A novel two-step strategy for in vitro culture of early-stage ovarian follicles in the mouse

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Objective: To develop an in vitro strategy to support the growth of early-stage follicles and produce mature oocytes competent for fertilization.

Design: Whole ovaries from 8-day-old mice were cultured for 4 days, and then secondary follicles were isolated and cultured for 12 days in a three-dimensional alginate or fibrin-alginate (FA) hydrogel matrix.

Setting: University-affiliated laboratory.

Animals: Mice.

Intervention(s): None.

Main Outcome Measures: Histologic evaluation of follicle development, steroid hormone production, and rates of oocyte maturation, oocyte fertilization, and embryo formation.

Result(s): Culture of 8-day-old mouse ovaries for 4 days resulted in transition of the follicle population from primordial and primary follicles to secondary follicles, similar to that seen in a 12-day-old ovary. Isolated secondary follicles cultured for 12 days showed larger increases in oocyte diameter and more frequent antrum formation and theca cell differentiation in the FA-hydrogel matrix compared with the alginate matrix. Steroid hormone secretion patterns were consistent with the changes in follicle morphology and cell differentiation observed in the cultured follicles. Compared with oocytes from alginate follicle cultures, a greater number of oocytes retrieved from the FA-based follicle cultures progressed to metaphase I, reached metaphase II, and could be fertilized and cleaved to two-cell embryos. The organ culture plus FA-hydrogel follicle culture strategy produced a very high rate of oocyte progression to metaphase II (88 ± 8.7% [mean ± SEM]) and formation of two-cell embryos (54 ± 4%).

Conclusion(s): A strategy combining whole ovary culture of early-stage follicles and subsequent FA hydrogel in vitro follicle culture produced a high percentage of oocytes competent for fertilization; this might provide new options for fertility preservation in women and prepubertal girls facing fertility-threatening diseases or treatments. (FertilSteril® 2009; ■■■■ – ■ ■ ■. ©2009 by American Society for Reproductive Medicine.)

Key Words: Follicle, alginate, fibrin, 3D, oocyte maturation, fertilization, in vitro culture

Recent advances in ovarian tissue or oocyte cryopreservation and transplantation or in vitro follicle culture and oocyte maturation may provide options for women who are facing fertility-threatening diseases or treatments (1, 2). A practical limitation to the success rate of these methods is the fact that the majority of follicles within the ovary, particularly within the cortex, are arrested at the primordial stage. Current methods limit the ability to dissect primordial follicles from ovarian tissue, leaving a significantly smaller population of secondary follicles available for collection and eventual in vitro oocyte maturation and/or IVF. Although culture of ovarian cortex tissue, with successful transition of primordial/primary follicles to secondary follicles in vitro, has been achieved in some species (mouse, bovine, baboon, and human) (3–6), in vitro follicle culture methods have been successful only with secondary or later stage follicles. Collection and culture of primordial and primary follicles in vitro to produce meiotically competent, fully mature oocytes would significantly expand the number of follicles available for future use, thus increasing the statistical odds of IVF success.

Successful in vitro culture of primordial/primary follicles has been hampered by the sheer complexity of recreating the multiple signals and cell-cell/cell-stromal interactions needed to support early follicle growth and selection into the growing follicle pool. In the past decade, important advances have been made in two-dimensional (2D) culture techniques for in vitro growth of preantral follicles (1, 7, 8). However, follicles cultured in these systems must attach to a flat culture surface on which somatic cells migrate away from the oocyte, thus altering the native three-dimensional (3D) structure of the follicle and disrupting the somatic cell–gamete interactions important for normal oocyte growth (9). Three dimensional (3D) in vitro culture systems that mimic the ovary’s internal architecture appear to be optimal for supporting follicle growth and oocyte maturation. A recently developed alginate follicle culture system provides a 3D scaffold matrix for supporting the growth and maturation of multilayered secondary follicles (10–13). This system has been shown to produce meiotically competent oocytes.
that are able to be fertilized and produce viable offspring (14). Subsequent studies have demonstrated that the concentration of the alginate hydrogel matrix can be modified to support the growth of earlier secondary follicles (15).

In this article, we describe the development of a two-step method of in vitro follicle culture involving ovarian tissue culture followed by secondary follicle culture in a modified alginate matrix, as described previously by our group (12–15). In this system, whole ovaries are cultured for 4 days to support early follicle growth and development. Secondary follicles recovered from the cultured ovaries are then cultured in interpenetrating fibrin-alginate (FA) beads (16) to support further follicle development and oocyte maturation. We hypothesize that the in-organ culture of primordial and primary follicles will produce a greater number of secondary follicles for subsequent in vitro culture, and that the FA gel is superior to alginate alone for producing mature oocytes competent for fertilization. The ability to culture follicles in vitro to produce mature oocytes competent for fertilization represents a major step forward in the development of fertility-sparing options for women and girls facing potentially gonadotoxic diseases or treatments.

MATERIALS AND METHODS

Animals

C57BL/6j x CBA/Ca F1 hybrid mice study were housed and bred in the Central Animal House of Northwestern University. Eight-day-old F1 female mice were used in this study. All animals were housed in a temperature- and light-controlled environment (12L:12D) and were provided with food and water ad libidum. Animals were treated in accordance with the National Institutes of Health Guide for the Care and Use of Laboratory Animals and the established Institutional Animal Care and Use Committee protocol at Northwestern University.

Organ Culture of 8-day-old Mouse Ovaries

As reported previously (17), ovaries were excised from the ovarian bursa and washed twice with culture medium: α-minimal essential medium (αMEM) supplemented with recombinant FSH (10 mIU/mL; A. F. Parlow, National Hormone and Peptide Program, National Institute of Diabetes and Digestive and Kidney Diseases), bovine serum albumin (3 mg/mL), bovine fetuin (1 mg/mL; Sigma-Aldrich, St. Louis, MO), insulin (5 mg/mL), transferrin (5 mg/mL), and selenium (5 mg/mL). Ovaries were transferred into 24-well plates with tissue culture well inserts (nonadhesive culture treated; