Effects of Low Oxygen Culture on Pluripotent Stem Cell Differentiation and Teratoma Formation

by

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Bachelor of Science in Chemical Engineering
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Pluripotent stem cells (PSC) hold promise for the study of embryonic development and the treatment of many diseases. Most pluripotent cell research is performed in incubators with a gas-phase oxygen partial pressure (pO₂) of 142 mmHg. However, embryonic cells in early development are exposed to a local pO₂ of 0-30 mmHg, and the effects of such conditions on differentiating PSC are poorly understood. Residual PSC within differentiated populations are problematic because of their potential to form tumors *in vivo*. This is a major safety issue that must be overcome before PSC-based therapies can be used in the clinic. In this study, we differentiated mouse and human embryonic stem cells and mouse induced pluripotent stem cells at different defined pO₂ on highly oxygen-permeable silicone rubber culture dishes and assessed differentiation to the three germ layers, endoderm, ectoderm, and mesoderm and to cardiomyocytes and assessed residual PSC within differentiated populations.

Low pO₂ drastically affects differentiation of PSC to the three germ layers and cardiomyocytes. Overall, differentiation was higher to endoderm, lower to ectoderm, and higher or the same to mesoderm. Differentiation to cardiomyocytes was greatly enhanced without the need for purification, possibly by lineage selection via increased Mesp1 and Mesp2 expression. Understanding the effects of pO₂ during differentiation is an important step towards the development of protocols for regenerative medicine.

Control of pO₂ to physiological levels typical of the developing embryo reduced the fraction of PSC within, and the tumorigenic potential of, differentiated populations. Culture under differentiating conditions at low pO₂ reduced measured pluripotency markers by up to four orders of magnitude. Upon implantation into immunocompromised mice, low pO₂-differentiated PSC either did not form tumors or formed tumors at a slower rate than high pO₂ PSC. Low pO₂ differentiation could be combined with cell sorting for improved benefits. Low pO₂ culture alone or in combination with other methods is a potentially straightforward method that could be applied to future cell therapy protocols to minimize the possibility of tumor formation.

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Chapter 1. Introduction

1.1. Pluripotent stem cells

Pluripotent stem cells, such as embryonic stem cells (ESC) and induced pluripotent stem cells (iPSC), have immense clinical potential due to their ability to form all somatic cells found in the body [1-3]. These cells can potentially make major advances in human health, for example, by use in drug screening, in modeling of human disease in vitro, and as an unlimited supple of replacement tissue to treat heart disease, diabetes, and Parkinson’s disease [3-6].

ESC are isolated from the inner cell mass of the developing blastocyst [7, 8] and are characterized by their ability to maintain their pluripotent state and differentiate. For mouse ESC (mESC), maintaining their pluripotent state is typically accomplished by the soluble factor leukemia inhibitory factor (LIF) and, for human ESC (hESC), is typically accomplished by the soluble factor basic fibroblast growth factor (bFGF) and coculture with mouse embryonic fibroblasts (MEFs) [7]. The molecular circuitry that regulates the pluripotent state of ESC is complex, involving transcription factors, chromatin modification, miRNA, and signaling pathways. The transcription factors Oct4, Nanog, and Sox2 are considered to be the master regulators of this network and lie within its core. ESC have been differentiated in vitro into a wide range of cell types, such as
cardiomyocytes [8], β-cells [9], and neurons [10] and then implanted into rodent models of cardiac infarct, diabetes, and Parkinson's disease to improve function.

iPSC are a promising alternative source of pluripotent stem cells that are derived from somatic cells, such as fibroblasts, via reprogramming by inserting, often by virus vectors, factors such as Oct4, Sox2, c-Myc, and Klf4 [11] or Oct4, Sox2, Nanog, and Lin28 [12]. iPSC have properties similar to those of ESC [13]. One major concern is the safety of these cells. c-Myc is a proto-oncogene that presents a tumorigenic risk. Mouse iPSCs (miPSC), derived with modified protocols without c-Myc, have reduced occurrence of tumor formation [14]. Viral insertions also possess a risk of tumor formation. Methods for generating iPSCs without viruses [15, 16], using small molecules in combination with Oct4 and Klf4 gene insertion [17], and without residual programming factors [18] have been explored. More recently, reprogramming has been achieved in mouse cells using only recombinant proteins [19].
1.2. The effects of low oxygen on self-renewal and differentiation of PSC

Human pluripotent cells are a renewable source of replacement tissue that have the potential to treat many diseases, including heart disease, type I diabetes, and Parkinson's disease. However, many hurdles remain before this new era of regenerative medicine can be realized. All cells of the developing embryo are exposed to oxygen levels in vivo far below that of atmospheric levels, and they differentiate and undergo organogenesis in a low oxygen environment [20-26]. The local oxygen environment is potentially an important variable for manipulation by researchers hoping to recapitulate specific differentiation pathways in vitro to produce replacement tissues from pluripotent stem cells for regenerative medicine. Here, we review recent reports investigating the effects of low oxygen on the undifferentiated phenotype and on the differentiation process, and we address the issue of control of pO$_2$cell during culture.

1.2.1. Undifferentiated phenotype

The ability to propagate pluripotent stem cells in their undifferentiated state is essential to produce sufficient cells for future applications, and the effects of oxygen levels much lower than that in a standard incubator on undifferentiated cells have been examined in a number of studies (Table 1-1). Throughout this review, we will refer to oxygen levels either as a percentage or a partial pressure in mmHg, as it was reported in the original papers, with standard incubator oxygen referred to as 20%, even though it is actually often 18.7% (142 mmHg).
1.2.1.1. Derivation of new embryonic stem cell lines

Derivation of new ES cell lines, both mouse [27, 28] and human [29], benefit from being performed in low oxygen levels. Studies examining the establishment of new ES cell lines from blastocysts under 5% oxygen have had greater success than under 20% more commonly used. The benefits of low oxygen conditions include an increase in the number of individual colonies exhibiting alkaline phosphatase activity [28], an increase in the number of proliferating cells [27], and removal of the need for undefined animal-derived products such as serum or matrices in the process [29], the last of which can be crucial in the applicability of these derived cells to clinical applications.

1.2.1.2. Pluripotency

Low oxygen affects the pluripotent phenotype of mESCs when they are cultured with leukemia inhibitory factor (LIF), a cytokine commonly used to prevent differentiation [30]. Normally, mESC colonies have smooth edges, but this morphological phenotype is lost with low oxygen exposure, instead causing the colonies to have fibroblast-like morphologies [31] or to form jagged edges [32], both signs of early differentiation despite the presence of LIF. These morphological changes are also associated with reduced levels of alkaline phosphatase activity and pluripotency gene (Oct4, Nanog, and Sox2) expression [31, 32]. One study [31] proposed that HIF-1α protein, the concentration of which
increases at low oxygen, mediates suppression of the LIF-STAT3 pathway as
evidenced by a reduction in the expression of the LIF receptor under hypoxia.
Interestingly, in one report [32], despite these observations, the cells maintained
some potential to differentiate into cells of all three germ layers.

In contrast to mESCs, expression of Oct4, Nanog, and Sox2 in hESCs are
apparently not affected by hypoxic culture (2% to 5% oxygen) [33, 34]. In one
study, however, Nanog and Notch1 mRNA expression is increased in hESCs
cultured over a four-week period at 5% oxygen compared to those cultured under
20% oxygen. This difference is not maintained after long term culture under the
hypoxic condition, since there is no significant difference in Nanog and Notch1
mRNA levels between cells cultured for 18 months under either 5% or 20%
oxygen. Consistent with earlier results, no differences in Oct4 expression are
observed under both oxygen conditions with either short or long term culture [35].

Despite not affecting gene expression of Oct4, Nanog, and Sox2 directly, hypoxic
culture results in wide-spread transcriptome differences in hESCs when
compared to culture under ambient oxygen conditions, including downstream
targets of unaffected genes Oct4, Nanog, Sox2 [34]. Amongst other affected
genes, Lefty2, a member of the TGFβ family of proteins, which is hypothesized to
play a role in preventing differentiation, is down-regulated at 20% oxygen
compared to culture at 4%. This may lead to a greater proclivity for the hESC to
differentiate. In studies on the transcriptome-wide effects of hypoxic culture on
hESCs kept under self-renewing conditions, culture at ambient pO$_2$ gas results in greater heterogeneity amongst different hESC lines than when cultured under hypoxic (2-4% oxygen) conditions [33]. This heterogeneity may arise from subpopulations of cells already undergoing early differentiation when cultured at 20% oxygen.

While low oxygen appears not to affect pluripotency gene expression directly, differentiated areas of each colony expressing SSEA-1, a marker of early differentiation for hESCs, are reduced when cultured under 3% and 1% oxygen compared to normoxic culture [36]. In a different study, however, culture at 2% oxygen did not affect expression of either SSEA-4 or SSEA-1 in hESCs when compared to ambient oxygen conditions [37].

HIF-1α has been investigated as an important gene mediating hESC hypoxic response. In one report, co-culture of hESCs with transgenic human fetal liver stromal cells ectopically expressing HIF-1α protein prevents hESC differentiation and even increases Oct4 and Nanog expression [38]. Another study [39] suggests a possible link connecting HIF-1α action with Notch signaling. In mouse embryonic teratocarcinoma cells and neural stem cells, hypoxia allows for the accumulation of HIF-1α, which stabilizes the Notch intracellular domain, in turn promoting the undifferentiated phenotype. The action of Notch in promoting self-renewal appears to be conserved in hESCs in view of a recent demonstration
that self-renewal is mediated by activation of the Notch pathway and that benefits of hypoxic culture are eliminated with inhibition of Notch signaling [35].

1.2.1.3. Proliferation

Many studies have found that low oxygen affects proliferation, but this effect varies between studies and cell lines. In one study [28], proliferation of mESCs increases under 5% compared to 20% oxygen, while in another [32], no difference is observed in one cell line and a small increase in another. Culture at 1% oxygen or lower, however, reduces the rate of proliferation compared to 20% [31, 32]. mESCs are able to adapt and continue to proliferate under anoxia, albeit with a longer doubling time compared to 20% oxygen, and have the ability to utilize anaerobic respiration exclusively for their energy requirements [32]. Similarly disparate results have been reported for hESCs: In one study, proliferation rates at 5% oxygen are reduced compared to 20% [35], in another, no differences in the rates are observed at 5% and 3% and only a slight reduction at 1% [36].

1.2.1.4. Clonal recovery

Low oxygen culture improves clonal recovery after passaging. mESCs at 5% oxygen [40] and hESCs at 2% [37, 41] have improved clonal survival. Cytogenetic instability at high passages for mESCs is not reduced in culture at
5% oxygen [27], and further reductions in oxygen to even 0% do not affect the amount of oxidative DNA damage incurred by the cells [32]. In contrast for hESCs, 2% oxygen reduces chromosomal abnormalities after multiple passages [37], suggesting a further benefit of hypoxic culture.

1.2.2. Differentiation

Differentiation is a major hurdle for the successful translation of stem cell research to clinical applications. Given that low partial pressures of oxygen (pO₂) is an important physiological cue during development [42], it is not surprising that low pO₂ influences differentiation to many cell types (Table 1-2).

1.2.2.1. Neurons

Low oxygen has different effects on specific stages of differentiation of pluripotent cells to dopaminergic neurons. While low oxygen reduces differentiation from mESCs to neural stem cells (NSCs) compared to 20% oxygen [43, 44], it increases subsequent differentiation to neurons. mESCs differentiated to NSCs at standard incubator oxygen followed by differentiation at low (3.5%) oxygen have a higher number of dopaminergic neurons (tyrosine hydroxylase (TH)-expressing cells) compared to 20% oxygen. Differentiation with a neuronal differentiation-promoting cytokine mixture increases the number of dopaminergic neurons at 20% oxygen to a value equal to that at low oxygen.
without the cytokine mixture, but, interestingly, combining the cytokine mixture with low oxygen culture does not further increase neural differentiation [43]. Knocking down HIF-1α expression with antisense nucleotides decreases differentiation to TH+ cells, indicating that low-oxygen enhancement of the second differentiation step is HIF-1α-mediated, which is not surprising since HIF-1α is necessary for proper brain development [45].

Even though low oxygen decreases differentiation of pluripotent cells to NSCs, differentiation of mESCs entirely at low oxygen can increase the yields of neurons compared to 20% oxygen. In one study [44], the number of beta III tubulin+ and MAP2+ (both neuron markers) cells are increased by factors of 55 and 114, respectively, when differentiated entirely at 2% compared to 20% oxygen [44]. This is related, in part, to a greater total cell number by a factor of 34 with differentiation at 2% compared to 20% oxygen.

1.2.2.2. Cardiomyocytes

Low oxygen increases the number of cardiomyocytes from both hESCs and mESCs in bioreactors. HIF-1α is necessary for proper heart development [46], and differentiation with a HIF-1α knockout mESC line results in no observable spontaneously-beating cells [47], suggesting oxygen is an important environmental parameter. Using a strategy to selectively kill non-cardiomyocytes, the yield of cardiomyocytes from mESCs is increased by a factor of 1.5 with
differentiation at 4% compared to 20% oxygen in one report [48]. However, no difference between the two oxygen levels occurs before genetic selection. With hESCs, differentiation at 4% oxygen increases the total cell number by 30-47% compared to 20% oxygen [49]. However, the fraction of spontaneously-beating EBs is the same under both oxygen conditions, and while some cardiac markers are increased, others are unaffected by low oxygen. From these reports, it is unclear if the increased cardiomyocyte number is due to preferential differentiation to cardiomyocytes or simply to increased total number of cells. Recent unpublished data from the authors' laboratory indicates that both phenomena occur when mESCs are differentiated to cardiomyocytes at low oxygen.

1.2.2.3. Hematopoietic progenitor and endothelial cells

Low oxygen works in a complicated network to influence differentiation of pluripotent cells to blood and endothelial fates [50]. Vascular endothelial growth factor (VEGF), a growth factor secreted by cells at low oxygen, increases differentiation of mESCs to hemogenic mesoderm. These cells continue to differentiate to endothelial cells with activation of Flk-1, a VEGF receptor. Without VEGF, which is the situation if cells are cultured at 20% oxygen, Flk-1 is inactivated, resulting in the formation of hematopoietic progenitor cells. Even at low oxygen, VEGF can be inhibited by the presence of soluble Flt-1, a protein that cells express with increasing differentiation by binding VEGF and making
VEGF unavailable to activate Flk-1. VEGF secretion and inhibition results in an oxygen-dependent competitive balance between free and bound VEGF that influences differentiation of mesoderm to hematopoietic progenitor and endothelial cells. This example illustrates how understanding the mechanism by which low oxygen affects lineage selection is important for successful applications in regenerative medicine.

1.2.2.5. Chondrocytes

Chondrocytes normally reside in a low oxygen environment [51], and the effects of low oxygen on differentiation of pluripotent cells to this cell type has been investigated. Low oxygen increases collagen synthesis and biomechanical functionality of hESC-derived chondrocytes [52]. Differentiation at 2% oxygen increases collagen I and II content, tensile modulus, and compressive properties compared to differentiation at 20%. However, when these low oxygen-derived chondrocytes are used to form tissue using a scaffold-less strategy, the reverse is true: tissues formed at 20% oxygen have higher collagen content and better physical properties. Low oxygen can have different effects on different aspects (e.g. differentiation and tissue engineering) of a bioengineering strategy, and these differences must be understood to fully utilize low oxygen culture in regenerative medicine.
1.2.3. Control of $\text{pO}_2\text{cell}$

Most of the studies reviewed here implicitly make the assumption that $\text{pO}_2\text{gas}$ is equal to $\text{pO}_2\text{cell}$. However, depending on the cell density, medium height, cellular oxygen consumption rate, and oxygen permeability of the culture substrate, $\text{pO}_2\text{cell}$ can differ drastically from $\text{pO}_2\text{gas}$ [53]. Furthermore, step changes in $\text{pO}_2\text{gas}$ due to removal of culture dishes from the incubators may result in transient changes in $\text{pO}_2\text{cell}$ with long time scales for equilibration. These problems are ameliorated by culturing the cells on the surface of a membrane with very high oxygen permeability, such as silicone rubber [53].

To illustrate this point, we estimate $\text{pO}_2\text{cell}$ in two published studies [43, 44] (Table 1-3) using oxygen consumption rate and transport parameters combined with a model we previously developed for mESCs in monolayer [53]. The differences in $\text{pO}_2\text{cell}$ and $\text{pO}_2\text{gas}$ are very small ($<2$ mmHg) for one study and large ($>20$ mmHg) for the other because the cell densities and media heights are substantially different. Even though $\text{pO}_2\text{gas}$ in one study ($26.6$ mmHg) is higher than in the other ($15.5$ mmHg), the order of the calculated $\text{pO}_2\text{cell}$ values is reversed, $5.2$ and $12.4$ mmHg respectively. These results demonstrate that the assumption that $\text{pO}_2\text{cell}$ is equal to $\text{pO}_2\text{gas}$ is not valid for high cell densities and media heights. Knowledge of $\text{pO}_2\text{cell}$ is important for successful interpretation of experimental results and comparisons between different studies.
Differences in $pO_{2\text{cell}}$ between different experiments carried out at the same $pO_{2\text{gas}}$ may be a possible explanation for the inconsistencies in data reported. For instance, one study [35] reports reduced proliferation of hESCs cultured at 5% oxygen, and another [36] reports no reduction in proliferation at the same oxygen condition. In the first study, there are four weeks between passages (initial and final cell numbers not reported). In the second study, $6 \times 10^5$ cells are plated per well of standard six-well tissue culture plates, and there are only two weeks between passages, with final cell numbers of $1.5-2.5 \times 10^6$ cells per well. It is possible that the $pO_{2\text{cell}}$ of the first study is significantly lower than the $pO_{2\text{gas}}$ because of higher cell densities, in particular after extended culture without passage, while $pO_{2\text{cell}}$ in the latter study is closer to $pO_{2\text{gas}}$. This would help explain the differences in results because the authors of the second study report a reduced proliferation at 1% gas phase oxygen level, which may be closer in cellular oxygen level to the 5% gas phase oxygen level of the first study that also demonstrated a reduced proliferation rate. Moreover, the first group use the Xvivo system (BioSpherix, Lacona, NY), which allows for continuous culture low $pO_{2\text{gas}}$ without interruption for media changes and passaging. The second group use a conventional sealed chamber from which cells are periodically removed for media changes and passaging, allowing for periods where $pO_{2\text{gas}}$ is at ambient levels, disrupting the low $pO_{2\text{cell}}$ conditions. Without information regarding the experimental conditions, especially the media height and cell density, it is impossible to draw any conclusions.
There are instances where oxygen levels higher than ambient may be beneficial for differentiation, but control and understanding of the oxygen level experienced by the cells is still essential. Mouse pancreatic bud explants cultured at 35% oxygen preferentially differentiate to β-cells [54]. This benefit is greatest when the cells are cultured on top of a membrane with enhanced oxygen permeability compared to culture on a micro-porous membrane completely immersed in culture medium, a standard method for explant culture. \( pO_{2\text{cell}} \) on the high permeability membrane is closer to \( pO_{2\text{gas}} \) than with the standard method, allowing for better control of \( pO_{2\text{cell}} \) because oxygen permeates across these membranes more rapidly than through culture media.

1.2.4. Conclusions

Although cells and tissues differentiated from pluripotent cells have the potential to treat many diseases, improvement in methods to produce these tissues is needed, and control of cell oxygen level is potentially important in addressing this issue. Under conditions maintaining the undifferentiated phenotype, low oxygen reduces spontaneous differentiation of hESCs but has the opposite effect on mESCs, although reports are conflicting. Similar disparities are reported for proliferation rates. Differentiation under low oxygen can increase yields of desired cell types through a combination of preferential differentiation and higher total cell numbers. Low oxygen can have different effects on different stages of differentiation, so detailed understanding of the phenomena and its mechanisms
is vital. One major problem in the field is the lack of recognition that $pO_{2\text{cell}}$ is often different from $pO_{2\text{gas}}$, sometimes by a substantial amount, which is exacerbated because values of cell density and media height necessary for retrospective estimation of $pO_{2\text{cell}}$ are often not reported. Knowledge of $pO_{2\text{cell}}$ is necessary for interpreting the results of studies and comparing them to data from other studies. Efforts to estimate or control $pO_{2\text{cell}}$ are essential for rational advancement of this field.
1.3. Differentiation to cardiomyocytes

Adapted from Horton, R.E., J.R. Millman, C.K. Colton, and D.T. Auguste,

The inability of the human heart to repair itself after trauma renders it vulnerable to ischemia, viral infection, or other pathologies. Significant loss of cardiomyocytes is irreversible and may progress to heart failure. Congestive heart failure is endemic; one in three American adults suffers from one or more types of cardiovascular disease [55]. Lack of a sufficient organ supply to fulfill demand coupled with donor compatibility and organ rejection leaves a dire need for new therapeutic alternatives. Cell replacement therapies may be the solution to this deficit.

Many regenerative efforts focus on ESCs as a possible method for organ repair. ESCs are derived from the inner cell mass of blastocysts and have the potential to regenerate indefinitely and differentiate into virtually any cell type. Because of these capabilities, hESCs may be a renewable source of differentiated cells for regenerative medicine. However, embryonic differentiation is poorly understood. Of particular interest is robust cardiomyocytes for cardiovascular repair.

Cardiomyocytes are an ideal cell type for implantation to replace diseased or necrotic heart tissue because of their unique electrical, mechanical, and biological characteristics. ESC-derived cardiomyocytes display pacemaker-, atrial-, and ventricular-like characteristics [56], and spontaneously beat in vitro. These cardiomyocytes can integrate in vivo into the myocardium and improve the function of ventricles after infarction in animal models [8, 57, 58]. For example, porcine hearts exhibiting complete blockage were injected with microsectioned
hESC-derived cardiomyocytes. Electocardiogram analysis revealed that the injected cardiomyocytes successfully coupled with native heart tissue and were able to conduct electrical activity [8, 57, 58]. Concerns regarding implantation remain, including beating synchrony between implanted cells and native cells and inflammation at the injection site.

In addition to integration concerns, sufficient hESC populations are needed to repair damaged tissue. It is estimated that on the order of $10^9$ replacement cells would be necessary to treat a cardiac infarct [59]. Attention has turned to the expansion of hESC and new strategies for their differentiation. Studies have focused on the use of signaling hESC through addition of morphogens and modifications to the microenvironment to increase cardiac cell generation. Here, we review strategies that give rise to increased cardiac differentiation based on controlling the stem cell microenvironment.

hESC are often induced to differentiate by generating three-dimensional suspended cellular aggregates called embryoid bodies (EBs) in the absence of MEF and bFGF [60, 61] (Figure 1-1). EBs recapitulate many aspects of embryonic development and can serve as a model for studying organ development. The complex three-dimensional environment provides temporal and spatial cues that control differentiation into all three germ layers.
Matrix-cell and cell-cell interactions are important in stem cell differentiation, which are affected by EB size. Conventional methods used to generate EBs often results in heterogeneous size and shape. Microcontact printing [62], forced aggregation [63-65], and microwells [66] are common methods for addressing heterogeneity.

Microcontact printing is a well established technique that restricts cell growth surface area, forcing cells to conform to the patterned protein regions. ECM proteins are 'inked' onto a polymer stamp containing an array of geometric designs and then printed onto a substrate. Areas without ECM protein are treated to inhibit cell adhesion, yielding a fixed array of protein islands. A cell suspension is washed over the substrate, and cells only attach to regions with protein. EBs are formed by detaching the resulting colonies from adherent surfaces and continuing culture in suspension. This technique allows for cell number control, which is dictated by the adhesive area, resulting in colony size control [67, 68].

Altering colony and EB size influences differentiation decisions to cardiomyocytes. Bauwens et al. demonstrated that hESC colony size, controlled using the microcontact printing technique, influences cellular gene expression. Using real-time polymerase chain reaction (PCR) quantification [69]. They show that the Gata6 (endodermal marker)/Pax6 (neural marker) ratio increases with smaller colony size and hESC with a high Gata6/Pax6 ratio preferentially differentiate to cardiomyocytes. The endoderm secretes procardiogenic factor
whereas neural signals, such as Wnt, inhibit embryonic heart development. hESC colony size and EB size are interrelated parameters that influence cell fate by altering gene expression and should be controlled in differentiation studies. In another study, Niebrugge et al. combined EB size and oxygen control. Using microcontact printing, uniform hESC-derived EBs were generated and exposed to hypoxic (4% oxygen) conditions, resulting in a 3.5-fold increase in overall cardiomyocyte output when compared to traditional EB formation methods [70].

Another technique for controlling EB size is forced aggregation, which typically entails the centrifugation of a defined number of cells into multiwell plates. Forced aggregation was used to deposit hESC into round bottom multi well plates to examine the effects of size on EB differentiation patterns. Ng et al. found that efficient blood cell formation required at least 500 hESC per well, whereas optimal erythropoiesis was achieved in wells containing 1000 hESC per well [65].

Control of colony size, the number of ESC comprising EBs, and the gene expression profiles of input cells within EBs are important in influencing cell fate decisions and is an important environmental parameter. Furthermore, controlling the size of EBs can allow for better control of biomolecular and oxygen gradients [71], which may affect differentiation mechanisms. Because many differentiation methods rely on factors to induce differentiation, molecular diffusion and gradient concentrations within cell colonies can produce heterogeneous differentiation.
Recent progress has been made differentiating hESC under two-dimensional, serum-free culture by adding exogenous factors to direct differentiation to cardiomyocytes, allowing better control of the microenvironment [8]. Passier et al. employs a 2D coculture method to induce differentiation of hESC towards cardiomyocytes. Replacing the typical MEF feeder layer with mitomycin C inactivated END-2 cells, which exhibit characteristics of visceral endoderm, in the presence or absence of serum led to cardiomyocyte formation. Furthermore, the absence of serum enhanced hESC differentiation towards cardiomyocytes [72].

Many studies have examined cellular responses to growth factors. In particular, studies have employed serum-free media, conditioned media, or used media additives, such as activin A [73], ascorbic acid [74], dimethyl sulfoxide (DMSO) [75], 5-azacytidine [76], retinoic acid [77], and vascular endothelial growth factor (VEGF) [29], to promote differentiation to cardiomyocytes. The in vitro functionality of these cardiomyocytes is assessed by visual observation of spontaneous contractions, measurement of the action potential, and measurement of intracellular Ca\textsuperscript{2+} oscillations [78]. Numerous signaling pathways, such as transforming growth factor beta (TGF\textbeta), bone morphogenic protein (BMP), Wnt, and Notch, are involved in cardiogenesis [6]. Behfar et al. reported that TGF\textbeta and BMP-2 activated cardiac promoters in mESC, leading to the expression of cardiac transcription factors NK2 transcription factor related, locus 5 (Nkx2.5) and myocyte enhancer factor 2C (MEF2C) and cardiomyocyte
differentiation [79]. mESC differentiation methods are often translated to hESC, but differentiation of cardiomyocytes from hESC is often less efficient. More recently, translation of differentiation strategies from hESC to iPSC has been investigated.

iPSC are a promising alternative source of autologous cells [11, 80] and are derived by reprogramming somatic cells to an ESC-like state by inserting, often by viral vectors, factors such as Oct4, Sox2, c-Myc, Klf4, Nanog, and Lin28 [81]. Cardiomyocytes derived from both mouse [82-84] and human [85] iPSC (miPSC and hiPSC) have nodal-, atrial-, and ventricular-like action potentials, similar cardiac gene and protein expression, and spontaneous contraction rates similar to ESC-derived cardiomyocytes. Some found the rate and efficiency of differentiation comparable to ESC [82], others slower [83] and less efficient [83, 85]. Persistent expression of the transgene c-Myc, a proto-oncogene that presents a tumorigenic risk, was observed with a cardiomyocyte differentiation protocol [85]. Other cardiovascular cell types reported include endothelial, smooth muscle, and hematopoietic cells [82, 84].

Methods for quantifying cardiomyocyte generation from ESC vary. The most basic approach is to count the fraction of resulting EBs containing cells that spontaneously contract, providing crude quantification of the effects of an experimental treatment on generating cardiomyocytes compared to a control. This method does not provide information on the cells within an EB. Better
quantification can be achieved by measuring the fraction of cardiomyocytes in the total cell population by immunostaining cells for cardiac-restricted proteins, such as sarcomeric myosin heavy chain, cardiac troponin T, α-sarcomeric actin, and cardiac-restricted alpha major histocompatibility complex. The fraction of cells expressing these proteins may be counted with a flow cytometer. This information can be combined with a measurement of the total cell number to calculate the number of cardiomyocytes or, when the number of initial ESCs is known, the number of cardiomyocytes generated per input ESCs, the most useful comparison. However, most or all of this information is seldom provided, limiting rigorous comparisons between different studies.

Numerous challenges need to be addressed before hESC can be used in clinical and therapeutic applications, such as the ability to generate cardiomyocytes in clinically relevant numbers, minimization of spontaneous differentiation, and elimination of tumor-formation risk. Differentiation typically yields a mixture of cell lineages of unsuitable purity for regenerative therapies. Residual undifferentiated hESC within differentiated populations have the potential to form tumors upon implantation. If these challenges can be overcome, hESC or iPSC-derived cardiomyocytes could be used to treat heart disease, opening a door to a new era of regenerative therapies.
1.4. The tumorigenicity of pluripotent stem cells

The tumorigenicity of PSC is a major obstacle for the safe translation of stem cell biology to regenerative medicine that must be overcome [86], and the presence of any number of PSC is a potential tumor formation risk. As few as 2 m ESC have formed tumors in immunocompromised mice [87]. There have been no reported cases of undifferentiated PSC being implanted into humans, but in one case study, fetal neural stem cells implanted into a human caused the formation of a glioneuronal neoplasm, derived from donor cells [88]. Implanted iPSC can also form tumors, with a propensity that varies depending on the tissue of origin [89]. iPSC have the additional concern that inserted c-MYC genes could become reactivated and cause tumor formation [90]. In addition to the health risks to the recipient by the presence of the tumor, tumor formation could also cause a reversal of the clinical benefits of the implanted differentiated cells [91]. Preclinical testing of any stem cell-derived product for tumor formation is very important [92, 93].

Differentiation of PSC reduces the risk of tumor formation [94]. However, residual PSC can still persist, even after being subjected to a differentiation protocol, and cause tumor formation upon implantation [9, 95]. Even though studies have reported that differentiation of PSC completely avoided tumor appearance [96, 97], these experiments were not designed to sensitively detect residual PSC. A small number of residual PSC may still exist and be a safety risk for humans,
which require larger cell numbers for treatment. The experimental design of other studies has allowed for the formation of tumors with low numbers of implanted PSC [87, 98]. The fraction of animals that form tumors and the rate of tumor formation depends on the location of the implanted cells, with subcutaneous and intramuscular injections being the most convenient, sensitive, and reliable [87, 98-100].

Several methods have been used to reduce residual PSC [101]. Genetic selection has been used to kill residual PSC while retaining the desired cell type [102]. However, genetic manipulations introduce an additional hurdle to clinical translation. Positive and/or negative selection by staining cells with an fluorescently-labeled antibody against surface markers, followed by fluorescence activated cell sorting (FACS), has been used [95, 103]. An antibody, termed mAb 84, has been identified that binds to the surface protein podocalyxin-like protein-1 (PODXL) and causes cytotoxicity to human ESC (hESC) via oncosis [104], reducing the viability of undifferentiated hESC populations by a factor of 3-5 [105].

1.5. Objectives

In order for pluripotent stem cells to create a new era of regenerative medicine, they must be successfully differentiated to clinically-relevant cell types of sufficient number and purity that can be implanted into patients without the risk of tumor formation. The central hypothesis of this thesis is that oxygen is an
important parameter that greatly affects pluripotent stem cell differentiation. The objectives of this research were to investigate the effects of low oxygen culture on differentiation to the three germ layers, with particular emphasis on cardiomyocytes, and on the number and fraction of potentially-tumorigenic residual undifferentiated pluripotent cells within differentiated populations. Results and experimental details of comprehensive studies on these two objectives are described in the subsequent chapters.

1.6. Overview

Chapter 2: Extended low oxygen culture of pluripotent stem cells reduces the fraction of residual tumor-forming cells in differentiated populations

This chapter describes the effects of low oxygen culture on residual PSC within differentiated populations. mESC, miPSC, and hESC were differentiated in vitro for up to 90 days at 142, 36, or 7 mmHg pO$_2$gas on highly-oxygen permeable silicone rubber culture plates. Residual PSC were assessed by real-time PCR measurements of Oct4 and Nanog mRNA and by flow cytometric measurements of Oct4-GFP expression. Cells were implanted into immunocompromised mice to assess tumor formation rate. Low oxygen culture methodology was combined with fluorescence activated cell sorting (FACS) of residual Oct4-GFP+, SSEA-1+, and SSEA-4+ cells. Residual Oct4-GFP+ cells were separated from differentiated populations using FACS and characterized by morphological assessment,
immunostaining for Oct4, Nanog, and SSEA-1 protein expression, and in vitro differentiation. The necessity of the HIF-1α gene was assessed through use of a HIF-1α homozygous knockout mESC line.

Chapter 3: Differentiation of pluripotent stem cells under low oxygen increases generation of cardiomyocytes and influences differentiation to ectoderm, mesoderm, and endoderm

This chapter describes work that was done with Dr. Daryl E. Powers on assessing the effects of low oxygen on the differentiation of PSC to derivatives of the three germ layers, ectoderm, mesoderm, and endoderm. mESC, miPSC, and hESC were differentiated in vitro for up to 13 days at 142, 36, or 7 mmHg pO2gas on highly-oxygen permeable silicone rubber culture plates. Real-time PCR, flow cytometry, and immunostaining for numerous markers of the three germ layers were used. The generation of cardiomyocytes was further assessed by measurements of spontaneously-beating area on the culture plate. A mathematical model was developed for the volumetric distribution of cardiomyocytes within differentiated cell aggregates. The necessity of the HIF-1α gene was assessed through use of a HIF-1α homozygous knockout mESC line.
Appendix A: Accurate control of oxygen level in cells during culture on silicone rubber membranes with application to stem cell differentiation

This appendix contains a published manuscript, of which I am a co-first author with Dr. Daryl E. Powers, demonstrating the use of silicone rubber membranes for controlling pO$_{2\text{cell}}$. Mass transport models of oxygen diffusion and consumption in 2- and 3-dimensional culture are described. Fibronectin and Matrigel were physically absorbed onto the membranes to promote cell attachment. mESC were differentiated 11 days in vitro at 142, 36, or 7 mmHg pO$_{2\text{gas}}$ on highly-oxygen permeable silicone rubber culture plates. Flow cytometry and measurements of spontaneously-beating area on the culture plate were used to quantitate the generation of cardiomyocytes.

Appendix B: Effects of oxygen on mouse embryonic stem cell growth, phenotype retention, and cellular energetics

This appendix contains a published manuscript, of which I am a co-author, describing the effects of oxygen on mESC growth, phenotype retention, and cellular energetics. mESC were cultured on polystyrene at pO$_{2\text{gas}}$ values ranging from 0-285 mmHg. Cell number and viability was determined with flow cytometry. Oxygen consumption rate was determined. A comet assay was used to determine oxidative DNA damage. Pluripotency and differentiation was assessed
with morphological characterization and measurements of the expression of markers with real-time PCR and flow cytometry.

Appendix C: Differentiation of hESC to β-cells

This appendix describes work performed at ViaCyte, Inc. demonstrating my ability to replicate a protocol developed by ViaCyte differentiating hESC to insulin-secreting cells. hESC were differentiated in vitro for up to 29 days at 142 mmHg pO$_{2\text{gas}}$ on polystyrene and highly-oxygen permeable silicone rubber culture plates. Real-time PCR and immunocytochemistry are used to measure the expression of pluripotency and differentiation markers. Oxygen consumption rate was determined. ELISA is used to measure c-peptide contain in the culture medium.

Appendix D: Experiments with the BioSpherix Xvivo

This appendix describes work performed with the Xvivo incubation system, fabricated by BioSpherix, Inc., demonstrating the effect of continuous oxygen exposure on VEGF secretion and cardiomyocyte differentiation and describing the use of oxygen sensors. mESC were cultured and differentiated in vitro for up to 10 days at 142 and 36 mmHg pO$_{2\text{gas}}$ on polystyrene and highly-oxygen permeable silicone rubber culture plates in flush boxes or Xvivo incubations systems. VEGF secretion was measured with ELISA. Differentiation to
cardiomyocytes was assessed with flow cytometry. Oxygen sensors were assessed with finite element modeling and used in conjunction with a PID controller under a variety of cell culture scenarios.
1.7. Tables and Figures
Table 1-1. Effects of hypoxic culture on pluripotent stem cells

<table>
<thead>
<tr>
<th>Species</th>
<th>Study</th>
<th>Cell Line(s)*</th>
<th>Culture Surface Modification</th>
<th>Oxygen Level (%)</th>
<th>Effects of low O₂ compared to standard O₂</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mouse</td>
<td>Wang 2008</td>
<td>Newlines from C57B1/6J mice</td>
<td>Gelatin</td>
<td>5, 20</td>
<td>With physiological glucose levels, enhanced proliferation of cells but did not reduce rate of apoptosis. No reduction in cytogenetic stability at high passages.</td>
</tr>
<tr>
<td></td>
<td>Gibbons 2006</td>
<td>Newlines from C57BL6 embryos</td>
<td>MEFs</td>
<td>5, 20</td>
<td>More blastocysts produced outgrowths and had AP activity. Increased proliferation.</td>
</tr>
<tr>
<td></td>
<td>Jeong 2007</td>
<td>CCE, R1, J1, E14TG2</td>
<td>Gelatin</td>
<td>1, 20</td>
<td>Reduced AP activity and Oct4 expression. Proposed mechanism whereby HIF-1α mediates suppression of the LIF-STAT3 pathway. Reduced proliferation rates.</td>
</tr>
<tr>
<td></td>
<td>Powers 2008</td>
<td>CCE, D3</td>
<td>Gelatin</td>
<td>0, 1, 5, 20, 40</td>
<td>Maintained pluripotency under a wide range of O₂ conditions, although high (40%) and low (0-1%) O₂ reduced Oct4/Nanog/Sox2 expression. Reduced proliferation rates at high (40%) and low (0-1%) O₂. No change for intermediate values (5-20%). No effects on oxidative damage to DNA under all conditions</td>
</tr>
<tr>
<td>Human</td>
<td>Ying 2008</td>
<td>E14TG2a</td>
<td>-</td>
<td>5, 21</td>
<td>Clonal propagation achieved without B27 supplement in serum-free culture.</td>
</tr>
<tr>
<td></td>
<td>Ludwig 2006</td>
<td>WA15 &amp; WA16</td>
<td>Matrigel or a combination of collagen IV, fibronectin, laminin, and vitronectin</td>
<td>5, 10, 20</td>
<td>Highest proliferation rate at 5% oxygen/10% CO₂</td>
</tr>
<tr>
<td></td>
<td>Forsyth 2006</td>
<td>H1, RH1, H9</td>
<td>Matrigel</td>
<td>2, 21</td>
<td>Reduced heterogeneity in transcriptome. No change in expression of Oct4/Nanog/Sox2.</td>
</tr>
<tr>
<td></td>
<td>Westfall 2006</td>
<td>H1, H9</td>
<td>MEFs or Matrigel</td>
<td>4, 20</td>
<td>Reduced heterogeneity in transcriptome. No change in expression of Oct4/Nanog/Sox2. Increased expression of Lef1/2.</td>
</tr>
<tr>
<td></td>
<td>Prasad 2009</td>
<td>CLS1, CLS2</td>
<td>HFF-1</td>
<td>1, 5, 10, 15, 21</td>
<td>No change in Oct4 expression, some increase in Nanog/Notch1 expression at 5% in culture for 4 weeks, no difference compared to 20% when cultured for 18 months. Reduced proliferation rates at 5%.</td>
</tr>
<tr>
<td></td>
<td>Ezashi 2005</td>
<td>H1</td>
<td>MEFs or Matrigel</td>
<td>1, 3, 21</td>
<td>Reduction of differentiated areas around colonies expressing SSEA-1 at 1% and 3%. No differences in proliferation rates at 3%, slight reduction at 1%.</td>
</tr>
<tr>
<td></td>
<td>Forsyth 2006</td>
<td>H1, H9, RH1</td>
<td>Matrigel</td>
<td>2, 21</td>
<td>No change in SSEA-1 or SSEA-4 expression, increased clonal recovery. Decreased chromosomal abnormalities.</td>
</tr>
<tr>
<td></td>
<td>Hewett 2006</td>
<td>H9</td>
<td>Matrigel</td>
<td>2, 21</td>
<td>Increased clonal recovery.</td>
</tr>
</tbody>
</table>

*Boldface indicates main cell line(s) used.

MEFs - Mouse Embryonic Fibroblasts; AP - Alkaline Phosphatase; HIF - Hypoxia Inducible Factor; LIF - Leukaemia Inhibitory Factor; HFF - Human Foreskin Fibroblasts

<table>
<thead>
<tr>
<th>Species</th>
<th>Study</th>
<th>Cell Lines</th>
<th>Oxygen Level (%)</th>
<th>Effects of low O$_2$ compared to standard O$_2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mouse</td>
<td>Kim 2008</td>
<td>D3</td>
<td>3.5</td>
<td>Differentiation to neurons. Number of neurons increased 35%. HIF-1α dependent.</td>
</tr>
<tr>
<td></td>
<td>Mondragon-Teran 2009</td>
<td>E14Tg2a</td>
<td>2</td>
<td>Differentiation to neurons. Number of neurons increased factor of 55-114.</td>
</tr>
<tr>
<td></td>
<td>Bauwens 2005</td>
<td>D3</td>
<td>4</td>
<td>Differentiation to cardiomyocytes. Total cell number increased 50%. Used bioreactor system.</td>
</tr>
<tr>
<td></td>
<td>Purpura 2008</td>
<td>R1</td>
<td>4</td>
<td>Differentiation to hemangioblast, endothelial cells, and hematopoietic progenitor cells. Low oxygen involved in complicated network via VEGF secretion to influence differentiation to these lineages.</td>
</tr>
<tr>
<td>Human</td>
<td>Niebruegge 2009</td>
<td>H9, HES2</td>
<td>4</td>
<td>Differentiation to cardiomyocytes. Total cell number increased 30-47%. Some cardiac markers increased.</td>
</tr>
<tr>
<td></td>
<td>Koay 2008</td>
<td>H9</td>
<td>2</td>
<td>Differentiation to chondrocytes. Collagen I and II, tensile modulus, and compressive properties increased.</td>
</tr>
</tbody>
</table>

# Table 1-3. Estimated values of pO$_{2\text{cell}}$ from published studies

<table>
<thead>
<tr>
<th>Study</th>
<th>Cell density (10$^3$ cells/cm$^2$)</th>
<th>Media Height (mm)</th>
<th>pO$_{2\text{gas}}$ (mmHg)</th>
<th>pO$_{2\text{cell}}$ (mmHg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kim 2008$^b$</td>
<td>150</td>
<td>3.8$^c$</td>
<td>26.6</td>
<td>5.2</td>
</tr>
<tr>
<td></td>
<td>150</td>
<td>3.8</td>
<td>142</td>
<td>99.7</td>
</tr>
<tr>
<td>Mondragon-Teran 2009$^a$</td>
<td>28</td>
<td>2.1</td>
<td>15.2</td>
<td>12.4</td>
</tr>
<tr>
<td></td>
<td>0.82</td>
<td>2.1</td>
<td>142</td>
<td>141.9</td>
</tr>
</tbody>
</table>

$^a$ Values from 8 days of differentiation  
$^b$ Values from the beginning of differentiation of NSC to neurons  
$^c$ Typical media height for coverslides  

ESCs aggregate into EBs via hanging drop, static suspension, convective suspension, micropatterned wells or by forced aggregation. ESCs may also be cultured in 2D plates on various ECM matrices or micropatterened substrates. ESCs may also be embedded in scaffolds. These microenvironments may be subjected to shear and morphogens, for example, to promote differentiation. During differentiation, expression of pluripotency markers, such as Oct4 and Nanog, decreases, early mesodermal markers, such as Mesp1 and Brachyury T, are transiently expressed and expression of late-stage markers, such as Nkx2.5, Mef2c, and α-MHC, increases. Reproduced from Horton, R.E., J.R. Millman, C.K. Colton, and D.T. Auguste, *Engineering microenvironments for embryonic stem*
Chapter 2. Extended Low Oxygen Culture of Pluripotent Stem Cells Reduces the Fraction of Residual Tumor-Forming Cells in Differentiated Populations
2.1. Abstract

Residual pluripotent stem cells (PSC) within differentiated populations are problematic because of their potential to form tumors. Simple methods to reduce their occurrence are needed. Here, we demonstrate that control of the oxygen partial pressure (pO$_2$) to physiological levels typical of the developing embryo reduces the fraction of PSC within, and the tumorigenic potential of, differentiated populations. Culture under differentiating conditions at low pO$_2$ reduced measured pluripotency markers by up to four orders of magnitude for mouse and human embryonic stem cells and mouse induced pluripotent stem cells. Upon implantation into immunocompromised mice, low pO$_2$-differentiated PSC either did not form tumors or formed tumors at a slower rate than high pO$_2$ PSC. Low pO$_2$ differentiation could be combined with cell sorting for improved benefits. Low pO$_2$ culture alone or in combination with other methods is a potentially straightforward method that could be applied to future cell therapy protocols to minimize the possibility of tumor formation.
2.2. Introduction

Pluripotent stem cells (PSC), such as embryonic stem cells (ESC) [106, 107] and induced pluripotent stem cells (iPSC) [11, 80], have immense clinical potential due to their ability to form many clinically-relevant cell types [1-3]. However, because of their pluripotency, ESC and iPSC have the potential to form teratomas [86], tumors composed of derivatives of the three germ layers [92], and are a serious safety risk if implanted into humans in their undifferentiated state [93].

The tumorigenicity of PSC is a major obstacle for the safe translation of stem cell biology to regenerative medicine that must be overcome [86], and the presence of any number of PSC is a potential tumor formation risk. As few as 2 mouse ESC (mESC) have formed tumors in immunocompromised mice [87]. There have been no reported cases of undifferentiated PSC being implanted into humans, but in one case study, fetal neural stem cells implanted into a human caused the formation of a glioneuronal neoplasm, derived from donor cells [88]. Implanted iPSC can also form tumors, with a propensity that varies depending on the tissue of origin [89]. iPSC have the additional concerns that inserted c-MYC genes could become reactivated and cause tumor formation [90] and that the process of reprogramming can cause genetic aberrations [108]. In addition to the health risks to the recipient by the presence of the tumor, tumor formation could also cause a reversal of the clinical benefits of the implanted differentiated cells [91].
Preclinical testing of any stem cell-derived product for tumor formation is very important [92, 93].

Differentiation of PSC reduces the risk of tumor formation [94]. However, residual PSC can still persist, even after being subjected to a differentiation protocol, and cause tumor formation upon implantation [9, 95]. Even though studies have reported that differentiation of PSC completely avoided tumor appearance [96, 97], these experiments were not designed to sensitively detect residual PSC. A small number of residual PSC may still exist and be a safety risk for humans, which require larger cell numbers for treatment. The experimental design of other studies has allowed for the formation of tumors with low numbers of implanted PSC [87, 98]. The fraction of animals that form tumors and the rate of tumor formation depends on the location of the implanted cells, with subcutaneous and intramuscular injections being the most convenient, sensitive, and reliable [87, 98-100].

Several methods have been used to reduce residual PSC [101]. Genetic selection has been used to kill residual PSC while retaining the desired cell type [102]. However, genetic manipulations introduce an additional hurdle to clinical translation. Positive and/or negative selection by staining cells with an fluorescently-labeled antibody against surface markers, followed by fluorescence activated cell sorting (FACS), has been used [95, 103]. An antibody, termed mAb 84, has been identified that binds to the surface protein podocalyxin-like protein-1
(PODXL) and causes cytotoxicity to human ESC (hESC) via oncosis [104], reducing the viability of undifferentiated hESC populations by a factor of 3-5 [105].

The oxygen partial pressure (pO₂) under which embryonic cells develop in vivo and PSC are cultured in vitro is an important parameter [2, 42]. During embryonic development, cells are exposed to pO₂ much lower than atmospheric (160 mmHg), which influences many aspects of embryogenesis, such as tracheal development, cardiovascular and bone morphogenesis, and adipogenesis, often via hypoxia inducible factors (HIF). Teratomas do not form normally, even though all differentiated cells are derived from pluripotent precursors. In addition, differentiation of PSC in vitro at low pO₂ affects the generation of several cell types, such as neurons, hematopoietic progenitors, endothelial cells, cardiomyocytes, and chondrocytes. pO₂ also affects many aspects of pluripotency, such as expression of the pluripotency genes Oct4, Nanog, and Sox2, in mESC and hESC. The reprogramming efficiency of mouse and human fibroblasts to iPSC is increased under low pO₂ [109, 110]. Female hESC derived under low pO₂ maintain two active X chromosomes and upon culture at atmospheric pO₂, irreversibly silences one X chromosome [111].

In this study, we show that differentiation and extended culture of mESC and mouse iPSC (miPSC), both containing a green fluorescence protein (GFP) transgene driven by an Oct4 promoter, and hESC at low pO₂ drastically reduced the fraction and number of residual PSC, which, upon implantation into nude
mice, slowed or eliminated the appearance of teratomas. Low pO₂ culture was combined with FACS removal of PSC based on GFP and PSC surface marker expression to reduce the fraction of residual PSC, more than either method alone. Residual Oct4-GFP+ mESC, after being subjected to the differentiation protocol, had in vitro characteristics similar to those of undifferentiated mESC.

2.3. Methods

2.3.1. Undifferentiated mESC and miPSC culture

Undifferentiated R1 mESC and miPSC, both with an inserted GFP coding sequence driven by an inserted Oct4 promoter (Oct4-GFP mESC) [112, 113] (generously donated by Professor Douglas Melton of Harvard University, Cambridge, MA, USA) and R1 mESC with a homozygous HIF-1α gene knockout (HIF-1α⁻/⁻ mESC) [114] (generously donated by Professor Peter Carmeliet of Katholieke Universiteit Leuven, Leuven, Belgium) were propagated as previously described [53] without mouse embryonic fibroblasts (MEFs) in Dulbecco’s modified Eagles medium (DMEM; SCRR-2010; American Type Culture Collection (ATCC), Manassas, VA, USA) supplemented with 10% (v/v) fetal bovine serum (FBS; SCRR 30-2020; ATCC), 2 mM L-alanyl-L-glutamine (SCRR 30-2115; ATCC), 50 units/ml penicillin and 50 μg/ml streptomycin (15070; Invitrogen, Carlsbad, CA, USA), 1x MEM non-essential amino acids (SCRR 30-2116; ATCC), 100 μM 2-mercaptoethanol (M7522; Sigma-Aldrich; St. Louis, MO,
USA), and 10³ units/mL leukemia inhibitory factor (LIF; LIF2005; Millipore, Billerica, MA, USA). Cells were passaged every two d by detaching with 0.25% trypsin (25-053-C1; Mediatech, Manassas, VA, USA) and plating at a density of 1.2x10⁴ cells cm⁻² on cell culture flasks (353014; BD Biosciences, Bedford, MA, USA).

2.3.2. Gas phase pO₂ control

pO₂ in the gas phase (pO₂gas) was maintained by flowing premixed gas containing 5% CO₂ and 20%, 5%, or 1% O₂ with a balance of N₂ (certified medical gas from Airgas, Hingham, MA, USA), which corresponded under humidified conditions to 142, 36, and 7 mmHg pO₂gas inside chambers maintained at 37 °C as previously described [53].

2.3.3. Silicone rubber membrane-based plate

Differentiation and extended culture was performed on silicone rubber membranes having high oxygen permeability as previously described [53]. In brief, the bottoms of 24-well polystyrene plates (353047; Becton Dickinson, Franklin Lakes, NJ, USA) were replaced with silicone rubber sheets (nonreinforced vulcanized gloss/gloss 127 μm thick; Specialty Manufacturing, Saginaw, MI, USA) and sterilized by filling with 70% (v/v) ethanol (111000200; Pharmco-AAPER, Brookfield, CT, USA) in water, incubating for 1 hr, and drying
overnight under a germicidal UV lamp in a biological safety cabinet (SterilGARD III Advance; The Baker Company, Sanford, MA, USA). In use, the corners of the plate were supported on top of a 150x15 mm Petri dish (35384-326; VWR, West Chester, PA, USA), and the silicone rubber membrane was exposed directly to the gas phase. By culturing on these silicone rubber membrane-based plates, pO$_2$ of the cells (pO$_{2\text{cell}}$) at the cell-membrane interface was maintained virtually equal to pO$_{2\text{gas}}$, and rapid equilibration of pO$_{2\text{cell}}$ occurred after a change in pO$_{2\text{gas}}$. Attachment of mESC and miPSC was achieved by incubating silicone rubber membranes at 37 °C with 2 µg/mL fibronectin (F1141; Sigma-Aldrich) in phosphate buffered saline (PBS; 21-040; Mediatech) for 24 hr, and of hESC by incubation with Matrigel (354234, Becton Dickinson) diluted 1:20 in serum-free medium for 1 hr, for hESC.

2.3.4. Differentiation of mESC and miPSC

mESC and miPSC were differentiated as previously described [53]. In brief, embryoid bodies (EBs) were formed in propagation medium without LIF with 500 cells per 20-µL hanging drops. After 2 d, 30 EBs per well were transferred to fibronectin-coated (F1141; Sigma) silicone rubber membrane-based 24-well plates filled with 2 mL of propagation medium without LIF. After 3 additional d, medium was replaced with 2.5 mL of serum-free medium, consisting of 50% (v/v) DMEM (90-133-PB; Mediatech) and 50% (v/v) Ham's F-12 (10-080-CV; Mediatech) supplemented with 0.75% (w/v) sodium bicarbonate (25-035-CI;
Mediatech), 9 mM glucose (G8769; Sigma-Aldrich), 5 μg/mL human insulin (I9278; Sigma-Aldrich), 50 μg/mL holo transferrin (T1283; Sigma-Aldrich), 31.2 nM sodium selenite (S9133; Sigma-Aldrich), 50 units/ml penicillin, and 50 μg/ml streptomycin. Cells were differentiated for a total of 10 d. As noted, pO2_{gas} was kept constant throughout differentiation in some experiments and was varied in others. In some experiments, 0.2 mM ascorbic acid (A4034; Sigma-Aldrich) was also included in the medium, as noted. At constant 142 and 7 mmHg, this protocol produced a final cell population of which approximately 10% and 30% of the cells were cardiomyocytes, respectively.

### 2.3.5. Extended culture of differentiated mESC and miPSC

After the 10-d differentiation protocol, cells were subjected to additional extended culture for up to 80 d. In some experiments, pO2_{gas} was identical to pO2_{gas} during differentiation, while in others, pO2_{gas} was changed. Cell samples were acquired for analysis by detachment with a 5-min incubation of 0.25% trypsin at 37 °C.

### 2.3.6. Flow cytometric analysis of Oct4-GFP expression

Oct4-GFP expression was measured with a flow cytometer (FACScan; Becton Dickinson) at the Koch Institute (MIT). A negative control of R1 mESC without the Oct4-GFP reporter was differentiated under the same conditions as the Oct4-GFP cells, and a gating threshold was set so that all of these cells were counted
as negative for Oct4-GFP. The fraction of cells expressing Oct4-GFP (Oct4-GFP+) was calculated as the number of cells above the threshold divided by the number of cells in the entire population at each specific measurement time and condition.

2.3.7. Real-time polymerase chain reaction (PCR)

Total RNA was isolated using the RNeasy Kit (74104, Qiagen, Valencia, CA), and RNase-Free DNase Set (79254, Qiagen), cDNA was synthesized using High Capacity cDNA Reverse Transcription Kit (4368814, Applied Biosystems, Foster City, CA). Real-time PCR was performed on a Fast Real-Time PCR System (7900HT, Applied Biosystems) using Power SYBR Green PCR Master Mix (4367659, Applied Biosystems), and 28S ribosomal RNA (rRNA) was used as an endogenous control. Primer sequences used for 28S rRNA, Oct4 (mouse), Nanog (mouse), Nestin, and Nkx2.5 have been previously reported [32]. Additional primers sequences used (in order of forward and reverse primer and 5’ to 3’) were Brachyury T TCCCGGTGCTGAAGGTAAAT and CCGTCAGCAGTCCAGCAA, Foxa2 TCAAGGCCTACGAACAGGTCAT and GCCCGCTTTGTTCGTGACT, cTnT AGATGCTGAAGGAGGTCCAGT and CACCAAGTGGGCATGAAGA, cardiac-α-Actin GCTTCCGCTGTCCAGAGACT and TGCCAGCAGATTCCATACCA, OCT4 (human) TGGGCTCGAGAAGGATGTG and GCATAGTCGCTGCTTGATCG, and NANOG (human) GCAATGTCTTTCTGCTGAG and
CCATGGAGGAGGGAAGAGGA. A standard calibration curve was constructed using serial dilutions of pooled cDNA from all conditions.

2.3.8. Cell enumeration

Detached cells or excised tumors were lysed to liberate nuclei by vigorous vortexing in deionized water containing 1% (v/v) Triton X-100 (T-9284; Sigma-Aldrich) and 0.1 M citric acid (0627; Mallinckrodt Specialty Chemicals Co., Paris, KY, USA). Nuclei were stained with Guava ViaCount (4000-0041; Millipore) and counted using a Guava PCA flow cytometer (Millipore) as previously described [53, 115].

2.3.9. Cell implantation into mice

Female nude mice, 43-56 d old (Charles River Laboratories International, Wilmington, MA, USA), were housed and cared for, and experiments were performed with a protocol (approved by the MIT Committee on Animal Care) based on one designed for biosafety testing of stem cell products, for which implantation of only two ES cells was sufficient to form detectable tumors [87]. Samples containing $10^5$ cells (for mESC and miPSC) and $10^6$ cells (for hESC) in 200 µl of 50% (v/v) Matrigel (354234; BD Biosciences) and 50% (v/v) serum-free differentiation medium were injected subcutaneously into the lower left flank of
isoflurane-anaesthetized mice using a 23G needle in a laminar-flow hood. Mice were visually observed every 3-4 d for the appearance of masses under the skin.

2.3.10. Histology

Tumors were excised from euthanized mice, fixed overnight in 4% (w/v) paraformaldehyde (36606; Alfa Aesar, Ward Hill, MA, USA) in PBS, paraffin embedded, sectioned, and stained with hematoxylin and eosin (H&E) at the Division of Comparative Medicine (DCM) at MIT. Stained histological sections were analyzed by an American College of Veterinary Pathologist board-certified pathologist.

2.3.11. Fluorescence activated cell sorting (FACS)

mESC and hESC that underwent 10 d differentiation and an additional 20 d extended culture were detached with trypsin and then sorted with an Aria 2 FACS (Becton Dickinson). mESC were sorted into Oct4-GFP+ and Oct4-GFP- populations or sorted based on surface marker expression of the mESC surface marker stage-specific embryonic antigen-1 (SSEA-1). hESC were sorted based on surface marker expression of the hESC marker SSEA-4. For SSEA-1 and SSEA-4 sorting, cells were treated with a phycoerythrin (PE)-conjugated anti-SSEA-1 antibody (12-8813; eBioscience, San Diego, CA, USA) or PE-conjugated
anti-SSEA-4 antibody (12-8843-71; eBioscience) before sorting into SSEA-1+, SSEA-1-, SSEA-4+, and SSEA-4- populations.

2.3.12. Flow cytometric analysis of immunostained cells

The fraction of Nestin+ and MF-20+ cells was measured with flow cytometry as previously described [53]. In brief, detached cells were fixed in 1% (w/v) paraformaldehyde in PBS for 20 min. Samples of 3x10^5 fixed cells were incubated 10 min in 0.5% (w/v) saponin (S-4521; Sigma-Aldrich) in PBS to permeabilize cell membranes and incubated in 2% (v/v) FBS in PBS for 30 min to block non-specific binding. Cells were incubated with primary antibody against Nestin (MAB2736; R&D Systems, Minneapolis, MN, USA) diluted 1:50 or sarcomeric myosin heavy chain (MF-20; MF-20 supernatant; Developmental Studies Hybridoma Bank (DSHB), Iowa City, IA, USA) diluted 1:10 in 2% FBS solution for 1 hr, washed 1x with PBS, and incubated with secondary goat anti-mouse PE-conjugated antibody (Jackson ImmunoResearch, West Grove, PA) diluted 1:250 in 2% FBS solution in the dark. Stained cells were washed thrice with PBS, and fluorescence intensity data were acquired using the Guava PCA flow cytometer with the Express software module.
2.3.13. Immunocytochemistry

Cells attached to silicone rubber or polystyrene were fixed in 4% paraformaldehyde in PBS for 20 min and then permeabilized and non-specific protein binding blocked with a 45-min incubation in 3% (v/v) donkey serum (DS; D9663; Sigma-Aldrich) and 0.1% Triton X-100 in PBS. Cells were then incubated overnight at 4 °C with primary antibodies against Oct4 (sc-9081; Santa Cruz Biotechnology, Santa Cruz, CA, USA) diluted 1:100, Nanog (IHC-00205; Bethyl Laboratories, Montogomery, TX, USA) diluted 1:100, SSEA-1 (MC-480; DSHB) diluted 1:10, Foxa2 (sc-6554; Santa Cruz Biotechnology) diluted 1:200, Nestin diluted 1:50, or cardiac troponin T (cTnT; MS-295-P1; NeoMarkers, Fremont, CA, USA) diluted 1:50 in 3% DS, 0.1% Triton X-100 solution, washed once with PBS, and incubated with secondary donkey Alexa Fluor 488 (green) or 594 (red) antibodies (Invitrogen) against the appropriate species diluted 1:250 in 3% DS, 0.1% Triton X-100 solution for 2 hr in the dark. DNA was stained with a 10-min incubation with 1 μg/mL 4′,6-diamidino-2-phenylindole (DAPI; 32670, Sigma-Aldrich) in PBS in the dark. Stained cells were then washed thrice with PBS and images acquired with an Axiovert 200 fluorescence microscope (Carl Zeiss, Oberkochen, Germany).
2.3.14. Differentiation of hESC

CyT49 hESC [9] (NIH Registration Number 0041; generously provided by ViaCyte, Inc, San Diego, CA, USA) were plated on Matrigel-coated silicone rubber plates at a density of $1.5 \times 10^5$ cells cm$^{-2}$ without MEFs, propagated 4 d in self-renewal media, and then switched to DMEM/F12 (15-090-CV; Mediatech) with 10% (v/v) Knockout serum replacement (A1099202; Invitrogen) for up to 90 d.

2.3.15. Statistics

Statistical analysis was performed using unpaired t-tests, with significant results requiring $p<0.05$. Data is presented as mean ± standard deviation.

2.4. Results

2.4.1. Low pO$_2$ reduces residual mESC

We differentiated and performed extended culture on Oct4-GFP mESC at various pO$_2$gas to examine the effect on residual mESC. Before differentiation, the fraction of mESC expressing Oct4-GFP (Oct4-GFP+), determined with flow cytometry, was 0.96 (Figure 1A). This fraction decreased rapidly during the 10-d differentiation, after which the fraction of total cells that were Oct4-GFP+ was
$3 \times 10^{-3}$, $6 \times 10^{-4}$ and $1 \times 10^{-3}$, for culture at 142, 36, and 7 mmHg, respectively. The residual Oct4-GFP+ subpopulation had GFP intensity values similar to that of undifferentiated mESC (data not shown). We hypothesized that these cells were residual PSC that could develop into tumors upon implantation into animals.

The differentiation process was largely complete by about 10 d of culture, on the basis that no more spontaneously-contracting cells formed after that time. After differentiation, culture at each constant $pO_{2\text{gas}}$ was extended for up to 80 additional d to further examine its effects on the reduction of Oct4-GFP+ cells. The fraction of Oct4-GFP+ cells did not change significantly at 142 mmHg but continued to decrease at 36 and 7 mmHg (Figure 1A). The greatest difference was observed at 7 mmHg after 90 total d of culture, at which point the fraction of Oct4-GFP+ cells was a factor of 1260 lower than at 142 mmHg.

Low oxygen culture also reduced the number of Oct4-GFP+ cells (Figure 1B). Each well was initially seeded with $1.44 \times 10^4$ Oct4-GFP+ cells. After 10 d of differentiation, this number remained unchanged at 142 mmHg but decreased to $3 \times 10^3$ and $4 \times 10^3$ at 36 and 7 mmHg, respectively. During the next 20 d of culture, Oct4-GFP+ cell number decreased to $5 \times 10^3$ at 142 mmHg and did not change significantly thereafter. This reduction was proportional with the total cell number (Figure 1C). In contrast, the number of Oct4-GFP+ cells decreased continuously at 36 and 7 mmHg, resulting in only 10 and 4 Oct4-GFP+ cells per well remaining by d 90, respectively. This reduction was substantially faster than the reduction in
total cell number. Gene expression of Oct4 and Nanog, measured by real-time PCR, decreased with time in a manner comparable to that observed with flow cytometry of Oct4-GFP+ cells (Figure 1D-E).

When observed with fluorescence microscopy, cells that expressed GFP after culture at 142 mmHg resided in clusters within aggregates (Figure 1F-G), whereas no cells expressing GFP were visually observed anywhere in the dish, including in aggregates, at 7 mmHg (Figure 1H-I) or at 36 mmHg (data not shown). The inability to visually observe Oct4-GFP+ cells at 36 or 7 mmHg with microscopy likely reflects the low fraction of cells expressing high enough levels of GFP to be detected by eye, rather than a complete absence of GFP-expressing cells.

Different pO$_{2\text{gas}}$ values during differentiation and extended culture were investigated by differentiating mESC for 10 d at either 142 or 7 mmHg, followed by 20 d extended culture at the same or the other pO$_{2\text{gas}}$ (Figure 2). After 10 d of differentiation at 142 mmHg, the fraction of Oct4-GFP+ cells was 1.4x10$^{-2}$ (Figure 2A). Extended culture for 20 d at 142 mmHg did not significantly change the fraction of Oct4-GFP+ cells. However, extended culture at 7 mmHg decreased the fraction to 5x10$^{-4}$. Differentiation for 10 d at 7 mmHg resulted in a fraction of Oct4-GFP+ cells of 6x10$^{-4}$ (Figure 2B), a factor of 22 lower than at 142 mmHg (Figure 2A). Extended culture for 20 d at 7 mmHg further decreased the fraction
to $3 \times 10^{-5}$. However, extended culture at 142 mmHg increased the mean fraction to $3 \times 10^{-3}$.

After 10 d of differentiation at 142 mmHg, the number of Oct4-GFP+ cells was $3.5 \times 10^5$ (Figure 2C). Extended culture for 20 d at 142 mmHg decreased the number of Oct4-GFP+ cells slightly to $1.6 \times 10^4$. However, extended culture at 7 mmHg decreased the number to $9.5 \times 10^2$. Differentiation for 10 d at 7 mmHg resulted in a number of Oct4-GFP+ cells of $1.1 \times 10^3$ (Figure 2D), a factor of 33 lower than at 142 mmHg. Extended culture for 20 d at 7 mmHg further decreased the number to 33. However, extended culture at 142 mmHg increased the mean number to $3.1 \times 10^3$. Extended culture at 142 and 7 mmHg for 20 d, at either pO$_{2_{gas}}$, decreased the total cell number by approximately same proportion (Figures 2E-F).

If differentiated cells that have been previously cultured under normoxic conditions are to be cultured under hypoxic conditions in order to reduce residual PSC, it is possible that desired cell types will die if they are unable to use anaerobic metabolism. To investigate this possibility, we differentiated mESC and cultured the cells for extended time with three different sequences of pO$_{2_{gas}}$: (A) Differentiate at 142 mmHg for 10 d, then extended culture at 142 mmHg for 20 d; (B) Differentiate at 142 mmHg for 10 d, then extended culture at 7 mmHg for 20 d; (C) Differentiate at 7 mmHg for 6 d, then differentiate at 142 mmHg for 4 d, then extended culture at 7 mmHg for 20 d. With all sequences, spontaneously-
contracting cells were observed. The total number of cells and the number of MF-20+ (cardiomyocyte marker) cells is tabulated in Table 1 for d 10 and 30 of culture, along with the relative decrease from each number. At d 10, sequence C maximized the number and fraction of cardiomyocytes. With all sequences, the total cell number and number of MF-20+ cells decreased from 10 to 30 d. The decreases were smallest with sequences B and C, which involved extended culture at 7 mmHg, compared to sequence A, which used extended culture at 142 mmHg.

2.4.2. Low pO₂ delays appearance of tumors after implantation of cultured mESC

pO₂gas culture conditions of differentiation affected the rate of tumor formation following subcutaneous injection into nude mice (Figure 3A). All animals injected with undifferentiated mESC (positive control) formed visible tumors by d 14. Differentiation for 30 d before implantation led to slower tumor formation rates and resulted in only 306, 7, and 2 Oct4-GFP+ cells being implanted per 10⁵ cell injection for cells cultured at 142, 36, and 7 mmHg, respectively. Visible tumors appeared in 7/8 animals by d 17 and 8/8 by d 21 for cells cultured at 142 mmHg. Culture at lower pO₂gas had a greater effect. Tumors were first observed in 3/8 animals by d 31, 5/8 by d 35, and 7/8 by d 38 for cells cultured at 36 mmHg and in 5/8 animals by d 35, 6/8 animals by d 38, and 7/8 animals by d 42 for 7 mmHg. Animals (0/6) injected with only Matrigel (negative control) did not form tumors.
Tumors were excised from euthanized mice, fixed, embedded, sectioned, and H&E stained, and the resulting tissue sections histopathologically analyzed (Figure 3B). The tumors contained derivatives of all three germ layers and were identified as teratomas. These in vivo data are consistent with the in vitro Oct4-GFP and real-time PCR data and demonstrate that extended hypoxic culture of mESC greatly reduces the rate of teratoma formation in vivo.

2.4.3. Combination of low pO\textsubscript{2} culture of mESC with FACS

In a separate experimental run, mESC after 30-d differentiation at either 142 or 7 mmHg were sorted for GFP expression with FACS into Oct4-GFP+ and Oct4-GFP- populations before implantation (Figure 4). With cells cultured at 142 mmHg, the fraction of residual Oct4-GFP+ cells remaining in the sorted Oct4-GFP- population decreased compared to the unsorted population by a factor of 50, as determined by subsequent flow cytometric analysis of the sorted population (Figure 4A). With culture at 7 mmHg, all Oct4-GFP+ cells were removed from the sorted Oct4-GFP- population, as determined by subsequent flow cytometric analysis. Low pO\textsubscript{2} culture combined with FACS also reduced the number of Oct4-GFP+ cells (Figure 4B). Before sorting, culture at 142 mmHg resulted in 4.1x10\textsuperscript{4} Oct4-GFP+ cells. Sorting into an Oct4-GFP+ population captured 3.8x10\textsuperscript{4} of the Oct4-GFP+ cells. However, 9.4x10\textsuperscript{2} Oct4-GFP+ cells were sorted into the Oct4-GFP- population. There are fewer Oct4-GFP+ cells when the Oct4-GFP+ and Oct4-GFP- populations are combined compared to the
unsorted Oct4-GFP+ cells, likely loss during sample handling or due to measurement error. For culture at 7 mmHg, each well contained 190 Oct4-GFP+ cells before sorting. Sorting into an Oct4-GFP+ population captured all the Oct4-GFP+ cells. FACS removal of Oct4-GFP+ cells decreased the total cell number by a very small fraction for both 142 and 7 mmHg cells, a factor of 1.07 and 1.0004, respectively (Figure 4C).

