Do Bioreactor Designs with More Efficient Oxygen Supply to Ovarian Cortical Tissue Fragments Enhance Follicle Viability and Growth In Vitro?

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Abstract: Background: Autotransplantation of cryopreserved ovarian tissue is currently the main option to preserve fertility for cancer patients. To avoid cancer cell reintroduction at transplantation, a multi-step culture system has been proposed to obtain fully competent oocytes for in vitro fertilization. Current in vitro systems are limited by the low number and health of secondary follicles produced during the first step culture of ovarian tissue fragments. To overcome such limitations, bioreactor designs have been proposed to enhance oxygen supply to the tissue, with inconsistent results. This retrospective study investigates, on theoretical grounds, whether the lack of a rational design of the proposed bioreactors prevented the full exploitation of follicle growth potential. Methods: Models describing oxygen transport in bioreactors and tissue were developed and used to predict oxygen availability inside ovarian tissue in the pertinent literature. Results: The proposed theoretical analysis suggests that a successful outcome is associated with enhanced oxygen availability in the cultured tissue in the considered bioreactor designs. This suggests that a rational approach to bioreactor design for ovarian tissue culture in vitro may help exploit tissue potential to support follicle growth.

Keywords: bioreactor; design; in vitro culture; ovarian tissue; oxygen; transport

1. Introduction

Ovarian tissue cryopreservation and autotransplantation is currently the only option to preserve the fertility of pre-pubertal girls or patients in need of immediate cancer therapy. To date, autotransplantation of frozen/thawed ovarian cortical tissue has yielded about 130 live births worldwide [1]. Clinical application of this procedure is limited by the possible transmission of cancerous cells on the re-implantation of tissue from patients with blood-borne or highly metastatic malignancies [2–4]. The fact that the follicular basal lamina hinders cancerous cell invasion of ovarian follicles [5] has favored the development of two safer alternative strategies. The first strategy is based on the in vitro culture of ovarian cortical tissue to activate the pool of primordial follicles and promote their growth to the secondary stage, (2) the isolation of secondary follicles from the ovarian stroma, (3) the in vitro culture of isolated follicles to promote the development of antral follicles, and (4) the final aspiration and maturation of cumulus-oocyte-complexes for the in vitro fertilization. Neither of these has yet made it to routine clinical practice. In fact, the isolation of
human primordial follicles is limited by the damage caused by the enzymatic digestion or mechanical disruption of the tough ovarian stromal cortex [10]. Current in vitro systems do not provide efficient support to the culture of viable and functional frozen/thawed ovarian tissue and promote the growth of only a limited number of secondary follicles.

Within this framework, considerable research efforts have been undertaken to optimize the culture media and the follicle encapsulating materials used in vitro with the intent of bringing the latter approach to the clinic. Reviewing the outcome of such research is beyond the scope of this paper. Detailed information may be found in the excellent reviews [5,11,12] and consensus papers [13] available in the literature.

Less attention has been devoted to the bioreactors used. The bioreactor most widely used for the culture of fragments of fresh or cryopreserved ovarian cortical tissue is still the conventional static culture dish (CD). In CDs, tissue fragments are cultured under a static layer of medium (at times with an oil overlay) under a controlled gaseous atmosphere containing oxygen (O\textsubscript{2}) and carbon dioxide (CO\textsubscript{2}). Various bioreactor designs have been explored to overcome the limitations associated with the use of CDs [14], which were intended to enhance the transport of dissolved gases (i.e., O\textsubscript{2} and CO\textsubscript{2}) from the gaseous atmosphere in the incubator to the follicles in the tissue. The inconsistent results reported [15–19] cast doubt on the fact that the bioreactor designs that enhance gas transport may significantly influence follicle viability and growth in the in vitro culture of ovarian cortical tissue. The assumption of this study is that the lack of a rationale in the design of the proposed bioreactors (i.e., the definition of their configuration, geometry, and operation) has thus far prevented exploitation of these bioreactors’ performance potential.

