A macrophage and theca cell-enriched stromal cell population influences growth and survival of immature murine follicles in vitro

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Abstract

Innovations in in vitro ovarian follicle culture have revolutionized the field of fertility preservation, but the successful culturing of isolated primary and small secondary follicles remains difficult. Herein, we describe a revised 3D culture system that uses a feeder layer of ovarian stromal cells to support early follicle development. This culture system allows significantly improved primary and early secondary follicle growth and survival. The stromal cells, consisting mostly of thecal cells and ovarian macrophages, recapitulate the in vivo conditions of these small follicles and increase the production of androgens and cytokines missing from stromal cell-free culture conditions. These results demonstrate that small follicles have a stage-specific reliance on the ovarian environment, and that growth and survival can be improved in vitro through a milieu created by pre-pubertal ovarian stromal cell co-culture.

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Introduction

The ability to culture ovarian follicles in vitro is an important step towards preserving fertility for women who would like to delay childbearing or for cancer patients, whose gametes can be damaged by cancer therapeutics (Jeruss & Woodruff 2009). In vitro follicle culture also provides an ideal environment to investigate the effects of extra-follicular hormones, gonadotropins, paracrine factors, intraovarian molecules, and non-follicle cells on the maturation of follicles and oocytes.

Current in vitro follicle culture systems vary widely; they can involve plating on a flat surface, encapsulation in a 3D matrix, or alternative culture conditions that prevent the follicle from adhering to a surface, such as culture with constant orbital rotation (reviewed in Picton et al. (2008) and Smitz et al. (2010)). Our group has pioneered a 3D culture environment that surrounds the follicles with alginate, a biologically inert molecule derived from seaweed that gels in the presence of calcium and can be degraded very quickly by alginate lyase (Pangas et al. 2003, Rinaudo 2008). This culture system has provided the context from which live births in the mouse have been achieved, starting from pre-antral follicles isolated and grown in vitro (Xu et al. 2006a).

More recently, we have successfully cultured follicles from both the non-human primate and human, from the secondary/pre-antral stage to the antral stage for both species (Xu et al. 2009a, 2009b). The alginate hydrogel culture system still has limitations, however, as primordial, primary, or small secondary follicles (≤120 μm in starting size) do not grow. These small stages have been the black box of follicle culture for the history of the process; the only successful technique for culture of these follicles involves the two-step processes of first culturing small follicles in either ovarian tissue chunks or liquid droplets before transferring the surviving, larger follicles to more long-term culture conditions (Liu et al. 2001, O’Brien et al. 2003, Adriaens et al. 2004, Lenie et al. 2004). Culturing the small follicle through antral follicle development in a single step, where conditions are not changed through the duration of culture, remains difficult. This apparent stage-specific reliance on the surrounding ovarian tissue, and inability to be cultured in vitro, is a clear indication that the ovarian environment may be required for culture of the smallest follicles in the 3D alginate system.

The influence and importance of the ovarian environment was elucidated in many new ways through the use
of the alginate in vitro follicle culture system, which demonstrated that both physical mechanics and molecular support of the ovary can have an effect on follicle development. It has been shown that, while survival is not impacted, follicle growth and antrum formation are affected by the physical rigidity of the in vitro environment (Xu et al. 2006a). Additionally, follicles cultured in a more rigid environment have smaller oocytes, decreased steroid hormone production, a lower maturation rate, lower fertilization rate, and lower blastocyst formation rate than follicles cultured in a less rigid matrix (Xu et al. 2006a, West et al. 2007, West-Farrell et al. 2009). These data led to the hypothesis that the permissive/non-permissive conditions of culture mimic the natural environment within the ovary (West et al. 2007) to contribute to the function of the individual follicles. Our group has also found that molecular components of the ovarian stroma, particularly extracellular matrix (ECM), can affect follicle growth. While alginate alone cannot support small secondary follicle growth, alginate that has been modified to include collagen matrix or RGD peptides (that serve as a scaffold for integrin binding) can support some growth of the follicle, even in absence of FSH (Kreeger et al. 2006). This effect could be due to an impact on cell survival or proliferation, steroidogenesis, cell signaling, or cell differentiation, all of which may be affected by ECM (Woodruff & Shea 2007). Our culture system is particularly useful in the growth and maturation of larger follicles, but the current alginate-only composition of the 3D matrix cannot support the growth of smaller follicles. It seems possible that a culture system that more closely resembles the native environment of the follicle could be a promising way to initiate and sustain the early growth of primordial, primary, and small secondary follicles.