After FACS, unsorted cells and sorted Oct4-GFP+ and Oct4-GFP- populations, each cultured at 142 or 7 mmHg, were implanted into nude mice (Figures 4D). Each unsorted cell injection contained, on average, 1770 and 13 Oct4-GFP+ cells at 142 and 7 mmHg, respectively, each sorted Oct4-GFP+ population injection contained 1729 and 13 Oct4-GFP+ cells at 142 and 7 mmHg, respectively, and each sorted Oct4-GFP- population injection contained 41 and 0 Oct4-GFP+ cells at 142 and 7 mmHg, respectively. For cells cultured at 142 mmHg, tumors appeared at approximately the same time in the unsorted and sorted GFP+ populations; The Oct4-GFP- population had an 11-d delay in tumor appearance. For 7 mmHg, there was a 6-d delay in initial tumor appearance for the sorted GFP+ population compared to the unsorted cells, but 50% of the GFP+ cells never formed tumors. Cells cultured at 7 mmHg then sorted into Oct4-GFP- populations formed no tumors during 210 d of observation.

The combination of low pO₂ culture with FACS using an Oct4 reporter cell line reduced residual PSC compared to low pO₂ culture alone. However, sorting
based on GFP expression required genetic modifications of the cells, which might be problematic for clinical application. Therefore, we investigated low pO₂ culture combined with FACS of mESC cell surface marker stained with a PE-conjugated anti-SSEA-1 antibody (Figure 5). mESC underwent, in a separate experimental run, a 30-d differentiation at either 142 or 7 mmHg and were sorted for SSEA-1 expression with FACS into SSEA-1+ and SSEA-1- populations before implantation. The SSEA-1- populations had a fraction of Oct4-GFP+ cells that were a factor of 31 and 16 lower than unsorted cells from 142 and 7 mmHg, respectively, as determined by subsequent flow cytometric analysis of the sorted population (Figure 5A). Culture at 7 mmHg combined with FACS removal of SSEA-1+ cells resulted in a reduction in the fraction of Oct4-GFP+ cells by a factor of 1020 compared to cells culture at 142 mmHg without sorting. Low pO₂ culture combined with immunostaining for SSEA-1 and FACS also reduced the number of Oct4-GFP+ cells (Figure 5B). Before sorting, culture at 142 mmHg resulted in 6.5x10³ Oct4-GFP+ cells. Sorting into an SSEA-1+ population captured 6.2x10³ of the Oct4-GFP+ cells. However, the SSEA-1- population contained 1.9x10² Oct4-GFP+ cells. For culture at 7 mmHg, each well contained 110 Oct4-GFP+ cells before sorting. Sorting into an Oct4-GFP+ population captured 95 Oct4-GFP+ cells. However, the SSEA-1- population contained 7 Oct4-GFP+ cells. There are fewer Oct4-GFP+ cells when the SSEA-1+ and SSEA-1- populations are combined compared to the unsorted Oct4-GFP+ cells, likely loss during sample handling or due to measurement error. FACS removal
of SSEA-1+ cells decreased the total cell number by a very small fraction for both 142 and 7 mmHg cells, a factor of 1.05 and 1.01, respectively (Figure 5C).

After FACS, unsorted, SSEA-1+, and SSEA-1- populations cultured at 142 or 7 mmHg were implanted into nude mice (Figure 5D). Each unsorted cell injection contained, on average, 443 and 6 Oct4-GFP+ cells at 142 and 7 mmHg, respectively, each sorted SSEA-1+ population injection contained 420 and 6 Oct4-GFP+ cells at 142 and 7 mmHg, respectively, and each sorted SSEA-1- population injection contained 13 and 0 Oct4-GFP+ cells at 142 and 7 mmHg, respectively. Unsorted cells and sorted SSEA-1+ populations produced tumors at the same time with cells cultured at 142 mmHg, whereas culture at 7 mmHg led to a delay of 4 to 12 d for the SSEA-1+ population. The 142 mmHg SSEA-1- population resulted in a 14-d delay in tumor appearance compared to unsorted cells, but the 7 mmHg SSEA-1- population resulted in no tumors during 210 d of observation.

2.4.4. Low-pO2 differentiation of HIF-1α−/− mESC does not reduce residual mESC and delay tumor formation

Since HIF-1α protein is present in PSC cultured at low pO2 and has been shown to affect differentiation [2], we differentiated HIF-1α−/− mESC for 30 d to determine what effect the absence of the HIF-1α gene would have on residual mESC and the delay in tumor appearance (Figure 6). The relative loss of Oct4 and Nanog
gene expression for HIF-1α−/− mESC at both 142 and 7 mmHg (Figure 6A) was similar to the relative loss of expression of wild type mESC at 142 mmHg (Figure 1E-F). In nude mice, undifferentiated HIF-1α−/− mESC formed tumors at d 21 (Figure 6B), 7 d later than wild type mESC (Figure 3A). Differentiated and cultured HIF-1α−/− mESC formed tumors at d 31 in all animals, regardless of culture pO_{2gas} (Figure 6B), 14 d later than wild type mESC cultured at 142 mmHg (Figure 3A). Although HIF-1α absence slowed tumor formation, the positive benefits of low pO_{2} culture were lost.

2.4.5. Residual Oct4-GFP+ mESC in vitro phenotype is the same as undifferentiated mESC

We further investigated residual Oct4-GFP+ mESC by removing Oct4-GFP+ cells from differentiated mixtures, culturing them as undifferentiated mESC for two weeks, and characterizing their in vitro phenotype. We compared three cases: (I) Oct4-GFP+ mESC that have never undergone differentiation, (II) Oct4-GFP+ cells obtained with FACS from mESC that have undergone 30-d differentiation at 142 mmHg, and (III) cells the same as case II, but with differentiation at 7 mmHg. The distribution of Oct4-GFP intensity per cell measured by flow cytometry was similar among the three cases (Figure 7A). For all three cases, essentially all of the cells were Oct4-GFP+ and, when subjected to differentiation, for the second time for cases II and III and for the first time for case I, the fraction decreased to 9×10^{-3} (data not shown). Oct4-GFP+ mESC colonies for all three cases were
morphologically similar and contained Oct4, Nanog, and SSEA-1 protein (Figure 7B-E).

All three cases expressed markers for all three germ layers upon differentiation for the second time for cases II and III and for the first time for case I (Figures 8-10). All three cases (1) expressed Brachyury T, Foxa2, Nestin, cardiac Troponin T (cTnT), cardiac-α-Actin, and Nkx2.5 mRNA (Figure 8), (2) stained positive for Foxa2, Nestin, and cTnT protein (Figure 9), and (3) contained the same fraction and number of Nestin+ and MF-20+ cells (Figure 10). For all of the in vitro assays carried out, residual Oct4-GFP+ cells sorted from differentiated populations have the same characteristics as mESC that have never been exposed to differentiation conditions.

2.4.6. Low pO₂ reduces residual hESC and delays the appearance of tumors after implantation of differentiated and cultured hESC

We extended our investigation to hESC by differentiating hESC at various pO₂gas for up to 90 d (Figure 11). Before differentiation, almost all hESC stained positive for OCT4 protein with immunocytochemistry (data not shown). During differentiation, OCT4 and NANOG gene expression decreased and was lowest at reduced pO₂ (Figure 11A-B). At 142 mmHg, OCT4 and NANOG gene expression decreased until d 30, then remained constant. At 36 and 7 mmHg, expression decreased continuously and was always lower than at 142 mmHg. After 90 d,
OCT4 expression was about 360 and 100 times lower at 36 and 7 mmHg, respectively, and NANOG expression was about 2800 and 460 times lower at 36 and 7 mmHg, respectively, compared to 142 mmHg. These results are qualitatively similar to data obtained with mESC, but the loss of OCT4 expression at all PO2gas is less extensive in differentiated hESC than in mESC (Figure 1F-G).

hESC differentiated for 30 d at different PO2gas and injected subcutaneously into nude mice formed tumors at different rates (Figure 11C). Undifferentiated hESC (positive control) formed visible tumors by d 17. Differentiated cells had slower tumor formation rates. Cells differentiated at 142 mmHg formed tumors in 2/6 animals by d 31 and 6/6 by d 38. Tumors were not observed until d 52 with cells cultured at 7 mmHg. Cells differentiated at 36 mmHg formed tumors in 2/6 animals by d 59 and 6/6 by d 66. Tumors were histopathologically identified as teratomas (data not shown). These in vivo data are consistent with the in vitro real-time PCR data and demonstrate that hypoxic culture of hESC greatly reduces the rate of teratoma formation in vivo.

2.4.7. Combination of low pO2 culture of hESC with FACS

We investigated low pO2 culture combined with FACS of hESC cell surface marker stained with a PE-conjugated anti-SSEA-4 antibody (Figure 12). hESC underwent, in a separate experimental run, a 30-d differentiation at either 142 or 36 mmHg and were sorted for SSEA-4 expression with FACS into SSEA-4+ and
SSEA-4- populations before implantation. The SSEA-4- populations had OCT4 gene expression that were a factor of 9 and 23 lower and NANOG gene expression that were a factor of 24 and 16 lower than unsorted cells from 142 and 36 mmHg, respectively, as determined by subsequent real-time PCR analysis of the sorted population (Figure 12A-B). Culture at 36 mmHg combined with FACS removal of SSEA-4+ cells resulted in a reduction in OCT4 and NANOG gene expression by a factor of $4 \times 10^3$ and $8 \times 10^3$, respectively, compared to cells culture at 142 mmHg without sorting.

After FACS, unsorted, SSEA-4+, and SSEA-4- populations cultured at 142 or 36 mmHg were implanted into nude mice (Figure 12C). Unsorted cells and sorted SSEA-4+ populations first produced tumors at the same time with cells cultured at 142 mmHg, whereas culture at 36 mmHg led to a 4-d delay for the SSEA-4+ population. The 142 mmHg SSEA-4- population resulted in a 21-d delay in tumor appearance compared to unsorted cells, but the 36 mmHg SSEA-4- population resulted in no tumors during 210 d of observation.

2.4.8. Differentiation at low pO$_2$ reduces residual miPSC

We extended our investigation to miPSC by differentiating Oct4-GFP+ miPSC at various pO$_2$gas (Figure 13). The fraction of miPSC that were Oct4-GFP+ was 0.97, assessed with flow cytometry, before differentiation (Figure 13A). This fraction decreased during the 10 d differentiation to $4 \times 10^{-2}$ at 142 mmHg and to $2 \times 10^{-4}$
and 3x10^{-4}, respectively, at 36 and 7 mmHg. The residual Oct4-GFP+ subpopulation had GFP intensity values per cell similar to that of undifferentiated miPSC (data not shown). After differentiation, culture at each constant pO2gas was extended for up to 80 additional d to further examine its effects on the reduction of Oct4-GFP+ cells. The fraction of Oct4-GFP+ cells did not change significantly at 142 mmHg but continued to decrease at 36 and 7 mmHg (Figure 13A). The greatest difference was observed at 7 mmHg after 90 d of culture, at which point the fraction of Oct4-GFP+ cells as a factor of 2.3x10^4 lower than at 142 mmHg. Though similar to data obtained with mESC, Oct4-GFP expression decreased even more dramatically at low pO_2 in miPSC (Figure 1A).

Low oxygen culture also reduced the number of Oct4-GFP+ cells (Figure 13B). Each well was initially seeded with 1.46x10^4 Oct4-GFP+ cells. After 10 d of differentiation, this number increased to 7x10^4 at 142 mmHg but decreased to 7x10^2 and 1x10^3 at 36 and 7 mmHg, respectively. During the next 20 d of culture, Oct4-GFP+ cell number decreased to 5x10^4 at 142 mmHg and did not change significantly thereafter. This reduction was proportional with the total cell number (Figure 13C). In contrast, the number of Oct4-GFP+ cells decreased continuously at 36 and 7 mmHg, resulting in only 10 and 5 Oct4-GFP+ cells per well remaining by d 90, respectively. This reduction was substantially faster than the reduction in total cell number. At all time points, the total cell number was higher at low pO2gas compared to 142 mmHg. Compared to mESC, more Oct4-GFP+ cells were present at 142 and approximately the same at 36 and 7 mmHg (Figure 1B).
Gene expression of Oct4 and Nanog, measured by real-time PCR, decreased with time in a manner comparable to that observed with flow cytometry of Oct4-GFP+ cells (Figure 13D-E). When observed with fluorescence microscopy, cells that expressed GFP after culture at 142 mmHg resided in clusters within aggregates (Figure 13F-G), albeit smaller aggregates than seen with mESC (Figure 1F and H), whereas no cells expressing GFP were visually observed anywhere in the dish, including in aggregates, at 7 mmHg (Figure 13H-I) or at 36 mmHg (data not shown).

2.4.9. Low pO2 delays appearance of tumors after implantation of cultured miPSC

pO2gas culture conditions of differentiation and extended culture affected the rate of tumor formation following subcutaneous injection of differentiated miPSC into nude mice (Figure 14A). All animals injected with undifferentiated miPSC (positive control) formed visible tumors by d 14. Differentiation and extended culture for 20 d before implantation led to slower tumor formation rates and resulted in only 4153, 1, and 0 Oct4-GFP+ cells being implanted per 10^5 cell injection for cells cultured at 142, 36, and 7 mmHg, respectively. Visible tumors appeared in 1/6 animals by d 14 and 6/6 by d 14 for cells cultured at 142 mmHg. Culture at lower pO2gas resulted in no tumors during 210 d of observation. Tumors were histopathologically identified as teratomas (data not shown). These in vivo data are consistent with the in vitro Oct4-GFP and real-time PCR data and
demonstrate that extended hypoxic culture of miPSC greatly reduces the rate of teratoma formation in vivo.

2.4.10. Number of implanted PSC correlates with the appearance of tumors

The time required for first appearance of tumors following implantation of differentiated cell populations is plotted versus the number of implanted Oct4-GFP+ mESC or hESC in each injected sample (Figure 15). The data for each species correlated exponentially, with $R^2 = 0.90$ and $R^2 = 0.99$, respectively. This data shows that our animal model is very sensitive to small numbers of and could be used in the estimation of residual pluripotent cells.

2.5. Discussion

We performed a quantitative study on the effects of low pO$_2$ on the risk of residual PSC to develop tumors. mESC, hESC, and miPSC were cultured under differentiating conditions at 142, 36, and 7 mmHg pO$_{2_{gas}}$ on highly oxygen-permeable silicone rubber dishes and assayed for residential PSC in vitro by measuring Oct4 and Nanog gene expression and, for mESC and miPSC, Oct4-GFP expression. The potential of differentiated cell populations to form tumors was assessed by subcutaneous implantation into nude mice. We also investigated the effects of low pO$_2$ culture combined with FACS removal of residual undifferentiated cells.
Low pO₂ culture during differentiation markedly decreased the fraction of PSC, measured with Oct4-GFP, and Oct4 and Nanog gene expression. The order of magnitude of this reduction ranged from $10^2$-$10^4$, depending on the duration of low pO₂ culture and the cell type (Figures 1, 11, 13). Residual Oct4-GFP+ cells expressed Oct4, Nanog, and SSEA1 protein and could be differentiated again in vitro to all three germ layers (Figures 7, 8, 9, and 10), all characteristics of PSC.

Combinations of high and low pO₂ culture could be performed and residual PSC reduced. For mESC, regardless of whether differentiation was carried out for the first 10 d at 142 or 7 mmHg, if the subsequent 20-d culture was performed at 7 mmHg, the fraction of Oct4-GFP+ cells decreased approximately by a factor of 25 compared to d 10 cells (Figure 2). Additionally, if the 20 d extended culture was performed at 142 mmHg, the fraction of Oct4-GFP+ cells either increased or stayed the same compared to d 10. These results demonstrate that differentiation can be performed at any pO₂, followed by an extended culture period at low pO₂ to reduce the fraction of Oct4-GFP+ cells without negative affecting cell number. This combination of pO₂ control strategies allows for the production of the desired differentiated cell type at the best pO₂ for producing that cell type while also reducing the fraction of potentially-tumorigenic residual PSC.
The reduction in PSC in vitro correlated with a reduction in the time required for tumor formation in vivo (Figures 3, 4, 5, 11, 12, 15). We were also able to form tumors with as few as 2 implanted mESC (Figure 2) and 100 implanted hESC (Figure 15). These values are similar to previous reports for mESC [87] and hESC [98]. The absence of tumor formation with Oct4-GFP- populations verified to contain no Oct4-GFP+ cells demonstrates that only Oct4-GFP+ mESC are likely responsible for the tumor formation in this system (Figures 4 and 5). However, if a sufficiently large number of cells had been injected, tumors would likely have formed. The possibility of other cell sources, such as multipotent progenitors, contributing to tumor or other type of undesired formation, especially with hESC, cannot be completely ruled out.

To further eliminate residual PSC, low pO2 culture was combined with FACS removal of cells based on surface marker expression. FACS removal of SSEA-1+ cells, for mESC, and SSEA-4+ cells, for hESC, reduced Oct4-GFP+ cells or OCT4 and NANOG gene expression by approximately a factor of 10 at both high and low pO2 (Figures 5 and 12). Combination of low pO2 culture with FACS reduced residual PSC more than either method alone. FACS removal of residual PSC after differentiation has been reported [101], but, to the best of our knowledge, detailed quantification of its effectiveness has not, especially in cell mixtures containing a low fraction of PSC. Low pO2 culture potentially could be used in conjunction with other selection methods, such as positive selection for
the cell type(s) of interest with FACS, selective killing of residual PSC, or
differentiation protocols that better remove residual cells.

We found that low pO₂ culture of HIF-1α⁺ mESC did not decrease residual
mESC (Figure 6), but the full mechanism of the phenomenon is not understood.
Low pO₂ could cause residual PSC to differentiate or die. In previous studies,
mESC cultured at low pO₂ display some signs of differentiation, but expression of
pluripotency genes for hESC are either not affected or increased [2]. However, in
those studies, cells were cultured under self-renewal conditions, which are very
different than the differentiation conditions used in this study. This area deserves
additional study.

We developed a model for tumor formation based on a general model of stem
cell behavior (Equation 7) that better describes the kinetics of PSC-derived tumor
formation than exponential (Equation 1) and Gompertz (Equation 2)
mathematical models (Figures 15 and 16). According to equation 7, for both
mESC and hESC implants, tumor growth progressed through two phases. At
early times, there were low numbers of differentiated cells, and the dynamics
were dominated by the behavior of PSC, and the rate of differentiation was
greater than effective cell growth. Loss of PSC occurred much faster with mESC
than hESC, consisted with in vitro published reports [116]. At later times, the
tumors consisted of mostly differentiated cells, and the kinetics were dominated
by the effective rate of cell growth of the differentiated cells. Our fitted effective
rate constant of cell growth for differentiated mESC is similar to our previously reported values for in vitro culture of differentiated mESC at low pO$_{2\text{gas}}$ [32]. Overall, our model suggests that behavior of PSC during tumor formation is similar to its behavior in vitro.

Our mathematical analysis of tumor growth has some limitations. While our general model (Equation 7) does well at predicting the time required for tumor formation within the range of implanted cells investigated, it likely does not quantitatively reflect the growth kinetics at early time points. The model could be greatly improved with the inclusion of data from early time points. For hESC, we assumed only 1% of the implanted cells survive or contribute to the tumor. Reports have observed hESC numbering in the range of $10^2$ to $10^5$ are necessary to form a tumor [98, 100] while as few as 2 mESC are required [87]. Understanding the critical number PSC required to form a tumor is worthy of further investigation and is extraordinarily valuable to fully assess the safety risk of any PSC therapy.

Our work shows that in vitro low pO$_2$ culture markedly reduces the fraction of cells that are pluripotent, as measured by Oct4-GFP and Oct4 and Nanog gene expression, in differentiated mESC, hESC, and miPSC populations. The reduction correlates with reduction in the time required for tumor formation and, when all PSC are removed from differentiated populations, tumors do not form. After differentiation, sorted residual Oct4-GFP+ mESC express Oct4, Nanog, and
SSEA-1 protein and can be differentiated again, all characteristics of undifferentiated mESC. We believe that low pO$_2$ culture alone or in combination with other methods is a potentially straightforward method that could be applied to future cell therapy protocols to minimize the possibility of tumor formation.
2.6. Tables and Figures
Table 2-1 Comparison of changes in pO$_{2}$gas on total cell number and number of MF-20+ cells

<table>
<thead>
<tr>
<th>pO$_{2}$gas (mmHg)</th>
<th>Ascorbic</th>
<th>Total cell number (x10$^8$)</th>
<th>Number MF-20+ cells (x10$^5$)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Acid</td>
<td>Time (days)$^a$</td>
<td>Change (%)</td>
</tr>
<tr>
<td>142</td>
<td>No</td>
<td>0</td>
<td>1.8±0.1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>20</td>
<td>1.4±0.1</td>
</tr>
<tr>
<td>142-7$^b$</td>
<td>No</td>
<td>0</td>
<td>1.8±0.1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>20</td>
<td>1.4±0.1</td>
</tr>
<tr>
<td>7-142-7$^c$</td>
<td>Yes</td>
<td>0</td>
<td>2.1±0.4</td>
</tr>
<tr>
<td></td>
<td></td>
<td>20</td>
<td>3.2±0.3</td>
</tr>
</tbody>
</table>

$^a$ Time in extended culture

$^b$ Differentiation 10 d at 142 mmHg, then 20 d extended culture at 7 mmHg

$^c$ Differentiation 6 d at 7 mmHg followed by 4 d at 142 mmHg, then 20 d extended culture at 7 mmHg
Figure 2-1. Expression of pluripotency markers in mESC having undergone differentiation and extended culture at various pO$_{2_{gas}}$. mESC were differentiated at 142, 36, or 7 mmHg pO$_{2_{gas}}$ then subjected to extended culture at the same pO$_{2_{gas}}$ used during differentiation for a total of 0, 10, 20, 30, or 90 days. (A) Fraction of cells that are Oct4-GFP+ as measured with flow cytometry (n=3). (B) The number of Oct4-GFP+ cells per well determined by
multiplying the fraction of Oct4-GFP+ cells and the total cell number (n=3). (C)
The total cell number per well as measured by nuclei count (n=3). (D-E) Relative
expression of Oct4 and Nanog mRNA measured with real-time PCR (n=3). (F
and H) En face bright field images of cell aggregates after 30 days culture. (G
and I) En face fluorescence images taken with a GFP filter corresponding with (F)
and (H), respectively.
Figure 2-2. Expression of Oct4-GFP in mESC having undergone differentiation and extended culture at the same or different pO$_2$gas.
mESC were differentiated at 142 (A, C, E) or 7 (B, D, F) mmHg pO$_{2\text{gas}}$ then subjected to extended culture at the same (black bar) or the other (white bar) pO$_{2\text{gas}}$ used during differentiation. (A and B) Fraction of cells that are Oct4-GFP+ as measured with flow cytometry (n=3). (C and D) The number of Oct4-GFP+ cells per well determined by multiplying the fraction of Oct4-GFP+ cells and the total cell number (n=3). (E and F) The total cell number per well as measured by nuclei count (n=3). * indicates significant difference compared to the black bar of the same time.
Figure 2-3. Time-course development of tumors in nude mice after subcutaneous implantation of mESC.
(A) Formation of tumors after implantation of undifferentiated mESC (n=6; positive control), mESC differentiated 10 days at 142, 36, or 7 mmHg followed by 20 days extended culture at the same pO$_{2\text{gas}}$ (n=8) or only Matrigel (n=6; negative control). Tumor formation was determined by visual observation of the appearance of masses underneath the skin. (B) Representative H&E-stained tissue section of a tumor derived from mESC differentiated and having undergone 20 days of extended culture at 7 mmHg pO$_{2\text{gas}}$, with (i) small glandular structure, (ii) hyaline cartilage, and (iii) neuroepithelial structures highlighted. All animals with no tumors were monitored for a total of 210 days after implantation, during which no tumors were observed.
Figure 2-4. Expression of Oct4-GFP and time-course development of tumors in nude mice after subcutaneous implantation of differentiated and cultured mESC sorted into GFP+ and GFP- populations. mESC underwent 10-day differentiation and 20-day extended culture at either 142 or 7 mmHg pO$_{2}$gas and cells before sorting (unsorted) and after being sorted into Oct4-GFP+ (GFP+) and Oct4-GFP- (GFP-) populations were analyzed. (A) Fraction of mESC that are Oct4-GFP+ as measured with flow cytometry (n=3). (B) The number of Oct4-GFP+ cells per well determined by multiplying the fraction of Oct4-GFP+ cells and the total cell number (n=3). (C) The total cell number per well as measured by nuclei count (n=3). (D) A sample of 10$^5$ cells were implanted
without sorting (unsorted; n=6) or $10^5$ cells were sorted into GFP+ and GFP-
populations (n=6), then implanted.
Figure 2-5. Expression of Oct4-GFP and time-course development of tumors in nude mice after subcutaneous implantation of differentiated and cultured mESC sorted into SSEA-1+ and SSEA-1- populations.

mESC underwent 10-day differentiation and 20-day extended culture at either 142 or 7 mmHg pO2gas, and cells before sorting (unsorted) and after being sorted into SSEA-1+ and SSEA-1- populations were analyzed. (A) Fraction of mESC that are Oct4-GFP+ as measured with flow cytometry (n=3). (B) The number of Oct4-GFP+ cells per well determined by multiplying the fraction of Oct4-GFP+ cells and the total cell number (n=3). (C) The total cell number per well as measured by nuclei count (n=3). (D) A sample of $10^5$ cells were implanted.
without sorting (unsorted; n=6) or $10^5$ cells were sorted into SSEA-1+ and SSEA-1- populations (n=6), then implanted.
Figure 2-6. Expression of pluripotency markers and time-course development of tumors in nude mice with HIF-1α<sup>c</sup> mESC cells differentiated 10 days at 142 or 7 mmHg followed by 20-days extended culture at the same pO<sub>2</sub>gas.

(A) Relative expression of Oct4 and Nanog mRNA measured with real-time PCR (n=3). (B) Formation of tumors after implantation of undifferentiated HIF-1α<sup>c</sup> mESC (n=6; positive control) and HIF-1α<sup>c</sup> mESC differentiated and cultured at 142 or 7 mmHg (n=6).
Figure 2-7. Characterization of (I) Oct4-GFP+ mESC that have never undergone differentiation and extended culture, (II) Oct4-GFP+ cells sorted with FACS from mESC that have undergone 10 days differentiation and 20 days extended culture.
at 142 mmHg, and (III) Oct4-GFP+ cells cultured similar to case II, but at 7 mmHg.

(A) Histogram of the distribution of Oct4-GFP fluorescence intensity per cell, determined with flow cytometry. (B-E) En face images of colonies from cell population II. (B) Bright field image. (C-E) Cells stained with DAPI (blue) and immunostained (red) for (C) Oct4, (D) Nanog, and (E) SSEA-1.
Figure 2-8. Differentiation of cases I, II, and III and assessment with real-time PCR.

Expression of (A) Brachyury T, (B) Foxa2, and (C) Nestin after 4 days of differentiation and (D) cTnT, (E) cardiac-α-Actin, and (F) Nkx2.5 after 10 days of differentiation (n=3). Expression is relative to case I before differentiation.
Figure 2-9. Differentiation of cases I, II, and III and assessment with immunocytochemistry.

En face images cells after (A-D) 4 days and (E-F) 10 days of differentiation stained with DAPI (A, C, E) and immunostained for (B) Foxa2, (D) Nestin, and (F) cTnT.
Figure 2-10. Differentiation of cases I, II, and III and assessment with flow cytometry.

(A) Fraction and (B) number of Nestin+ cells after 4 days of differentiation. (C) Fraction and (D) number of MF-20+ cells after 10 days of differentiation. (n=3).
Figure 2-11. Expression of pluripotency markers and time-course development of tumors of hESC cultured under differentiation conditions at 142, 36, or 7 mmHg pO$_{2\text{gas}}$ up to 90 days.

(A-B) Relative expression of OCT4 and NANOG mRNA measured with real-time PCR (n=3). (C) Formation of tumors after subcutaneous implantation of
undifferentiated hESC (positive control; n=6) and hESC differentiated 30 days at 142, 36, or 7 mmHg (n=6).
Figure 2-12. Gene expression of OCT4 and NANOG and time-course development of tumors in nude mice after subcutaneous implantation of differentiated and cultured hESC sorted into SSEA-4+ and SSEA-4- populations. hESC underwent 30-day differentiation at either 142 or 36 mmHg pO$_{2\text{gas}}$, and cells before sorting (unsorted) and after being sorted into SSEA-4+ and SSEA-4- populations were analyzed. (A-B) Relative expression of OCT4 and NANOG
mRNA measured with real-time PCR (n=3). (C) A sample of $10^6$ cells were implanted without sorting (unsorted; n=6) or $10^6$ cells were sorted into SSEA-4+ and SSEA-4- populations (n=6), then implanted.
Figure 2-13. Expression of pluripotency markers in miPSC having undergone differentiation and extended culture at various pO$_{2gas}$.

miPSC were differentiated at 142, 36, or 7 mmHg pO$_{2gas}$ then subjected to extended culture at the same pO$_{2gas}$ used during differentiation for a total of 0, 10, 20, 30, or 90 days. (A) Fraction of cells that are Oct4-GFP+ as measured with flow cytometry (n=3). (B) The number of Oct4-GFP+ cells per well determined...
multiplying the fraction of Oct4-GFP+ cells and the total cell number (n=3). (C) The total cell number per well as measured by nuclei count (n=3). (D-E) Relative expression of Oct4 and Nanog mRNA measured with real-time PCR (n=3). (F and H) En face bright field images of cell aggregates after 30 days culture. (G and I) En face fluorescence images taken with a GFP filter corresponding with (F) and (H), respectively.
Figure 2-14. Time-course development of tumors in nude mice after subcutaneous implantation of miPSC.

Formation of tumors after implantation of undifferentiated miPSC (n=6; positive control), mipSC differentiated 10 days at 142, 36, or 7 mmHg followed by 20 days extended culture at the same $pO_2_{gas}$ (n=6). Tumor formation was determined by visual observation of the appearance of masses underneath the skin.
Figure 2-15. The time required for tumor appearance after implantation of different numbers mESC and hESC.

The number of implanted residual Oct4-GFP mESC, having undergone differentiation and extended culture under various conditions (Figures 1A, 4A, 5A), is plotted against the corresponding tumor formation data (Figures 3A, 4D, and 5D) (circles; n=79). $10^6$, $10^4$, and $10^2$ undifferentiated hESC were implanted and the appearance of tumors observed (squares; n=6). The mouse (solid line) and human (dashed line) data fit the form $N_0 = Ae^{Bt}$, where $N_0$ is the number of implanted pluripotent cells, t is time, and $A = 1.22 \times 10^4$ and $3.03 \times 10^8$ and $B = -0.209$ and -0.278, respectively.
Chapter 3. Differentiation of Pluripotent Stem Cells
Under Low Oxygen Increases Generation of
Cardiomyocytes and Influences Differentiation to
Ectoderm, Mesoderm, and Endoderm
3.1. Abstract

Pluripotent stem cells (PSC) hold promise for the study of embryonic development and cell replacement therapy, but efficient differentiation to desired cell types remains a major obstacle. Most PSC research is performed in incubators with high, non-physiological partial pressures of oxygen (pO₂). However, embryonic cells in early development are exposed to much lower local pO₂, and the effects of such conditions on differentiating PSC are poorly understood. Here, low pO₂ drastically affects differentiation of mouse and human embryonic stem cells and mouse induced pluripotent stem cells to the three germ layers and cardiomyocytes. Overall, differentiation was higher to endoderm, lower to ectoderm, and higher or the same to mesoderm. Differentiation to cardiomyocytes was greatly enhanced, with our best protocol resulting in 35% cardiomyocytes and 60 cardiomyocytes being generated per input mouse embryonic stem cell without the need for purification. Understanding the effects of pO₂gas during differentiation is an important step towards the development of protocols for regenerative medicine.
3.2. Introduction

Pluripotent stem cells (PSC), including embryonic stem cells (ESC) and induced pluripotent stem cells (iPSC), have the potential to differentiate into any cell type found in a mature organism. The clinical potential for such PSC is immense [3], including treatments for heart disease, diabetes, Parkinson's disease, and leukemia [4-6]. For example, PSC could be differentiated to cardiomyocytes and implanted into infarcted hearts to potentially replace necrotic tissue after a heart attack [117]. However, before PSC-based therapies can be used effectively in clinical applications, the ability to generate the desired differentiated cells in sufficient purity and quantity for use in humans must be developed [118].

Differentiation of PSC to the three germ layers in vitro is often induced with soluble signaling molecules. High and low levels of nodal signaling specify endoderm and mesoderm, respectively, while FGF signaling specifies ectoderm [119]. Numerous molecules can be used to further direct cardiomyocyte differentiation, including vascular endothelial growth factor, dickkopf homolog 1, ascorbic acid, and 5-azacytidine [120-123].

Low partial pressure of oxygen (pO₂) has been shown to increase PSC differentiation to derivatives of the ectodermal and mesodermal lineages, such as neurons, cardiomyocytes, hematopoietic progenitors, endothelial cells, and chondrocytes in separate studies [2]. In most studies, the pO₂ experienced by the
cell (pO$_2_{\text{cell}}$) is not controlled or understood [53]. Low pO$_2$ is generally thought to affect cells by decreasing exposure to reactive species generated by cellular respiration [124] and by regulating the stability of the oxygen-responsive transcription factor hypoxia inducible factor-1 (HIF-1) [125], but other mechanisms may also be involved [126].

Even though pO$_2$ is an important culture parameter, PSC are usually cultured at relatively high pO$_2$, typically in a humidified atmosphere consisting of 95% air/5% CO$_2$, resulting in a gas phase pO$_2$ (pO$_2_{\text{gas}}$) of 142 mmHg. This value is higher than that found in most mammalian tissues and in utero, such as rat microvasculature (25 mmHg) [20], rhesus monkey reproductive tract (15-45 mmHg) [127], and fetal human venous blood (25-30 mmHg) [21, 22]. The pO$_2$ in early developing embryos is likely very low because circulation is not yet occurring [128, 129].

In this study, we quantitatively investigated the effects of pO$_2$ on the differentiation of mouse ESC (mESC) to the three germ layers and mESC, mouse iPSC (miPSC), and human ESC (hESC) to cardiomyocytes. Cells were cultured on highly oxygen permeable silicone rubber membrane plates, which allowed us to set pO$_2_{\text{cell}}$ at the membrane-cell interface virtually to the value of pO$_2_{\text{gas}}$. Overall, differentiation of mESC at low pO$_2$ increased endoderm, decreased ectoderm, and either increased or didn’t change mesoderm differentiation compared to 142 mmHg. The expression of many differentiation
genes was delayed at low compared to high pO₂. Constant low pO₂ enhanced
differentiation of mESC, miPSC, and hESC to cardiomyocytes, compared to 142
mmHg without the need for purification and additional soluble molecules beyond
serum, but the largest cardiomyocyte yields occurred with temporal variation of
pO₂gas. pO₂cell of cardiomyocytes within resulting cellular aggregates after
differentiation was calculated.