To test such an assumption, in this paper a theoretical analysis is proposed of O\textsubscript{2} transport to fragments of ovarian cortical tissue in some exemplary bioreactor designs investigated as an alternative to CDs. The analysis is used to review and seek for a rationale regarding the reported experimental results of an in vitro culture of human ovarian cortex fragments. Albeit in an approximate fashion, the theoretical analysis describes the complex interplay between the physical transport of O\textsubscript{2} in the bioreactor and inside the tissue stroma, and its metabolic consumption inside the tissue fragment. Attention was focused on O\textsubscript{2} for a number of reasons. O\textsubscript{2} is important because ovaries have a high oxygen consumption rate per unit of tissue mass (i.e., 1.4 mL/(min 100g\textsubscript{tissue})) [20] lower only than heart (i.e., 10.7 mL/(min 100g\textsubscript{tissue})), kidney and liver (i.e., ca. 5 mL/(min 100g\textsubscript{tissue})), and brain (i.e., 3.5 mL/(min 100g\textsubscript{tissue})) tissue [21,22]. In vivo, during normal growth, the outer layers of the follicle (e.g., the theca layers) are vascularized, which ensures adequate oocyte oxygenation. Poor oxygenation has been associated with cytoplasmic and nuclear abnormalities [23]. Ex vivo, grafting of bovine cortical fragments beneath the chorioallantoic membrane of gonadectomized chick embryos has shown that the strip vascularization (and the consequent enhancement of O\textsubscript{2} transport) allowed for the activation of primordial follicles and progression to the secondary stage [24]. In vitro, the dissolved O\textsubscript{2} concentration has been reported to affect the development of human embryos [25] and follicles, both in cortical strips [26] and isolated [27–29], as well as the viability of oocytes [30,31], sometimes with inconsistent species-dependent indications. In the in vitro culture of fragments of human ovarian cortical tissue, it has been reported that gaseous O\textsubscript{2} concentrations higher than in air ensure adequate oxygenation and promote follicle viability long-term, as well as follicle activation and progression [32]. Recently, it has been shown that too low or too high dissolved O\textsubscript{2} concentrations in tissue are detrimental to both follicle viability and development [19]. In this analysis, the effect of gaseous CO\textsubscript{2} transport was not considered because in Talevi et al. [19], it was shown to be less relevant than O\textsubscript{2}, at least for the bioreactors and the conditions investigated therein.

The proposed theoretical analysis associates the culture outcome with oxygen availability in tissue and provides a rationale for the understanding of both the successful and unsuccessful outcome of in vitro culture experiments with the considered bioreactor designs. This suggests that a rational approach to the development of novel bioreactors, ensuring that ovarian cells are cultured in vitro
under controlled dissolved $O_2$ concentrations in fragments of human ovarian cortical tissue, might have a significant impact on the exploitation of the full fertility potential of such tissue.

2. Materials and Methods

2.1. Bioreactor Designs

In this review and the analysis herein reported, only papers were included which reported on bioreactor designs with an $O_2$ supply to follicles in human ovarian cortical tissue that were intended to be more efficient than CDs, where the information reported permitted the estimation of dissolved $O_2$ transport and metabolic parameters, as well as the comparison of the experimental outcome with CDs.

Information on the considered bioreactor designs is reported in Figure 1 and Table 1. In the conventional dish, taken as the control, $O_2$ is transported from the above gaseous atmosphere to the upper surface of the tissue fragments across a static layer of medium (Figure 1a). To reach follicles and stromal cells inside the avascular cortex fragment, $O_2$ has to: (i) partition between the gas and the medium at the gas–medium interface; (ii) be transported across the layer of medium between the gas phase and the upper fragment surface; and (iii) be transported across the fragment thickness, mainly by passive diffusion, while it is consumed by follicular and stromal cells [14]. The poor solubility of $O_2$ in medium, the resistance to $O_2$ transport of medium and stromal tissue, and the $O_2$ metabolic consumption of stromal and follicular cells all contribute to making the dissolved $O_2$ concentration inside the fragments much lower than in air.

![Figure 1](image-url)  
*Figure 1*. Schemes of the bioreactor configurations used for the in vitro culture of fragments of ovarian cortical tissue in the literature reports analyzed in this study: (a) conventional dish (CD), (b) CD with medium mixed by holding the CD on an orbital shaker during culturing [17], (c) CD with medium mixed by recirculating medium above the tissue fragments with an external pump [18], (d) submerged fragment culture on a permeable membrane at the bottom of an insert suspended in a well of a 24-well plate [15,16,33], (e) fragment culture on a permeable membrane at the bottom of an insert suspended in a well of a 24-well plate under a drop of medium [16], and (f) dish with a gas-permeable bottom (PD) [19]. Gas supply portrayed as green arrows.
Table 1. Culture conditions in the literature reports analyzed in this study. Legend for supplements: AA—ascorbic acid; Am—amphotericin; BrG—8-bromo-guanosine; BSA—bovine serum albumin; EGF—epidermal growth factor; FCS—fetal calf serum; FSH—follicle stimulating hormone; Glu—glutamine; ITS—insulin-transferrin-selenium; IHS—inactivated human serum; Ins—insulin; LH—luteinizing hormone; PS—Pen-Strep; Pyr—pyruvate. Abbreviations: Morph—morphological; N—no; Y—yes.