Because there is clearly a stage-specific reliance on the physical mechanics and molecular support of the ovarian environment, it may be hypothesized that cells of the ovarian stroma themselves are instrumental in follicle development, particularly early follicle development that cannot be recapitulated by our current in vitro conditions. The ovarian stroma is a diverse mix of cell types and adhesion molecules that includes theca–interstitial cells, immune cells, endothelial cells of the blood vessels, smooth muscle cells, and several types of extracellular matrix proteins (Kent & Ryle 1975, Paranko & Pelliniemi 1992, Brannstrom et al. 1993, Best et al. 1996, Magoffin 2002, Berkholtz et al. 2006). Both theca–interstitial cells and immune cells are present in large numbers in the stroma, and both are known to be highly secretory cells (Nathan 1987, Brannstrom et al. 1993, Best et al. 1996, Magoffin 2002); we hypothesize that these cells could be key modulators of folliculogenesis. Of the ovarian immune cells, macrophages are present in the greatest numbers in the rat and human (Brannstrom et al. 1993, Best et al. 1996).

In this study, we investigated what effect a co-culture of ovarian stroma cells may have on in vitro growth of immature follicles. We find a significant increase in growth and survival of primary and early secondary follicles in the presence of an ovarian stromal population that consists mainly of thecal–interstitial cells and macrophages. We identify several factors produced by these stromal cells that may influence this growth and survival, and demonstrate that the co-culture recapitulates the in vivo ovarian environment in a way that is missing from previous culture systems. The work further accentuates the importance of the ovarian environment for culture of small follicles and provides the basis for improved culture outcome for these follicles.

Results

Co-culture of pre-pubertal follicles with stromal cells

Isolated small growing follicles (90–120 μm) from day 16 animals were co-cultured with a feeder layer of ovarian stromal cells isolated from days 22 to 26 animals.

Culture of early secondary follicles with stromal cells

In the 3D culture system used, small secondary follicles (starting size of ~120 μm) from the CD1 mouse strain do not grow in the absence of stromal cells or FSH (Fig. 1A). In fact, after 6 days in culture in the absence of FSH, the follicles cultured without stromal cells begin to die at a high rate, and only about 30% survive to day 10 of culture; even those that survive do not grow (Fig. 1A and B). However, when these follicles are cultured in the presence of stromal cells, they increase in size from 120 μm in diameter to ~190 μm in diameter by day 10 and 75% of follicles survive the entire culture period (Fig. 1A and B). This phenomenon of superior growth and survival in the presence of stromal cells is further exaggerated when FSH is added to the culture media. While control follicles grown with 10 mIU/ml FSH still do not grow significantly by day 10 of culture and have a low survival rate (~40%), follicles grown with stromal cells and 10 mIU/ml FSH reach over 270 μm in diameter by day 10 and have a 98% survival rate (Fig. 1C and D).

Culture of late primary and primary follicles with stromal cells

Encouraged by the superior growth and survival of early secondary follicles, the experiments above were repeated for follicles with an average starting size of both 100 μm, which characterizes the late primary stage (Lenie et al. 2004), and 90 μm. Because FSH was found to be necessary for optimum growth, 10 mIU/ml FSH was used for the duration of the culture in both
Figure 1 Co-culture experiments with stromal cells and follicles from pre-pubertal animals. Follicles of starting size of 120 \( \mu \)m (small secondaries) were cultured either with no FSH in the media (A and B) or with 10 mIU/ml FSH (C and D). Follicles with a starting size of 100 \( \mu \)m (E and F) and 90 \( \mu \)m (G and H) were cultured with 10 mIU/ml FSH for the entire duration of culture. Follicle growth (A, C, E, and G) and survival (B, D, F, and H) were measured. Black lines with triangle icons represent follicles cultured alone, while gray lines with square icons show follicles cultured with stromal cells. N=60 follicles for each group, 20 follicles in each group for triplicate experiments. * \( p \leq 0.05 \), ** \( p \leq 0.01 \), *** \( p \leq 0.001 \).

Physiological mimicry of FSH levels in late primary/stromal cell co-culture

In an attempt to recapitulate in vivo conditions seen by the follicles, late primary follicle culture conditions were created, wherein no FSH was added to media for the first 6 days of culture and 10 mIU/ml FSH was added for all media changes from day 6 until the end of culture. The addition of 10 mIU/ml FSH only to replaced media allowed a gradual increase in FSH level, as might be experienced by small growing follicles in the ovary (these follicles saw a concentration of 8.75 mIU/ml FSH by day 12 of culture). As with all follicles described thus far, late primary follicles cultured in this condition in the absence of stromal cells did not grow (Fig. 2A). Additionally, only about 40% of these follicles survived until day 12 of culture (Fig. 2B). In contrast, late primary follicles cultured with stromal cells grew to an average size of nearly 175 \( \mu \)m by day 12, a 175% increase in size, and over 70% of follicles survived throughout culture (Fig. 2A and B). A second set of culture conditions also recapitulated a low FSH/high FSH environment by having 5 mIU/ml FSH added to media for the first 6 days of culture and then 10 mIU/ml FSH added for all media changes from day 6 until the end of culture (FSH concentration in the media was 9.375 mIU/ml by day 12 of culture). The late primary follicles cultured in these conditions had a significant increase in growth and survival compared with follicles cultured without stromal cells. Follicle co-cultured with stromal cells grew to an average size of \( \sim 270 \mu \)m and had an incredible 93% survival rate, as compared to controls, which never grew and had only a 15% survival rate (Fig. 2C and D). Lower survival of control follicles in cultures with FSH added early, compared with FSH absent early, could be due to increased demands for, or attempts at, growth that are unsupported by the stromal cell influence (i.e. response to cues for growth rather than simply survival or maintenance). Examples of control or stromal cells co-cultured follicles grown from a starting size of 100 \( \mu \)m in low FSH/high FSH conditions are shown in Fig. 2E and F respectively. By day 12 of culture, the control follicle has clearly begun to die, as the surrounding granulosa cells have lost contact with the oocyte in one hemisphere, and the oocyte has begun to be extruded from the follicle (Fig. 2E). In contrast, the follicle co-cultured with stromal cells has a visible antrum (asterisks) and the oocyte is asymmetrically located with surrounding somatic cells (arrow head, Fig. 2F).

Characterization of stromal cells

At the beginning of culture, stromal cells appeared morphologically heterogeneous. Nearly all cells at days 0, 2, and 4 of culture were flat and spread out on the culture dish. Many of these cells had apparent lipid
droplets surrounding the nucleus that resembled thecal cells in morphology (Fig. 3A), though some cells lacked the nucleus-adjacent lipid droplets and appeared similar to fibroblasts. Indeed, alkaline phosphatase staining, used as a marker for theca cells, revealed that over 80% of cells were theca-like at day 0 of culture (Fig. 3D). The heterogeneity of the cellular morphology increased over the culture period, as rounded cells with long cytoplasmic processes were seen frequently by day 6 of culture (Fig. 3B). Because these cells resembled peritoneal macrophages in culture (Fig. 3C), we examined the expression of CD11b using ICC and adherent cytometer counts. Counts revealed that ~18% of cells are macrophages at day 4, ~37% are macrophages by day 8, and the majority of cells (83%) are macrophages by day 12 (Fig. 3E). At the same time, the number of alkaline phosphatase-positive cells decreases significantly over the culture period, so that by days 8 and 12 of culture, the stromal cell feeder layer consists of <15% AP-positive theca cells (data not shown). Therefore, the stromal cell population transitions from being mostly constituted by theca cells at day 0 of culture to being mostly ovarian macrophages by day 12 of culture.

**Macrophone number and localization during the pre-pubertal period**

Because the stromal cell population contained a large number of macrophages by the end of culture, the *in vivo* localization of ovarian macrophages in the pre-pubertal ovary was investigated to determine if culture conditions were physiologically relevant. No quantitative analysis of macrophage localization has ever been performed in the pre-pubertal ovary. To accomplish this, immunohistochemistry (IHC) for the presence of macrophages was performed on ovaries from day postnatal 6, 10, 19, and 26 mice. Ovaries were sectioned and stained for f4/80, a specific antibody used to identify macrophages and their immediate precursors (monocytes), which gives clear staining in paraffin-embedded sections (Austyn & Gordon 1981). Concomitant follicle counts were performed with the macrophage cell counts. We find that macrophages are associated with a heterogeneous population of follicles at every time point during the pre-pubertal period.
examined. They are found with only a portion of each follicle class at every stage, with only atretic follicles always being associated with macrophages (Fig. 4 and Supplementary Figure 1, see section on supplementary data given at the end of this article). At day 6, there is a significant association of macrophages with primordial and primary follicles (Supplementary Figure 1A). By day 10, there is no longer a significant association of macrophages with primordial/primary follicles (Supplementary Figure 1B). Instead, days 10 and 19 counts show a strong association with healthy secondary and antral follicles (Supplementary Figure 1B and C). Primordial and primary follicles at day 26 show an interesting re-association with macrophages (Fig. 4A and B); day 26 also, like day 19, shows a strong association with atretic follicles (Supplementary Figure 1D). In summary, macrophages seem to show a strong association with primordial/primary follicles on day 6 (Fig. 4A and B), which changes to a more significant association with secondary/antral follicles on days 10 and 19 (Fig. 4C and D), and are-association with primordial/primary follicles on day 26 (Fig. 4A and B). Macrophages are also always seen with atretic follicles when such follicles are present (Fig. 4E). Additionally, the number of macrophages associated per follicle is drastically increased depending on follicle class and day of examination. For example, each secondary and antral follicle associated with macrophages on day 19 is associated with an average of 204 macrophages, while each primordial or primary follicle is associated with an average of only six macrophages (Supplementary Table 1B and C, see section on supplementary data given at the end of this article). Representative histological images depicting macrophage association with primordial/primary follicles, healthy secondary/antral follicles, and atretic follicles are shown in Fig. 4Fi–Fiii. Macrophage localization profiles by day, macrophage association as a percentage of total, numbers in the stroma, and total numbers of ovarian macrophages by day are all represented in Supplementary Figures 1 and 2 (see section on supplementary data given at the end of this article) respectively.

