in 5% oxygen was higher than that for cubes containing cells maintained in 20% oxygen. In most instances, four cubes were implanted per animal for each of the two oxygen conditions.

There was, however, no qualitative histologic difference between the two sets of cubes. In either case, osseous material was initially deposited along the inner walls of bone-containing pores by cuboidal cells presumed to be osteoblasts (Fig. 6A). Some osteoblasts became enclosed within the bony matrix (becoming osteocytes in the process) as deposition of bone progressed uniformly toward the center of the pore. Varying amounts of fibrous tissue and vascular elements were also located within bone-containing pores. In some cases, entire pores were filled with cartilage, identified by large round to cubodial cells surrounded by matrix which stained metachromatically with toluidine blue (Fig. 6B). Cartilage-containing pores represented a small portion of pores identified as bone- or cartilage-positive. Pores which contained neither bone nor cartilage were filled with vascularized fibrous tissue.

**DISCUSSION**

In this study, rat marrow-derived MSCs were cultivated in standard tissue culture conditions and in an atmosphere of reduced oxygen in order to determine whether the two conditions differ in their effect on cell attachment and proliferation and on osteochondrogenesis. The results suggest, first, that attachment and/or survival of rMSCs in primary culture is enhanced in a reduced oxygen environment, as indicated by cell colonies which were significantly higher in number than for those maintained in control oxygen. Secondly, a greater number of cells per dish were harvested at first passage from cultures maintained in 5% oxygen, suggesting that the rate of cell proliferation in primary culture was higher in low oxygen. While it is possible that the increase in cell number resulted solely from the greater number of colonies, examination of the cultures by phase contrast microscopy suggests that colonies in cultures maintained in low oxygen are, on average, larger in size and not just greater in number than those in control oxygen, although the difference in colony size was not quantified.

**TABLE 2. Effect of oxygen concentration in primary culture on bone formation in implanted rMSC-loaded porous ceramic cubes**

<table>
<thead>
<tr>
<th>Cell preparation (1–5); host animal (A or B)</th>
<th>Cells cultured in control oxygen</th>
<th>Cells cultured in low oxygen*</th>
</tr>
</thead>
<tbody>
<tr>
<td>1A</td>
<td>9.6</td>
<td>35.8</td>
</tr>
<tr>
<td>1B</td>
<td>27.0</td>
<td>43.8</td>
</tr>
<tr>
<td>2A</td>
<td>51.2</td>
<td>61.4</td>
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</tr>
<tr>
<td>4A</td>
<td>36.4</td>
<td>70.9</td>
</tr>
<tr>
<td>4B</td>
<td>38.3</td>
<td>75.4</td>
</tr>
<tr>
<td>5A</td>
<td>64.1</td>
<td>81.1</td>
</tr>
<tr>
<td>5B</td>
<td>44.3</td>
<td>69.1</td>
</tr>
</tbody>
</table>

*Rat MSCs from five separate preparations (indicated by numbers 1–5) cultured in control or low oxygen were loaded into porous ceramic cubes which were implanted into two syngeneic host animals (designated by the letter A or B) per preparation. Cubes were harvested after six weeks and evaluated for the presence of bone and cartilage. The scores illustrate the number of bone- or cartilage-containing pores as a percentage of the total number of pores and are expressed as the mean for four cubes per oxygen condition per host animal. *Significantly higher (P < 0.0003) than scores for cells in control oxygen when the data are analyzed with the paired t-test.
Moreover, elevated rates of proliferation, independent of any influence of colony number, continued in first passage for rMSCs which had been in 5% oxygen in primary culture and first passage, as indicated by higher DNA values for these cells in both control and OS medium. Changing oxygen concentration at first passage almost always resulted in less of an increase in DNA, over time, for cells switched from low to control oxygen, and in a greater increase in DNA for cells transferred from control to low oxygen (compared, in each case, with those for which oxygen tension was not changed).