| Authors          | Medium Used            | Growth-Affecting Supplements | Culture System | Support Coating | Culture Mode | No Strips Per Well | Medium Volume | Medium Change | pO2 | Tissue | Data on Fresh Tissue | Culture Time Days | Viability | Stage | Grade | Hormones | Reference |
|------------------|------------------------|------------------------------|----------------|-----------------|--------------|-------------------|---------------|---------------|-----|--------|----------------------|-------------------|-----------|-------|-------|----------|-----------|
| Hovatta et al., 1997 | -MEM, Earle’s          | FSH, IHS, Ins, LH, Pyr       | Inserts in well in 24 well plate | none, Matrigel<sup>1</sup> | static, double medium layer | 1–3 | NA | 2nd day | air | Fresh & thawed | Y | up to 21 | Morph | Y | N | N | [15] |
| Wright et al., 1999 | -MEM,                   | FSH, HSA, ITS-G              | Inserts in well in 24 well plate | Matrigel<sup>1</sup> | static, double medium layer | NA | 100 ul. + 400 ul. | 150 ul. @ 2nd day | air | Fresh & thawed | Y | up to 14 | Morph | Y | N | N | [16] |
| Isachenko et al., 2006 | Iskove’s modified       | EGF, FCS, FSH, ITS          | 200 mL dishes | none | static vs. mixed on orbital shaker | 20 | 30 mL | NO | air | Fresh & thawed | N | up to 21 | Morph | Y | N | N | [17] |
| Liebenthron et al., 2013 | McCoy’s 5a             | AA, BSA, Glu, ITS           | 6 well plate | none | static vs. medium recirculation | NA | 4.5 mL | NO | air | Fresh | Y | up to 6 | Morph calcein AM | Y | N | Y | [18] |
| Talevi et al., 2018   | -MEM                   | AA, Am, BSA, Glu, ITS, PS    | 50 mm dishes | none | static, conventional vs. permeable bottom | 10 | 5 mL | half every other day | air | Fresh | Y | up to 9 | Morph dead/live | Y | Y | N | [19] |
To enhance the \( \text{O}_2 \) transport from the gas/medium interface to the upper fragment surface, it has been proposed that the medium be mixed either by holding CDs on an orbital shaker (Figure 1b) [17], or by periodically recirculating medium above the fragments with an external pump (Figure 1c) [18] during culturing. Alternatively, it has been that the \( \text{O}_2 \) be supplied from both the upper and the lower surface of the tissue fragments. This has been pursued either by culturing the fragments on permeable inserts suspended in a well of a multi-well plate submerged in static medium (Figure 1d–e) [15,16], or in dishes with a gas-permeable bottom (PD) (Figure 1f) [19]. In both cases, minimization of the resistance to \( \text{O}_2 \) transport external to the fragments has been pursued using a culture under medium layers of decreasing thicknesses. In a few cases, \( \text{O}_2 \) transport to follicles has been enhanced by decreasing the thickness of the tissue fragments [15,33].

2.2. Models of Oxygen Transport

To investigate possible associations between culture outcome and the level of oxygen at which ovarian cells are cultured inside a fragment, estimates of the dissolved oxygen concentration anywhere in the tissue fragment are needed for varying bioreactor designs and operating conditions. In this analysis, such estimates were obtained by developing mathematical models describing the transport of dissolved oxygen from the gaseous oxygen source across the medium and inside the ovarian cortical tissue for the bioreactors and the operating conditions used for the considered papers. In the transport models, the considered bioreactors were schematically described, as shown in Figure 2. The tissue fragments were approximated as a single tissue slab, entirely covering the bottom of the dish or the insert, with uniform transport and metabolic properties. The latter assumption implies that follicles are uniformly distributed in the tissue, are all at the same developmental stage, and consume \( \text{O}_2 \) at the same rate. \( \text{O}_2 \) was assumed to be supplied only through the upper (in some cases also the lower) fragment surface at a steady rate. According to the film theory [34], the resistance to \( \text{O}_2 \) transport at any interface between the medium and tissue is concentrated in a stagnant layer of medium adherent to the tissue surface, the thickness of which depends on the bioreactor geometry, the volume of medium used, and the degree of medium mixing. In statically operated bioreactors, the stagnant layer thickness equals the height of medium under which tissue is cultured. In bioreactors in which the medium is mechanically or fluid-mechanically mixed, the resistance to \( \text{O}_2 \) transport was estimated in terms of an \( \text{O}_2 \) transport coefficient, \( k_L \), from available semi-empirical correlations for the given bioreactor geometry and fluid dynamic conditions. The model assumes that an anoxic zone forms farthest from the \( \text{O}_2 \) source if \( \text{O}_2 \) is supplied only from the upper fragment surface, or in the middle of the fragment thickness if \( \text{O}_2 \) is supplied from both the upper and the lower fragment surfaces.

The \( \text{O}_2 \) transport model for each bioreactor design was based on previously published models for dissolved \( \text{O}_2 \) transport in ovarian cortical tissue cultured under a layer of stagnant medium in CDs and PDs [19]. For this reason, each model’s description may resemble, occasionally verbatim, that reported in Talevi et al. [19]. Briefly, in bioreactors in which \( \text{O}_2 \) is supplied only through the upper fragment surface (Figure 2a), the mass balance equations for \( \text{O}_2 \) in tissue were obtained in analogy to purely diffusive heat transport in a homogeneous slab when the resistance to transport from the liquid in contact with the slab to the slab outer surface is not negligible [34]. A reference system was used with a z-axis with the origin at the lower fragment surface, oriented towards the medium above. The mass balance equations were expressed in dimensionless form to minimize the bias caused by the lack of reliable estimates of the geometric, transport, and metabolic properties of the ovarian cortical tissue fragments. This also suggests how such properties combine in the dimensionless groups to determine the dissolved \( \text{O}_2 \) concentration profile across the tissue fragment (and the bioreactor performance). The dimensionless dissolved \( \text{O}_2 \) concentration in tissue, \( C_T^* \), and the dimensionless space coordinate, \( z^* \), were defined with respect to the dissolved \( \text{O}_2 \) concentration that equilibrates with the gaseous \( \text{O}_2 \) pressure at the gas-medium interface, \( C_B \), and the fragment thickness, \( h_T \), respectively, as follows:
$C_T^* = C_T/C_B$ and $z^* = z/\delta_T$. The one-dimensional steady state mass balance of O$_2$ about an infinitesimal tissue control volume $Adz$ yields:

$$\frac{d^2 C_T^*}{dz^*} = \frac{G'''}{D_T C_B} \delta_T^2 \phi^2$$

subject to the following boundary conditions:

$$BC1: \frac{d}{dz^*} \left. \left( \phi C_T^* \right) \right|_{z^*=1} = \frac{k_L}{D_T} \delta_T \left( 1 - C_T^*(z^* = 1) \right) = B_i m \left( 1 - C_T^*(z^* = 1) \right)$$

$$BC2: z^* = \delta_{i,U}/\delta_T \quad C_T^* = 0$$

where $\delta_{i,U}$ is the distance from the fragment bottom at which the anoxic zone begins. BC1 states that the oxygen transported across the medium diffuses into the tissue at the same rate.

\begin{figure}
\centering
\includegraphics[width=\textwidth]{figure2.png}
\caption{Bioreactor schemes used for the transport models and qualitative dissolved oxygen concentration profile in the medium and inside the fragment: (a) conventional or mixed dish in which oxygen is supplied only through the upper fragment surface, and (b) submerged fragment culture on a permeable membrane at the bottom of an insert suspended in a well of a 24-well plate. Captions: C—dissolved oxygen concentration; C$_B$—dissolved oxygen concentration in the medium at equilibrium with pO$_2$gas; pO$_2$gas—gaseous oxygen tension above the medium surface; $\delta_m$ and $\delta_{m,U}$ or $\delta_{m,B}$—thickness of the medium layer above or below the strip, respectively; $\delta_T$—thickness of the tissue fragment; $\Delta$CO$_2$ext—external dissolved oxygen concentration drop in the medium; $\Delta$CO$_2$int—internal dissolved oxygen concentration drop in the fragment. Other symbols are defined in the text. Please note that drawings are not to scale.}
\end{figure}

Integration of Equations (1) and (2) yields the dimensionless dissolved O$_2$ concentration $C_T^*$ at any dimensionless distance $z^*$ from the bottom of the fragment, as follows [19]:

$$C_T^* = \frac{1}{2} \phi^2 \left[ \left( \frac{\delta_{i,U}}{\delta_T} \right)^2 + \left( \frac{z}{\delta_T} \right)^2 - 2 \frac{\delta_{i,U}}{\delta_T} \frac{z}{\delta_T} \right]$$

with

$$\frac{\delta_{i,U}}{\delta_T} = 1 - \left[ \frac{1}{\phi} \sqrt{\frac{\phi^2}{B_i m^2} + 2 - \frac{1}{B_i m}} \right]; \quad \phi^2 = \frac{G'''}{D_T C_B} \delta_T^2; \quad B_i m = k_L \frac{\delta_T}{D_T}$$

where: $D_m$ and $D_T$ are the O$_2$ diffusivity in the medium and tissue, respectively; $G'''$ is the cellular O$_2$ consumption rate per unit tissue volume; $\delta_{i,U}$ and $\delta_T$ is the thickness of the stagnant medium layer above the fragment and the fragment thickness, respectively; $B_i m$ is the dimensionless mass Biot number at
the upper fragment surface; and \( \phi \) is the dimensionless Thiele modulus. \( \phi \) compares the maximal rate of \( \text{O}_2 \) cellular consumption to its diffusion in tissue. \( \text{Bi}_m \) compares the dissolved \( \text{O}_2 \) concentration drop inside the tissue to that in the stagnant medium layer adhering on the given fragment surface through which \( \text{O}_2 \) is supplied to the ovarian cells in the fragment.

In bioreactor designs in which \( \text{O}_2 \) is also supplied through the lower fragment surface (Figure 2b), the formation of an anoxic zone in the middle of the fragment permits us to obtain the dissolved \( \text{O}_2 \) concentration profile in the lower part of the fragment independent of the \( \text{O}_2 \) supply from the upper surface. The mass balance equations for the dissolved \( \text{O}_2 \) concentration in the lower part of the tissue fragment were written with reference to a \( z' \)-axis with origin \( O' \) at the upper fragment surface and oriented towards the medium below. In bioreactor designs in which \( \text{O}_2 \) is transported across a layer of stagnant medium in contact with the lower fragment surface, the boundary conditions were assumed to be identical to Equation (2), but for the \( \text{Bi}_m \) at the lower fragment surface. Integration of the mass balance equations and boundary conditions yields the dissolved \( \text{O}_2 \) concentration profiles and the distance at which the anoxic zone begins to form from the bottom, \( \delta_{i,B} \), similar to Equations (3) and (4). The transformation \( z' = \delta_T - z \) yields the actual profiles in terms of \( z \) and \( z' \). In bioreactors in which the tissue fragment is in direct contact with a gas-permeable membrane exposed to the same gaseous atmosphere as the medium above the tissue, Equation (2a) was replaced by the condition stating that the dissolved \( \text{O}_2 \) concentration at the lower tissue surface equals \( C_B \). The integration of the mass balance equation and the new boundary conditions yields [19]:

\[
C_T' = \left[ 1 + \frac{1}{2} \phi^2 \left( \frac{z}{\delta_T} \right)^2 - \sqrt{2} \phi \left( \frac{z}{\delta_T} \right) \right]^{3/2}
\] (3')

Equations (3), (4), and (3') show that the dissolved \( \text{O}_2 \) concentration in tissue depends on the \( \text{O}_2 \) transport properties in medium and tissue, on the cellular \( \text{O}_2 \) metabolic consumption, and on the fragment geometry, as well as on the bioreactor geometry and fluid dynamics described by the dimensionless Thiele modulus, \( \phi \), and the mass Biot number, \( \text{Bi}_m \). Model predictions were obtained for \( C_B = 0.2 \text{ mol}/\text{m}^3 \) (corresponding to a \( \text{pCO}_2 \) of about 21%), \( D_T = 2.8 \times 10^{-9} \text{ m}^2/\text{s} \), \( D_m = 3.5 \times 10^{-9} \text{ m}^2/\text{s} \), \( C'' = 2 \times 10^{-2} \text{ mol/(s·m}^3 \) [19], assuming a density \( \rho = 9.97 \times 10^2 \text{ kg/m}^3 \) [35] and a dynamic viscosity \( \mu = 8.2 \times 10^{-4} \text{ kg/(m·s)} \) for the medium [36].

The efficiency of \( \text{O}_2 \) transport to the tissue was expressed in terms of the oxygen availability in the tissue (i.e., the volume-averaged dissolved \( \text{O}_2 \) concentration in the fragment), \( \overline{C}_{T,avg} \), and the width of the anoxic zone, \( w_{az} \). \( w_{az} \) was defined as \( w_{az} = \delta_{i,LI} \) in bioreactors in which \( \text{O}_2 \) is supplied only through the upper fragment surface, and as \( w_{az} = \delta_{i,LI} - (\delta_T - \delta_{i,B}) \) in bioreactor designs in which the \( \text{O}_2 \) is supplied through both fragment surfaces.

2.3. Estimation of the Oxygen Transport Coefficient, \( k_L \)

2.3.1. Static Bioreactors

In static bioreactors, \( \text{O}_2 \) is transported by pure diffusion across the layer of stagnant medium between the gas-medium interface and the fragment surface. Hence, \( k_L \) was estimated as the ratio of the \( \text{O}_2 \) diffusivity in the medium to the stagnant medium layer thickness, \( k_L = D_m/\delta_m \). In bioreactors in which \( \text{O}_2 \) is supplied only across the upper fragment surface, \( \delta_m \) was estimated using the medium volume used for the culture for a dish diameter of 14.5 cm [17] or 3.5 cm [18]. In the bioreactors used for the culture on an insert, and in which \( \text{O}_2 \) is supplied across both the upper and the lower fragment surface, the diameter of a well in a 24-well plate was assumed to be equal to 1.56 cm [15,16,33]. The thickness of the upper medium layer, \( \delta_{m,LI} \), was estimated from the medium volume by neglecting both the insert and membrane volumes and by assuming a 1 mm distance between the bottom of the insert and the well [15,33,36,37] or it was assumed to be equal to 500 \( \mu \text{m} \) (as for experiment III in Wright et al. [16]). The thickness of the lower medium layer, \( \delta_{m,B} \), was estimated from the medium
volume assuming a straight diffusive path for O₂ from the gaseous phase to the membrane bottom in the insert, as shown in Figure 2b.

2.3.2. Mixed Bioreactors

In mixed bioreactors, the O₂ transport coefficient in medium, \( k_L \), was estimated using available semi-empirical correlations for the given bioreactor type, geometry, and fluid dynamic conditions. In large dishes held on an orbital shaker [17], the increase of the gas exchanging surface area caused by the shaking may be neglected [38]. Hence, \( k_L \) was estimated by adapting the following semi-empirical correlation for shaking flasks to circular dishes [39]:

\[
k_L a = 0.5 \frac{d^7}{36} n d_0^{1/4} V_m^{-8/9} D_m^{1/2} \nu^{3/54} g^{-7/54}
\]  

where \( a \) is the gas exchanging medium surface-to-volume ratio in the dish, \( d \) is the inner dish diameter, \( d_0 \) is the shaking orbit diameter, \( n \) is the shaking frequency, \( V_m \) is the medium volume, \( \nu = \rho/\mu \) is medium kinematic viscosity, and \( g \) is the gravitational acceleration. Model predictions were obtained for an orbital diameter \( d_0 = 1.9 \text{ cm} \).