**Secretion of cytokines and hormones in stromal cell-follicle co-culture**

The co-culture of stromal cells with follicles resulted in increased follicle growth and survival for primary, late primary, and small secondary follicles, in the absence or presence of FSH. Because the culture system used encapsulates follicles within a surrounding 3D alginate bead, no follicle cells are ever in direct contact with the adherent stromal cells cultured on the bottom of the well. Any effects exerted by the stromal cells must therefore be due to a secreted factor, either produced by, or in response to, the stromal cells. Because theca cells and macrophages are the major constituents of the stromal cell population, the culture media were analyzed for presence of cytokines and androgens. The media from day 2 of culture (either with or without FSH) was tested for the presence of 40 different secreted cytokines. In media from cultures with no FSH, many cytokines were upregulated when follicles were co-cultured with stromal cells, including IL6, KC, M-CSF, JE, MIP-1α, and MIP-2, though only M-CSF, MIP-1α, and TIMP1 reached statistical significance (Fig. 5A). Interestingly, several cytokines such as IL1α and IL1β were significantly downregulated in stromal cell/follicle co-cultures versus follicles cultured by themselves. The pattern of regulation in the 5 mL/ml FSH culture media was similar to that of the culture with no FSH in that IL6, KC, M-CSF, JE, MIP-1α, and MIP-2 were still upregulated (though only IP-10, JE, TNF, and TREM1 reached statistical significance), and IL1α and IL1β were still significantly downregulated when follicles were co-cultured with stromal cells (Fig. 5B). That said, the degree of upregulation was significantly higher in the culture with FSH than that without.
Effect of androstenedione and cytokines on follicle development

Several of the cytokines, as well as androgen, that were found to be differentially produced in stromal cell/follicle co-culture were added to follicle culture in the absence of any co-culturing. Follicles of 100 μm starting size were cultured with 10 mIU/ml FSH and either 1) 0.1, 1, 10, 50, or 100 ng/ml M-CSF, 2) combinations of 50/0.05, 300/0.1, 500/1 ng/ml of IP-10/IL6, 3) combinations of 10, 50, or 100 ng/ml each of MIP-1α, JE, KC, and M-CSF, or 4) increasing levels of androstenedione (1 ng/ml on day 0, 4 ng/ml on days 2, and 10 ng/ml on day 4-end of culture). None of these isolated or combined cytokines or androgens were able to recapitulate the improved growth and survival of small follicles, as seen with stromal cell co-culture, suggesting that the milieu created by these cells is more complex than these few chosen components.

Discussion

We have found that both small secondary and late primary follicles isolated from the pre-pubertal ovary have increased follicle growth, granulosa cell proliferation, and follicle survival when cultured in the presence of ovarian stromal cells. This is the first study, to our knowledge, to examine direct co-culture of pre-pubertal stromal cells and isolated follicles in vitro, as is made possible by the 3D alginate culture system. We have demonstrated that small follicles have a stage-specific reliance on stromal cell co-culture, that our co-culture system recapitulates in vivo conditions through the presence of theca cells and the physiologically relevant association of macrophages, and that the stromal cells present produce androgens and cytokines that may be important for the growth and survival effects observed.

Macrophages are associated with growing follicles in the pre-pubertal ovary

We find that the stromal cell population is predominantly made up of macrophages by the end of culture. In order to determine whether the association of these immune cells with growing follicles is physiologically relevant, we examined the localization of macrophages with each follicle class across the prepubertal period.