3.3. Methods

3.3.1. Culture of undifferentiated mESC and miPSC

Undifferentiated J1 mESC (American Type Culture Collection (ATCC),
Manassas, VA, USA), miPSC [113] and mESC harboring a Sox17-dTomato
reporter [130] (Sox17-dTomato mESC) (both generously donated by Professor
Douglas Melton of Harvard University, Cambridge, MA, USA), mESC with a
homozygous HIF-1α, a subunit of HIF-1, gene knockout (HIF-1α−/− mESC) [114]
generously donated by Professor Peter Carmeliet of Katholieke Universiteit
Leuven, Leuven, Belgium), and mESC harboring a Brachyury T-GFP reporter (T-
GFP mESC) [131] (generously donated by Professor Gordon Keller of McEwen
Centre for Regenerative Medicine, Toronto, ON, Canada) were cultured without
mouse embryonic fibroblasts as previously described [53] in Dulbecco’s Modified
Eagles Medium (DMEM; SCRR-2010; ATCC) supplemented with 10% (v/v) fetal
bovine serum (FBS; SCRR 30-2020; ATCC), 1x L-alanyl-L-glutamine (SCRR 20-
2115; ATCC), 1x penicillin streptomycin (P/S; 15070; Invitrogen, Carlsbad, CA, USA), 1x MEM non-essential amino acids (SCRR 20-2116; ATCC), 100 µM 2-mercaptoethanol (M7522; Sigma-Aldrich; St. Louis, MO, USA), and 10³ units/mL leukemia inhibitory factor (LIF; LIF2005; Millipore, Billerica, MA, USA). Cells were passaged by treatment with 0.25% trypsin (25-053-Cl; Mediatech, Manassas, VA, USA) every two days and seeded at 1.2x10⁴ cells cm⁻² on cell culture flasks (353014; Becton Dickinson, Franklin Lakes, NJ, USA).

3.3.2. Control of pO₂gas

pO₂gas was controlled by placing cells inside chambers maintained at 37 °C with premixed gas flown in containing 5% CO₂ and 20%, 5%, or 1% O₂ and a balance of N₂ (certified medical gas from Airgas, Hingham, MA, USA), which corresponded, under humidified conditions, to 142, 36, and 7 mmHg pO₂gas, respectively, as previously described [53].

3.3.3. Silicone rubber membrane-based plate

Differentiation was performed on custom-made highly oxygen permeable silicone rubber plates, as previously described [53]. In brief, 24-well polystyrene plate bottoms (353047; Becton Dickinson) were replaced with silicone rubber sheets (nonreinforced vulcanized gloss/gloss 127 µm thick; Specialty Manufacturing, Saginaw, MI, USA) and sterilized by 1 hr treatment with 70% (v/v) ethanol
(111000200; Pharmco-AAPER, Brookfield, CT, USA) in water and overnight drying under a biological safety cabinet germicidal UV lamp (SterilGARD III Advance; The Baker Company, Sanford, MA, USA). When in use for cell culture, a silicone rubber plate sat on top of a 150x15 mm Petri dish (35384-326; VWR, West Chester, PA, USA) with only the plate corners being supported by the Petri dish, allowing the silicone rubber membrane to be directly exposed to the gas phase. Silicone rubber membranes maintained pO2cell at the cell-membrane interface at virtually the same value as pO2gas, and allowed for rapid equilibration of pO2cell after a change in pO2gas. Treatment of silicone rubber membranes at 37 °C with 2 μg/mL fibronectin (F1 141; Sigma-Aldrich) in phosphate buffered saline (PBS; 21-040; Mediatech) for 24 hr for mESC and miPSC or Matrigel (354234, Becton Dickinson) diluted 1:20 in serum-free medium for 1 hr for hESC permitted cell attachment.

3.3.4. Differentiation of mESC and miPSC

mESC and miPSC differentiation was performed as previously described [53]. In brief, embryoid bodies (EBs) were formed in 20-μL hanging drops, consisting of 2.5x10^4 cells ml⁻¹ in propagation medium without LIF. After 2 days, 30 EBs per well were transferred to fibronectin-coated silicone rubber wells filled with 2 mL of propagation medium without LIF, which was replaced with 2.5 mL of serum-free medium after 3 additional days, consisting of 50% (v/v) DMEM (90-133-PB; Mediatech) and 50% (v/v) Ham’s F-12 (10-080-CV; Mediatech) supplemented
with 0.75% (w/v) sodium bicarbonate (25-035-CI; Mediatech), 9 mM glucose (G8769; Sigma-Aldrich), 5 µg/mL human insulin (I9278; Sigma-Aldrich), 50 µg/mL holo transferrin (T1283; Sigma-Aldrich), 31.2 nM sodium selenite (S9133; Sigma-Aldrich), and 1x P/S. Cells were differentiated for up to a total of 13 days. 

$pO_{2gas}$ was kept constant throughout differentiation in some cases and in others, was varied, as noted. In addition, 0.2 mM ascorbic acid (A4034; Sigma-Aldrich) was also included in culture medium in some cases, as noted. Cell samples at various time points were acquired for analysis by dissociation with a 5-min exposure to 0.25% trypsin at 37 °C.

3.3.5. Real-time polymerase chain reaction (PCR)

Total RNA was extracted with the RNeasy Kit (74104, Qiagen, Valencia, CA) and RNase-Free DNase Set (79254, Qiagen), cDNA created using High Capacity cDNA Reverse Transcription Kit (4368814, Applied Biosystems, Foster City, CA), real-time PCR conducted on a Fast Real-Time PCR System (7900HT, Applied Biosystems) at the Center for Biomedical Engineering (MIT) using Power SYBR Green PCR Master Mix (4367659, Applied Biosystems) or Taqman Gene Expression Master Mix (4369016, Applied Biosystems), as appropriate, and 28S ribosomal RNA (rRNA) was measured for an endogenous control. Primer sequences used for 28S rRNA, Sox17, Nestin, and Nkx2.5 have been previously reported [32]. Additional primers sequences used (in order of forward and reverse primer and 5' to 3') were Foxa2 TCAAGGCCTACGAACAGGTCAT and
GCCCGCTTTTGTTCGTGACT, Hnf4a CAGACGTCCCTCCTTTTCTTGTGATA and TGTTTGGTGTTGAGGTAGTATGATA, Pdx1 CGCGTCCAGCTCCCTTTT and CCTGCCACTGGCCTTT, Sox1 TGCACAGGTCGCCCTGAGT and GCACAAAAACTTGCCCAAGAG, Pax6 TGGAAACAAACGCCCTAGCT and GGACAGGAACACACCAACTTT, Brachyury T TCCCCGTGCTGAAGGTAAAT and CCGTCACGAAGTCCGCAA, Mesp2 TTTGGGCTGCCTTTGGAAGT and AGCTAAGCAGACCTCAAATGTCA, Tbx20 CCAGCGAAGAGATGGCTAAAA and TGTCCTAGCTCTCCTTGTT, Mef2c TGCTGGTCACCTGTGTAAC and ATCCTTTGATTCACTGATGGCAT, cardiac-α-Actin
GCTTCCGCTGTCCAGAGACT and TGCCAGCAGATTCCATACCA, and cardiac Troponin T (cTnT) AGATGCTGAGAGGTCCAGTAGAG and CACCAAGTTGGGCATGAAGAGA. Premade primers for Mesp1 (Mm00801683_g1, Applied Biosystems), Tbx5 (Mm00803518_m1, Applied Biosystems), Titin (Mm00621005_m1, Applied Biosystems), human cTNT (Hs00165960_m1, Applied Biosystems), human MEF2C (Hs00231149_m1, Applied Biosystems), and human NKKX2.5 (Hs00231763_m1, Applied Biosystems) were also used. A standard curve was constructed with pooled cDNA from all time points and PO2gas serial diluted.

3.3.6. Immunocytochemistry

Attached cells were fixed with 4% (w/v) paraformaldehyde (36606; Alfa Aesar, Ward Hill, MA, USA) in PBS for 20 min, permeabilized and non-specific protein
binding blocked in a PBS solution containing 3% (v/v) donkey serum (DS; D9663; Sigma-Aldrich) and 0.1% (v/v) Triton X-100 (T9284; Sigma-Aldrich) for 45 min, and incubated overnight at 4 °C with primary antibodies against Foxa2 (sc-6554; Santa Cruz Biotechnology, Santa Cruz, CA, USA) diluted 1:200, Nestin (MAB2736; R&D Systems, Minneapolis, MN, USA) diluted 1:50, cTnT (MS-295-P1; NeoMarkers, Fremont, CA, USA) diluted 1:50, or Nkx2.5 (sc-8697; Santa Cruz Biotechnology) in 3% DS 0.1% Triton X-100 solution. After washing 1x with 3% DS 0.1% Triton X-100 solution, cells were incubated for 2 hr at room temperature in the dark with Alexa Fluor 488 (green) or 594 (red) secondary antibodies (Invitrogen, Carlsbad, CA, USA) against the appropriate primary antibody species diluted 1:200 in 3% DS 0.1% Triton X-100 solution. After washing 1x, cells were incubated with 1 μg/mL 4’,6-diamidino-2-phenylindole (DAPI; 32670, Sigma-Aldrich) in PBS for 10 min in the dark to stain DNA (blue), after which cells were then washed 3x with PBS and visualized with an Axiovert 200 fluorescence microscope (Carl Zeiss, Oberkochen, Germany).

### 3.3.7. Flow cytometric analysis of immunostained cells

The fraction of Nestin+ and MF-20+ cells in dispersed cells was acquired with flow cytometry, as previously described [53]. In brief, samples were fixed by 20 min incubation in 1% (w/v) paraformaldehyde in PBS. Fixed cells numbering $3 \times 10^5$ were permeabilized in 0.5% saponin (S-4521; Sigma-Aldrich) for 10 min, non-specific binding blocked 30-min incubation in 2% (v/v) FBS in PBS, and
incubated at room temperature with primary antibody against Nestin diluted 1:50 or sarcomeric myosin heavy chain (MF-20; MF-20 supernatant; Developmental Studies Hybridoma Bank, Iowa City, IA, USA) diluted 1:10 in 2% FBS solution for 1 hour. Samples were then incubated with goat anti-mouse phycoerythrin-conjugated secondary antibody (115-116-146; Jackson ImmunoResearch; West Grove, PA) diluted 1:250 in 2% FBS solution for 30 min at room temperature in the dark, washed 3x in 2% FBS solution, and staining data were measured using a Guava PCA flow cytometer (Millipore).

3.3.8. Nuclei enumeration

Dispersed cells were exposed to a lysis solution of 1% (v/v) Triton X-100 and 0.1 M citric acid (0627; Mallinckrodt Specialty Chemicals Co., Paris, KY, USA) and vortexed to obtain nuclei samples. Samples were stained with Guava Viacount (4000-0041; Millipore) and data acquired with a Guava PCA flow cytometer, as previously described [53].

3.3.9. Flow cytometric analysis of T-GFP expression

T-GFP expression was acquired with a flow cytometer (FACScan; Becton Dickinson) at the Koch Institute (MIT). Gating was set so that all of a negative control, J1 mESC without T-GFP differentiated under the same conditions as the T-GFP mESC, were counted as negative for T-GFP, and the fraction of cells
expressing T-GFP (T-GFP+) was calculated as the number of cells above this gate divided by the total number of cells at each time and condition.

3.3.10. Immunostaining of histological sections

Histological specimens obtained from cells at differentiation day 11 were fixed with 4% paraformaldehyde overnight and subsequently embedded in paraffin and sectioned by the Joslin Diabetes Center histology core to yield 5-μm sections. Samples were deparaffinized and rehydrated by 7 min rinses with xylene (8671; Mallinckrodt) (twice), 5 min rinses with 100% ethanol (twice), a 3 min rinse with 95% ethanol, a 10 min rinse with 70% ethanol, and 5 min rinses with deionized water (25-055-CM; Mediatech) (twice). Antigen retrieval was performed by boiling sections for 10 min in a 10 mM sodium citrate (0754; Mallinckrodt) solution at pH 6. Endogenous peroxidase activity was quenched by incubating the slides at room temperature for 10 min in 0.3% hydrogen peroxide (386790; Calbiochem, La Jolla, CA, USA), and slides, blocked with 1 hr incubation in 1% FBS. Samples were incubated with MF-20 (1:10) or anti-cTnT primary antibody overnight at 4 °C and, after 3 washes, were incubated with goat anti-mouse IgG (115-035-062; Jackson ImmunoResearch) diluted 1:50 in 1% FBS for 3 hr at room temperature. After 3 additional washes, samples were incubated with mouse peroxidase-anti-peroxidase complex (223-005-024; Jackson ImmunoResearch) diluted 1:500 in 1% FBS for 1 hr at room temperature. Visualized was achieved by incubating 2 min in 2 mM diaminobenzidine (D5637; Sigma-Aldrich) containing 0.015%
hydrogen peroxide, and slides were counterstained by incubating 15 sec in haematoxylin (HHS16; Sigma-Aldrich), followed by 3 washes in 30 mM sodium borate (B10267-34; EMD Chemicals, Gibbstown, NJ, USA). Slides were then dehydrated following the reverse procedure for rehydration and preserved using Permount (SP15; Fisher Chemical, Fair Lawn, NJ, USA).

3.3.11. Semi-quantitative assessments

The fractional surface coverage of spontaneously-contracting cells within 15 randomly arranged 2 mm² circles per well was estimated by visual observation and the average of which taken to be the fractional surface coverage of cells for that well, as previously described [53]. Histological sections of J1 mESC differentiated for 11 days with 0.2 mM ascorbic acid at 142, 36, and 7 mmHg pO₂ gas immunostained with MF-20 were manually counted for the number of MF-20+ and MF-20- cells.

3.3.12. Calculated volumetric pO₂ cell within MF-20+ regions

We previously reported calculating the pO₂ cell distribution within aggregates [53]. Calculations similar to those were used to estimate the pO₂ cell for cells positively immunostained with MF-20. For these calculations, the aggregate dimensions were measured, and MF-20+ regions were identified in the 5-μm tissue sections. The finite element model for oxygen transport was used to predict the pO₂ profile
within the entire aggregate, and the integration to determine the volume of tissue between surfaces of constant pO$_2$ was only performed for simulated MF-20+ regions. The volume fraction of tissue in MF-20+ regions within specific limits of pO$_2$ was determined by:

$$
\Phi_{ab,\text{MF20}} = \frac{\sum_{j=1}^{n} V_{ab,j,\text{MF20}}}{\sum_{j=1}^{n} V_{\text{aggregate},j,\text{MF20}}},
$$

(1)

where $V_{ab,j,\text{MF20}}$ is the volume of tissue with a pO$_2$ between $a$ and $b$ in the $j^{th}$ MF-20+ region and $V_{\text{aggregate},j,\text{MF20}}$ is the total volume of the $j^{th}$ MF-20+ region.

Because no MF-20+ cells were observed in the sheet-like morphology, $\phi_{\text{sheet}}$ was set to 0 and does not appear in Eq. 1.

### 3.3.13. Differentiation of hESC

EBs were formed with 500 CyT49 hESC [9] (NIH Registration Number 0041; generously provided by ViaCyte, Inc, San Diego, CA, USA) per microwell of an AggreWell plate (27845; Stemcell Technologies, Vancouver, BC, Canada) for 1 day, transferred to suspension culture on silicone rubber plates, previously treated with 1% (w/v) pluronic F-127 (P2443; Sigma-Aldrich) to prevent attachment for 7 days, and 30 EBs transferred to each well of a Matrigel-coated 24-well silicone rubber plate for attachment culture or 12 days. Differentiation was performed entirely in mESC and miPSC propagation medium without LIF with 0.2 mM ascorbic acid.
3.3.14. Statistics

Statistical analysis was performed with unpaired two-tailed t-tests, with \( p<0.05 \) being considered statistically significant. All data presented as mean ± standard deviation of biological replicates.

3.4. Results

3.4.1. The effect of PO2 on endodermal differentiation of mESC

We differentiated J1 mESC at various, constant PO2gas to examine differentiation to the endodermal lineage (Figure 1). Different PO2gas values changed the timing and magnitude of endodermal gene expression, measured with real-time PCR. Expression overall was higher at low compared to high PO2gas. Highest Foxa2 expression, an early endoderm marker [132], occurred on days 4, 7, and 13 for 142, 36, and 7 mmHg, respectively, with expression at 36 mmHg being 2.7 times higher than and at 7 mmHg being statistically the same as 142 mmHg (Figure 1A). Highest Sox17 expression, an early endoderm marker [133], occurred on days 8, 8, and 9 for 142, 36, and 7 mmHg, respectively, with expression at 36 mmHg being 1.9 times higher than and at 7 mmHg being statistically the same as 142 mmHg (Figure 1B). Highest Hnf4a expression, a primitive gut tube marker [134], occurred on days 7, 7, and 13 for 142, 36, and 7 mmHg, respectively, with
expression at 36 mmHg being 2.3 times higher than and at 7 mmHg being 3.0 times higher than as 142 mmHg (Figure 1C). Highest Pdx1 expression, a posterior foregut marker [135], occurred on days 12, 10, and 12 for 142, 36, and 7 mmHg, respectively, with expression at 36 mmHg being 4.5 times higher than and at 7 mmHg being 4.1 times higher than as 142 mmHg (Figure 1D).

More Foxa2+ and Sox17-dTomato+ cells were observed with microscopy at low compared to high pO$_2$gas (Figure 1E-H), qualitatively correlating with our real-time PCR data (Figure 1A-B). However, with this protocol, the fraction of Foxa2+ and Sox17-dTomato+ cells was less than 1%, assessed with visual observation, at all pO$_2$gas, and we were unable to accurately quantitate the numbers with flow cytometry.

Different pO$_2$gas values slowed or delayed endodermal gene expression, as in expression was higher at some pO$_2$gas at earlier times and higher at other pO$_2$gas at later times. Gene expression was delayed 1-2 days for Sox17 and Hnf4a at 36 mmHg and for Foxa2, Sox17, and Hnf4a at 7 mmHg, compared to 142 mmHg (Figure 1A-C). However, the trend with Pdx1 was different, with expression being first significantly higher than undifferentiated mESC on day 9 for 36 mmHg, day 10 for 142 mmHg, and day 12 for 7 mmHg (Figure 1D). We are uncertain of the cause of the difference between Pdx1 and the earlier markers.
3.4.2. The effect of pO$_2$ on ectodermal differentiation of mESC

We differentiated J1 mESC at various, constant pO$_2$gas to examine differentiation to the ectodermal lineage (Figure 2). Expression overall was higher at high compared to low pO$_2$gas. Highest Sox1 expression, an early ectodermal marker [136], occurred on days 13, 7, and 7 for 142, 36, and 7 mmHg, respectively, with expression at 142 mmHg being 3.2 and 8.6 times higher than 36 and 7 mmHg, respectively (Figure 2A). Highest Nestin expression, a neural stem cell marker [137], occurred on days 12, 4, and 4 for 142, 36, and 7 mmHg, respectively, with expression at 142 mmHg being 1.4 times higher than at 7 mmHg and statistically the same as 36 mmHg (Figure 2B). Highest Pax6 expression, a neurogenic marker [138], occurred on days 13, 12, and 12 for 142, 36, and 7 mmHg, respectively, with expression at 142 mmHg being 7.8 and 4.3 times higher than at 36 and 7 mmHg, respectively (Figure 2C). The fraction and number of Nestin+ cells was also lower with differentiation at low pO$_2$gas (Figure 2D-E), correlating with our real-time PCR data. After 12 days of differentiation, the fraction of Nestin+ cells was 32, 20, and 8.6% for 142, 36, and 7 mmHg, resulting in 95, 60, and 18 Nestin+ cells generated per each input mESC. More Nestin+ cells were also observed with microscopy at high compared to low pO$_2$gas (Figure 2F-G).
3.4.3. The effect of pO$_2$ on mesodermal differentiation of mESC

We differentiated J1 mESC at various, constant pO$_2$gas to examine mesodermal differentiation (Figure 3). pO$_2$ changed the timing and magnitude of mesodermal gene expression. Expression of Brachyury T, an early mesoderm marker [139], was higher at 142 mmHg on days 3 and 4 compared to 36 and 7 mmHg. Maximal Brachyury T expression was the same between 142 and 36 mmHg and slightly higher at 7 compared to 142 mmHg. The first large increase in gene expression occurred with Mesp1 and Mesp2, cardiac mesoderm markers [140], being a factor of 2 higher at 36 and 7 mmHg compared to 142 mmHg for Mesp1 and a factor of 3 and 2 higher at 36 and 7 mmHg, respectively, compared to 142 mmHg for Mesp2 on day 5.

Because of the large increase in Mesp1 and Mesp2 expression at low pO$_2$gas, we focused on differentiation to cardiomyocytes. Differentiation at 7 mmHg resulted in a delay in the expression of Tbx20, Tbx5, Nkx2.5, Mef2c, Foxh1, cardiac-α-Actin, cTnT, and Titin, cardiac markers [141-145], compared to 142 mmHg. Differentiation at 36 mmHg only delayed Foxh1 expression. After days 7-8, expression was higher at most time points at 7 compared to 142 mmHg for Tbx20, Tbx5, Nkx2.5, Mef2c, and Foxh1, and after day 9 for cardiac-α-Actin, cTnT, and Titin. Expression at 36 mmHg for most times was lower than 7 and the same or higher than 142 mmHg.
Differentiating T-GFP and J1 mESC at various constant pO$_{2\text{gas}}$ delayed the appearance of large numbers of T-GFP+ and MF-20+ cells and increased the generation of MF-20+ cells (Figure 4). Maximal fraction of T-GFP+ cells occurred on days 4, 5, and 6 at 142, 36, and 7 mmHg, respectively but was approximately 70% at all pO$_{2\text{gas}}$ (Figure 4A). The appearance of MF-20+ cells from J1 mESC was delayed by 2 days with differentiation at 7 compared to 142 mmHg (Figure 4B). By day 8 and later, the fraction of MF-20+ cells was higher at 36 and 7 compared to 142 mmHg. The maximal fraction of MF-20+ cells was 12, 18, and 31%, which occurred on days 7, 9, and 11, at 142, 36, and 7 mmHg, respectively. Up to day 9, total cell number was the same for most time points between 7 and 142 mmHg and higher at 36 mmHg (Figure 4C). After day 9, total cell number was approximately the same between 36 and 7 mmHg, both of which had a substantially higher cell number than 7 mmHg. Differentiation at 36 and 7 mmHg resulted in more MF-20+ cells than 142 mmHg (Figure 4D). A steady-state number of MF-20+ cells was reached by day 7 for 142 mmHg and day 8 for 36 and 7 mmHg, after which the number of MF-20+ cells was the same for most times for 36 and 7 mmHg. Differentiation at low pO$_{2\text{gas}}$ produced 2-3x more MF-20+ cells compared to high pO$_{2\text{gas}}$.

We further focused our investigation on differentiation to cardiomyocytes with the addition of ascorbic acid, which has been shown to increase differentiation of mESC to cardiomyocytes [74]. Differentiating J1 mESC at various, constant pO$_{2\text{gas}}$ for 11 days with 0.2 mM ascorbic acid resulted in a large increase in the
fraction of MF-20+ cells with decreasing values of pO$_{2gas}$ that was reproducibly observed in 7 additional independent experimental runs (Figure 5A). The highest number fraction of MF-20+ cells was 31% with a pO$_{2gas}$ of 7 mmHg, compared to 23% and 9% at 36 and 142 mmHg, respectively. Culture at 7 mmHg resulted in a substantial decrease in total cell number relative to 36 and 142 mmHg, respectively, both of which had comparable cell numbers (Figure 5B). As a result, the number of MF-20+ cells was higher at 36 mmHg by a factor of 2.3 and 2.9 compared to 142 and 7 mmHg, respectively, resulting in 55 MF-20+ cells being generated per initial mESC at 36 mmHg, compared to 24 and 28 at 142 and 7 mmHg, respectively (Figure 5C). Variations of this protocol without ascorbic acid, without serum-free media, with different initial cell densities in the hanging drops, and with different time periods in hanging drop also showed an enhancement in cardiomyogenesis at low compared to high pO$_{2gas}$ (data not shown). The inclusion of ascorbic acid increased the mean fraction and number of MF-20+ cells by a factor of 2.1 and 2.4, respectively, for 142 mmHg, 1.2 and 1.3, respectively for 36 mmHg, and had no effect on 7 mmHg compared to day 11 data on Figure 4D, albeit these increases were not statistically significant. The expression of the cardiac marker genes cTnT, Mef2c, and Nkx2.5 were also higher at low compared to high pO$_{2gas}$ (Figures 5D-F). Low pO$_{2}$-differentiated mESC immunostained positive for cTnT, which were striated, demonstrating sarcomeric organized (Figure 5G), and for Nkx2.5 (Figure 5H).
3.4.5. Verification of cardiomyocyte assessment methodology

We compared MF-20 immunostaining of histological sections, counting with flow cytometry, immunostaining of cTnT on histological sections, and the area covered with spontaneously contracting cells in each well to further assess mESC-derived cardiomyocytes (Figure 6). Immunostaining of serial sections with MF-20 and anti-cTnT antibodies showed excellent co-localization in all examined sections (Figure 6A). The fraction of MF-20+ cells assessed with histology and flow cytometry data correlated highly, with a correlation coefficient of 0.7 (Figure 6B). There was also a strong correlation (correlation coefficient of 0.8) between the number of MF-20+ cells assessed with flow cytometry and the area covered by spontaneously contracting cells (Figure 6C).

3.4.6. pO\textsubscript{2cell} of cardiomyocytes after differentiation

We used MF-20-immunostained histological sections to estimate pO\textsubscript{2cell} of MF-20+ cells within cellular aggregates after 11 days of differentiation with 0.2 mM ascorbic acid of J1 mESC (Figure 7). With histological examination, mESC differentiated at 7 mmHg preferentially formed thin cell sheets and smaller aggregates than were found at 36 or 142 mmHg (Figure 7A), with 90 ± 7%, 75 ± 9%, 53 ± 6% of the tissue volume being present in aggregate regions at 142, 36, and 7 mmHg, respectively. MF-20+ cells were found together in clusters at all pO\textsubscript{2gas}.  

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We previously reported a theoretical model of oxygen consumption and diffusion to estimate the volumetric distribution of $pO_{2cell}$ within aggregations after 11 days of differentiation [53]. Combining these results with immunostaining showing the location of MF-20+ cells (Figure 7A), we calculated the $pO_{2cell}$ of the MF-20+ cells. Even though silicone rubber dishes were used in this study, oxygen gradients still existed within the aggregates. A comparison of the volumetric distribution of $pO_{2cell}$ of all of the tissue and that of MF-20+ cells at 142 mmHg is shown in Figure 7B. This analysis only considered the aggregate regions (90% of the total tissue volume), since no MF-20+ cells were observed in sheet-like areas at any $pO_{2gas}$. MF-20+ cells represented approximately 10% of the total aggregate volume and were preferentially located in regions of moderately low $pO_{2cell}$ (20 – 30 mmHg), with relatively fewer cardiomyocytes at high and low $pO_{2cell}$ extremes.

3.4.7. Effect of temporal variation of $pO_{2gas}$ on differentiation of mESC to cardiomyocytes

The effects of differentiating J1 mESC at 142 and 7 mmHg $pO_{2gas}$ for different time periods in different orders on the fraction of MF-20+ cells, total cell number, and number of MF-20+ cells after 10 days total is shown in Figure 8. The fraction of cardiomyocytes was greater when the cells were initially started at 7 mmHg. Increased cardiomyocyte fraction was observed if days 0-6 were carried out at 7 mmHg and was not affected by increasing $pO_{2gas}$ to 142 mmHg after day 6.
Changing pO$_{2\text{gas}}$ from 142 to 7 mmHg after day 2 did not increase the cardiomyocyte fraction. The total cell number was greatest for cultures in which the pO$_{2\text{gas}}$ at days 6-10 was 142 mmHg, regardless of the pO$_{2\text{gas}}$ used prior to day 6. Therefore, the total number of cardiomyocytes present at day 10 was highest if the pO$_{2\text{gas}}$ was 7 mmHg during the first 6 days of culture and then increased to 142 mmHg for the final 4 days. Using these pO$_{2\text{gas}}$ conditions, 35% of cells were MF-20+ and 60 MF-20+ cells were generated for each initial mESC, both of which represent a 3x increase relative to constant culture at 142 mmHg. The mean total MF-20+ number in this case was slightly higher than that obtained using constant culture at a pO$_{2\text{gas}}$ of 36 mmHg in the same experiments, but the difference was not statistically significant.

For cultures started at 7 mmHg, the increase in cardiomyocyte and total cell number that occurred during days 6-10 at 142 mmHg was accompanied by significant morphological changes (data not shown). The cells that were in sheet-like and small aggregate structures on day 6 proliferated and migrated to form large aggregates similar in size to those present for cultures maintained at 142 mmHg for all 10 days. These aggregates contracted vigorously and contained numerous cardiomyocytes.
3.4.8. Low pO₂ differentiation of HIF-1α⁺⁻ mESC does not increase yield of cardiomyocytes

HIF-1α⁺⁻ mESC were differentiated at various, constant pO₂gas to assess the necessity of HIF-1α in the low pO₂-mediated increase in cardiomyocytes (Figure 9). Expression of Mesp1 and Mesp2 was delayed at 36 and 7 mmHg compared to 142 mmHg (Figure 9A-B) with HIF-1α⁺⁻ mESC, similar to J1 mESC (Figure 3). However, maximal expression, which occurred on day 4 for 142 and day 5 for 36 and 7 mmHg for both genes, was the same at all pO₂gas for HIF-1α⁺⁻ mESC (Figure 9A-B), contrasting with J1 mESC, which had markedly enhanced expression at low compared to high pO₂gas (Figure 3). By day 11, spontaneously-contracting cells were observed with HIF-1α⁺⁻ mESC at all pO₂gas. However, the number and fraction were the same at all pO₂gas, averaging 17 MF-20+ cells per original HIF-1α⁺⁻ mESC and 8% of the final population, similar to the low values obtained with J1 mESC differentiated at 142 mmHg (Figure 4B and D).

3.4.9. Low pO₂ increases differentiation of hESC to cardiomyocytes

We extended our study to investigate the effects of pO₂ on differentiation of hESC to cardiomyocytes by differentiating CyT49 hESC for 21 days at constant 142, 36, or 7 mmHg pO₂gas (Figure 10). The gene expression of cTNT, MEF2C, and NKX2.5, measured with real-time PCR, was higher at 36 and 7 compared to 142 mmHg (Figure 10A-C). The average increase in cTNT, MEF2C, and NKX2.5
expression from 142 to 7 mmHg was similar between hESC and mESC but from 142 to 36 mmHg was higher with hESC compared to mESC (Figure 5D-F). More hESC-derived cTNT+ cells were observed with microscopy at low compared to high pO$_2$gas (Figure 10D-E). However, with this protocol, the fraction of cTNT+ cells was less than 1%, assessed with visual observation, at all pO$_2$gas, and we were unable to accurate quantitate the number with flow cytometry. A small amount of spontaneously-beating cells were also observed at 7 mmHg, but none were observed at 142 or 36 mmHg.

3.4.10. Low pO$_2$ increases differentiation of miPSC to cardiomyocytes

We extended our study to investigate the effects of pO$_2$ on differentiation of miPSC to cardiomyocytes. Differentiating miPSC at various, constant pO$_2$gas for 11 days with 0.2 mM ascorbic acid resulted in a large increase in the fraction of MF-20+ cells at low compared to high pO$_2$gas (Figure 11A). The highest fraction of MF-20+ cells was 11% with a pO$_2$gas of 36 and 7 mmHg, compared to 142 mmHg, values which were 2-3 times lower than J1 mESC (Figure 5A). Culture at 36 and 7 mmHg resulted in a substantial increase in total cell number relative to 142 mmHg (Figure 11B), unlike J1 mESC, which had the opposite trend (Figure 5B). As a result, the number of MF-20+ cells was highest at 7 mmHg by a factor of 5.2 and 1.4 compared to 142 and 36 mmHg, respectively, resulting in 34 MF-20+ cells being produced per each initial miPSC at 7 mmHg, compared to 25 and 7 at 36 and 142 mmHg, respectively (Figure 11C). The number of MF-20+ cells was higher at 142 and 36 mmHg and lower for 7 mmHg with J1 mESC compared to
miPSC (Figure 5C). The expression of the cardiac marker genes cTnT, Mef2c, and Nkx2.5 were also higher at low compared to high pO$_{2}$gas (Figures 11D-F). Low pO$_{2}$-differentiated miPSC immunostained positive for cTnT and Nkx2.5 (Figure 11G-H).

3.5. Discussion

We studied the effects of pO$_{2}$ on mESC differentiation to the three germ layers and mESC, miPSC, and hESC differentiation to cardiomyocytes. Throughout our studies, we used highly oxygen-permeable silicone rubber dishes to set pO$_{2}$cell at the membrane-cell interface. Real-time PCR was used to measure relative gene expression of cardiomyocyte and germ layer markers, and immunocytochemistry was used to verify the presence of marker protein. Flow cytometry was used to measure the fraction and number of Nestin+, T-GFP+, and MF-20+ cells. Histological analysis was used to quantitatively verify MF-20 staining and quantification with flow cytometry and to provide geometric measurements for estimations of pO$_{2}$cell with finite element simulations. Temporal variations in pO$_{2}$gas were performed to determine the time period over which a low pO$_{2}$cell was required to promote cardiomyocyte differentiation and to further increase MF-20+ cell fraction and number.

Low pO$_{2}$ during differentiation of mESC changed the timing and magnitude of temporal gene expression for most markers of all three germ layers. Overall, we
observed higher gene expression of endodermal genes (Figure 1), lower expression of ectodermal genes (Figure 2), and approximately the same expression of mesodermal genes at low compared to high pO₂gas (Figure 3-4). To our knowledge, there have been no previous reports on the effects of pO₂ on differentiation to early endoderm, although high pO₂ has been shown to affect later stages of pancreatic development [54]. Our protocol was not designed to produce large amounts of endodermal cells, so further investigating low pO₂ with a better protocol is of interest. Low pO₂ has been previously reported to affect different stages of ectodermal differentiation differently, with differentiation of PSC to neural stem cells (NSC) being reduced at low pO₂ while differentiation of NSC to neurons being enhanced [2]. pO₂ has been previously implicated in changing the timing and magnitude of the expression of the mesodermal marker Brachyury T, but this has not been shown statistically [146].

We demonstrated that control of pO₂ can result in a factor of 3 increase in cardiomyocyte fraction and number from mESC above levels typically obtained at 142 mmHg without purification (Figure 8). The best protocol reported in this study involved the addition of ascorbic acid and removal of serum after 5 days of culture. Nonetheless, the positive effect that low pO₂ culture had on mESC differentiation into cardiomyocytes was found using several variations in the differentiation protocol, such as with or without ascorbic acid or serum removal, and with miPSC and hESC, suggesting the benefits of low pO₂ are broadly applicable and not simply the artifact of using a specific differentiation protocol.
Previous studies have investigated the effects of low pO$_2$ on differentiation of PSC to cardiomyocytes in bioreactors. Bauwens et al. [48] reported no significant difference between 4% and 20% oxygen from mESC, likely due to the low cardiomyocyte yields without genetic selection. Only after combining low pO$_2$ with genetic selection to purify cardiomyocytes, 3.8 cardiomyocytes were generated per input mESC at 4% oxygen, a factor of 1.5 higher than 20% oxygen with genetic selection. This low oxygen result is a factor of 16 lower than what we achieved with our temporal modulations in pO$_2$gas protocol (Figure 8). We likely were able to achieve higher yields because our protocol consistently obtained a cardiomyocyte fraction of 5-10% from mESC at 142 mmHg, comparable to other optimized differentiation protocols [118, 147, 148], which allows for easier measurement of the effects of pO$_2$. Additionally, with hESC, 4% oxygen has reported to increase the expression of some cardiac markers and the total cell number compared to 20% oxygen. However, the fraction of beating EBs is the same between 20% and 4% and no information on fraction or number of cardiomyocytes was reported [49].

The mechanism by which oxygen exerted its effects in our experiments is not known. The oxygen-sensitive transcription factor HIF-1 is active in wild type PSC cultured at low pO$_2$ [149] and is seen in hypoxic regions of the developing chick heart [150]. We demonstrated that differentiation of HIF-1α$^{-/}$ mESC to cardiomyocytes was insensitive to pO$_2$ (Figure 9). HIF-1 targets hypoxic
responsive elements within the promoter regions of and increases expression of the cardiac-associated transcription factors Mef2c and Tbx5 and the sarcomere organization protein Titin [46]. HIF-1 stabilization and activity at low pO$_2$ therefore seems likely to be responsible for at least some of the effects observed in our experiments.

Oxygen exerted its strongest effects on cardiomyocyte differentiation during the first 6 days (Figure 8), the time period that immediately precedes the emergence of the first spontaneously-contracting and MF-20+ cells and coincides with increases Mesp1 and Mesp2 expression. Thus, we infer that pO$_2$ affected differentiation into cardiomyocyte progenitor cells and/or induced an increase in proliferation of such progenitor cells relative to other cells types. The increase in total cardiomyocyte number suggests a direct positive effect of reduced oxygen on differentiation or proliferation along the cardiomyocyte lineage, but the exact mechanism and cell types through which this process is occurring is not yet known. After the appearance of cardiomyocytes, high pO$_2$gas caused a higher total cell number, presumably by minimizing the oxygen or other nutrient limitations as the cell number increased.

The use of silicone rubber culture dishes to control pO$_2$cell at the membrane-cell interface was crucial for the success of our experiments. We previously reported that simply differentiating mESC on oxygen-impermeable polystyrene dishes at low pO$_2$gas does not result in more cardiomyocytes compared to high pO$_2$gas [53].
Furthermore, when polystyrene and silicone rubber dishes were compared directly, the highest cardiomyocyte fraction and number occurred with differentiation at low $pO_2_{gas}$ on silicone rubber dishes. Without control or understanding of $pO_2_{cell}$, new important knowledge may remain hidden and apparent inconsistencies between published studies may arise [2].

Even though the cells were cultured on oxygen-permeable silicone rubber, differentiating cells preferentially formed aggregates that contained $pO_2$ gradients (Figure 7) [53]. The size of the aggregates was smaller with decreasing $pO_2_{gas}$ values, which may have been due to limitations on oxygen and nutrient availability, especially given that cells initially cultured at 7 mmHg would form large aggregates within 4 days of transfer to 142 mmHg (data no shown). MF-20+ cells were exclusively present in aggregates and preferentially occupied regions of moderately low local $pO_2$ after 11 days of differentiation at 142 mmHg (Figure 7B). However, the $pO_2_{cell}$ during differentiation (before day 11) of these cells is unknown. Oxygen gradients were large within day-11 aggregates but are certainly smaller during the first 6 days of differentiation, since the aggregates are smaller. Experiments investigating histological sections from earlier times during differentiation immunostained with antibodies to markers of cardiomyocyte progenitor cells would be worthwhile to further explore whether cardiomyocytes preferentially form in regions within a certain range of $pO_2_{cell}$. 

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Our work shows that PO₂<sub>cell</sub> profoundly affects differentiation of PSC into cardiomyocytes and the three germ layers, which has major ramifications. One potential application of PSC is to better understand development and disease, and this work suggests that controlling PO₂ to levels experienced by the developing embryo is necessary to accurately explore this. The markedly increased yield of cardiomyocytes may enhance prospects for their therapeutic use in treating heart disease. Control of dissolved oxygen is inexpensive compared to growth factors and small molecules and can be accomplished with current technology, making our results applicable to commercial development. Reduced PO₂ culture combined with other proteins and small molecules may further enhance differentiation to cardiomyocytes and other desired cell types. Because the cells in the early embryo are exposed to PO₂<sub>cell</sub> that lie within the range that we observed largest effects on differentiation, PO₂<sub>cell</sub> may play a more important role in early embryonic development than heretofore appreciated.
3.6. Figures
Figure 3-1. pO₂ affects differentiation to the endodermal lineage.

J1 mESC were differentiated for up to 13 days at 142, 36, or 7 mmHg pO₂gas without ascorbic acid. (A-D) Temporal gene expression of Foxa2, Sox17, Hnf4a,
and Pdx1 acquired with real-time PCR (n=3). (E-F) En face fluorescence images of mESC after 6 days of differentiation at 142 (E) or 36 (F) mmHg stained with DAPI (blue) and against Foxa2 (green). (G-H) En face phase contrast images of Sox17-dTomato mESC after 6 days of differentiation at 142 (G) or 36 (H) mmHg with respective Sox17-dTomato (red) fluorescence images superimposed.
Figure 3-2. pO₂ affects differentiation to the ectodermal lineage.

J1 mESC were differentiated for up to 13 days at 142, 36, or 7 mmHg pO₂gas without ascorbic acid. (A-C) Temporal gene expression of Sox1, Nestin, and Pax6 acquired with real-time PCR (n=3). (D-E) Number and fraction of Nestin+ cells after 12 days of differentiation acquired with flow cytometry (n=3). (F-G) En face fluorescence images of mESC after 10 days of differentiation at 142 (F) or 7 (G) mmHg stained with DAPI (blue) or against Nestin (green). * indicates statistical difference (p<0.05) compared to 142 mmHg (black bar), and # indicates statistical difference compared to 36 mmHg (grey bar).
Figure 3-3. Temporal gene expression of the mesodermal and cardiac lineage markers Brachyury T, Mesp1, Mesp2, Tbx20, Tbx5, Nkx2.5, Mef2c, Foxh1, cardiac α-Actin, cTnT, and Titin for J1 mESCs differentiated for up to 13 days at 142, 36, or 7 mmHg pO$_2$gas without ascorbic acid.

All data acquired with real-time PCR (n=3).
mESC were differentiated for up to 13 days at 142, 36, or 7 mmHg pO$_{2gas}$ without ascorbic acid. (A) Fraction of T-GFP+ cells from T-GFP mESC acquired with flow cytometry (n=3). (B) Fraction of MF-20+ cells from J1 mESC acquired with flow cytometry (n=3). (C) The total cell number per well from J1 mESC acquired with nuclei counts (n=3). (D) The number of MF-20+ cells from J1 mESC calculated by multiplying the fraction of MF-20+ cells (B) and the total cell number (C) (n=3).
Figure 3-5. Low pO₂ increases mESC differentiation to cardiomyocytes.

J1 mESC were differentiated 11 days culture at 142, 36, or 7 mmHg pO₂gas with ascorbic acid. (A) Fraction of MF-20+ cells from mESC acquired with flow cytometry (n=7). (B) The total cell number per well from mESC acquired with nuclei counts (n=7). (C) The number of MF-20+ cells from mESC calculated by multiplying the fraction of MF-20+ cells (A) and the total cell number (B) (n=7). (D-F) Gene expression of Nkx2.5, Mef2c, and cTnT acquired with real-time PCR for (n=4). (G-H) En face fluorescence images of mESC differentiated at 7 mmHg pO₂gas and stained against cTnT (green) and with DAPI (blue) and against Nkx2.5 (green). * indicates statistical difference (p<0.05) compared to 142 mmHg.
(black bar), and # indicates statistical difference compared to 36 mmHg (grey bar).
Figure 3-6. Comparison of different cardiomyocyte assessment methodologies. 

(A) Consecutive 5-μm tissue sections of J1 cells differentiated 11 days at 36 mmHg with ascorbic acid immunostained with a primary antibody to sarcomeric myosin heavy chain (MF-20 - top), cardiac troponin T (cTnT - middle), or no primary antibody (control - bottom). (B) Comparison of the fraction of MF-20+ cells counted from stained 5-μm sections to that measured with flow cytometry of trypsin dispersed cells. Open, gray filled, and black filled symbols represent samples taken from pO$_{2gas}$ of 142, 36, and 7 mmHg, respectively, and each
shape (circle, triangle, upside-down triangle) represents an independent experimental run. Results for flow cytometry are the mean ± standard deviation (n=3), while the histological results are from a single section that was obtained in each experiment and pO_{2gas}. (C) Comparison between the total number of MF-20+ cells counted using flow cytometry and the total surface area of the culture dish covered with spontaneously contracting cells estimated visually. The fraction of MF-20+ cells ranged from 3 to 36% in these samples. The best-fit line was determined using linear regression, R^2 = 0.80.
Figure 3-7. Modeling of the volumetric distribution of $pO_2$ within MF-20+ and all regions.

(A) Representative images of 5-µm tissue sections of J1 cells immunostained with MF-20 (brown) and counter stained with hematoxylin. (B) Volumetric distribution of $pO_{2cell}$ values in MF-20+ regions at 142 mmHg. Filled bars represent the distribution of $pO_{2cell}$ within MF-20+ regions, and open bars represent the distribution of $pO_{2cell}$ in the entire volume of aggregates in the culture dish (including MF-20+ regions).
Figure 3-8. Temporal modulation of oxygen culture at 7 and 142 mmHg affects differentiation to cardiomyocytes.

J1 mESC were differentiated with ascorbic acid starting at either 142 or 7 mmHg and then switched to the other condition on day 2, 4, 6, or 8 for a total of 10 days.
Fraction MF-20+, total cell number, and number of MF-20+ cells assessed with flow cytometry after 10 days of differentiation are shown (n as shown at the top of each). The 10 different pO$_{2}$gas histories and day of the pO$_{2}$gas change are shown schematically on the x-axis.
Figure 3-9. HIF-1α is necessary for low pO₂-enhancement of cardiomyocyte generation.

HIF-1α⁺⁻ mESC were differentiated up to 10 days at 142, 36, or 7 mmHg pO₂gas without ascorbic acid. (A-B) Time-dependent gene expression of Mesp1 and Mesp2 acquired with real-time PCR (n=3). (C-D) The fraction and number of MF-20+ cells acquired with flow cytometry (n=6).
Figure 3-10. Low pO₂ increases hESC differentiation to cardiomyocytes.

CyT49 hESC were differentiated 20 days at 142, 36, or 7 mmHg pO₂gas. (A-C) Gene expression of cTNT, MEF2c, and NKX2.5 acquired with real-time PCR (n=4). (D-E) En face fluorescence images of hESC differentiated at 142 (D) or 7 (E) mmHg and stained with DAPI and against cTNT. * indicates statistical difference (p<0.05) compared to 142 mmHg (black bar), and # indicates statistical difference compared to 36 mmHg (grey bar).
Figure 3-11. Low pO₂ increases miPSC differentiation to cardiomyocytes.

miPSC were differentiated 11 days at 142, 36, or 7 mmHg pO₂gas. (A) Fraction of MF-20+ cells acquired with flow cytometry (n=4). (B) The total cell number per well acquired with nuclei counts (n=4). (C) The number of MF-20+ cells calculated by multiplying the fraction of MF-20+ cells (A) and the total cell number (B) (n=4). (D-F) Gene expression of cTnT, Mef2c, and Nkx2.5 acquired with real-time PCR for (n=4). (G-H) En face fluorescence images of miPSC differentiated at 7 mmHg pO₂gas and stained with DAPI (blue) and against cTnT and Nkx2.5 (green). * indicates statistical difference (p<0.05) compared to 142 mmHg (black bar), and # indicates statistical difference compared to 36 mmHg (grey bar).
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Appendix

A. Accurate control of oxygen level in cells during culture on silicone rubber membranes with application to stem cell differentiation

Accurate Control of Oxygen Level in Cells During Culture on Silicone Rubber Membranes with Application to Stem Cell Differentiation

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Oxygen level in mammalian cell culture is often controlled by placing culture vessels in humidified incubators with a defined gas phase partial pressure of oxygen ($P_{O_2}\text{gas}$). Because the cells are consuming oxygen supplied by diffusion, a difference between $P_{O_2}\text{gas}$ and that experienced by the cells ($P_{O_2}\text{cell}$) arises, which is maximal when cells are cultured in vessels with little or no oxygen permeability. Here, we demonstrate theoretically that highly oxygen-permeable silicone rubber membranes can be used to control $P_{O_2}\text{cell}$ during culture of cells in monolayers and aggregates much more accurately and can achieve more rapid transient response following a disturbance than on polystyrene and fluorinated ethylene-propylene co-polymer membranes. Cell attachment on silicone rubber was achieved by physical adsorption of fibronectin or Matrigel. We use these membranes for the differentiation of mouse embryonic stem cells to cardiomyocytes and compare the results with culture on polystyrene or on silicone rubber on top of polystyrene. The fraction of cells that are cardiomyocyte-like increases with decreasing $P_{O_2}$ only when using oxygen-permeable silicone membrane-based dishes, which contract on silicone rubber but not polystyrene. The high permeability of silicone rubber results in $P_{O_2}\text{cell}$ being equal to $P_{O_2}\text{gas}$ at the tissue-membrane interface. This, together with geometric information from histological sections, facilitates development of a model from which the $P_{O_2}$ distribution within the resulting aggregates is computed. Silicone rubber membranes have significant advantages over polystyrene in controlling $P_{O_2}\text{cell}$, and these results suggest they are a valuable tool for investigating $P_{O_2}$ effects in many applications, such as stem cell differentiation. © 2009 American Institute of Chemical Engineers Biotechnol. Prog., 26: 805-818, 2010

Keywords: embryonic stem cells, oxygen, hypoxia, differentiation, silicone rubber

Introduction

Laboratory animal cell culture experiments are usually performed in glass or polystyrene dishes having zero or low oxygen permeability in an incubator having a humidified atmosphere consisting of 95% air/5% CO$_2$ that results in a gas oxygen partial pressure ($P_{O_2}\text{gas}$) of 142 mm Hg. Because oxygen consumption by the cells induces oxygen concentration gradients in the medium, $P_{O_2}$ at the cell surface ($P_{O_2}\text{cell}$) is lower than $P_{O_2}\text{gas}$ and the difference ($\Delta P_{O_2}$), which is usually unknown, can be large, recognized using theoretical models of oxygen transport, and experiments with microelectrodes. These gradients have been exploited in static culture systems to measure oxygen consumption rate (OCR) of cells in monolayer culture and in embryoid bodies (EBs). In addition to affecting steady-state values of $P_{O_2}\text{cell}$, the long time scales associated with transient oxygen diffusion through the static medium make it nearly impossible to accurately study effects of intermittent hypoxia.

To reduce $\Delta P_{O_2}$, cells can be cultured in gas-permeable dishes, in dishes that are mechanically mixed, or in a perfused system to induce convective transport. Mechanical mixing is preferred for oxygenation of large-scale bioreactors but introduces cumbersome equipment requirements for small-scale research. Additionally, mixing in culture vessels induces shear stresses on cells, which may affect their function. Perfusion systems can avoid this problem but are complex and susceptible to contamination. In contrast,
culture on a gas-permeable membrane at the bottom of a dish requires no additional equipment and no changes in the culture environment, other than the surface on which the cells are growing. Membranes previously examined include fluorinated ethylene-propylene copolymer (FEP) and silicone-poly carbonate membrane copolymer.\textsuperscript{11} FEP was selected because of its optical clarity and strength, not because it is optimal for oxygen transfer. Dishes made with this membrane are available (Lumox dishes, Greiner Bio-One, Munich) with either a hydrophobic or plasma gas-treated hydrophilic surface suitable for cell culture.\textsuperscript{1,2} The oxygen permeability of silicone rubber is two orders of magnitude higher than that of FEP (Table 1).\textsuperscript{5,6} Silicone rubber is preferred for oxygen delivery and is used in membrane-aerated bioreactors.\textsuperscript{16-19} High density culture of hybridoma cells\textsuperscript{20,21} and cell aggregates\textsuperscript{21} Small-scale dishes suitable, e.g., for stem cell research that incorporate these membranes are not commercially available.

Silicone rubber surfaces must be modified prior to culture of adherent cells.\textsuperscript{22,23} Treatment with plasmas of various compositions have been used\textsuperscript{22,23} but surfaces modified this way are not stable for extended periods of time due to migration of functional groups from the surface into the bulk material.\textsuperscript{24-26} Modifications in which functional coatings are grafted onto the silicone rubber are more stable but infrequently used.\textsuperscript{27,28} Recently, polyelectrolyte multilayers (PEMs) have been used for surface modification of silicone rubber because this method is inexpensive, easy to apply, and has flexible chemistry.\textsuperscript{29-33}

Here we show theoretical justification for the superiority of silicone rubber membranes for accurate control of the $P_{O_2}^{cell}$ during culture of both cell monolayers and aggregates. We also show that physical adsorption of fibronectin or Matrigel to silicone rubber is sufficient for robust cell adhesion of the cell line employed, mouse embryonic stem (mES) cells. These materials are used to study the effects of oxygen on the differentiation of mES cells into cardiomyocytes using an 11-day differentiation protocol, the last 9 days of which occur with the cells attached to silicone rubber membrane-based dishes. Finally, we used histology to determine the geometry of differentiated aggregates of mES cells for use in a theoretical model for predicting the $P_{O_2}^{cell}$ distribution in these aggregates.

### Materials and Methods

#### Silicone rubber membrane-based plate

The bottom surface of eight central wells of 24-well tissue culture treated plates (353047, Becton Dickinson, Franklin Lakes, NJ) was removed using a $3/8 \times 3$-in fixed handle nutdriver (Cooper Hand Tools, Apex, NC) heated in a Bunsen burner, and edges of the holes trimmed with sterile scalpel to be flush with the rest of the plate bottom. A very thin layer of silicone adhesive (Henkel Loctite Corp., Rocky Hill, CT) was spread on the base of the plate around each hole. Silicone rubber sheets (nonreinforced vulcanized gloss/gloss 0.005 in thick, Specialty Manufacturing, Saginaw, MI), cut to $8.5 \times 4.5$ cm and sterilized by autoclaving for 30 min at $121^\circ$C, was stretched so that it was flat (no wrinkles) and pressed onto the uncut bottom surface. After allowing the adhesive to set for 24 hr, the plates were filled with 70% (v/v) ethanol in water for 1 hr and dried overnight under a germicidal UV lamp in a biological safety cabinet. In use, the silicone rubber membrane at the bottom of the plate was placed on four layers of dry paper towels to provide access to ambient $P_{O_2}^{gas}$.

#### Coating of silicone rubber using layer-by-layer PEM film growth

PEMs were prepared by adapting a published method.\textsuperscript{29} Solutions in Dulbecco's phosphate buffered saline (DPBS) (Mediatech, Herndon, VA) were prepared containing 3 mg/mL poly-sodium 4-styrene-sulfonate (PSS), 2 mg/mL poly-ethyleneimine (PEI), 0.5 mg/mL poly-d-lysine hydrobromide (PDL), 1 mg/mL type A gelatin, 1 mg/mL type B gelatin and 50 $\mu$g/mL fibronectin (all from Sigma-Aldrich, St. Louis, MO). Coating a silicone rubber membrane at the bottom of a well was carried out at room temperature using 1 mL of solution for each step. PSS solution (polyanion) was added to an untreated plate and incubated for 1 hr, followed by a 10-min wash with DPBS, an 1-hr incubation with PEI (polycation), and a 10-min wash with DPBS. Two additional incubations with both PSS and PEI (30 min each) with intervening washes (10 min) were done to build a (PSS-PEI)$_n$ PEM. An additional three layers each of type B gelatin (polyanion) and PDL (polycation) were formed using 30-min incubations with the polyelectrolyte solution and 10-min washes with DPBS to yield (PSS-PEI)$_n$-(type B gelatin-PDL)$_n$, after which an additional 20-hr incubation with type A gelatin, type B gelatin, or fibronectin was performed. Plates were washed and incubated for 24 hr with DPBS prior to emptying the wells and adding culture medium and cells.

#### Coating of silicone rubber membrane by physical adsorption of proteins

Plates to be coated were incubated at 37°C in a 0.1% (w/v) solution of type A gelatin in tissue culture water for 24 hr (Mediatech, Herndon, VA), 2 $\mu$g/mL fibronectin (Sigma) in DPBS for 24 hr, or Matrigel (354234, Becton Dickinson) diluted 1:20 in serum-free medium for 1 hr. Immediately before plating cells, the solution was removed and replaced with mES cell medium.

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### Table 1. Oxygen Diffusivity and Solubility Parameters at 37°C

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<th>Continuous Phase</th>
<th>$D$ ($10^{-5}$ cm$^2$ sec$^{-1}$)</th>
<th>Source</th>
<th>$\alpha$ (10$^{-9}$ mol cm$^{-3}$ mm Hg$^{-1}$)</th>
<th>Source</th>
<th>$D_{O_2}/h_0^*$ (10$^{-14}$ mol sec$^{-1}$ cm$^{-2}$ mm Hg$^{-1}$)</th>
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<td>Culture medium</td>
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<td>1.27</td>
<td>Dionne 1990$^b$</td>
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<td>Silicone rubber</td>
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<td>Average value$^c$</td>
<td>12.2</td>
<td>Average value$^d$</td>
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<td>Fluorinated ethylene-propylene copolymer</td>
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<td>DuPont 1996</td>
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<td>Hodge et al. 2001</td>
<td>8.60</td>
<td>Chen et al. 1994</td>
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<td>Dionne 1990$^b$</td>
<td>1.02</td>
<td>Dionne 1990$^b$</td>
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$^a$Thickness $h$ of 5.0,127,0.025, and 1 mm was used for culture medium, silicone rubber, fluorinated ethylene-propylene copolymer, and polystyrene, respectively. $^b$Mean of published values. $^c$Reitlinger et al. 1956; Robb 1968. $^d$Barrer et al. 1965; Reitlinger et al. 1956; Robb 1968.
Table 2. Composition of Undifferentiated mES Cell Maintenance and mES Cell Differentiation Mediuma

<table>
<thead>
<tr>
<th>Component</th>
<th>Manufacturer and Catalog Number</th>
<th>Volume per Liter of Medium (mL)</th>
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<td>Dulbecco’s modified eagles medium (DMEM)</td>
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<td>Fetal bovine serum</td>
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<td>ES cell qualified</td>
</tr>
<tr>
<td>L-lysine</td>
<td>ATCC SCRR 20-2115</td>
<td>10</td>
<td>200 mM stock solution</td>
</tr>
<tr>
<td>MEM nonessential amino acid solution</td>
<td>ATCC SCRR 20-2116</td>
<td>1</td>
<td>10 mM stock solution prepared in DMEM</td>
</tr>
<tr>
<td>2-mercaptoethanol solution</td>
<td>Sigma-Aldrich M7522</td>
<td>10</td>
<td>100 mM stock solution prepared in DMEM</td>
</tr>
<tr>
<td>Leukemia inhibitory factor</td>
<td>Chemicon ESG 1106</td>
<td>1</td>
<td>100 unit/mL stock solution</td>
</tr>
<tr>
<td>L-ascorbic acid solution</td>
<td>Sigma-Aldrich A4034</td>
<td>1</td>
<td>200 mM stock solution prepared in DPBS</td>
</tr>
</tbody>
</table>

a Leukemia inhibitory factor was added only to undifferentiated ES cell maintenance medium. Ascorbic acid was added only to ES cell differentiation medium.

Table 3. Formulation of Serum-Free ITS Medium

<table>
<thead>
<tr>
<th>Component</th>
<th>Manufacturer and Catalog Number</th>
<th>Volume per Liter of Medium (mL)</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dulbecco’s modified eagles medium (DMEM)</td>
<td>Mediatech 90-133-PB</td>
<td>485</td>
<td>8.4 mg/mL stock solution prepared in deionized water</td>
</tr>
<tr>
<td>Sodium bicarbonate</td>
<td>Invitrogen 37165-035</td>
<td>10</td>
<td>7.5% (w/v) solution</td>
</tr>
<tr>
<td>F12 nutrient mixture</td>
<td>Mediatech 25-035-CI</td>
<td>496</td>
<td></td>
</tr>
<tr>
<td>25 M glucose solution</td>
<td>Sigma-Aldrich G8769</td>
<td>1.6</td>
<td>Final medium concentration of 9 mM</td>
</tr>
<tr>
<td>Human insulin solution</td>
<td>Sigma-Aldrich B278</td>
<td>0.5</td>
<td>10 mg/mL solution</td>
</tr>
<tr>
<td>Holo transferrin solution</td>
<td>Sigma-Aldrich T1283</td>
<td>5</td>
<td>10 mg/mL stock solution prepared in DPBS</td>
</tr>
<tr>
<td>Sodium selenite solution</td>
<td>Sigma-Aldrich S9133</td>
<td>0.26</td>
<td>0.12 mM stock solution prepared in DPBS</td>
</tr>
<tr>
<td>L-ascorbic acid solution</td>
<td>Sigma-Aldrich A4034</td>
<td>1</td>
<td>200 mM stock solution prepared in DPBS</td>
</tr>
</tbody>
</table>

Gas phase \( \text{pO}_2 \) control

As previously described,\(^{34}\) cell culture vessels were placed in sealed polystyrene chambers (Billups-Rothenburg, Del Mar, CA) inside a standard incubator (Queue Systems, Parkersburg, WV) at 37°C. Gas was bubbled through a bottle of water open only to the gas cylinder and chamber in the incubator; an open dish of deionized water in each chamber provided additional humidification. The flow rate to the chambers of premixed gas containing 20%, 5%, or 1% \( \text{O}_2 \), 5% \( \text{CO}_2 \), and the remainder \( \text{N}_2 \) (certified medical gas, Airgas, Hingham, MA) was set to 2 L/min for 15 min following closure of the chambers (after cell medium exchange) and to 30 mL/min otherwise. The corresponding \( \text{pO}_2 \text{gas} \) in the chambers was 142, 36, or 7 mm Hg, respectively.

Measurement of transient \( \text{pO}_2 \) in culture dishes

To examine the transient response of polystyrene and silicone rubber dishes to a change in \( \text{pO}_2 \text{gas} \), measurements were made in an Xvivo Incubation System (BioSpherix, Lacona, NY), an oxygen-controlled incubator, using the included 2-mm diameter fiber optic oxygen-sensing probes (Model F2 OxyValidator, Kyodo International, Kawasaki, Japan). The probe was placed vertically in each of 2 wells of a 24-well plate, one well with a polystyrene and one with a silicone rubber bottom filled with 2 mL of DPBS (medium height 10.5 cm) so that the sensing tip was in contact with the upper surface of the plate bottom inside of a sealed incubator set at \( \text{pO}_2 \text{gas} \) = 100 mm Hg and 37°C. At \( t = 0 \), the incubator gas controller was set to \( \text{pO}_2 \text{gas} \) = 7 mm Hg, and the \( \text{pO}_2 \) of each of the sensors monitored for 20 hr. Plates with silicone rubber bottoms were placed on four layers of paper towels in the same configuration as for normal cell culture.

Culture of undifferentiated mES cells

Undifferentiated J1 mES cells ATCC (SCRC 1010, ATCC, Manassas, VA) were grown in 25 cm\(^2\) cell culture T-flasks (Becton Dickinson) using 4 mL of medium (Table 2) exchanged daily. Cells were detached with 0.25% trypsin every two days and placed in a new culture flask at a density of \( 1.2 \times 10^4 \) cell/cm\(^2\).

Enumeration of undifferentiated cells

Unattached cells (suspended in the supernatant medium) were mixed with an equal volume of a lysis solution containing 1% Triton X-100 (Sigma) and 0.1 M citric acid (Sigma) in deionized water and stored at 4°C for up to 1 day before analysis. Nuclei from attached cells were obtained by adding 0.5 mL of lysis solution to the well. After 10 min, solution was removed, and 0.5 mL of DPBS was added, pipetted up and down to remove any remaining nuclei, and saved for analysis. Nuclei were counted as previously described using a Guava PCA flow cytometer (Guava Technologies, Hayward, CA).\(^{34}\) Viable cell counts were performed by detaching cells with 0.25% trypsin (ATCC), resuspending in 0.4% trypan blue solution (Sigma), and counting membrane-intact cells with a hemacytometer.

Differentiation of mES cells to cardiomyocytes

Differentiation. Detached undifferentiated mES cells were centrifuged, supernatant medium removed, resuspended in differentiation medium without LIF (Table 2), and diluted to 25,000 membrane-intact cells/mL. Single 20-μL drops of this cell suspension were placed on the inside surface of the lids of 10 × 10 cm Petri dishes (Becton Dickinson) using an 8-channel pipette. The lids were inverted and placed onto dish bottoms filled with 15 mL of (37°C) solution containing 75% DPBS, 25% water, and 0.002% (w/v) type A gelatin. After 2 days, the resulting EBs were manually pipetted off the lids using wide-orifice tips (Molecular BioProducts, San Diego, CA) and placed in wells with a fibronectin-coated bottom consisting of either polystyrene, a silicone rubber membrane, or a silicone rubber membrane on top of polystyrene. Exactly thirty EBs were placed in each well, and
medium was mixed by pipetting to distribute EBs uniformly across the bottom. EBs attached and spread within one day of transfer to silicone rubber. Medium was changed every 48 hr after transfer and replaced with 1 mL of serum-free differentiation medium. After 1 day, medium was replaced with 1 mL of serum-free differentiation medium (Table 3), which was exchanged daily for the next 4 days.

**Cell Processing for Nuclei Counts and Flow Cytometry.** Medium was removed from each well, attached cells were washed with DPBS, and 200 μL of trypsin solution was added. After 5 min at 37°C, 800 μL of differentiation medium was added, and the contents were vigorously pipetted up and down using a 1-mL pipette to dislodge and disperse the cells, which were transferred to a 1 mL tube and allowed to settle for 2 min. The bottom 50 μL, which contained any remaining large clumps of cells and extra-cellular matrix, was removed and discarded (or saved for later analysis). The cell sample was mixed by vortexing, and an aliquot saved for nuclei counting. The remaining cell sample was centrifuged, and the supernatant discarded. Cells were resuspended in 750 μL DPBS, 250 μL of 4% (w/v) paraformaldehyde (Alfa Aesar, Ward Hill, MA) in DPBS was added to fix the cells, and each sample was incubated for 20 min at room temperature. Samples were then centrifuged, supernatant removed, and 1 mL of DPBS added. Samples of nuclei and of fixed cells were stored at 4°C before analysis.

**Flow Cytometry.** Samples containing 3 x 10^5 fixed cells were incubated for 10 min with an equal volume of 1% (w/v) saponin to permeabilize cell membranes, washed, and resuspended in 50 μL of 2% (v/v) fetal bovine serum (FBS, ATCC) in PBS for 30 min, then 5 μL of antisarcomeric myosin heavy chain antibody (MF-20, MF-20 supernatant, Developmental Studies Hybridoma Bank, Iowa City, IA) was added. Samples were incubated for 1 hr, 0.5 mL of 2% FBS solution was added to each tube, and the samples centrifuged. Supernatant was discarded, and the cells resuspended in 50 μL of goat anti-mouse PE-conjugated secondary antibody (Jackson ImmunoResearch, West Grove, PA) diluted 1:250 in 2% FBS. Samples were incubated 30 min in the dark, washed twice with 0.5 mL of PBS, and fluorescence intensity data were acquired using the Guava PCA flow cytometer with the Express software module.

**Semiquantification of Spontaneously Contracting Areas.** The fractional area of each well containing spontaneously-contracting cells was estimated by visual observation. The average beating area of fifteen random 2 mm² circles in each well was taken to be the fraction of beating cells for that well.

**Cell Processing for Histology.** Nine days after plating EBs, medium was removed from wells, and cells were washed with DPBS and incubated for 30 min with 4% (w/v) paraformaldehyde. After washing twice with 1 mL of DPBS, the sheet of cells was released from the well bottom by stretching the silicone rubber membrane and transferred into a 1.5 mL microtube. Each tube was centrifuged at 300g, all but 100 μL of the DPBS was removed, and 900 μL of 1% (w/v) agarose (Life Technologies, Gaithersburg, MD) in deionized water at 95°C was added and mixed by vortexing. Samples were centrifuged for 1 min at 20,000g, cooled for 1 hr at 4°C, and 500 μL of 4% (w/v) paraformaldehyde was added to the tubes, which were stored overnight. Agarose was removed from the tube the following day and trimmed with a razor blade so that only the pellet and a small additional amount of agarose remained. Pellets were placed in histology cassettes and embedded in paraffin. Five micrometer sections were cut, deparaffinized, and rehydrated by two 7-min rinses with xylene, two 5-min rinses with 100% ethanol, a 3-min rinse with 95% ethanol, a 10-min rinse with 70% ethanol, and two 5-min rinses with deionized water. Sections were stained by incubating for 15 sec in filtered hematoxylin (Sigma), followed by three washes in 30 mM sodium borate (EMD, Gibbstown, NJ), then dehydrated following the reverse procedure for rehydration and preserved using Permount (Fisher Chemical, Fair Lawn, NJ).

**Theory**

Mathematical models were developed to estimate pO2cell for (1) a monolayer of cells, common for undifferentiated cells, and (2) cells that have combined into three-dimensional aggregates, typical of cells undergoing differentiation (Figure 1). We used c_i = x_i pO_2, where c_i is oxygen concentration, x_i is the Bunsen solubility coefficient, and i denotes the phase. OCR was assumed to depend on the local pO2, according to Michaelis-Menten kinetics,

\[ V = \frac{V_{\text{max}} pO_{2\text{cell}}}{K_m + pO_{2\text{cell}}} \]  

where \( V_{\text{max}} \) is the maximal OCR (amol/cell sec), and \( K_m \) is the Michaelis constant, taken to be 0.44 mm Hg.\(^{35}\) We previously measured \( V_{\text{max}} \) of undifferentiated CCE mouse mES cells that had been cultured at various pO2gas from 0 to 285 mm Hg for a day or more\(^{34}\) under conditions such that the difference between pO2gas and pO2cell was small. Data were fitted to a hyperbolic expression of the form

\[ V_{\text{max}} = V_{\text{max,0}} + (V_{\text{max,∞}} - V_{\text{max,0}}) \left( \frac{pO_2}{K_v + pO_2} \right) \]  

where \( V_{\text{max,0}} = 10 ± 2 \) and \( V_{\text{max,∞}} = 30 ± 2 \) amol/cell sec, and the fitting constant \( K_v = 16 ± 1 \) mm Hg. Preliminary measurements with EBs after two days culture and with tissue differentiated for 10 days\(^{36}\) both with J1 cells, gave results that were similar to those obtained with undifferentiated CCE cells. For analysis related to mES cells, the OCR as a function of local pO2 was taken as the product of Eqs. 1 and 2 using the same parameters.

**Cell Monolayer.** We assume the cells are uniformly distributed on the surface at a density for which a one-
dimensional approximation in the medium is valid. At steady state, the rate of oxygen consumption by the cells is balanced by the sum of the rates of oxygen diffusion through the stagnant medium and support layer:

\[ \frac{D_m \alpha_m}{h_m} + \frac{D_s \alpha_s}{h_s} (pO_{2\text{gas}} - pO_{2\text{cell}}) = \rho V \] (3)

where \( D_i \) and \( \alpha_i \) are oxygen diffusion coefficient and solubility in phase \( i \), respectively, \( h_i \) is thickness of phase \( i \), \( \rho \) is cell surface density, and subscripts \( m \) and \( s \) refer to medium and support layer phases, respectively. For all calculations, \( h_s \) was taken to be actual or typical values (Table 1).

If \( V \) is represented by the product of Eqs. 1 and 2, and all variables are made dimensionless, Eq. 3 becomes

\[ (1 - \theta) = \left[ F_V + (1 - F_V) \left( \frac{\theta}{\beta_V + \theta} \right) \right] \left( \frac{\theta}{\beta_M + \theta} \right) f(D_{a,i}) \] (4)

where

\[ f(D_{a,i}) = \left( \frac{1}{D_{a,m}} + \frac{1}{D_{a,s}} \right)^{-1} \] (5)

and \( \theta = \frac{pO_{2\text{cell}}}{pO_{2\text{gas}}} \), \( F_V = V_{\text{max,0}} \), \( \beta_V = K_V/pO_{2\text{gas}} \), \( \beta_M = K_M/pO_{2\text{gas}} \), and \( D_{a,i} = \rho V_{\text{max,0}} \alpha_i h_i / D_i \alpha_i pO_{2\text{gas}} \), which is a Damkohler number of the second kind for phase \( i \). If the support is impermeable (1/Da = 0), \( f(D_{a,i}) \) reduces to \( D_{a,i} \). Equation 4 applies to the most general situation, and is solved implicitly for \( \theta \) as a function of the dependent variables. In addition, there are several simpler cases of interest (Table 4). 38

The transient response of \( pO_2 \) at the support-medium interface following a step change in \( pO_{2\text{gas}} \) for the situation with no cells present as examined by solving the species conservation equations for the idealized situation of a one-dimensional composite slab with equality of \( pO_2 \) and oxygen flux in each layer at the interface and \( pO_{2\text{gas}} \) specified at both external boundaries. The finite element package COMSOL Multiphysics (COMSOL, Burlington, MA) was used; an analytical solution is also available. 39

Cell Aggregates. Steady state oxygen transport within the medium, supporter layer, and cell aggregate was modeled using the oxygen species conservation equation

\[ D_i \alpha_i \nabla^2 pO_2 = \frac{V_i}{v_{\text{cell}}} \] (11)

where \( V_i \) is OCR of the cell aggregate and is zero elsewhere. The boundary conditions for the model were \( pO_2 = pO_{2\text{gas}} \) at the gas/liquid and gas/support interfaces, and continuity of flux and \( pO_2 \) at the support/medium, support/tissue, and medium/tissue interfaces. The conservation equations and boundary conditions were solved by the finite element method (COMSOL Multiphysics).

\( v_{\text{cell}} \), the specific volume of a cell estimated from light microscopic measurements of EB diameters with a calibrated reticule and nuclei counts from dispersed EBs, was 10.1 ± 1.1, 11.2 ± 1.4, and 12.7 ± 1.5 × 10⁻¹⁰ cm³/cell for day 2 EBs at \( pO_{2\text{gas}} \) of 7, 36, and 142 mm Hg, respectively. Regions of the 5-μm thick histological sections encompassing aggregated cells were examined by light microscopy, and the number of nuclei whose center point was within the volume of tissue contained in the region was estimated. Assuming that the nuclei were spheres with a diameter of 7 μm, \( v_{\text{cell}} \) was estimated to be 15 ± 4, 14 ± 6, and 14 ± 7 × 10⁻¹⁰ cm³ for cells cultured for 11 days at \( pO_{2\text{gas}} \) of 7, 36, and 142 mm Hg, respectively. Cell size varied significantly within different regions of the tissue, and this heterogeneity was not considered in the models. Additional details are available. 36

Geometry Used for Finite Element Model. The day 2 EB was modeled as a sphere of radius 99, 115, and 114 μm for \( pO_{2\text{gas}} \) of 7, 36, and 142 mm Hg, respectively, resting on the surface of a silicone rubber membrane, \( h_s = 127 \mu \), and with \( h_m = 5 \mu \). Each EB sphere or aggregate was modeled as being in the center of a cylindrical well with a 1.5 mm radius. Parameters for day 11 aggregates were obtained using light microscopy and stereological point counting of at least 300 points in the 5-μm tissue sections from three independent experiments, such as the one shown in Figure 2. The volume fraction of tissue that was present in aggregate and sheet-like morphologies was determined. Aggregates accounted for 53, 75, and 90% of the total tissue volume for cells cultured at \( pO_{2\text{gas}} \) of 7, 36, and 142 mm Hg, respectively. Sheets were modeled as a thin slab with a thickness of 15 μm. The distance from the basal to apical surface (tissue thickness) and the length of each aggregate was recorded, and results are summarized in Table 5. Each aggregate was approximated as an oblate hemispheroid with major and minor axes that corresponded to the measured height and half width, respectively.

Distribution of \( pO_2 \) in Tissue. The numerical solution of Eq. 11 yielded the distribution of \( pO_2 \) throughout each of the individual spherical hemispheroidal and planar tissue geometries analyzed. For each geometry simulated, surfaces of constant \( pO_2 \) were determined, from which contours were plotted. In addition, numerical integration was performed to determine the volume of tissue between the surfaces of constant \( pO_2 \). For the average contribution of aggregates and sheets in an actual culture dish as determined from the 5-μm sections, the volume fraction of tissue within specific limits of \( pO_2 \) was determined from

\[ \Phi_{ab} = \left[ (\Phi_{\text{sheet}}) (\Phi_{\text{aggregate}}) \right] + \phi_{\text{aggregate}} V_{\text{aggregate},n} \] (12)

where \( \Phi_{ab} \) is the total volume fraction with a \( pO_2 \) between \( a \) and \( b \), \( \Phi_{\text{sheet}} \) and \( \Phi_{\text{aggregate}} \) are the volume fractions of tissue in the total preparation in the sheet and aggregate geometry, respectively, \( V_{\text{aggregate},n} \) is the volume of tissue with a \( pO_2 \) between \( a \) and \( b \) in the \( j \)th aggregate of cells, and \( V_{\text{aggregate},n} \) is the total volume of the \( n \) aggregates analyzed.

Statistics

Statistical comparisons of data were performed using unpaired t-tests. Results were deemed significant if the two-tailed \( P \)-value was 0.05 or less.

Results

Theoretical comparison of materials for monolayer culture

Steady State. The effect of oxygen diffusion limitations in the medium and the support layers on \( pO_{2\text{cell}} \) relative to
Table 4. Solutions to Estimate \( \text{pO}_2 \text{cel} \) in Cell Monolayer

<table>
<thead>
<tr>
<th>Case</th>
<th>Limits</th>
<th>Expression</th>
<th>Equations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Eq. 2: ( \text{pO}<em>2 )-dependent ( V</em>{\text{max}} )</td>
<td>Eq. 1: Michaelis-Menten kinetics</td>
<td>( (1 - \theta) = \left[ F_{\text{V}} + (1 - F_{\text{V}}) \left( \frac{\theta}{\beta_{\text{V}} + \theta} \right) \right] \left[ \frac{\theta}{\beta_{\text{M}} + \theta} \right] f(Da) )</td>
<td>(4)</td>
</tr>
<tr>
<td>Eq. 2: Constant ( V_{\text{max}} )</td>
<td>Eq. 1: Michaelis-Menten kinetics</td>
<td>( \theta = -\frac{1}{2} \left{ (\beta_{\text{M}} + f(Da)) - 1 \right} + \left[ (\beta_{\text{M}} + f(Da)) - 1 \right]^2 + 4\beta_{\text{M}} } )</td>
<td>(6)</td>
</tr>
<tr>
<td>Zero order ( \beta_{\text{M}} \ll 0 )</td>
<td></td>
<td>( \theta = 1 - f(Da) )</td>
<td>(7)</td>
</tr>
<tr>
<td>First order ( \beta_{\text{M}} \gg 0 )</td>
<td></td>
<td>( \theta = \frac{\beta_{\text{M}}}{f(Da) + \beta_{\text{M}}} )</td>
<td>(8)</td>
</tr>
<tr>
<td>Eq. 2: ( \text{pO}<em>2 )-dependent ( V</em>{\text{max}} )</td>
<td>Eq. 1: Zero order kinetics</td>
<td>( \theta = -\frac{1}{2} \left{ (\beta_{\text{V}} + f(Da)) - 1 \right} + \left[ (\beta_{\text{V}} + f(Da)) - 1 \right]^2 + 4\beta_{\text{V}} } )</td>
<td>(9)</td>
</tr>
<tr>
<td>Zero order ( \beta_{\text{V}} \ll 0 )</td>
<td></td>
<td>( \theta = 1 - f(Da) )</td>
<td>(7)</td>
</tr>
<tr>
<td>First order ( \beta_{\text{V}} \gg 0 )</td>
<td></td>
<td>( \theta = \frac{\beta_{\text{V}}}{1 + (1 - F_{\text{V}}) f(Da)} )</td>
<td>(10)</td>
</tr>
</tbody>
</table>

\(^*\)The solution for this case with Da = 0 and \( f(Da) = Da_{\text{m}} \) was previously described.\(^{38}\)

Figure 2. Representative hematoxylin-stained histological section used to estimate tissue geometry of cell aggregates for numerical solution of oxygen species conservation equation, Eq. 11.

This section shown is from tissue cultured at a \( \text{pO}_2 \text{gas} \) of 36 mm Hg.

\( \text{pO}_2 \text{gas} \) is shown in Figure 3 for three values of \( \text{pO}_2 \text{gas} \). Da for each phase increases with increasing cell density, maximal V, and height and with decreasing oxygen diffusivity and solubility and \( \text{pO}_2 \text{gas} \). As Da of either phase increases, \( f(Da) \) decreases, and the ratio \( \theta = \text{pO}_2 \text{cel} / \text{pO}_2 \text{gas} \) decreases. The various cases begin to diverge from each other for a \( f(Da) \) in the range of about 0.1-0.4. The most general case, Eq. 4, which includes both Michaelis-Menten kinetics and \( \text{pO}_2 \)-dependent \( V_{\text{max}} \), drops most slowly because it entails the largest decrease in V with decreasing \( \text{pO}_2 \). If \( V_{\text{max}} \) is constant, Eq. 6, the curve has similar shape but lower magnitude than Eq. 4. For both cases, if Michaelis-Menten is replaced by zero order kinetics, Eqs. 7 and 9, the curves drop precipitously as \( \theta \) approaches zero. Equation 7 represents global zero-order kinetics.

The rates of oxygen diffusion through the medium and support relative to the rate of oxygen consumption is embodied in the function \( f(Da) \), which depends on the individual Da of each phase. The phase-dependent properties are summarized by the group \( D_{\text{eff}} h_t / h_t \) (the inverse of the

Table 5. Data from Sections Used for Finite Element Models

<table>
<thead>
<tr>
<th>( \text{pO}_2 \text{gas} ) (mm Hg)</th>
<th>7</th>
<th>36</th>
<th>142</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of aggregates analyzed</td>
<td>21</td>
<td>19</td>
<td>12</td>
</tr>
<tr>
<td>Fraction of tissue in aggregates (%)</td>
<td>53</td>
<td>75</td>
<td>90</td>
</tr>
<tr>
<td>Aggregate height (( \mu \text{m} ))</td>
<td>Mean</td>
<td>120</td>
<td>190</td>
</tr>
<tr>
<td>Std Dev</td>
<td>60</td>
<td>100</td>
<td>150</td>
</tr>
<tr>
<td>Aggregate width (( \mu \text{m} ))</td>
<td>Mean</td>
<td>450</td>
<td>630</td>
</tr>
<tr>
<td>Std Dev</td>
<td>290</td>
<td>320</td>
<td>370</td>
</tr>
</tbody>
</table>

5-\( \mu \text{m} \) tissue sections from three experiments were analyzed to obtain morphology estimates that could be used in finite element models. Stereological point counting was performed to determine the volume fraction of tissue that was present in aggregates and in cell sheets. The height of each aggregate was measured as the distance from its basal to apical surface using a calibrated reticle, and the length was measured in a similar manner. The basal surface was determined by the shape of the aggregate and the surface which was continuous with the cell sheet.
by value for medium, oxygen is supplied almost exclusively through the medium, and \( f(Da) \) is of the order of 1. The most permeable material is silicone rubber, and \( D_{\text{polystyrene}}/h_p \) is about \( 10^2 \) higher than medium. Consequently, \( f(Da) \) is typically less than \( 10^{-1} \) and \( \theta \) is close to 1, so that virtually all of the oxygen is supplied by diffusion through the membrane.

The effects of the different support materials in determining actual values of \( p_{O_2,\text{cell}} \) calculated from Eqs. 4 or 6 are shown in Figure 4 as a function of cell density for medium heights ranging from 1 to 5 mm. Panels A, B, and C on the left apply to the OCR parameters of CCE mES cells for which \( V_{max} \) decreases with \( p_{O_2,\text{cell}} \); panels D, E, and F on the right are illustrative of typical adult mammalian cells, which have a higher \( V_{max} \) that is independent of \( p_{O_2,\text{cell}} \) and leads to even greater effects of oxygen diffusion limitations. Predicted values of \( p_{O_2,\text{cell}} \) are very sensitive to medium height and cell surface density for culture on polystyrene, less so for FEP copolymer, and virtually not at all with silicone rubber if realistically large medium heights are used.

**Transient Response.** The predicted response of \( p_{O_2} \) at the medium-support interface of an idealized one-dimension system following a step change in \( p_{O_2,\text{gas}} \) was determined for each of the three support materials (Figure 5A). The equilibration time was quite long on a polystyrene dish and was highly dependent on the medium height: approximately 1, 4, and \( 25 \times 10^3 \) sec were required to reach 90% of the steady state value at medium depths of 1, 2, and 5 mm, respectively, in a polystyrene dish. The time required was about 100 and 500 sec at depths of 1 and 5 mm, respectively, on an FEP-teflon membrane, and about 10 seconds in a silicone rubber dish with any medium height.

To verify that silicone rubber dishes set up as for normal cell culture reached steady-state very rapidly compared to polystyrene, \( p_{O_2} \) at the surface of silicone rubber membrane-based and polystyrene dishes was measured in an acellular system in response to a change in \( p_{O_2,\text{gas}} \) (Figure 5B) with a fiberoptic sensor in contact with each bottom surface. The incubator changed \( p_{O_2,\text{gas}} \) from 160 to 7 mm Hg in \( \sim 15 \) min; \( p_{O_2} \) at the silicone rubber surface lagged \( p_{O_2,\text{gas}} \) by several minutes. The polystyrene dish required significantly longer time, \( \sim 20 \) hr, to reach steady-state.

**Theoretical comparison of materials for aggregate culture**

In order to explore the relationships between oxygen diffusion in the medium, tissue, and different support materials, the \( p_{O_2} \) profile within wells containing hypothetical 100-\( \mu \)m radius hemispherical cell aggregates spaced 1 mm apart (2.3 \( \times 10^5 \) cell/cm\(^3\)) was calculated for three different \( p_{O_2,\text{gas}} \) (Figure 6). In contrast to the behavior with monolayers, \( p_{O_2} \) varied within the aggregate. At a \( p_{O_2,\text{gas}} \) of 142 mm Hg (Figure 6A,B), there was a 55 mm Hg drop in \( p_{O_2} \) from the gas to the upper surface of the aggregate (\( \Delta p_{O_2,\text{gas}} \)) on polystyrene dishes, which was slightly larger than the \( \Delta p_{O_2} \) of 40 mm Hg for a cell monolayer at the same overall average density on the support. \( \Delta p_{O_2,\text{gas}} \) was reduced to 24 and 4 mm Hg on FEP and silicone rubber membranes, respectively. The maximum \( p_{O_2} \) decrease within the aggregate was 45, 30, and 27 mm Hg on polystyrene, FEP and silicone rubber, respectively. The total \( p_{O_2} \) drop from the gas to the region of the aggregate with the lowest local \( p_{O_2} \) was

![Image](https://example.com/image.png)

Figure 3. Steady-state reduction in \( p_{O_2,\text{cell}} \) relative to \( p_{O_2,\text{gas}} \) as a result of oxygen diffusion limitations in the support layer (s) and the medium (m) for cells in a monolayer.

\( \theta \) is plotted as a function of \( f(Da) \), defined by Eq. 5, for fixed values of parameters \( F_a = 1/3, k_a = 16 \) mm Hg, \( k_b = 0.44 \) mm Hg at different values of \( p_{O_2,\text{gas}} \): (A) 142 mm Hg, \( \beta_a = 0.113, \beta_b = 3.1 \times 10^{-3} \); (B) 36 mm Hg, \( \beta_a = 0.444, \beta_b = 0.012 \); (C) 7 mm Hg, \( \beta_a = 2.29, \beta_b = 0.063 \). Equation numbers refer to equations in Table 4.

---

mass transfer resistance), which are tabulated in Table 1 together with their value relative to that of the medium. Because of its low permeability and large thickness, \( D_{\text{polystyrene}}/h_p \) for polystyrene is on the order of \( 10^2 \) lower than the value for medium, oxygen is supplied almost exclusively by diffusion through the medium, and \( f(Da) \) is large, typically well in excess of 1, which is associated with low values of \( \theta \) (Figure 3). The mass transfer resistances of FEP are comparable to that of medium, and \( f(Da) \) is of the order of 1. The most permeable material is silicone rubber, and \( D_{\text{polystyrene}}/h_p \) is about \( 10^2 \) higher than medium. Consequently, \( f(Da) \) is typically less than \( 10^{-1} \) and \( \theta \) is close to 1, so that virtually all of the oxygen is supplied by diffusion through the membrane.

The effects of the different support materials in determining actual values of \( p_{O_2,\text{cell}} \) calculated from Eqs. 4 or 6 are shown in Figure 4 as a function of cell density for medium heights ranging from 1 to 5 mm. Panels A, B, and C on the left apply to the OCR parameters of CCE mES cells for which \( V_{max} \) decreases with \( p_{O_2,\text{cell}} \); panels D, E, and F on the right are illustrative of typical adult mammalian cells, which have a higher \( V_{max} \) that is independent of \( p_{O_2,\text{cell}} \) and leads to even greater effects of oxygen diffusion limitations. Predicted values of \( p_{O_2,\text{cell}} \) are very sensitive to medium height and cell surface density for culture on polystyrene, less so for FEP copolymer, and virtually not at all with silicone rubber if realistically large medium heights are used.

**Transient Response.** The predicted response of \( p_{O_2} \) at the medium-support interface of an idealized one-dimension system following a step change in \( p_{O_2,\text{gas}} \) was determined for each of the three support materials (Figure 5A). The equilibration time was quite long on a polystyrene dish and was highly dependent on the medium height: approximately 1, 4, and \( 25 \times 10^3 \) sec were required to reach 90% of the steady state value at medium depths of 1, 2, and 5 mm, respectively, in a polystyrene dish. The time required was about 100 and 500 sec at depths of 1 and 5 mm, respectively, on an FEP-teflon membrane, and about 10 seconds in a silicone rubber dish with any medium height.

To verify that silicone rubber dishes set up as for normal cell culture reached steady-state very rapidly compared to polystyrene, \( p_{O_2} \) at the surface of silicone rubber membrane-based and polystyrene dishes was measured in an acellular system in response to a change in \( p_{O_2,\text{gas}} \) (Figure 5B) with a fiberoptic sensor in contact with each bottom surface. The incubator changed \( p_{O_2,\text{gas}} \) from 160 to 7 mm Hg in \( \sim 15 \) min; \( p_{O_2} \) at the silicone rubber surface lagged \( p_{O_2,\text{gas}} \) by several minutes. The polystyrene dish required significantly longer time, \( \sim 20 \) hr, to reach steady-state.

**Theoretical comparison of materials for aggregate culture**

In order to explore the relationships between oxygen diffusion in the medium, tissue, and different support materials, the \( p_{O_2} \) profile within wells containing hypothetical 100-\( \mu \)m radius hemispherical cell aggregates spaced 1 mm apart (2.3 \( \times 10^5 \) cell/cm\(^3\)) was calculated for three different \( p_{O_2,\text{gas}} \) (Figure 6). In contrast to the behavior with monolayers, \( p_{O_2} \) varied within the aggregate. At a \( p_{O_2,\text{gas}} \) of 142 mm Hg (Figure 6A,B), there was a 55 mm Hg drop in \( p_{O_2} \) from the gas to the upper surface of the aggregate (\( \Delta p_{O_2,\text{gas}} \)) on polystyrene dishes, which was slightly larger than the \( \Delta p_{O_2} \) of 40 mm Hg for a cell monolayer at the same overall average density on the support. \( \Delta p_{O_2,\text{gas}} \) was reduced to 24 and 4 mm Hg on FEP and silicone rubber membranes, respectively. The maximum \( p_{O_2} \) decrease within the aggregate was 45, 30, and 27 mm Hg on polystyrene, FEP and silicone rubber, respectively. The total \( p_{O_2} \) drop from the gas to the region of the aggregate with the lowest local \( p_{O_2} \) was
Figure 4. Predicted $pO_{2cell}$ of cell monolayers as a function of cell density $\rho$ and medium height $h_m$ for culture on polystyrene (P), fluorinated ethylene-propylene copolymer (FEP) membrane, and silicone rubber (SR) membrane surfaces at $pO_{2gas}$ of (A and D) 142, (B and E) 36, and (C and F) 7 mm Hg.

The panels on the left (A, B, and C) are for mES cells for which OCR parameters $^3$ are given in conjunction with Eq. 2, oxygen diffusivity and solubility parameters are from Table 1, and the $pO_{2cell}$ was determined using Eq. 4. The panels on the right (D, E, and F) are for cells having constant $V_{\text{mm}} = 50$ amol/sec cell, which is typical for mammalian fibroblasts. $^4$ $pO_{2cell}$ was determined using Eq. 6. A single line is shown for SR because the prediction is virtually independent of $h_m$.

Greater than 98% of undifferentiated mES cells were attached to uncoated polystyrene wells after a day in culture (Table 6). Coating polystyrene with fibronectin or to a lesser extent gelatin caused the cells to have a more widely spread morphology but did not affect cell proliferation (Figure 7). Cells did not attach to well bottoms made of silicone rubber that were untreated or coated with gelatin. When mES cells were placed into such wells, they formed freely floating EBs after 24 hr in culture. Exposure of the membrane to 2 $\mu$g/mL fibronectin in DPBS (or a serum-free ITS medium) for a day before addition of the cells was sufficient for attachment of 99% of the cells. There was a small but significant decrease in the total cell number on the silicone rubber treated with fibronectin relative to the polystyrene tissue culture plastic. Cells would not attach to the silicone rubber surface when...
Figure 5. (A) The fractional approach of $P_{O_2}$ at the medium-support interface to steady state after a step change in $P_{O_2\text{gas}}$ as estimated using numerical simulations in a cell-free system with a polystyrene (P), fluorinated ethylene-propylene copolymer (FEP), and silicone rubber (SR) bottom. Numbers refer to medium height in mm. A single line is shown for SR because the prediction is virtually independent of $h_m$. (B) Transient response of the $P_{O_2}$ measured with a fiberoptic sensor in contact with the culture surface in acellular SR and P wells of a 24-well plate filled with 2 mL DBPS ($h_m = 10.5$ mm) following a change in $P_{O_2\text{gas}}$ setting from 160 to 7 mm Hg.

Figure 6. Theoretical predictions for $P_{O_2}$ profiles within wells containing hypothetical 100-μm radius hemispherical aggregates spaced 1 mm apart (overall average on support $2.3 \times 10^6$ cells/cm²) on polystyrene, FEP membranes, or silicone rubber membranes at the bottom for $P_{O_2\text{gas}}$ of 142, 36, and 7 mm Hg. The $P_{O_2}$ profile shown is on a line passing through the center of an aggregate and perpendicular to the bottom of the well. The dashed lines on the plots indicate the locations of the upper and lower aggregate surfaces. Panel (A) shows the entire distance from the top of the medium to the bottom of the culture vessel, (B), (C), and (D) are expanded to show regions within 100 μm of the top and bottom surfaces of the aggregate. Distances greater than 2 mm represent areas within the support, and the plotted lines showing the local $P_{O_2}$ profiles terminate at the lower surface of the culture vessel, the location of which is different for each of the three different support materials. Parameters used in the calculations are for mES cells.

The 2 μg/mL fibronectin solution was made in DMEM with 10% FBS (data not shown). This may have resulted from competitive adsorption of other serum proteins onto the silicone surface, which limited fibronectin adsorption. Additionally, poor cell attachment was observed when cells were cultured in mES cell differentiation medium (Table 2) with 2% instead of 10% FBS, which could be rectified by absorbing Matrigel instead of fibronectin (data not shown).

Deposition of a (PSS-PEI)$_4$-(PDL-gelatin)$_3$ PEM onto silicone rubber before cell addition could be used to promote cell adhesion (Table 6). Cells adhered better to silicone rubber with a PEM than they did to unmodified silicone, but the total number of attached cells was lower on PEM-treated silicone rubber than any of the polystyrene surfaces tested. The PEM-gelatin structures did not support attachment as well as the single layer of fibronectin adsorbed.
Differentiation to cardiomyocytes on silicone rubber and polystyrene surfaces

Differentiation of mES cells to cardiomyocytes is an oxygen-sensitive cell culture process. We investigated if culture on gas-permeable (silicone rubber) or gas-impermeable (polystyrene alone and silicone rubber membrane glued with silicone adhesive on top of polystyrene) surfaces would lead to different results, since the cells would be exposed to different PO2gas conditions. To illustrate the difference in PO2cell between the gas-permeable and gas-impermeable substrates, approximate estimates of PO2cell were calculated using Eq. 4 by assuming that all of the attached cells measured at each condition were distributed in a uniform layer (Table 7). Culture on gas-permeable silicone rubber resulted in values of PO2cell within 2 mm Hg of PO2gas. In contrast, culture on the gas-impermeable substrates resulted in very small values of PO2cell (<1 mm Hg) at all PO2gas, pO2cell was slightly higher for cells on silicone rubber on polystyrene compared to polystyrene alone because of slightly higher numbers of attached cells for the latter case.

Spontaneous contraction was not observed at any PO2gas condition on polystyrene in multiple experiments (Table 8), even though cells stained positive with MF-20. Contractions were observed on silicone rubber membranes alone and on silicone rubber on top of polystyrene. The largest fractional area of beating cells occurred on silicone rubber alone at a PO2gas of 36 mm Hg, and the smallest was on silicone rubber on top of polystyrene at 7 mm Hg. Although there was substantial variation between beating area assessments, with biological replicates at the same condition, the average fraction of the entire well area covered by spontaneously contracting cells for the latter case.

Table 6. Cell Attachment to Polystyrene and Modified Silicone Rubber Surfaces

<table>
<thead>
<tr>
<th>Surface Treatment</th>
<th>Cell Number (% of Total on Polystyrene)</th>
<th>Fraction Attached (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Total</td>
<td>Attached</td>
</tr>
<tr>
<td>Tissue culture polystyrene</td>
<td>100 ± 7</td>
<td>98 ± 7</td>
</tr>
<tr>
<td>Unmodified*</td>
<td>95 ± 5</td>
<td>95 ± 5</td>
</tr>
<tr>
<td>Gelatin A&quot;</td>
<td>98 ± 2</td>
<td>97 ± 2</td>
</tr>
<tr>
<td>Gelatin B&quot;</td>
<td>110 ± 16</td>
<td>109 ± 16</td>
</tr>
<tr>
<td>Silicone rubber</td>
<td>69 ± 9</td>
<td>39 ± 18</td>
</tr>
<tr>
<td>Untreated*</td>
<td>71 ± 0</td>
<td>46 ± 3</td>
</tr>
<tr>
<td>Gelatin A only*</td>
<td>91 ± 1</td>
<td>90 ± 1</td>
</tr>
<tr>
<td>Gelatin B only*</td>
<td>88 ± 14</td>
<td>85 ± 14</td>
</tr>
<tr>
<td>PEM-gelatin A</td>
<td>80 ± 5</td>
<td>76 ± 5</td>
</tr>
<tr>
<td>PEM-gelatin B</td>
<td>64 ± 5</td>
<td>61 ± 5</td>
</tr>
</tbody>
</table>

*Conditions shown in Figure 7.

Approximately 2 x 10⁵ viable cells were placed into each well of a 24-well plate and cultured for about 24 hr before assessing cell attachment. Data are from two separate experiments each with an untreated polystyrene control condition. Results are reported as the mean ± SD for triplicate wells for each condition, and are normalized to the total cell number in the untreated polystyrene control for the experiment from which the data were taken.

Figure 7. Adhesion of mES cells to polystyrene and silicone rubber in their native states or after incubation with PBS, 0.1% gelatin in water, or 2 µg/mL fibronectin in PBS for 24 hr before cell addition.
The fraction of MF-20+ cells was determined by flow cytometry of trypsin-dispersed cell samples. Data presented were obtained by flow cytometry of trypsin-dispersed cell samples. Essentially identical data was obtained when no adhesive was used. All experiments were started with 30 EBs/well (0.15 x 10^3 cells/well). Data are shown as mean ± SD for four replicate wells in a single experiment. * * indicates statistical difference (P < 0.05) between value on SR compared to P and SR-P at the same pO_{2gas}.

**Discussion**

We studied silicone rubber membrane as a support for the precise control of the pO_{2cell} during culture with cell monolayers and aggregates. We also investigated surface modifications to promote cell attachment to the silicone rubber membrane and further proliferated on the silicone rubber membrane; the idealized spherical geometry was lost, and approximations of aggregate shapes were used in the model based on measurements from sectioned tissue (Figure 2). Cells cultured at a high pO_{2gas} grew to form larger aggregates, with a mean height after 11 days in culture of 120, 190, and 280 μm at pO_{2gas} of 7, 36, and 142 mm Hg, respectively (Table 5). The axially-symmetric three-dimensional pO_{2cell} distribution within the aggregates having mean dimensions tabulated in Table 5 was obtained by numerical solution of Eq. 11, from which the pO_{2cell} contours in Figure 10 were evaluated. Calculated pO_{2} profiles in sheet-like tissue and all aggregates were used to determine the volumetric distribution of pO_{2cell} in Figure 11 for the entire tissue mass at each condition.

In spherical EBs after 2 days culture, about 60% of the tissue was exposed to pO_{2cell} less than 1 mm Hg during culture at pO_{2gas} of 7 mm Hg, whereas more than 80% of tissue cultured at 36 mm Hg was at pO_{2cell} greater than 7 mm Hg, and all of the tissue cultured at 142 mm Hg was at pO_{2cell} greater than 94 mm Hg (Figure 11). After 11 days of culture, about 28%, 26%, and 15% of the total volume of tissue was at a pO_{2cell} of less than the K_{m} for oxygen consumption (0.44 mmHg) at 7, 36, and 142 mmHg, respectively. However, a large portion of the tissue remained at a pO_{2cell} that was relatively near that of the gas phase (Figures 10 and 11). In contrast, if aggregates with similar dimensions and cell numbers had been cultured on a polystyrene dish at 142 mm Hg, ~80% of the aggregate volume would have been at a pO_{2cell} less than 0.44 mm Hg. Oxygen effects would therefore have been impossible to accurately study on polystyrene, because all pO_{2gas} conditions would have had extreme oxygen starvation.

pO_{2cell} during cardiomyocyte differentiation on silicone rubber membranes

After 2 days in hanging drops, the differentiating cells were in the form of spherical EBs of radius about 100 μm and the predicted oxygen gradients were relatively small. At times later than 2 days, the EBs were allowed to attach the fibronectin-coated silicone rubber surface. The cells spread and further proliferated on the silicone rubber membrane; the idealized spherical geometry was lost, and approximations of aggregate shapes were used in the model based on measurements from sectioned tissue (Figure 2). Cells cultured at a high pO_{2gas} grew to form larger aggregates, with a mean height after 11 days in culture of 120, 190, and 280 μm at pO_{2gas} of 7, 36, and 142 mm Hg, respectively (Table 5). The axially-symmetric three-dimensional pO_{2cell} distribution within the aggregates having mean dimensions tabulated in Table 5 was obtained by numerical solution of Eq. 11, from which the pO_{2cell} contours in Figure 10 were evaluated. Calculated pO_{2} profiles in sheet-like tissue and all aggregates were used to determine the volumetric distribution of pO_{2cell} in Figure 11 for the entire tissue mass at each condition.

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Table 7. Calculation of \( pO_{2\text{cell}} \) for Day 11 Differentiated mES Cells

<table>
<thead>
<tr>
<th>Material</th>
<th>( pO_{2\text{gas}} ) (mm Hg)</th>
<th>Cell Density (10^5 cells/cm^2)</th>
<th>( pO_{2\text{cell}} ) (mm Hg)</th>
<th>( pO_{2\text{gas}} - pO_{2\text{cell}} ) (mm Hg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>P</td>
<td>142</td>
<td>7.7</td>
<td>0.830</td>
<td>141</td>
</tr>
<tr>
<td>P</td>
<td>36</td>
<td>6.2</td>
<td>0.125</td>
<td>35.9</td>
</tr>
<tr>
<td>P</td>
<td>7</td>
<td>4.8</td>
<td>0.027</td>
<td>6.97</td>
</tr>
<tr>
<td>SR-P</td>
<td>142</td>
<td>7.3</td>
<td>0.921</td>
<td>141</td>
</tr>
<tr>
<td>SR-P</td>
<td>36</td>
<td>4.5</td>
<td>0.193</td>
<td>35.8</td>
</tr>
<tr>
<td>SR</td>
<td>7</td>
<td>3.7</td>
<td>0.035</td>
<td>6.97</td>
</tr>
<tr>
<td>SR</td>
<td>142</td>
<td>15.1</td>
<td>1.40</td>
<td>1.96</td>
</tr>
<tr>
<td>SR</td>
<td>36</td>
<td>15.9</td>
<td>34.3</td>
<td>1.73</td>
</tr>
<tr>
<td>SR</td>
<td>7</td>
<td>5.4</td>
<td>6.62</td>
<td>0.377</td>
</tr>
</tbody>
</table>

P, polystyrene; SR-PS, silicone rubber on polystyrene; SR, silicone rubber.

*Calculated with Eq. 4.

Table 8. Comparison of Beating Areas on Different Culture Surfaces

<table>
<thead>
<tr>
<th>Material</th>
<th>( pO_{2\text{gas}} ) (mm Hg)</th>
<th>Fractional Beating Area (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>P</td>
<td>142</td>
<td>0</td>
</tr>
<tr>
<td>P</td>
<td>36</td>
<td>0</td>
</tr>
<tr>
<td>P</td>
<td>7</td>
<td>0</td>
</tr>
<tr>
<td>SR-P</td>
<td>142</td>
<td>1.7 ± 1.6</td>
</tr>
<tr>
<td>SR-P</td>
<td>36</td>
<td>2.5 ± 1.8</td>
</tr>
<tr>
<td>SR-P</td>
<td>7</td>
<td>0.8 ± 0.4</td>
</tr>
<tr>
<td>SR</td>
<td>142</td>
<td>3.3 ± 2.6</td>
</tr>
<tr>
<td>SR</td>
<td>36</td>
<td>9.6 ± 3.5</td>
</tr>
<tr>
<td>SR</td>
<td>7</td>
<td>2.6 ± 1.2</td>
</tr>
</tbody>
</table>

P, polystyrene; SR-P, silicone rubber on polystyrene; SR, silicone rubber, \( n = 4 \).

*Fraction of entire well area.

![Figure 9](image_url)
number of cardiomyocyte-like cells in $p_{O_2_{gas}}$ conditions less than 142 mm Hg using a gas-impermeable dish. Extensive cell contraction was observed on silicone rubber alone or on top of polystyrene, but not on polystyrene alone, for which there are at least two possible explanations: (1) the $p_{O_2}$ underneath cell layers with sheets morphology (Figure 2) is higher than under adjacent aggregates, which contain cardiomyocyte-like cells. $p_{O_2}$ affects the beating phenotype of mES cell-derived cardiomyocytes, and slight differences in $p_{O_2_{cell}}$ could explain the difference in the appearance of beating areas on polystyrene and silicone rubber on polystyrene. (2) silicone rubber is much more elastic than polystyrene, and elasticity affects the beating phenotype of embryonic cardiomyocytes; cells on more rigid substrates stop beating while cells on more elastic substrates continue to beat. This issue deserves further investigation. In this experimental system, the use of nearly gas-impermeable dishes to study oxygen effects gives misleading results as a consequence of the large differences between the $p_{O_2_{gas}}$ and $p_{O_2_{cell}}$.