In culture dishes in which the medium is periodically recirculated above the tissue fragments, the actual fluid dynamic depends on the position and geometry of the influent and effluent cannulas. In the absence of such information, transport efficiency was estimated by assuming that the medium continuously flows as a plug above the fragments located at the bottom of the dish. The local \( k_L \) value corresponding at any given chord length normal to medium flow in the dish was estimated from the following semi-empirical correlation [40]:

\[
k_L z/D_m = 0.323 \left( \frac{v(z)z}{v} \right)^{1/2} \left( \frac{\nu}{D_m} \right)^{1/3}
\]  

where \( z \) is the distance from the inlet section and \( v(z) \) is the current medium velocity at the distance \( z \). A \( k_L \) value averaged over the dish surface, \( k_{L,avg} \), was used to obtain model predictions.

3. Results and Discussion

Equations (3) and (4) show that in static CDs, the actual dissolved O₂ concentration inside the tissue depends on the gaseous O₂ pressure (i.e., \( C_B \)), on the resistance to O₂ transport of the medium layer and tissue (i.e., \( 1/k_L \) and \( \delta_T/D_T \)), and on the metabolic rate at which the cells in the ovarian cortex consume O₂ (i.e., \( G''' \)). Optimization of bioreactor design to provide ovarian cells with an adequate O₂ supply has often been approached by trying to optimize some of the above variables one at a time (e.g., either \( C_B \), or \( k_L \), or else \( \delta_T \)) directly in culture experiments. Such an approach has often yielded inconsistent results. For instance, an increase of the O₂ transport coefficient, \( k_L \), obtained either by increasing the degree of medium mixing [17,18] or by decreasing the thickness of the upper medium layer (i.e., between the gaseous atmosphere and the upper fragment surface) [15,16,33] has not always enhanced follicle viability.

Theoretical models of O₂ transport may help in understanding the interplay among contributing variables and parameters and the reported culture outcome. Models have been proposed to describe dissolved O₂ transport in pre-antral and antral follicles [41–44] that account to varying degrees for the changing position and geometry of the oocyte in the growing follicle when the dissolved O₂ concentration outside the follicle equals that in the blood vessels surrounding it. Such models have been valuable for understanding the role of antrum formation and of follicular cells’ metabolism in ensuring adequate oxygenation to the oocyte in the preovulatory follicle in vivo. The use of such models to estimate O₂ availability to follicles in fragments of ovarian cortical tissue cultured in vitro is impractical. In fact, they do not account for the effects on O₂ transport from an external gaseous source of the stromal extracellular matrix and the cells in cortical tissue, nor for bioreactor geometry and fluid dynamics.
The models discussed here describe O\textsubscript{2} transport in fragments of ovarian cortical tissue cultured in vitro in various bioreactor designs, and account, albeit in an approximate fashion, for tissue properties and bioreactors geometry and fluid dynamics. Ideally, the models ought to describe the extent to which the different cells and stroma components present in tissue determine its transport and metabolic properties. Bioreactor geometry and fluid dynamics should also be described in detail. The scarcity of reliable quantitative information on the composition and structure of the ovarian cortical tissue, the metabolic functions of the ovarian cells, and the bioreactors used makes it challenging to build reliable complex transport models. To obtain qualitative predictions of the dissolved O\textsubscript{2} distribution inside a tissue fragment, a less detailed but effective description of ovarian cortical tissue was used in terms of uniform lumped transport and metabolic characteristics, the value of which could be estimated from literature information for human ovarian cortex or similar tissue. Results and model predictions obtained for different bioreactor designs were generally compared when reported in the same paper, or in different papers by the same research group, to minimize the bias caused by the broad variability of medium composition and supplements, and of the methods used to assess follicle viability and stage. The broad variability of the follicle number and stage in ovarian cortical tissue from different species and age, and even in tissue harvested from different regions of the same ovary [45], contributes to making data interpretation even more problematic.

The model-predicted O\textsubscript{2} availability and width of the anoxic zone inside the tissue fragment for the considered bioreactor configurations and operating conditions are reported in Table 2. The value of \( \phi \), consistently greater than unity, suggests that, under all conditions, the dissolved O\textsubscript{2} concentration in tissue steeply decreases towards the innermost regions of the fragment as a result of the large diffusional resistance of tissue, the more so the greater the \( \phi \) [46]. The value of \( Bi_{in} \) varies by two orders of magnitude and suggests that the interplay of transport and metabolic phenomena is rather different in the various bioreactor configurations yielding broadly different dissolved O\textsubscript{2} concentrations at the tissue–medium interfaces. In bioreactors in which O\textsubscript{2} is supplied only from the upper fragment surface, Equations (3) and (4) suggest that the combination of \( \phi \) and \( Bi_{in} \), rather than each single group, determines the dissolved O\textsubscript{2} concentration profile inside tissue and the width of the anoxic zone, \( \omega_{az} \). In bioreactors in which O\textsubscript{2} is also supplied through the lower fragment surface, the dissolved O\textsubscript{2} profile inside tissue and \( \omega_{az} \) is determined by the combination of the values of \( \phi \) and \( Bi_{in} \) at both medium-fragment interfaces.