Our study clearly demonstrates that macrophages are associated with healthy growing follicles in the pre-pubertal ovary, in addition to the association with dying follicles that may be expected. The presence of macrophages in the pre-pubertal ovary is not surprising, as macrophages have been observed in the adult ovary of the mouse, pig, rat, and human (Bulmer 1964, Gillim et al. 1969, Paavola 1979, Hume et al. 1984).
In our examination of macrophages in the pre-pubertal ovary, we propose a model by which to explain the macrophage localization data observed. There is a significant association of macrophages with the smallest class of follicle when this cohort begins to grow around day 6 (Fig. 6, Box A). As these follicles continue to grow, the macrophages remain associated with them, therefore showing a higher association with larger follicles on days 10 and 19 (Fig. 6, Box B). At the same time, macrophages are attracted to atretic follicles (Fig. 6, Box B). As the first pre-ovulatory class of follicles prepares to ovulate (around day 28), the ovary must begin recruiting the next cohort of pre-ovulatory follicles. A reassociation of macrophages with small follicles is therefore seen around day 26 (Fig. 6, Box C). It is expected that the pattern of macrophage association seen earlier will be cyclical so that by day 30, macrophages would again be more significantly associated with secondary follicles (Fig. 6, Box D).

Though no study has performed quantitative analysis of macrophage number concomitant with follicle counts in the pre-pubertal ovary, our results, and our model proposed, are supported by macrophage counts in the adult ovary and qualitative analysis in the pre-pubertal ovary. In studies of the adult ovary in which the association of macrophages with certain classes of follicle has been followed, there is a correlation between macrophage association and follicle maturity in both the mouse and the human (Petrovska et al. 1996, Katabuchi et al. 1997). These data confirm the physiological relevancy of our culture system, which has an increasing number of macrophages as the follicle grows. In contrast to our data in the pre-pubertal ovary, the studies in the adult ovary of both humans and mice show no macrophages associated with primordial and primary follicles (Petrovska et al. 1996, Katabuchi et al. 1997, Takaya et al. 1997, Gaytan et al. 1998, Li et al. 1998).

In light of our examination of macrophages in the pre-pubertal mouse ovary did no counts of primordial or primary follicles, but do corroborate our data in finding an association of macrophages with healthy growing follicles, in addition to atretic ones (Gaytan et al. 1998). Lastly, though the study is very qualitative and performs no actual counts of macrophages or follicles, our data are nicely corroborated by a second study of pre-pubertal ovarian macrophages, which states that macrophages are seen among primary follicles on day 7, in the theca of growing follicles on day 14, and heavily populating the interstitium by day 21 (Li et al. 1998).

Our demonstration of macrophage association with healthy, growing follicles indicates that phagocytosis of dead cells may not be the only, or even the main role of macrophages in the ovary. This idea is supported by data that shows that ovarian macrophages are more weakly phagocytic than peritoneal or testicular macrophages (Itoh et al. 1999), and that ovarian macrophages surrounding developing non-atretic follicles in the adult ovary do not contain vacuoles or granules present in the cytoplasm that would indicate phagocytic activity (Katabuchi et al. 1997). Because macrophages secrete over 100 cytokines, growth factors, ECM components, and bioactive lipids and peptide products that can act locally (Nathan 1987), our method of counting macrophages that are not only within a follicle, but also closely associated with it (within one cell layer of the outer layer of the follicle), provides a much clearer understanding of the contribution of both secretory and phagocytic (two activities that are not always mutually inclusive (Butler et al. 1982)) macrophages to follicle development.

**Influence of theca cells on early follicle development**

In contrast to the ovarian macrophages that predominate in the stromal cell feeder layer later in culture, theca cells predominate in the beginning of culture. In culture, follicles that have a preserved theca cell layer grow much better than follicles with only a few theca cells when only LH is added to the media, demonstrating the importance of theca cells in in vitro cultures (Cortvrindt et al. 1998). It is possible that the stromal cell co-culture system developed provides theca cells that are normally associated with the follicle in vivo, but that become lost due to the follicle isolation procedure. Surprisingly,
though theca cell number as percentage of total cells decreases over the culture period, androgen production continues to increase throughout culture. This apparent discrepancy is likely due to the proliferation of cells in the feeder layer over time that seems to favor macrophage proliferation, thereby giving a fairly stable number of theca cells over the culture duration. The theca cells that are present could also gain an increased steroidogenic ability over time; current studies are examining the theca cells in the feeder layer in more detail to investigate their full steroidogenic activity and interactions between the theca cells and other cells within the feeder layer.

**Production of cytokines and hormones in the stromal cell co-culture and their possible role in small follicle growth**

We find that several cytokines and growth factors are differentially regulated by the addition of stromal cells to follicle culture; these cytokines and growth factors are possible products of the macrophages present in the stromal cell population or products of the follicle produced in response to the presence of the macrophage-enriched stromal cells in culture. Indeed, previous observations have shown direct effects of macrophages and their secreted cytokines on follicular cells and the process of ovulation. *In vitro*, co-culturing macrophages with granulosa cells causes an increase in granulosa cell proliferation and an increase in progesterone production (Kirsch *et al.* 1983, Polan *et al.* 1989, Fukumatsu *et al.* 1992); this steroidogenic effect has also been seen in ovarian luteal cells co-cultured with macrophages (Halme *et al.* 1985). Peritoneal macrophages and ovarian macrophages both increase the progesterone production of granulosa and granulosa-luteal cells when they are added to culture (Kirsch *et al.* 1981, 1983, Halme *et al.* 1985). In addition, it has been shown that cytokine production by granulosa cells is increased by co-culture with macrophages and that granulosa cell-conditioned media causes migration of immune cells towards the source (Polec *et al.* 2009). Reproductive effects can be seen in models of macrophage depletion, further demonstrating a role of macrophages in ovarian processes. A model of specific ovarian macrophage depletion demonstrates a negative effect on the number of ovulations (Van der Hoek *et al.* 2000), while systemic loss of macrophages and the expression of CSF-1 lead to several reproductive effects, including fewer mature follicles and fewer ovulations (Araki *et al.* 1996, Cohen *et al.* 1997). Finally, many of the cytokines we found to be highly regulated by stromal cell co-culture and/or FSH, namely M-CSF, IL6, and JE/MCP-1, have been previously shown to be regulated by FSH and highly purified urinary menotropin in isolated cultured mouse follicles (Foster *et al.* 2010).

In addition to the large array of cytokines and growth factors measured, we also measured high levels of androgens produced by the stromal cells, likely through the specific actions of the theca-like cells in the population. Androgens can be aromatized to estrogens by the mature granulosa cells (Short 1962, Liu & Hsueh 1986); androgens have been shown to have a positive effect on follicle growth in mice, cows, and non-human primates (Murray *et al.* 1998, Vendola *et al.* 1998, Yang & Fortune 2006). In bovines, cortical strips cultured *in vitro* contained a higher number of secondary follicles when cultured in the presence of testosterone, while primordial and primary follicle numbers were not significantly affected (Yang & Fortune 2006). The mouse model demonstrates similar results: an androgen receptor antagonist or anti-androgen serum slows the growth of isolated *in vitro* cultured follicles, while culturing with dihydrotestosterone (DHT) increases the growth of large follicles (Murray *et al.* 1998). In the non-human primate, *in vivo* treatment with testosterone increased the number of healthy primary, secondary and small antral follicles. Granulosa and thecal cell proliferation were increased in these groups over controls, and follicle atresia and granulosa cell apoptosis were decreased (Vendola *et al.* 1998). Ovarian apoptosis was also suppressed by androgen in cultures of human ovarian tissue *in vitro* (Olta *et al.* 2004). This anti-atretogenic effect of androgens is in conflict with data from hypophysectomized rats, which demonstrates that DHT treatment *in vivo* caused a large amount of follicle atresia in primary, secondary, and tertiary follicles (Bagnell *et al.* 1982, Billig *et al.* 1993). This discrepancy could be due to the particular model and the lack of endogenous gonadotropin control.

Despite these large differences in cytokine and androgen production, neither several cytokines (M-CSF, IP-10, IL6, MIP-1α, KC and JE) nor androgen (androstenedione) could fully recapitulate the improved growth and survival of primary and early secondary follicles seen with ovarian stromal cell co-culture. This suggests that the co-culture system developed provides a complex milieu of hormones and cytokines, as well as possible inhibitors and growth factors, which is ideally suited for optimal growth of small follicles. It is also possible that the cytokines and other factors produced in the stromal cell/follicle co-culture are the result of the interaction between the feeder layer and the follicle itself, and that a feedback loop exists where both must be present for the optimal degree of factor regulation required for increased follicle growth and survival.

In conclusion, we have developed a stromal cell co-culture system for follicles that recapitulates the *in vivo* environment of the ovary by providing theca and macrophage cell influence. This stromal cell support is required in a stage-specific manner for the growth and survival of small follicles. Further work in characterizing...
the molecular and biochemical support provided by each constituent of the stromal cell population will optimize the system and even further expand our ability to successfully mature small follicles in vitro.