Our work shows that culture on silicone rubber membrane offers substantial improvements in tissue oxygenation over polystyrene and FEP membrane-based culture vessels, and that physical adsorption of fibronectin or matrigel to silicone rubber surfaces can be used to promote prolonged mES cell attachment. These findings strongly suggest that silicone rubber membrane-based dishes should be used whenever possible if oxygen is a variable of interest.

**Acknowledgment**

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**Literature Cited**


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B. Effects of oxygen on mouse embryonic stem cell
growth, phenotype retention, and cellular energetics

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Effects of oxygen on mouse embryonic stem cell growth, phenotype retention,
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**Editors' Choice**

**Biotechnology & Bioengineering**

**Effects of Oxygen on Mouse Embryonic Stem Cell Growth, Phenotype Retention, and Cellular Energetics**

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**ABSTRACT:** Most embryonic stem (ES) cell research is performed with a gas phase oxygen partial pressure (pO₂) of 142 mmHg, whereas embryonic cells in early development are exposed to pO₂ values of 0–30 mmHg. To understand effects of these differences, we studied murine ES (mES) growth, maintenance of stem cell phenotype, and cell energetics over a pO₂ range of 0–285 mmHg, in the presence or absence of differentiation-suppressing leukemia inhibitory factor (LIF). With LIF, growth rate was sensitive to pO₂ but constant with time, and expression of self-renewal transcription factors decreased at extremes of pO₂. Subtle morphological changes suggested some early differentiation, but cells retained the ability to differentiate into derivatives of all three germ layers at low pO₂. Without LIF, growth rate decreased with time, and self-renewal transcription factor mRNA decreased further. Gross morphological changes occurred, and overt differentiation occurred at all pO₂. These findings suggested that hypoxia in the presence of LIF promoted limited early differentiation. ES cells survived oxygen starvation with negligible cell death by increasing anaerobic metabolism within 48 h of anoxic exposure. Decreasing pO₂ to 36 mmHg or lower decreased oxygen consumption rate and increased lactate production rate. The fraction of ATP generated aerobically was 60% at or above 142 mmHg and decreased to 0% under anaoxia, but the total ATP production rate remained nearly constant at all pO₂. In conclusion, undifferentiated ES cells adapt their energy metabolism to proliferate at all pO₂ between 0 and 285 mmHg. Oxygen has minimal effects on undifferentiated cell growth and phenotype, but may exert more substantial effects under differentiating conditions.

**KEYWORDS:** embryonic stem cells; oxygen; hypoxia; proliferation; differentiation; cellular energetics

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**Introduction**

Embryonic stem (ES) cells have the potential to generate any type of cell for transplantation to treat diseases such as diabetes, leukemia, and Parkinson’s disease (Bonner-Weir et al., 2000; Kaufman et al., 2001; Kim et al., 2002; Soria, 2001). Development of efficient techniques for expansion of undifferentiated cells and subsequent directed differentiation requires an understanding of how ES cells interact with their microenvironment and methods to control these interactions. The oxygen partial pressure (pO₂) to which the cell is exposed (pO₂cell) is potentially one important environmental parameter.

Mammalian tissues are usually exposed to pO₂ values far below atmospheric levels, such as 25 mmHg in the rat microvasculature (Intaglietta et al., 1996), 15–45 mmHg in the reproductive tract of rhesus monkeys (Fischer and Bavister, 1993), and 25–30 mmHg in venous human fetal blood in the second and third trimester (Sigaard-Andersen and Huch, 1995; Soothill et al., 1986). Lower values occur in the gestational sac during the first trimester (Jauniaux et al., 2003b), which also provides an environment containing antioxidant molecules to further reduce oxidative damage (Jauniaux et al., 2003a). pO₂ values within early developing embryos are likely to be even lower because circulation does not occur until a well-formed vascular network is present (Malpepe and Simon, 1998; Palis et al., 2001), and there are hypoxic regions even after a well-developed vascular network exists (Land, 2004; Lee et al., 2001).

Stem cell culture is usually performed at pO₂ levels much higher than that of the embryo in vivo, typically in a humidified atmosphere consisting of 95% air/5% CO₂ that results in a gas phase pO₂ (pO₂gas) of 142 mmHg. Most experiments with ES cells are carried out in polystyrene culture dishes, which have very low oxygen permeability, and oxygen is supplied by diffusion through the medium. Oxygen concentration gradients exist in the medium that cause pO₂cell to be different from pO₂gas (Metzen et al., 1995;

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PO2cellu depends on several factors, including medium depth, cell density, cellular oxygen consumption rate, and PO2gas, which are usually not taken into account (Csete, 2005; Metzen et al., 1995). As a consequence, the value of PO2cell during ES cell culture is usually not controlled to be in the range experienced by the early embryo. The implications of this poor control are not well understood. Published data obtained with ES and other cell types suggest that cell survival and proliferation, maintenance of the stem cell phenotype, and cellular energetics are three important parameters that can be significantly affected by the magnitude of PO2cell.

Cell proliferation can be adversely affected by too high or too low a PO2. Culture at a PO2 higher than needed exposes cells to higher concentrations of reactive oxygen species (ROS), which damage lipids, proteins, and nucleic acids (Lee and Wei, 2005; Saretzki et al., 2004) and can lead to senescence and cell death (Parrinello et al., 2003). Many cell types have a higher growth rate at PO2gas conditions less than 142 mmHg, including fibroblasts (Bradley, 1978; Parrinello et al., 2003), hematopoietic cells (Bradley, 1978; Hevehan et al., 2000; Mostafa et al., 2000), neural progenitors (Milosevic et al., 2005; Morrison et al., 2000; Studer et al., 2000), muscle satellite cells (Chakravarthy et al., 2001; Csete et al., 2001), CHO and hybridoma cells (Goetz, 1975; Miller et al., 1987), as well as early embryos of both humans and mice (Dumoulin et al., 1999; Orsi and Leese, 2001; Quinn and Harlow, 1978). Providing too little oxygen can also be harmful and lead to apoptotic or necrotic cell death (Brunelle and Chandel, 2002).

In addition to growth, PO2gas can affect whether stem or progenitor cells remain undifferentiated. Culture of mouse and human hematopoietic progenitor (Cipolleschi et al., 1997; Hevehan et al., 2000; Mostafa et al., 2000; Ramirez-Bergeron et al., 2004), rat CNS progenitor (Studer et al., 2000), and rat neural crest stem cells (Morrison et al., 2000) at reduced PO2gas favors self-renewal, while increased PO2gas favors differentiation and maturation. Some studies suggest that reduced PO2gas increases the growth rate of mouse and human ES cells and may be beneficial for maintaining the undifferentiated phenotype, but usually only two or three values of PO2gas were examined, and results are often not in agreement with one another (Carmeliet et al., 1998; Ezashi et al., 2005; Forsyth et al., 2006; Gibbons et al., 2006; Hopfl et al., 2002; Iyer et al., 1998; Kurosawa et al., 2006; Ludwig et al., 2006; Ording et al., 2005; Peura et al., 2005; Wang et al., 2006).

Cellular energetics are affected by PO2cell (Miller et al., 1987), and survival at low oxygen requires anaerobic glycolysis for ATP generation. Mouse, bovine, and human blastocysts have high glycolytic capacity and can readily convert glucose into lactic acid (Devreker and Englert, 2000; Harvey et al., 2002; Hewitson and Leese, 1993; Houghton et al., 1996). Nonetheless, 75–85% of the ATP production in embryos occurs through oxidative phosphorylation at high PO2gas (Houghton, 2006; Houghton et al., 1996; Thompson et al., 1996). The high levels of glycolytic enzymes in ES cells during culture at a PO2gas of 142 mmHg are further increased at reduced PO2gas conditions through the action of HIF-1α (Iyer et al., 1998; Wenger, 2002). Quantitative oxygen consumption and lactate production measurements with ES cells have not been reported, and it is unknown how these parameters are affected by the culture PO2cell.

In this study we performed a systematic quantitative investigation of the effects of PO2 on growth rate, stem cell phenotype, and energetics of murine ES (mES) cells. Varying PO2gas in the range from 0 to 285 mmHg had an effect on the growth rate of undifferentiated mES cells, both with and without inclusion of the differentiation-suppressing cytokine leukemia inhibitory factor (LIF) in the culture medium. High and low PO2gas reduced mRNA levels for transcription factors associated with the maintenance of the undifferentiated phenotype during culture with LIF, but only modest changes characteristic of early differentiation occurred in morphology, and cells retained the ability to differentiate into derivative of all three germ layers. Aerobic metabolism generated approximately 60% of the energy required by mES cells at high oxygen conditions, but substantially smaller fractions if cells were oxygen starved. This shift from aerobic to anaerobic respiration occurred rapidly with minimal cell death.

**Materials and Methods**

**Cells and Media**

Two mES cell lines were used, passage 22 CCE (Keller et al., 1993; Robertson et al., 1986) from Stem Cell Technologies (Vancouver, BC) and unknown passage number D3 (Doetschman et al., 1985) from ATCC (CRL-1934, Manassas, VA). Undifferentiated cells were grown in high glucose DMEM (30-2002, ATCC) supplemented to 10% (v/v) ES cell qualified fetal bovine serum (FBS) (06905, Stem Cell Technologies, or SCRR 30-2020, ATCC). The medium was further supplemented to a final concentration of 1,000 U/mL leukemia inhibitory factor (LIF, ESG1106, Chemicon, Temecula, CA) and 0.1 mM 2-mercaptoethanol (M-7522, Sigma-Aldrich, St. Louis, MO). Differentiation was carried out in an otherwise identical medium without LIF.

**Cell Culture**

Unless otherwise stated, cells were grown with 4 mL of medium in 25 cm² cell culture flasks (353109, Becton Dickinson, Franklin Lakes, NJ) that were treated for 30 min with a sterile 0.1% (w/v) solution of gelatin (G-2500, Sigma-Aldrich) in tissue culture water (25-055-CM, Mediatech, Herndon, VA). Cells were plated at a density of about 12,000 cells/cm². Medium was exchanged daily, and cells were detached with 0.25% trypsin (30-2101, ATCC)
every 2 days. Split fractions were chosen so that cells were always plated at 12,000 cells/cm². Medium was incubated in the appropriate pO₂gas condition for 24 h prior to use.

ES Cell Differentiation

ES cells were differentiated using the culture procedure described above (monolayer differentiation), except that medium without supplemental LIF was used. For one series of experiments, as indicated in the text and figure captions, differentiation was carried out in embryoid bodies (EBs) that were formed in 20 µL hanging drops of DMEM with 10% FBS containing 500 ES cells per drop. After 2 days, 30 EBs were transferred to each well of a gelatin-coated 24-well plate and cultured for another 3 days.

Gas Phase pO₂ Control

Cell culture vessels were placed inside polystyrene chambers (MIC-101, Billups-Rothenburg, Del Mar, CA) contained within a standard incubator (OW12720A, Queue Systems, Parkersburg, WV) maintained at 37°C. An open dish of deionized water in each chamber provided humidification. The desired pO₂gas was attained using premixed gas containing 5% CO₂ and 40%, 20%, 5%, 1%, or 0% O₂ (certified medical gas from Airgas, Hingham, MA), which corresponded under humidified conditions to 285, 142, 36, 7, and 0 mmHg. The flow rate of this gas to the chambers was 2 L/min for 15 min for an initial purge following closure of the chamber (after cell medium exchange or passage); and was 30 mL/min at all other times. To examine the response to a change in pO₂ under simulated culture conditions, a fiber optic sensor from the Micro Oxygen Monitoring System (FO/SYS2-T250, Instech Labs, Plymouth Meeting, PA) that measures oxygen fluorometrically was attached along the bottom of a 25 cm² flask through a hole drilled in the flask with the sensing face of the tip perpendicular to the culture surface. A volume of 4 mL of water equilibrated to room air was added, and the chamber was sealed. The gas flowing into the chamber was changed to nitrogen, and the pO₂ was measured as a function of time.

Immunocytochemistry for ES Cell-Specific Markers Analyzed by Flow Cytometry

Samples with 10⁵ cells were centrifuged for 3 min at 300g, supernatant was removed and replaced with 750 µL of Dulbecco's phosphate buffered saline (DPBS), the cells were resuspended, and 250 µL of 4% (w/v) paraformaldehyde (Alfa Aesar, Ward Hill, MA) in DPBS was added. Cells were fixed for 20 min, washed with 1 mL DPBS, and resuspended in 100 µL of DPBS. For Oct4 staining, 100 µL of 1% (w/v) saponin (S-4521, Sigma-Aldrich) in DPBS was added to the sample, the cells were incubated for 10 min for permeabilization, then washed and resuspended in 50 µL of 1% (v/v) FBS in PBS. Samples for SSEA-1 staining were centrifuged and resuspended in 50 µL of 1% FBS in PBS without saponin treatment. All samples were incubated in the 1% FBS solution for 30 min, then 5 µL of a 1:10 dilution of either anti-Oct4 (611202, BD Transduction Laboratories, Franklin Lakes, NJ) or anti-SSEA-1 (MC-480 ascites, Developmental Studies Hybridoma Bank, Iowa City, IA) was added and the samples incubated for 1 h. To each tube 1 mL of 1% FBS solution was added, samples were centrifuged, supernatant was discarded, and the cells were resuspended in 50 µL of an appropriate PE conjugated secondary antibody (115-116-075 or 115-116-146, Jackson Immunoresearch, West Grove, PA) diluted 1:500. Samples were incubated 30 min in the dark, then washed three times with 1 mL of PBS, and fluorescence intensity data were acquired with a flow cytometer (Guava Technologies with express software module, Hayward, CA). A negative control was prepared using cells stained only with the secondary antibody, and gating for the flow cytometry was set so that 98% or more of the unstained cells were counted as negative. The fraction of cells expressing SSEA-1 and Oct4 was measured with flow cytometry and expressed as the fraction of cells above this threshold. All steps were performed at room temperature.

Real-Time Polymerase Chain Reaction (PCR)

Total RNA was isolated using the RNeasy Kit (74104, Qiagen, Valencia, CA) and RNase-Free DNase Set (79254, Qiagen), cDNA was synthesized using High Capacity cDNA Reverse Transcription Kit (4368814, Applied Biosystems, Foster City, CA), and real-time PCR was performed on a Fast Real-Time PCR System (7900HT, Applied Biosystems), using Power SYBR Green PCR Master Mix (4367659, Applied Biosystems). 28S ribosomal RNA was used as an oxygen insensitive endogenous control (Zhong and Simons, 1999). Primer sequences used to assess gene expression are shown in Table I. A standard calibration curve was constructed using undifferentiated mES cells.

<table>
<thead>
<tr>
<th>Target Gene</th>
<th>Direction</th>
<th>Sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>28S rRNA</td>
<td>Forward</td>
<td>GAATCCGCTAAGGAGTTGTGAACA</td>
</tr>
<tr>
<td></td>
<td>Reverse</td>
<td>CTCCAGCGCCATCCATTT</td>
</tr>
<tr>
<td>Oct4</td>
<td>Forward</td>
<td>CAGCACTGGAAGAAACCTCAGA</td>
</tr>
<tr>
<td></td>
<td>Reverse</td>
<td>TCTCCACATCTCAGGGATG</td>
</tr>
<tr>
<td>Nanog</td>
<td>Forward</td>
<td>CCTGTACCTCTCAACCTGCCA</td>
</tr>
<tr>
<td></td>
<td>Reverse</td>
<td>GCCCGTGAGAGAACACAGTCC</td>
</tr>
<tr>
<td>Sox2</td>
<td>Forward</td>
<td>GACAGCTACGGCCACATGA</td>
</tr>
<tr>
<td></td>
<td>Reverse</td>
<td>GGTGCTACGTGGTCCATGC</td>
</tr>
<tr>
<td>Nkx2.5</td>
<td>Forward</td>
<td>CAGTTGAGCTGGCAAAGG</td>
</tr>
<tr>
<td></td>
<td>Reverse</td>
<td>TTAGGAGCGTTCTGGAAGCA</td>
</tr>
<tr>
<td>Sox17</td>
<td>Forward</td>
<td>GCTGGCGGCGTCTGGAAGTC</td>
</tr>
<tr>
<td></td>
<td>Reverse</td>
<td>TAATGCTGATGACCATCTT</td>
</tr>
<tr>
<td>Nestin</td>
<td>Forward</td>
<td>GGGGAGGAGAGAGATGGA</td>
</tr>
<tr>
<td></td>
<td>Reverse</td>
<td>CCAGGATCTGACCGATCTG</td>
</tr>
</tbody>
</table>

Sequences reported 5'-3'.

Powers et al.: Effects of Oxygen on ES Cell Growth and Energetics

Biotechnology and Bioengineering
Cell Enumeration, Membrane Integrity, and Apoptosis Assessment

After detachment with trypsin, the total number of cells and the number of viable cells (assayed by membrane integrity measurements) in a sample were evaluated by staining cells using a Guava Viacount assay kit (Guava Technologies) and acquiring data with the Guava PCA flow cytometer. Apoptotic cell fraction was determined using an Annexin V assay kit (Guava Technologies).

Oxygen Consumption Rate

Cells were detached with trypsin and resuspended at a density of between 6 and 12 x 10^6 viable cells/mL in fresh culture medium equilibrated to 37°C and ambient oxygen, leading to a reoxygenation of hypoxic cells. The oxygen consumption rate (OCR) of this suspension was measured with a Micro Oxygen Monitoring System (FO/SYS2-T250, Instech Labs) as described previously (Papas et al., 2007). Briefly, a 200 μL aliquot of the cell suspension was sealed within the titanium chamber stirred with a glass-coated magnetic stirring bar. The time-dependent PO2 within the chamber was recorded with a fluorescence-based oxygen sensor, and the data at PO2 values greater than 30 mmHg were fit to a straight line by linear regression analysis. The oxygen consumption rate (OCR) was evaluated from the following equation:

\[
OCR = \frac{V_{ch} \alpha \Delta P_{O2}}{\Delta t}
\]

where \(V_{ch}\) is the chamber volume (200 μL), \(\alpha = 1.19 \times 10^{-9}\) mol/cm^3/mmHg is the Bunsen solubility coefficient for oxygen in medium (Avgoustiniatos, 2002), and \(\Delta P_{O2}/\Delta t\) is the slope of the fitted line. OCR/cell was determined by dividing the total OCR by the total number of viable cells that were loaded into the chamber.

Lactate Concentration

Assay reagent, prepared fresh for each test, consisted of 1 U/mL lactate oxidase (L-0638, Sigma-Aldrich), 0.025 U/mL peroxidase (P-6782, Sigma-Aldrich), and 0.1 mg/mL of o-dianisidine (F-5803, Sigma-Aldrich) in DPBS. Samples of supernatant medium from cell cultures were collected, and lactate concentration was measured by adding 100 μL of assay reagent to 50 μL of diluted sample (final concentration 0.1–0.3 mM lactate) in wells of a 96-well plate. The plate was incubated for 30 min at 37°C, 100 μL of 12 N sulfuric acid was added to stop the reaction, absorbance was measured at 540 nm using a microplate reader (Molecular Devices, Sunnyvale, CA), and lactate concentration was determined with a standard curve prepared for each test using lactic acid (L-1750, Sigma-Aldrich). DPBS (21-030-CM, Mediatech) was used as a diluent throughout.

Lactate Production Rate

We assumed that the lactate production rate (LPR) is proportional to the number of cells (N) at any instant of time (t), which can be represented by

\[
LPR = V_m \frac{dC}{dt} = \lambda N
\]

where \(V_m\) is the medium volume, \(C\) is the lactate concentration in the medium, and the coefficient \(\lambda\) is the LPR per cell, which is assumed to be constant during an experiment. Rearranging and integrating yields

\[
V_m (C_t - C_0) = \lambda \int_0^t N dt
\]

where subscripts 0 and f represent the initial and final conditions, and \(t_f\) is the duration of the experiment. When cells are in exponential growth, which applied in all of our experiments, we can write

\[
N = N_0 \exp (\mu t)
\]

and integrating leads to

\[
\lambda = \frac{V_m (C_t - C_0)}{(N_0/\mu) \exp(\mu t_f) - 1}
\]

from which \(\lambda\) is evaluated, since all quantities in Equation (5) were known. In our experiments, we measured final cell number \(N_f\) at \(t = t_f\), which was used to estimate \(N_0\) with Equation (4).

Lactate Dehydrogenase (LDH) Activity

A 100-μL aliquot containing 10^5 cells was added to a 1.5 mL tube containing 1 mL of DPBS and centrifuged for 3 min at 300g to obtain a cell pellet. To lyse the cells, the supernatant was removed, and 1 mL of 1% (v/v) Triton X-100 (T-9284, Sigma-Aldrich) in DPBS was added to each sample. The tubes were then briefly mixed using a vortex mixer and kept at room temperature for 1 h. LDH activity in the samples was measured with an Ektachem Vitros DT 60 II analyzer (Johnson and Johnson Clinical Diagnostics, Rochester, NY), which monitored the kinetics of the change in reflection density on a test slide as pyruvate and NAD were enzymatically converted into lactate and NAD by LDH in the test sample.
Estimation of $pO_2_{\text{cell}}$

$pO_2_{\text{cell}}$ was estimated by equating the rate of diffusion of oxygen through the stagnant medium to the rate of oxygen consumption:

$$\frac{D \alpha}{L} [pO_{2_{\text{gas}}} - pO_{2_{\text{cell}}}] = \rho \left( \frac{V_{\text{max}} pO_{2_{\text{cell}}}}{K_m + pO_{2_{\text{cell}}}} \right)$$

(6)

where $D$ and $\alpha$ are the diffusivity and solubility of oxygen in cell culture medium, respectively, $L$ is the medium depth, $pO_{2_{\text{gas}}}$ and $pO_{2_{\text{cell}}}$ are the gas phase and cell surface $pO_2$ values, respectively, $\rho$ is the viable cell density on the surface, $K_m$ is the Michaelis constant for oxygen consumption, and $V_{\text{max}}$ is the experimentally measured OCR/cell. The presence of a small $pO_2$ drop across the layer of cultured cells was ignored. For our calculations we used $D = 2.97 \times 10^{-5} \text{cm}^2/\text{s}$, $\alpha = 1.19 \times 10^{-9} \text{mol/cm}^2/\text{mmHg}$, $L = 0.15 \text{cm}$, and $K_m = 0.44 \text{mmHg}$ (Avgoustiniatos, 2002; Wilson et al., 1979). Equation (6) was solved for $pO_{2_{\text{cell}}}$ by iterative solution (Excel solver).

Estimation of Specific Growth Rate

The cell specific growth rate, $\mu$, was approximately constant during culture of undifferentiated cells. The specific growth rate was determined for these cultures using linear regression of the cumulative cell number versus time plotted on a semi-log arithmetic scale (as in Fig. 2A). The specific growth rate varied with time in culture when ES cells were differentiating (as in Fig. 3A). The instantaneous specific growth rate at a given time $t$ was determined by numerically differentiating the data using the average of the forward, backward, and central differences, according to the following equation:

$$\mu = \frac{\ln(N/N_0)}{3(t-t_0)} + \frac{\ln(N_1/N)}{3(t_1-t)} + \frac{\ln(N_1/N_0)}{3(t_1-t_0)}$$

(7)

where $N_0$, $N$, and $N_1$ are the cumulative cell numbers at successive times $t_0$, $t$, and $t_1$, respectively.

Assessment of Oxidative DNA Damage

A single cell gel electrophoresis (comet) assay (Tice et al., 2000) was used to detect DNA damage. This assay was done with and without the use of formamidopyrimidine DNA glycosylase (FPG, F-3174, Sigma-Aldrich), which creates strand breaks at sites of 8-oxoguanine (Collins and Dusinska, 2002), a well-documented product of oxidative damage (Helbock et al., 1999). Undifferentiated mES cells were cultured at specified $pO_{2_{\text{gas}}}$ for 72 h, then detached with trypsin and resuspended at a concentration of 200 cell/$\mu$L in 37°C, 1% (w/v) low melting point agarose (A-6560, Sigma-Aldrich) in DPBS. Methods for alkaline single-cell gel electrophoresis were subsequently followed (Collins and Dusinska, 2002). Comets were stained with 50 nM Sytox Orange dye (S11368, Molecular Probes, Eugene, OR), images were acquired with a 10× objective on a Zeiss epifluorescence microscope, and image analysis was performed using NIH image (Helma and Uhl, 2000). Samples with oxidative DNA damage (positive controls) were prepared by treating cells with 25, 85, and 150 $\mu$M hydrogen peroxide for 30 min immediately before embedding them in low melting point agarose.

Statistics

Statistical comparisons of data were performed using t-tests (paired or unpaired as appropriate), and results were deemed significant if the two-tailed $P$-value was 0.05 or less.

Results

Characterization of $pO_2$ Control in Experimental Apparatus

The time-dependent response of the liquid phase $pO_2$ to a step change in $pO_2$ of the gas flowing into the flask under simulated culture conditions is shown in Figure 1. The $pO_2$ in the region near the bottom of the water layer dropped from 160 to about 10 mmHg within 1 h and to less than 1 mmHg after 100 min. The $pO_2$ continued to drop at longer times, but it could not be measured when it was below the resolution of the sensor. The presence of oxygen-consuming cells at the bottom of the flask under actual culture conditions would have accelerated the decrease of $pO_{2_{\text{cell}}}$ and would have brought it to values substantially smaller than 1 mmHg in less than 2 h. Thus, the response time for liquid phase $pO_2$ was much smaller than the time of the

Figure 1. Response of liquid $pO_2$ to a change in ambient $pO_{2_{\text{gas}}}$. The $pO_2$ at the bottom of 4 mL of water contained within a 25 cm² tissue culture flask is plotted as a function of time. The flask was initially equilibrated with room air. At $t=0$, the gas flowing into the flask was changed to nitrogen. The time dependence of the inlet gas flow rate was identical to that used in all cell culture experiments.
experiment in all experiments carried out. Cells cultured at low \( pO_2 \) would be exposed to higher \( pO_2 \) for a period of at most 3 h once every two days as the result of passaging.

### Cellular \( pO_2 \)

Estimates of \( pO_2_{cell} \) calculated from Equation (6) for cell densities ranging from 0.1 to \( 2.0 \times 10^5 \) cell/cm\(^2\), corresponding to the lowest and highest cell density present before and after passaging cells, respectively, are tabulated in Table II. Although the magnitude of the \( pO_2 \) drop across the culture medium was small in all cases, the difference between \( pO_2_{gas} \) and \( pO_2_{cell} \) was large relative to \( pO_2_{gas} \) at very low oxygen concentrations, for example, cells cultured using a \( pO_2_{gas} \) of 7 mmHg were often exposed to \( pO_2_{cell} \) values that approached 0.

### Table II. Experimental \( pO_2_{gas} \) conditions and the corresponding \( pO_2_{cell} \) for undifferentiated mES cell monolayers.

<table>
<thead>
<tr>
<th>% Oxygen</th>
<th>( pO_2_{gas} ) (mmHg)</th>
<th>Maximal OCR ( b ) ( V_{max} ) (amol/sec cell)</th>
<th>Cellular ( pO_2_{cell} ) (mmHg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>40</td>
<td>285</td>
<td>28</td>
<td>262–284</td>
</tr>
<tr>
<td>20</td>
<td>142</td>
<td>29</td>
<td>118–141</td>
</tr>
<tr>
<td>5</td>
<td>36</td>
<td>23</td>
<td>18–35</td>
</tr>
<tr>
<td>1</td>
<td>7</td>
<td>17</td>
<td>0.5–6.5</td>
</tr>
<tr>
<td>0</td>
<td>0</td>
<td>10</td>
<td>0</td>
</tr>
</tbody>
</table>

\( a \) The oxygen content in the pre-mixed gas cylinders (containing 5% CO\(_2\)) was specified, and the \( pO_2_{gas} \) was determined from \( pO_2_{gas} = (\% \text{ oxygen}) \times (P_{\text{atm}} - P_{\text{H}_2\text{O}}) \), where \( P_{\text{atm}} \) is the atmospheric pressure and \( P_{\text{H}_2\text{O}} = 47 \text{ mmHg} \) is the vapor pressure of water at 37°C.

\( b \) OCR measured experimentally (Fig. 9).

\( c \) \( pO_2_{cell} \) calculated with Equation (6) for extreme low and high densities of 0.1 and \( 2.0 \times 10^5 \) cell/cm\(^2\) (about 2–50% confluence), respectively.

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**Cell Growth**

Undifferentiated CCE and D3 mES cells at \( pO_2_{gas} \) of 0, 7, 36, 142, and 285 mmHg grew exponentially for up to 25 days (Fig. 2A). There were no temporal changes that suggested senescence, differentiation, or long-term conditioning to the culture conditions. The fraction of cells with intact cell
membranes was between 94% and 98%, which further suggested that undifferentiated mES cells survived and proliferated in all conditions tested. Both CCE and D3 cells had roughly the same specific growth rate (Fig. 2B) and were tolerant of a wide range of oxygen conditions. Growth was significantly suppressed under anoxia, but the cells continued to grow exponentially with a doubling time between 24 and 27 h. Elevated pO_{2gas} of 285 mmHg also caused a decrease in growth rate in both cell lines relative to that at 142 mmHg. D3 cells consistently grew faster at a pO_{2gas} of 36 mmHg compared to 142 mmHg. The specific growth rate \( \mu \) in four experiments was 1.05 \( \pm 0.12 \) day\(^{-1} \) at 142 mmHg and 1.15 \( \pm 0.10 \) day\(^{-1} \) at 36 mmHg. This small, but statistically significant difference (\( P < 0.01 \), paired t-test) would lead to a threefold difference in cell number after 10 days of culture. No difference in growth at these same two pO_{2gas} conditions was observed with CCE cells.

Without LIF in the culture medium, CCE cells readily differentiated, and the specific growth rate (Fig. 3) decreased with increasing time after LIF withdrawal. After a time lag of about 2 days, the specific growth rate, on average, decreased at a similar rate for pO_{2gas} conditions ranging from 7 to 142 mmHg, and reached a minimum after about 10 days (Fig. 3F). Changing pO_{2gas} in this range did not significantly delay or accelerate the onset or rate of differentiation. Individual experiments were reproducible at 285, 36, and 7 mmHg, less so at other conditions. The specific growth rate at 285 mmHg was suppressed to a greater extent, relative to that at other values of pO_{2gas} than was observed in the presence of LIF. A small number of experiments with D3 cells suggested similar trends (data not shown), which occurred more slowly due to a decreased rate of spontaneous differentiation.

### Oxidative DNA Damage

Comet assays run with and without FPG using cells cultured for 72 h at each pO_{2gas} produced similar values of tail length (Table III) and tail moment, both measures of the relative amount of DNA damage, and there was no trend of increased damage with increased pO_{2gas}. Cells treated with hydrogen peroxide displayed dose-dependent increases in mean tail length and moment, and a further dose-dependent increase occurred when the assay was performed with FPG. These results suggest that pO_{2gas} did not significantly affect the amount of oxidative DNA damage detectable with a comet assay, a result consistent with growth rate data (Fig. 2A).

#### Retention of Undifferentiated Stem Cell Phenotype at Different pO_{2gas}

**Protein and Gene Expression**

When using culture medium containing LIF, which suppresses mES cell differentiation (Viswanathan et al., 2002), pO_{2gas} did not affect the fraction of cells that expressed SSEA-1 and Oct4 using flow cytometry. Both proteins were expressed by 85–95% of all cells over a 25-day period at all pO_{2gas}. When gene expression of several transcription factors (Oct4, Sox2, and Nanog) typically associated with the undifferentiated ES cell phenotype was measured with real-time PCR, there were no changes after 8 days from the undifferentiated stem cells with cells cultured at 142 and 36 mmHg. However, other values of pO_{2gas} led to differences in the amount of mRNA present (Fig. 4). After 8 days in culture (3 passages), both high (285 mmHg) and low (7 and 0 mmHg) pO_{2gas} resulted in significant decreases of Oct4, Sox2, and Nanog mRNA to levels ranging from 10% to 40% of those in the undifferentiated stem cells at day 0, and the levels of each transcription factor were sequentially lower at 7 and 0 mmHg.

Without LIF in the culture medium, the fraction of CCE cells expressing Oct4 protein, as measured by flow cytometry, also decreased with time except for pO_{2gas} of 285 mmHg and some experiments at 142 mmHg, at which the fraction decreased and then rebounded (Fig. 5A and B). Individual experiments were reproducible at 7 and 285 mmHg but less so at 142 and 36 mmHg, similar results were observed at 0 and 7 mmHg. The averaged data (Fig. 5F) demonstrate a decrease in the fraction of Oct4 positive cells with decreasing pO_{2gas}. Because of the scatter in the data, this trend was only significant (\( P < 0.04 \)) when 142 and 7 mmHg were compared at day 6, and 285 and 7, 36, or 0 mmHg were compared on day 16. After about 16 days, the average fraction of Oct4 positive cells converged to about 30% at 0, 7, and 36 mmHg and was about 55% and 70% at 142 and 285 mmHg, respectively.

With the exception of inconsistent behavior at 285 mmHg, the data for gene expression of Oct4, Nanog, and Sox2 (Fig. 4) displayed several trends when LIF was absent. (1) The relative gene expression of all transcription factors decreased as pO_{2gas} decreased from 142 to 0 mmHg. The relative magnitude of the decline decreased in the order Nanog > Oct4 > Sox2. The overall trend for Oct4 was consistent with that of the flow cytometry measurements in

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**Table III.** Relative tail length from a Comet assay.

<table>
<thead>
<tr>
<th>pO_{2gas} (mmHg)</th>
<th>H_{2}O_{2} (\mu M)</th>
<th>Without FPG</th>
<th>With FPG</th>
</tr>
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<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>1.4</td>
<td>1.8</td>
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<tr>
<td>7</td>
<td>0</td>
<td>1.3</td>
<td>1.7</td>
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<td>36</td>
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<td>1.4</td>
<td>1.8</td>
</tr>
<tr>
<td>142</td>
<td>0</td>
<td>1.0</td>
<td>1.6</td>
</tr>
<tr>
<td>285</td>
<td>0</td>
<td>1.2</td>
<td>1.1</td>
</tr>
<tr>
<td>142</td>
<td>25</td>
<td>0.9</td>
<td>1.9</td>
</tr>
<tr>
<td>142</td>
<td>85</td>
<td>1.8</td>
<td>2.9</td>
</tr>
<tr>
<td>142</td>
<td>150</td>
<td>2.3</td>
<td>4.2</td>
</tr>
</tbody>
</table>

*Assay was carried out with and without FPG, which induces DNA strand breaks at 8-oxoguanine sites and increases comet tail lengths. Relative tail length is the measured tail length divided by the value at 142 mmHg. Data tabulated are from a single experiment.
Figure 4. Real-time PCR analysis of ES cell markers Oct4, Sox2, and Nanog in CCE cells cultured for 8 days with and without LIF in the culture medium. Results reported are mean ± SD of data obtained with four different cultures carried out with the same undifferentiated stem cells starting at day 0. All data is normalized to that for the cells at day 0. Gene expression for conditions with LIF that are significantly different from the day 0 undifferentiated cells are indicated with an asterisk (*). For conditions without LIF, the X denotes a significant difference in gene expression between cultures with LIF and without LIF at identical \( P_{\text{O}_2}\) conditions, and the plus sign (+) denotes significant difference between \( P_{\text{O}_2}\) conditions indicated by the brackets, all without LIF.