First, the models were used to analyze the inconsistent results obtained with bioreactors in which O\textsubscript{2} transport is enhanced with respect to CDs by increasing \( k_{L} \) (bioreactor #3 vs. #4 and bioreactor #5 vs. #6 in Table 2). The transport models predicted a very poor O\textsubscript{2} availability and an anoxic zone well in excess of 95% of the fragment thickness when the fragments are cultured on an insert in a well, both under an upper medium layer about 2.3 mm thick and a much thinner layer of about 0.5 mm (i.e., bioreactors #5 vs. #6 in Table 2) [16]. Similar predictions were obtained for fragments cultured in static CDs or in CDs in which the medium was mixed by recirculating it above the fragments with an external pump (i.e., bioreactor #3 vs. #4 in Table 2) [18]. The models’ predictions suggest that the bioreactor designs used did not ensure a good O\textsubscript{2} supply to the ovarian cells, irrespective of how they were operated. Under such conditions, cells likely starved for O\textsubscript{2} and struggled for survival. It is no surprise that follicle viability was not found to be any different between each pair of bioreactors.

Second, the models were used to analyze the successful results obtained with bioreactors in which O\textsubscript{2} transport is enhanced by increasing \( k_{L} \) or by decreasing the fragment thickness, \( \delta_{T} \). In bioreactors in which \( k_{L} \) is enhanced with respect to static CDs by holding the CDs on an orbital shaker during culture [17] (i.e., bioreactors #1 vs. #2 in Table 2), the models predicted a much higher O\textsubscript{2} availability and an anoxic zone about 20% smaller in mixed CDs than in static CDs (Table 2). Correspondingly, the number of viable follicles was about 2.6 times higher, with a higher proportion of morphologically intact follicles, than in CDs. Also, in bioreactors in which tissue fragments were statically cultured under a layer of medium at the bottom of gas permeable dishes (PDs), in direct contact with an O\textsubscript{2} source [19] (i.e., bioreactor #8 vs. #9 in Table 2) the models predicted a higher O\textsubscript{2} availability in PDs.
than in CDs with a gas-impermeable bottom. Correspondingly, the width of the anoxic core in PDs was one third of that in CDs. Histological and viability analysis showed that culture in PDs for up to 9 days enhanced the follicle viability and progression to the secondary stage over cultures in CDs. In one case, the tissue resistance to oxygen transport was reduced by culturing tissue fragments of decreasing thickness (i.e., 2 vs. 0.3 mm) on an insert placed in a well of a 24-well plate (i.e., bioreactors #5 vs. #7 in Table 2) [15,16]. The models predicted an O$_2$ availability two times higher and a 15% smaller anoxic zone in the bioreactors with thinner tissue fragments. Correspondingly, the number of atretic follicles in thinner fragments was about one fourth of that in thicker fragments (4% vs. 17%, respectively).

That which is reported above shows that models of O$_2$ transport in the ovarian cortex and bioreactors provide rational grounds for understanding the experimental results reported for all considered bioreactors, some of which would otherwise appear inconsistent. This lends support to this study’s assumption that the rational optimization of bioreactor design for the in vitro culture of fragments of ovarian cortical tissue may ensure adequate oxygenation to the ovarian cells and may enhance follicle viability and growth. A reliable transport model that predicts the dissolved O$_2$ concentration anywhere in the tissue fragment as a function of its value in the medium bulk for a given fragment and bioreactor geometry and fluid dynamics would also provide a means to indirectly monitor the dissolved O$_2$ concentration in tissue from measurements of the dissolved O$_2$ concentration in the medium bulk, the only measurable parameter. This would make it possible for an operator to change the operating conditions and adjust the supply of O$_2$ (and also of nutrients and metabolic effectors) to the current cell metabolic requirements so as to maintain the follicles viability and possibly guide their progression during culture.

A model, however, is as good as its assumptions and its parameters estimates. To increase the reliability of a model’s predictions, it would be necessary to quantitatively characterize the structure of the ovarian cortical tissue and the metabolic requirements of the ovarian cells at any developmental stage. Flow and mass transport in the bioreactor ought to be based on a more detailed description of bioreactor and fragment geometry. CO$_2$ transport and the effect of pericellular pH changes on ovarian cell metabolism should also be accounted for because they might also play a role in bioreactor configurations with a high tissue-to-medium volume ratio.