Materials and Methods

Animals

CD1 mice were maintained in accordance with the policies of the Northwestern University’s Animal Care and Use Committee. Mice were housed and bred in a controlled barrier facility within Northwestern University’s Center for Comparative Medicine (Chicago, IL, USA) and were provided with food and water ad libitum. Temperature, humidity, and photoperiod (14 h light:10 h darkness) were kept constant. Animals were fed Teklad Global (Madison, WI, USA) irradiated 2919 or 2916 chow, which does not contain soybean or alfalfa meal and contains minimal phytoestrogens.

Tissue processing and histological staining

Mouse ovaries (from days 6, 10, 19, and 26 animals) were placed in 4% formaldehyde (Sigma) fixative for 8–24 h, depending on size. The tissue was then dehydrated and paraffin embedded. Sections (5 μm) were cut with a microtome and mounted on Superfrost-Plus slides (Vector Laboratories, Inc., Burlingame, CA, USA) or Surgipath Snowcoat Extra slides (Surgipath, Richmond, IL, USA). Hematoxylin and eosin staining were performed using a Leica Autostainer XL (Leica Microsystems, Wetzlar, Germany). All tissue processing was performed by the Northwestern University Center for Reproductive Sciences Histology Core.

Antibodies

The antigen f4/80 was detected by IHC using the rat monoclonal anti-f4/80 (Clone C1:A3-1, Cat. #ab6640, Abcam, Inc., Cambridge, MA, USA) at a primary antibody concentration of 1:50. The antigen CD11b was detected by immunocytochemistry (ICC), using the FITC-conjugated rat monoclonal anti-mouse CD11b (recognizes the 170 kDa z[M] chain of CD11b, Cat. #553310, BD Biosciences, San Jose, CA, USA) at a concentration of 0.5 μg/ml.

Immunohistochemistry

IHC for f4/80 was performed as previously described (Bristol-Gould et al. 2006a), except that no antigen retrieval was performed. Adult mouse spleen tissue was used as a positive control, while protocol lacking primary antibody served as negative control. Staining of bursal macrophages was also determined to be an internal positive control at all time points.

ICC and cell counting

ICC for CD11b was performed as follows: live adhered ovarian stroma cells at day 4, 8, or 12 of cell culture were washed three times with 100 μl staining buffer (1 × PBS with 2% FCS and 0.1% sodium azide), then incubated in the dark for 30 min with 0.5 μg/ml FITC-z-CD11b and 5 μg/ml Hoechst 33422 prepared in staining buffer. Following incubation, cells were washed three times with 100 μl staining buffer and then imaged within 1 h. Imaging and cell counts were performed using the Cyntellect Celigo Adherent Flow Cytometer with two-channel detection (Hoechst- Excitation: 377, Emission: 447 and FITC- Excitation: 483, Emission: 536) (Cyntellect, Inc., San Diego, CA, USA). Analysis was performed using a modified live/dead protocol with intensity gating of 2 and size gating of 10 or 15 for Hoechst and FITC channels respectively.

Alkaline phosphatase staining

Alkaline phosphatase (AP) staining was performed as previously described (Cortvrindt et al. 1998). Briefly, adherent cultured stromal cells were washed three times with PBS, fixed for 5 min with 4% paraformaldehyde, and washed three more times with PBS. Cells were then incubated in the dark in histochemical stain solution (0.25 mg/ml naphthol AS/BI phosphate and 0.75 mg/ml fast blue salt prepared in 0.2 M Tris–HCl, pH 9.3) for 45 min. Following incubation, cells were washed three times with double-deionized water (ddH2O) and counterstained with 0.5% nuclear Fast Red for 10 min. Following three washes with ddH2O, cells were imaged. Cell counts were performed in three random parts of each well (n = 5 wells for each day examined) and represented as percent AP positive out of total cells. If individual cells could not be examined due to a cell mass, a calculated estimate was made on percent positive for AP staining.

Macrophage and follicle counting

Macrophages were counted in every 10th section as being associated with primordial/primary follicles (follicles with only a single granulosa cell surrounding them), healthy secondary/antral follicles (follicles with multiple granulosa cell layers with complete absence of pyknotic nuclei), atretic follicles (follicles characterized morphologically by the presence of at least one pyknotic nuclei), or stroma tissue. A macrophage was considered to be associated with a follicle if it was within one cell layer of the outermost portion of the follicle (the granulosa layer in monolaminar follicles and the outermost theca cell layer in larger follicles), and was considered to be in the stroma if it was not in the immediate vicinity of any follicles. A macrophage that was close to two follicles of different classes was counted in the numbers of both classes, but only counted once in the total. Because macrophages have long pseudopodia-like extensions that could traverse multiple sections, double counting of cells was avoided by counting only stained cells, where the nucleus, as stained by hematoxylin counterstaining, could be detected. Macrophage nuclei were small enough not to be double counted in multiple sections, so the average number of cells per section was multiplied by the total number of sections obtained to find the number of macrophages associated with each type of follicle class per ovary. Macrophage numbers are presented as per ovary counts, unless otherwise noted. Representative images of
macrophage association with each class of follicle counted are shown in Fig. 4F and Supplementary Figure 3, see section on supplementary data given at the end of this article.