Evidence for the notion that in vitro osteogenesis is enhanced in an atmosphere of reduced oxygen tension is found in data regarding calcium content and von Kossa staining of bone nodules. For each of the five preparations of rMSCs tested, calcium content was elevated in all cultures which were maintained in low oxygen during at least one passage in vitro, relative to those in control oxygen throughout their time in culture. Moreover, von Kossa staining of selected rMSC cultures from each cell preparation revealed that more nodules are present in those maintained continuously in low oxygen than in control oxygen. Bellows and Aubin (1989) have demonstrated that individual nodules are derived from single osteoprogenitors in cultures of fetal rat calvarial cells. Assuming that this observation also applies to rMSCs, it is conceivable that they may be affected by the low oxygen tension in primary culture and first passage, as indicated by higher calcium values were higher than for rMSCs in the same medium conditions but in control oxygen in both passages. It is important to note that calcium values were low in all oxygen conditions in the absence of osteogenic supplements in the culture medium. Thus, low oxygen alone may predispose rMSCs to commit to the osteogenic lineage, but, in the absence of the appropriate stimuli, elevated expression of osteogenic markers is not observed.

On the other hand, elevated calcium and alkaline phosphatase levels and increased von Kossa-positive staining were also observed in rMSCs maintained in control oxygen in primary culture but switched to low oxygen and OS medium at first passage. Thus, even in the absence of putative pre-induction by reduced oxygen tension in primary culture, the osteogenic expression of rMSCs is enhanced in the presence of low oxygen in first passage. It may be that low oxygen favors both commitment to osteogenic lines and differentiation along those lines, but also favors osteogenic differentiation even in the absence of prior enhanced commitment.

Other investigators have examined the effect of reduced oxygen tension on osteoblasts (Tuncay et al., 1994; Reilly et al., 1998), pericytes (Brighton et al., 1992; Reilly et al., 1998), and periosteal cells (Deren et al., 1990). Like rMSCs, periosteal cells and pericytes are thought to have the potential to differentiate into osteoblasts given the proper environmental cues. It is interesting to note, therefore, that the rates of proliferation and the alkaline phosphatase activity of these cells are elevated in a low oxygen environment. On the other hand, alkaline phosphatase activity of rat calvarial osteoblasts was higher in 90% oxygen than in either 10 or 21% oxygen (Tuncay et al., 1994). Additionally, levels of osteocalcin, thought to be a late marker of osteoblastic expression, were higher in cultures of pericytes at 7 weeks of culture in 21 and 60% oxygen than in lower oxygen tensions (Brighton et al., 1992; Reilly et al., 1998). While osteocalcin levels were not examined in the present study, calcium content, bone nodule formation, and in vivo osteogenesis, other late markers of osteogenic expression, were found to be elevated in reduced oxygen in rMSCs. These parameters, however, were not examined as late as seven weeks of culture, and it is difficult to make comparisons among these various studies because of differences in cell type, species, medium, serum, and the timing of the assays.

The low oxygen mixture employed in this study differs from the atmosphere customarily used in cell culture not only in oxygen concentration, but also in elevated nitrogen tension and in the absence of argon, neon, and other gases present in the atmosphere in trace quantities. While it is possible, though we believe unlikely, that these non-oxygen related differences affected the experimental results, to our knowledge, no data are available regarding the impact of these atmospheric constituents on cell culture. While human MSCs were not examined under the conditions described in this paper, it is conceivable that they may be affected by lower oxygen tension in the same manner as rMSCs. If that were the case, cultivation of human MSCs in low oxygen could play an important role in possible clinical application of these cells.

Finally, a review of the literature reveals a broad range of cell types (beyond the brief list presented in the Introduction) for which cultivation in physiologic oxygen has been found to be superior to that in conditions normally found in vitro. Perhaps the oxygen concentration to be employed in the culture of a given cell type should be accorded as much consideration as is given to the choice of culture medium or serum. It seems fair to say that, while the use of 95% air (20% oxygen) is sufficient to observe osteogenic events in vitro, this oxygen level is neither optimal nor physiologic.

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LITERATURE CITED