In spite of its limitations, the model-based review and analysis of reports on the in vitro culture of fragments of ovarian cortical tissue suggests that bioreactor designs enhancing the O$_2$ supply to the ovarian cells have the potential to maintain about 65% [19] of follicles as viable for longer than a week, and to activate up to about 20% of these [8,15,19] to progress from the primordial to the secondary stage, without significant differences between fresh and frozen-thawed tissue. However, in the long-term, in vitro culture follicle quality decreases (with only 45% grade I follicles after 9 days of culture [19], and the proportion of viable follicles decreases as well. Correspondingly, the proportion of degenerating/atretic follicles increases from a few units to 14–50% [8,15] after a week of culture. This suggests that there is still quite some room to further optimize bioreactor design to enhance O$_2$ transport to ovarian cells and to adjust it to their changing requirements as follicles grow, as a possible means to guide their progression in vitro.
Table 2. Model-predicted oxygen availability, CT, avg, width of anoxic zone, wac, and O$_2$ concentration at tissue medium interface, CT, m, for the bioreactor geometry and fluid dynamic conditions used in the literature reports analyzed in this study. Other symbols are defined in the text.

| Bioreactor # | Bioreactor Type       | Fragment Side of O$_2$ Supply | $\delta_T$ m | $\delta_m$ m | $k_L$ m/s | Thiele Number $\phi$ | Biot Number $Bi_m$ | Oxygen Availability | Anoxic Core | Reference |
|--------------|-----------------------|-------------------------------|---------------|---------------|------------|----------------------|---------------------|---------------------|-------------|-----------|
| 1            | Static CD             | Upper                         | $1.00 \times 10^{-3}$ | $8.18 \times 10^{-4}$ | $4.28 \times 10^{-6}$ | 6.0 | 1.53 | 3.07 | 3.07 $\times 10^{-3}$ | 95.85 [17]   |
| 2            | CD on orbital shaker  | Upper                         | $1.00 \times 10^{-3}$ | NA            | $1.13 \times 10^{-4}$ | 6.0 | 40.37 | 81.14 | 3.15 $\times 10^{-1}$ | 78.68 [17]   |
| 3            | Static CD             | Upper                         | $1.00 \times 10^{-3}$ | $3.69 \times 10^{-3}$ | $9.49 \times 10^{-7}$ | 6.0 | 0.34 | 0.16 | 1.60 $\times 10^{-4}$ | 99.05 [18]   |
| 4            | CD w/ periodic medium flow | Upper                          | $1.00 \times 10^{-3}$ | NA            | $9.94 \times 10^{-7}$ | 6.0 | 0.36 | 0.18 | 1.76 $\times 10^{-4}$ | 99.01 [18]   |
| 5            | Culture on insert in well | Upper                          | $2.00 \times 10^{-3}$ | $2.26 \times 10^{-3}$ | $1.55 \times 10^{-6}$ | 12.0 | 1.10 | 0.42 | 5.44 $\times 10^{-4}$ | 98.82 [16]   |
| 6            | Culture on insert in well | Upper                          | $2.00 \times 10^{-3}$ | $5.00 \times 10^{-4}$ | $7.00 \times 10^{-6}$ | 12.0 | 5.00 | 7.49 | 1.69 $\times 10^{-5}$ | 95.61 [16]   |
| 7            | Culture on insert in well | Upper                          | $3.00 \times 10^{-4}$ | $1.33 \times 10^{-3}$ | $2.63 \times 10^{-6}$ | 1.8  | 0.28 | 1.20 | 1.53 $\times 10^{-3}$ | 84.25 [15]   |
| 8            | Static CD             | Upper                          | $5.00 \times 10^{-4}$ | $1.40 \times 10^{-3}$ | $2.50 \times 10^{-6}$ | 2.5  | 0.7  | 3.6  | 5.30 $\times 10^{-2}$ | 89.3 [19]    |
| 9            | Static PD             | Lower                          | $5.00 \times 10^{-4}$ | $1.40 \times 10^{-3}$ | $2.50 \times 10^{-6}$ | 2.5  | 0.7  | 3.6  | 1.25 | 32.6 [19]    |
|              |                       | Lower                          | $5.00 \times 10^{-4}$ | $1.40 \times 10^{-3}$ | $2.50 \times 10^{-6}$ | 2.5  | 0.7  | 3.6  | 1.25 | 32.6 [19]    |
4. Conclusions

The critical review reported in this paper was aimed at investigating whether the lack of a rationale to the design of the proposed bioreactor designs for the in vitro culture of fragments of cortical ovarian tissue has prevented the full understanding of the effect of O2 transport on the culture outcome, and from fully exploiting the tissue fertility potential. Theoretical models were developed for O2 transport from a gaseous atmosphere to cells in ovarian cortical tissue fragments in some exemplary bioreactors proposed to enhance O2 transport in in vitro culture. The models were used for predicting the dissolved O2 concentration anywhere in the fragment. The model-based analysis evidenced an association between higher O2 availability in tissue and an enhanced follicle viability and progression. It also provided a rationale to help understand the unsuccessful reported culture outcomes. This suggests that bioreactor designs enhancing O2 transport to cells in fragments of cortical ovarian tissue may significantly influence the outcome of the in vitro culture. It also suggests that future strategies to maintain follicle viability long-term and possibly guide follicle activation and growth are better based on bioreactors for the in vitro culture of ovarian tissue designed and operated according to rational engineering criteria to control the perifollicular culture microenvironment. Within this methodological framework, models describing the transport of medium flow and relevant dissolved species in bioreactors and tissue in detail may be valuable developmental tools, provided that their predictions are validated against experimental measurements.

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