Differentiation to Germ Layer Derivatives

To examine the pluripotency of mES cells cultured with LIF at reduced \( P_{\text{O}_2}\) for 8 days, cells were subsequently subjected to a differentiation protocol involving EB formation followed by attached culture. Gene expression measurements (Fig. 7) after 5 days of culture for Nkx2.5, Sox17, and Nestin, demonstrated a 1–2 order of magnitude increase in these markers for mesoderms, endoderm, and ectoderm, respectively, relative to undifferentiated mES cells.

Continuous Culture Without Passaging

In experiments described to this point, medium was exchanged daily, and cells were passaged every 2 or 3 days. Fresh medium was pre-equilibrated to the desired \( P_{\text{O}_2}\) so that there was minimal disturbance associated with medium exchange; however, during passaging the mES cells were exposed to ambient oxygen for about 1 h and to the stress of trypsin detachment, which may have selectively harmed either differentiated or undifferentiated cells. To study effects of \( P_{\text{O}_2}\) on mES cell growth and differentiation without these potential artifacts, D3 and CCE cells were
plated at a density of 100 cells/well in 6-well plates containing 3 mL of culture medium (three wells for each condition) both with and without supplemental LIF. The plates were maintained for 8 days without passaging the cells or changing the medium. Plates were kept at a pO2gas of 142 or 7 mmHg continuously (24 h/day) or at a pO2gas of 7 mmHg for 23 h/day and 142 mmHg for 1 h/day. With and without LIF, exposure of D3 and CCE ES cells to a pO2gas of 7 mmHg for 23 h and 142 mmHg for 1 h each day did not significantly affect specific growth rate or the fraction of cells expressing SSEA-1 and Oct4 protein (measured by flow cytometry), relative to culture at 7 mmHg continuously (data not shown), although it is possible that a more sensitive method such as real-time PCR might detect differences.

**Temporal Response to Anoxia**

The ability of mES cells to grow in anoxia with a doubling time of about 24 h indicated that mES cells are well suited to, or can readily adapt to, low oxygen conditions. To further explore this behavior, undifferentiated mES cells were subjected to a change in pO2gas from 142 to 0 mmHg. The fraction of cells with intact membranes, the fraction of apoptotic cells, OCR/cell immediately after reoxygenation, and LDH activity were measured at various times up to 50 h after the transition to anoxia (Fig. 8).

**Cellular Energy Metabolism**

The data in Figure 8 provide evidence that mES cells increased anaerobic metabolism when exposed to hypoxia or anoxia. To quantify this further, we estimated the rate of ATP produced aerobically and anaerobically under each of the pO2 conditions tested. OCR/cell was used as a measure of oxidative phosphorylation and LPR/cell as a measure of anaerobic glycolysis. ATP production rate was estimated by assuming production of 6 mol of ATP per mol of oxygen consumed.
Figure 6. Phase contrast photomicrographs of CCE cells cultured (A) for up to 72 h with 1,000 U/mL LIF at pO$_{2}$gas of 142 mmHg (top panels) or 7 mmHg (middle panels) or (B) 8 days without LIF (bottom two panels) at a pO$_{2}$gas of 142 or 7 mmHg.

Figure 7. Real-time PCR analysis of the differentiation markers Nkx2.5, Sox17, and Nestin of CCE cells cultured at the indicated pO$_{2}$gas for 8 days with 1,000 U/mL LIF and then differentiated without LIF in hanging drop EBs for 2 days and in attached culture for an additional 3 days at the same pO$_{2}$. Nkx2.5, Sox17, and Nestin are markers for mesoderm, endoderm, and ectoderm, respectively. Results reported are mean ± SD of data obtained with four different cultures carried out with the same undifferentiated stem cells starting at day 0, and all of the results were significantly different from the control (undifferentiated ES cells at 142 mmHg).

Discussion

In order to determine whether oxygen level affected proliferation and maintenance of undifferentiated phenotype in differentiation-suppressing and -permissive conditions, we measured the growth rate, Oct4, Sox2, and Nanog mRNA expression, the potential to differentiate into derivatives of all three germ layers, and the cell and colony consumed and 1 mol of ATP per mol of lactate formed (Lindqvist et al., 2002; Miller et al., 1987).

A relatively high level of anaerobic metabolism occurred in the mES cells, even at high pO$_{2}$gas at which there was a plentiful supply of oxygen to the cells (Fig. 9). Nearly 40% of the ATP production was provided by anaerobic metabolism of glucose into lactate at 142 mmHg or greater. When pO$_{2}$gas was reduced to 36 mmHg, a small but significant decrease in OCR and increase in LPR was observed. This shift to increased anaerobic metabolism became more pronounced as the pO$_{2}$gas decreased to 0 mmHg, a condition in which no significant aerobic ATP production could occur. Overall, the total estimated ATP production rate remained nearly constant as long as oxygen was present, and decreased about 10% at 0 mmHg.
and using real-time PCR revealed a decrease in gene expression of Oct4, Sox2, and Nanog at pO2gas of 285, 7, and 0 mmHg (Fig. 4). Compared to 142 mmHg, photomicrographs of cells grown at pO2gas of 7 mmHg revealed subtle changes indicating initiation of differentiation (Fig. 6A). Nonetheless, cells cultured at low pO2gas in the presence of LIF retained the ability to subsequently differentiate into derivatives of all three germ layers (Fig. 7).

When LIF was removed from the culture medium, the cells differentiated. The growth rate decreased with time and was not affected by pO2gas, from 7 to 142 mmHg (Fig. 3), but there were differences at the extremes of 0 and 285 mmHg as there were for growth in the undifferentiated state. Differentiation occurred much faster in the CCE cell line than in the D3 cell line. Such variations between ES cell lines have been observed by others in many different aspects of culture (Ward et al., 2004). The rate at which CCE cells lost Oct4 protein expression increased with decreasing pO2gas in the range of 0–285 mmHg (Fig. 5). For 0–142 mmHg pO2gas, gene expression of Oct4, Nanog, and Sox2 decreased with decreasing pO2gas and was always lower than when LIF was present at the same pO2gas (Fig. 4). At pO2gas of 7 and 0 mmHg, the effect on gene expression of reduced oxygen was comparable to or greater than the

morphology of undifferentiated mES cells cultured with LIF, and a subset of these characteristics for mES cells cultured without LIF, at pO2gas in the range of 0–285 mmHg. We also studied mES cell energetics over the same range of pO2gas to gain further insight into the extent to which mES cells adapt to different environmental oxygen conditions.

When LIF was present in the culture medium, growth of undifferentiated mES cells was sensitive to pO2 (Fig. 2), but the growth rate was constant at any fixed pO2gas up to 24 days. When compared to culture at a pO2gas of 142 mmHg, the specific growth rate (Fig. 2B) was similar at 36 mmHg, moderately reduced at 7 and 285 mmHg as others have found for mES cells (Iyer et al., 1998; Saretzki et al., 2004) and hES cells (Ezashi et al., 2005; Ludwig et al., 2006) and further reduced at 0 mmHg. D3 cells had a slightly higher growth rate at a pO2gas of 36 relative to 142 mmHg, while growth rates for CCE cells were the same. The fraction of cells expressing Oct4 and SSEA-1 protein as quantified with flow cytometry was not affected by pO2gas, but measurements of mRNA levels using real-time PCR revealed a decrease in gene expression of Oct4, Sox2, and Nanog at pO2gas of 285, 7, and 0 mmHg (Fig. 4). Compared to 142 mmHg, photomicrographs of cells grown at pO2gas of 7 mmHg revealed subtle changes indicating initiation of differentiation (Fig. 6A). Nonetheless, cells cultured at low pO2gas in the presence of LIF retained the ability to subsequently differentiate into derivatives of all three germ layers (Fig. 7).
separate effect of LIF removal (Table IV). Gross changes in cell and colony morphology were evident when LIF was not present (Fig. 6B).

Taken together, our observations suggest that hypoxic culture of mES cells causes initiation of differentiation, even in the presence of LIF. This finding is consistent with that of Jeong et al. (2007), who observed that hypoxia inhibits mES cell self-renewal and induces early differentiation. Their data suggested that hypoxia-induced in vitro differentiation of mES cells is triggered at least in part by the HIF-1α-mediated suppression of LIF-STAT3 signaling. Jeong et al. observed a 50% reduction in Oct4 gene expression after CCE cells were cultured for 48 h with PO2gas of 7 mmHg, which is consistent with our data. They also observed much more dramatic morphological changes after 48 h than we did, which may be due to their use of a defined serum replacement in place of FBS used in our study. Niwa et al. (2001) used conditional expression to reduce Oct4 by a factor of 2 in mES cells. With cells under normoxia, they observed cell compaction at 48 h, as did we, but found extensive flattening with enlarged nuclei at 96 h, which we did not.

Although we observed substantial reduction in gene expression of transcription factors associated with self-renewal after culture at 7 or 0 mmHg in the presence of LIF, all other measurements—constancy of growth rate with time, morphological changes, pluripotency—demonstrated only modest or no departure from the behavior of undifferentiated stem cells under the conditions we studied. Such paradoxical behavior suggests that the hypoxia-induced early differentiation was self-limiting. We hypothesize that it could result from the simultaneous presence of self-renewal promoting characteristics of hypoxia that counterbalance the HIF-1α-mediated suppression of LIF-STAT3 signaling. Because this signaling pathway is important for maintenance of mES but not hES cells in an undifferentiated state (Daheron et al., 2004), our speculation draws support from a previous study (Ezashi et al., 2005) that found pO2gas of 36 or 21 mmHg promoted self-renewal of hES cells, although other studies have not reported significant effects on hES cell self-renewal (Forsyth et al., 2006; Ludwig et al., 2006; Peura et al., 2005).

Further research is needed to explore the effects of hypoxic culture on mES cells in the presence of LIF on subsequent differentiation. There are situations where ES cell growth and survival are enhanced at moderately low PO2 conditions, such as during the establishment of new mES cell lines (Gibbons et al., 2006; Wang et al., 2006) and during culture of hES cells at very low cell densities or sub-optimal culture conditions (Forsyth et al., 2006; Ording et al., 2005). Reduced pO2gas conditions may also reduce the occurrence of chromosomal abnormalities in hES cells (Forsyth et al., 2006), but similar results have not been observed with mES cells (Wang et al., 2006). We found no effect of culture PO2 in the range of 0–285 mmHg on the amount of oxidative DNA damage in undifferentiated mES cells using the comet assay.

The limited effects of culture pO2 on ES cell proliferation, particularly under anoxia, led us to further investigate ES cellular energetic (Figs. 8 and 9). We measured the rates of lactate production and oxygen consumption (Fig. 9A) as indicators of the contributions of aerobic and anaerobic metabolism to the total cellular ATP production rate at all pO2gas conditions (Fig. 9B). With excess oxygen, 40% of the ATP demand of ES cells was satisfied through anaerobic glycolysis, which is higher than the 15–25% reported for early embryos (Houghton, 2006; Houghton et al., 1996; Thompson et al., 1996). The reliance on anaerobic metabolism increased further with oxygen depletion, and all of the ATP was produced anaerobically under anoxia. The total ATP production rate remained nearly constant during this change, and ES cells could complete this adaptation in less than 2 days without significant loss of viability (Fig. 4).

Use of anaerobic metabolism of glucose and the ability of ES cells to adapt to reduced pO2 conditions including anoxia is logical from a developmental viewpoint, since fetal development occurs in a very low pO2 environment. The pO2 in the maternal endometrium supporting an early embryo is about 40 mmHg, which in turn produces an oxygen concentration of about 0.05 mM in the surrounding tissue (Rodesch et al., 1992). Compared to oxygen, glucose is in substantial excess, with a fasting plasma concentration of approximately 3–4 mM (Weijers et al., 2002). An early embryo can therefore maximize its energy production rate by using both aerobic and anaerobic metabolism of glucose. The fact that mES cells utilize anaerobic metabolism to a substantial extent even at high pO2 may also partially explain their ability to survive hyperoxia, since such a metabolic profile reduces the generation of harmful ROS (Andreyev et al., 2005). This profile is also observed in tumor cells (Gatenby and Gawlinski, 2003), and it has been shown that the glycolytic and the cancer phenotypes are very closely linked to one another (Kondoh et al., 2005; Ramanathan et al., 2005).

Many other cells increase anaerobic metabolism when confronted with hypoxia, which was first observed by Pasteur during yeast fermentation (Berg et al., 2002). The Pasteur effect has been particularly well documented in CHO and hybridoma cultures (Lin and Miller, 1992; Miller et al., 1987). Although the Pasteur effect is commonly observed, not all cells can adapt equally well to survive hypoxic/anoxic insult. As ES cells differentiate into mature cells, a shift to aerobic metabolism will occur for many lineages (Sharma et al., 2006), and some of the resulting cells will become sensitive to oxygen deprivation. Study of the energetics of differentiated cells obtained from ES cells could potentially be used as an evaluative tool for determination of cell maturation.

Our work shows that for bioprocess development with cells having the properties of undifferentiated mES cells, pO2 control is not likely to be a critical parameter. The ideal pO2cell is about 40 mmHg, which is likely to maximize growth without leading to excessive lactate formation and might also provide some protection from DNA damage.
However, both lower and higher pO2\textsubscript{cell} conditions can be tolerated without excessive cell death or undesired changes in cell phenotype, although there may be effects on subsequent differentiation with LIF removal. We have observed a large effect of reduced pO2 during the directed differentiation of ES cells to cardiomyocytes (Powers, 2007). Under these conditions pO2 control is a much more important area that warrants further study.

This study was supported in part by grants from the NIH (NCRR ICR U41PR 16606 and ROI-5056108 OIA) and the JDRF Center for Islet Transplantation at Harvard Medical School. Helpful discussions and technical assistance with aspects of the work were provided by Dr. Susan Bonner-Weir (Joslin Diabetes Center, Harvard Medical School), Anna Pisansia, and Michael J. Rappel (MIT, Chemical Engineering), and Dr. Yuh-Shin Chang (Massachusetts General Hospital).

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C. Differentiation of hESC to β-cells

C.1. Introduction

Pluripotent stem cells are a potential source of insulin-secreting β-cells that has been explored for the past decade with both mouse and human cells [151-153]. ViaCyte, Inc. has recently developed a four-stage protocol for differentiating hESC in vitro in successive stages that mimic in vivo pancreatic organogenesis, inducing (1) definitive endoderm (DE), (2) primitive gut tube (PGT), (3) posterior foregut (PF), and (4) pancreatic endoderm (PE) and endocrine precursor (EP) [154, 155] and was the first to show that implantation of hESC-derived PE into mice resulted in the formation of insulin-secreting cells that can cure diabetic mice [9]. The Melton laboratory is also following an approach similar to ViaCyte but with the addition of small molecules, which are identified with a high-throughput screening assay at each stage. These small molecules result in an increased fraction of DE cells derived from mESC and hESC [130].

In 2009, the Colton and Melton laboratories along with ViaCyte, Inc. were awarded a research grant from the Juvenile Diabetes Research Foundation to increase the efficiency of differentiating human pluripotent stem cells into β-cells to cure type 1 diabetes and to improve characteristics of these cells for implantation. As part of this grant, I worked at ViaCyte, Inc. for one month to learn hESC culture and β-cell differentiation protocols and to begin to merge
these protocols with our low oxygen culture methodologies. The following are the results from my time at ViaCyte, Inc.

C.2. Differentiation of CyT49 hESCs with ViaCyte protocol

The ViaCyte protocols employed culture at 142 mmHg pO$_{2\text{gas}}$ on standard polystyrene culture dishes, as described in Kroon et al [9]. When expanded in the pluripotent state, almost all CyT49 hESC cultured remained undifferentiated (Figure C-1). Two CyT49 clones, 304d and 511b, were differentiated. Assays were carried out for all markers and with all methods normally used by ViaCyte and cells were periodically observed by light microscopy. During stage 1, expression of BRACHYURY T, CER, SOX17, AND FOXA2 were detected with real-time PCR, and by the end of stage 1, most cells expressed FOXA2 and few expressed OCT4 (Figure C-2). HNF4A expression was detected during stage 2 (Figure C-3). PDX1 expression was detected during stage 3 (Figure C-4). During stage 4, expression of NKX6.1, PTF1A, NGN3, NKX2.2 was detected with qPCR (Figure C-5). Many cells expressed PDX1, and a subset of these cells coexpressed NKX6.1 (Figure C-6). After 16 days of differentiation, insulin-, glucagon-, and somatostatin-expression cells were observed (Figure C-7); some of these cells expressed more than one of these hormones, and c-peptide was detected in the culture medium (Figure C-8). These results are consistent with previous results from ViaCyte Inc and demonstrate that all aspects of the ViaCyte protocol were replicated, which is a necessary and critical first step in completing
all subsequent aims and milestones in the proposal. Cells from clone 304d were maintained in extended culture under stage 4 conditions for an additional 17 d. OCT4 expression decreased by four orders of magnitude during the 12-day differentiation and by another two orders of magnitude after 29 d (Figure C-9).

C.3. OCR of differentiating CyT49 hESC

We measured the OCR of cells differentiated with the ViaCyte protocol after 0, 2, 8, and 14 days of differentiation (Table C-1). The OCR for undifferentiated CyT49 cells was 21 amol cell\(^{-1}\) s\(^{-1}\), a value about 2/3 that of undifferentiated mouse embryonic stem cells, which we reported in Powers et al. [32], and the OCR remained unchanged with differentiation time.

C.4. Culture and differentiation of CyT49 cells on silicone rubber

One of the major aspects of the JDRF grant is to perform studies in well-defined oxygen environments by culturing in dishes having a silicone rubber membrane culture surface. Because of rapid progress made with the standard ViaCyte protocol, there was some time available to begin studies with these dishes. We plated undifferentiated CyT49 cells on silicone rubber membranes pretreated with fibronectin, matrigel, human serum, or with no treatment. After 24 hr, cells attached with fibronectin and matrigel coating, but poor attachment was observed with human serum treatment or without treatment (Figure C-10). After 3 days,
cells were confluent on fibronectin- and matrigel-coated silicone rubber (Figure C-11). Cells were subsequently differentiated for 7 days, at which time many cells stained positive for PDX1 (Figure C-12). Throughout culture, cell detachment was not observed with fibronectin or Matrigel treatment. This data demonstrates that the ViaCyte protocol can be performed on silicone rubber plates, and we do not expect problems to arise due to the use of these plates.
### Table C-1. Oxygen consumption rate of CyT49 hESC during differentiation

<table>
<thead>
<tr>
<th>Stage</th>
<th>Differentiation Time (days)</th>
<th>Oxygen Consumption Rate (amol cell$^{-1}$ sec$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>21.0 ± 2.6</td>
</tr>
<tr>
<td>1</td>
<td>2</td>
<td>18.5 ± 1.5</td>
</tr>
<tr>
<td>3</td>
<td>8</td>
<td>20.9 ± 1.0</td>
</tr>
<tr>
<td>4</td>
<td>14</td>
<td>19.2 ± 4.3</td>
</tr>
</tbody>
</table>

Cells differentiated on polystyrene at pO$_{2}$=$142$ mmHg with Novocell protocol.

Data presented as average ± stdev.

n=2 for all except 0 days, in which n=7.
Figure C-1. En face immunofluorescence images of undifferentiated CyT49 cells.
Figure C-2. Expression of stage 1 markers in differentiating CyT49 cells.

(A-D) Relative expression of BRACHYURY T, CER, SOX17, AND FOXA2 mRNA measured with real-time PCR (n=4). (E-G) En face images of cells differentiated 5 days and stained with DAPI or immunostained against FOXA2 or OCT4 protein.
Figure C-3. Relative expression of the stage 2 marker HNF4A mRNA measured with real-time PCR in differentiating CyT49 cells (n=4).
Figure C-4. Relative expression of the stage 3 marker PDX1 mRNA measured with real-time PCR in differentiating CyT49 cells (n=4).
Figure C-5. Relative expression of the stage 4 markers NKX6.1, PTF1A, NGN3, NKX2.2 mRNA measured with real-time PCR (n=4).
Figure C-6. En face images of cells differentiated 12 days and stained with DAPI and immunostained against PDX1 or NKX6.1 protein.
Figure C-7. En face images of cells differentiated 16 days and stained with DAPI and immunostained with INSULIN, SOMATOSTATIN, or GLUCAGON protein.
Figure C-8. Concentration of c-peptide in medium determined by ELISA for CyT49 cells differentiated with ViaCyte protocol (n=1)
Figure C-9. Real-time PCR analysis of OCT4 expression of CyT49 hESCs during differentiation with ViaCyte protocol on polystyrene at pO_{2gas}=142 mmHg. After the 12-day protocol was completed, cells from clone 304d were maintained for another 17 days under conditions used to reach stage 4 (n=2).
Figure C-10. En face phase contrast images 24 hr after CyT49 cells were plated at a density of $1.3 \times 10^5$ cells cm$^{-2}$ on silicone treated with fibronectin, Matrigel, human serum, or with no treatment.
Figure C-11. En face phase contrast images 3 days after CyT49 cells were plated at a density of $1.3 \times 10^5$ cells $\text{cm}^{-2}$ on silicone treated with fibronectin or Matrigel.
Figure C-12. En face immunofluorescence image of CyT49 cells differentiated 7 days to stage 3 with ViaCyte protocol on Matrigel-coated silicone rubber at 142 mmHg pO\textsubscript{2}\textsubscript{gas}.

(A) DAPI. (B) PDX1. 20x zoom.
D. Experiments with the BioSpherix Xvivo

D.1. Introduction

The Xvivo incubation system is fabricated by BioSpherix, Inc and allows for the continuous cellular culture under reduced oxygen. It combines incubators with interconnecting, sealed glove box and microscope chambers in which pO$_2$ can be continuously controlled electronically. When the cells must be removed from the incubator and manipulated, sudden transient spikes in the oxygen concentration are avoided. We hypothesize that this feature is important for low oxygen culture because HIF-1α is rapidly degraded at atmospheric oxygen, and it takes hours to re-accumulate under low oxygen culture, which results in cells having depressed HIF-1α content for up to 8 hr [156]. Use of the Xvivo incubation system avoids these issues and allows for uninterrupted expression of HIF-1α and subsequent target genes. Combination of silicone rubber dishes [53] to the Xvivo system would provide a system to monitor and control the pO$_2$ to which the cells are exposed, a capability that does not now exist. The Xvivo incubation system can dynamically control the oxygen concentration of the incubator atmosphere. For example, when the cells are initially seeded at low density, a specific concentration of oxygen can be used. However, as the cell density increases, pO$_2$$_{\text{cell}}$ decreases. To compensate for this, the system could gradually increase pO$_2$$_{\text{gas}}$ according to an algorithm or based on sensor measurements to keep pO$_2$$_{\text{cell}}$ constant.
I performed several pilot studies to investigate the technical advantages of the Xvivo with and without silicone rubber membrane culture plates described hereafter.

D.2. The effect of low oxygen control methodologies on mESC VEGF secretion

D.2.1. Introduction

We investigated the secretion of VEGF protein by mESC cultured on silicone rubber membranes on top of polystyrene (SR-P) or silicone rubber alone (SR) using a flush box or Xvivo system. VEGF is an important signaling molecule that is secreted by cells at low oxygen and is involved in embryonic development and stem cell differentiation [42]. We hypothesized that VEGF expression during experiments involving low oxygen culture of cells in standard low oxygen incubators is likely affected by periodic high-oxygen exposure when the cells are handled (e.g. media changes) and that this problem is exacerbated if cell culture is being performed on oxygen-impermeable polystyrene dishes, since on the order of 20 hours can be required for the cells to return to low oxygen levels upon placement in a low oxygen environment [157]. We found that culture in the Xvivo and on SR dishes resulted in more VEGF production compared to the flush box.
system and SR-P dishes because of their ability to better maintain pO₂ at low levels.

D.2.2. Methods

R1 mESC numbering 2.4x10⁴ were plated in a 24-well plate with culture surfaces consisting of either fibronectin-coated SR-P or SR, placed in a flush box or Xvivo incubator, and cultured in DMEM supplemented with 10% FBS and 10³ units/mL of LIF at 142 or 7 mmHg pO₂gas. For SR-P, having silicone rubber on top of polystyrene controlled for differences in surface properties, while maintaining the oxygen transport characteristics of polystyrene. After 48 hours, the media was replaced with fresh media. Media exchange for the plates in the flush box was performed in an open biological safety cabinet at atmospheric pO₂gas (160 mmHg) and for cells in the Xvivo, was performed in the closed Xvivo system at 7 mmHg pO₂gas. After 24 additional hours of culture, media and cells were collected. The VEGF content of the media was assayed with ELISA (Calbiochem), and the viable cell number was determined with a membrane-integrity assay (Guava ViaCount).

D.2.3. Results

With incubation in the flush box, the VEGF content was higher under low oxygen compared to culture at 142 mmHg (Figure D-1). Among all 7 mmHg culture
conditions, culture on SR-P in the flush box had the smallest increase in VEGF content compared to 142, a factor of 1.9. Culture on SR in the flush box resulted in an even larger increase in VEGF content compared to 142, a factor of 5.1. VEGF content at low oxygen in the flush box is higher with SR compared to SR-P because pO$_{2cell}$ rapidly reaches steady-state after exposure to atmospheric oxygen, requiring only minutes, while polystyrene requires on order 20 hours.

Incubation in the Xvivo system at 7 mmHg resulted in the higher VEGF content compared to 7 mmHg in the flush boxes for both SR-P and SR (Figure D-1). Culture in the Xvivo increased VEGF content by a factor of 2.8 and 1.2 on SR-P and SR, respectively, compared to culture in the flush box. There was no statistical difference between SR-P and SR at 7 mmHg in the Xvivo. In the Xvivo, cells were never exposed to atmospheric oxygen during media exchange, since the Xvivo allows media exchange to occur without changing pO$_{2gas}$. This allowed the cells to be at low pO$_{2cell}$ during the entire duration of the experiment, longer than with the flush box.

**D.2.4. Conclusions**

With incubation at low pO$_{2gas}$, culture on SR and in the Xvivo system decreases and eliminates, respectively, periodic high-oxygen exposure that occurs with conventional culture in flush boxes. Because of this more consistent exposure to low oxygen with SR and the Xvivo, VEGF content was higher. These results
demonstrate the importance of oxygen-control methodologies, as they have large effects on cell phenotype.
Figure D-1. VEGF content in culture medium for mESC cultured 24 hours at 142 or 7 mmHg pO$_2$gas determined with ELISA.

R1 mESC were culture on either fibronectin-coated polystyrene or silicone rubber in a flush box or Xvivo incubator. * p<0.05 between the two indicated conditions. n=3.
D.3. Experiments with oxygen sensors

D.3.1. Introduction

The pO\textsubscript{2} at the surface of cell culture dishes can differ from that in the gas phase, in particular if a culture vessel without a silicone rubber membrane bottom is used [157]. Mathematical models are available to predict pO\textsubscript{2\text{cell}}; however, these require measurements of the OCR, which represents an added burden to stem cell researchers. Even when OCR is known, prediction of pO\textsubscript{2} within aggregates is more reliable if the pO\textsubscript{2} profile at the surface of the aggregate is measured.

The Xvivo culture system comes with 2-mm diameter oxygen sensors that can be inserted into open culture vessels to measure pO\textsubscript{2} in real time. The Xvivo can also change pO\textsubscript{2\text{gas}} in response to sensor measurements, e.g., to keep pO\textsubscript{2} of the sensor (pO\textsubscript{2\text{sensor}}) at a constant, set value. However, the presence of the oxygen sensor may significantly disrupt the oxygen gradient within culture medium normal to the cell surface that is present in static cell culture.

In the following pilot study, I investigated the disturbance of the oxygen gradient by the oxygen sensor and attempted to use oxygen sensor measurements coupled with Xvivo control of pO\textsubscript{2\text{gas}} to maintain pO\textsubscript{2\text{sensor}} at a set value in mESC culture.
D.3.2. Methods

Finite element modeling was used to simulate pO$_2$ profiles and measurements for oxygen sensors ranging 0.1-1 mm in diameter in mESC culture using previously published oxygen permeability values [157]. To investigate oxygen sensor-based pO$_2$ control, 5x10$^4$ mESC cm$^{-2}$ were seeded on fibronectin-coated polystyrene or silicone rubber 24-well plates with a 2 mm medium height. After 24 hr, the medium was replaced with medium equilibrated at 76 mmHg pO$_2$ and a 2 mm-diameter oxygen sensor placed 0.5 mm above the cells (t=0). The cells were cultured for up to an additional 30 hr with either no process control or processor control where the set point was pO$_2^\text{sensor}$=76 mmHg.

D.3.3. Effects of oxygen sensors on oxygen gradients

The presence of oxygen sensors disrupted the oxygen gradient and caused predicted pO$_2$ measurements lower than the actual pO$_2$ (Figure D-2). Sensors with a 1 and 0.5 mm diameter resulted in predicted measurements that were approximately 20 and 10 mmHg, respectively, lower than the actual pO$_2$ for all heights. The difference between the predicted measured and actual pO$_2$ increased the closer the sensors were to the cells. Small-diameter sensors had small (<10 mmHg) differences between predicted measured and actual pO$_2$ at all heights except near 0 µm, which had a difference greater than 10 mmHg.
Radial oxygen gradients were present on the surface of the oxygen sensor. Depletion of oxygen becomes increasingly severe towards the center of the probe surface with large diameter probes (0.5 and 1.0 mm), virtually reaching 0 mmHg. Smaller probes (0.1 mm and 0.2 mm) resulted in a less dramatic pO₂ drop between the bulk and the region directly underneath the sensor.

D.3.4. Control of pO₂sensor

Measurements of pO₂sensor coupled with the Xvivo control of pO₂gas were used to control the value of pO₂sensor to 76 mmHg (Figure D-3). Without process control, pO₂sensor reached 0 mmHg by 18 hr on polystyrene (Figure D-3A). With process control at early times, large oscillations in the values of pO₂gas and pO₂sensor were seen with polystyrene culture, with pO₂sensor oscillating around 76 mmHg (Figure D-3B). By 30 hrs, pO₂gas had reached the maximum the system is capable of (660 mmHg), and the value of pO₂sensor dropped to 0 mmHg (Figure D-3C). With silicone rubber culture plates, the value of pO₂sensor was maintained at 76 mmHg both with and without process control (data not shown).

D.3.5. Conclusions

Physical insertion of oxygen sensors to measure pO₂cell is often not practical and disrupts the oxygen gradient, resulting in incorrect measurements of the actual pO₂cell in the absence of the probe. Sensors with larger diameters result in
diffusion limitations under the surface. Oxygen sensors with diameters smaller than 200 μm theoretically could avoid this problem but are likely too fragile. The development of practical microsensors or other alternative oxygen measurement methodologies is desirable.

In addition, coupling of these measurements to the Xvivo oxygen control system would provide researchers a way to precisely control the cell surface pO₂ in any commercially available culture dish. However, experiments coupling large 2 mm-diameter probes with the Xvivo control system were unable to control pO₂ in polystyrene dishes, presumably because of the large size of the probe. Use of silicone rubber dishes avoids the need for process control but are not commercially available.
Figure D-2. Numerical simulations of oxygen sensors within oxygen gradient.

(Top) Numerical simulation of pO₂ measurements by probes 0.1, 0.2, 0.5, and 1.0 mm in diameter as a function of height above respiring cells in 2 mm of media. The dashed line is the calculated pO₂ without any probe. (Bottom) Numerical simulation of the pO₂ profile across the surface of probes of different diameters placed 20 μm above the surface of respiring cells.
Figure D-3. Control of $pO_2_{sensor}$ by controlling $pO_2_{gas}$.

mESC plated on polystyrene were cultured for 30 hr with either (A) no process control or (B) process control. The controller attempted to keep $pO_2_{sensor}$ at a set point of 76 mmHg by changing $pO_2_{gas}$. The data plotted in (C) is the same data as in (B) but with a different scale.
D.4. The effect of continuous oxygen control on mESC differentiation to cardiomyocytes

D.4.1. Introduction

Chapter 2 demonstrates that low-pO$_2$ differentiation increases differentiation of mESC to cardiomyocytes. However, the incubation system employed was a flush box, which only controls pO$_2$gas while the cells are inside, but pO$_2$gas is 160 mmHg when the cells are removed for observation and medium exchange. We hypothesized that the continuous low oxygen culture environment afforded by the Xvivo would yield superior yields of cardiomyocytes compare to the period high oxygen exposure of the flush box. This periodic high oxygen exposure is of interest because continuous uninterrupted low oxygen culture is not currently available in most laboratories, and periodic high oxygen exposure is unavoidable.

D.4.2. Methods

J1 mESC were differentiated for 10 days with the differentiation protocol described in Chapter 2 on highly-oxygen permeable silicone rubber membranes with media supplemented with 0.2 mM ascorbic acid inside either flush box and Xvivo culture systems at 142 and 36 mmHg pO$_2$gas. Cells cultured in the flush box were removed and placed in a biological culture cabinet for 1 hour at room temperature and atmospheric O$_2$ and CO$_2$ every 24 hours, after which they were
returned to the flush box. Cells cultured in the Xvivo were cultured continuously at the indicated pO\textsubscript{2}gas and 5% CO\textsubscript{2} or subjected to changes in pO\textsubscript{2}gas or CO\textsubscript{2} as indicated. MF-20+ cells were counted with flow cytometric and cell number counted with nuclei counts, as described in Chapter 2.

**D.4.3. Results**

The fraction and number of MF-20+ cells was the same between cells cultured in the flush box and the Xvivo, regardless of the changes in pO\textsubscript{2}gas and CO\textsubscript{2} (Table D-1). Based on these results, low-pO\textsubscript{2} differentiation of mESC to cardiomyocytes on silicone rubber membranes appears insensitive to these types of periodic high oxygen and low CO\textsubscript{2} perturbations. However, more acute processes, such as VEGF secretion (Appendix D.2), are sensitive to periodic high oxygen exposure. Moreover, mESC VEGF secretion was more sensitive to period high oxygen exposure when the cells were cultured on polystyrene culture plates, which required significantly more time to equilibrate back to low pO\textsubscript{2}gas compared to silicone rubber membranes (Appendix A). Taken together, low-pO\textsubscript{2} differentiation of mESC to cardiomyocytes may be sensitive to period high oxygen exposure if the cells were instead cultured on polystyrene.
D.4.4. Tables

Table D-1. Comparison of different gas control strategies on mESC differentiation to cardiomyocytes

<table>
<thead>
<tr>
<th>Culture System</th>
<th>po2gas (mmHg)</th>
<th>O2 Constant?</th>
<th>CO2 Constant?</th>
<th>Fraction MF-20+ (%)</th>
<th>Number MF-20+ (x10^6 cells)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Flush Box</td>
<td>142</td>
<td>No</td>
<td>No</td>
<td>8.3 ± 3.2</td>
<td>1.0 ± 0.53</td>
</tr>
<tr>
<td></td>
<td>36</td>
<td>No</td>
<td>No</td>
<td>16.2 ± 1.7</td>
<td>3.7 ± 0.068</td>
</tr>
<tr>
<td>Xvivo</td>
<td>142</td>
<td>Yes</td>
<td>Yes</td>
<td>8.0 *</td>
<td>1.5 *</td>
</tr>
<tr>
<td></td>
<td>36</td>
<td>Yes</td>
<td>Yes</td>
<td>16.5 ± 3.5</td>
<td>3.8 ± 1.3</td>
</tr>
<tr>
<td></td>
<td>36</td>
<td>No*</td>
<td>Yes</td>
<td>16.2 ± 1.4</td>
<td>4.2 ± 1.3</td>
</tr>
<tr>
<td></td>
<td>36</td>
<td>Yes</td>
<td>No b</td>
<td>16.3 *</td>
<td>3.8 *</td>
</tr>
</tbody>
</table>

n=2 for all samples except where designated by *, where n=1 due to infection

a po2gas was changed to 160 mmHg for 1 hr every 24 hr

b CO2 was changed to 0% for 1 hr every 24 hr