Follicle counts were also performed in the ovaries of prepubertal animals, as previously described (Bristol-Gould et al. 2006b, Tingen et al. 2009).

Ovarian stromal cell isolation and plating

Ovaries isolated from day 22 to day 26 animals with the bursa removed were placed in media containing L15, 0.5% Pen-Strep and 1 mg/ml BSA (‘wash media’). Two insulin gauge needles were used to poke each ovary 100 times to release granulosa cells. The remaining ‘husk’ was cut into eight pieces, placed into dissociation media (wash media with 0.1% collagenase and 0.01% DNase I), and pipetted with a 1 ml tip gently. Tissue was then incubated for 1 h at 37 °C in a 95% air/5% CO2 chamber, pipetting every 15 min with successively smaller-orificed pipette tips to break up the tissue. Tissue was placed back in incubator between pipettings. Solution was allowed to settle and supernatant was transferred to eppendorf tubes. Solution was then pelleted (1485 g for 5 min) and washed once each with 37 °C wash media and 37 °C plating media (RPMI 1640 containing 25 mM HEPES and 2 mM L-glutamine (Cat. #22400-071 Gibco Invitrogen) with 10% heat-inactivated FCS (Cat. # 10082-139 Gibco Invitrogen) and 1% Pen-Strep). Cells were then resuspended in 1–2 ml plating media and plated in 96-well polystyrene plates at an approximate density of 150 000 cells per well. Cells were incubated at 37 °C for 3 h in a 95% air/5% CO2 chamber, and then all non-adhered cells were removed by washing plates three times with 37 °C PBS. Cells were then allowed to ‘rest’ in plating media at 37 °C in a 95% air/5% CO2 chamber overnight. All wells were washed three times with 37 °C PBS the following day before follicles and follicle growth media (as described in ‘follicle isolation, encapsulation, culture, and diameter measurement’) were added in order to remove all of the plating media containing serum.

Follicle isolation, encapsulation, culture, and diameter measurement

Follicle isolation, encapsulation, culture, and measurement of follicle diameter were performed as previously described (Xu et al. 2006a), with following modifications: alginate concentration used was 0.5% instead of 1.5% and ovaries were not incubated in αMEM/FBS/collagenase/DNAseI prior to mechanical isolation. Starting sizes of follicles were also roughly ~90, 100–105, or 120 μm, as described for each experiment in ‘Results’, FSH was added as described in ‘Results’, and cultures were of a 10- or 12-day duration. Follicles were isolated from day 16 animals and all culture experiments had 20 follicles in each follicle group, with each experiment performed in triplicate, for a total of 60 follicles per group. As previously described (Xu et al. 2006b), follicles were considered dead if the oocyte was no longer surrounded by a granulosa cell layer, if the granulosa cells had become dark and fragmented, and the follicle had decreased in size, or if the oocyte had fragmented.

Cytokine and hormone analysis from conditioned media

Media collected after media change (every other day during culture) was placed in 96-well plates and frozen at −80 °C until use. For each cytokine array, five wells worth of media were pooled (~250 μl total) and 240 μl of media were used in the Proteome Profiler Mouse Cytokine Array/Panel A Array Kit (Cat. #ARY006, R&D Systems, Minneapolis, MN, USA), as instructed by the manufacturer, except that the streptavidin-HRP was diluted at 1:200. For each hormone assay, five wells worth of media were pooled (~250 μl total) and 25 μl of media were used for each condition in the Active Androstenedione Enzyme Immunoassay Kit (Cat. #DSL-10-3800, Diagnostic Systems Laboratories, Webster, TX, USA), as instructed by the manufacturer.

Statistical analysis

Data were analyzed by one-way ANOVA between groups, followed by a Tukey test of significance for macrophage localization. For follicle sizes and survival, an unpaired, two-tailed t-test was performed between macrophage and control groups at each day. A one-sample t-test was used to compare fold change of cytokine levels to a theoretical value of 1. Significance was determined for P<0.05. All statistical analyses were performed using GraphPad Prism 4 (GraphPad Software, Inc., San Diego, CA, USA).

Supplementary data

This is linked to the online version of the paper at http://dx.doi.org/10.1530/REP-10-0483.

Declaration of interest

The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.

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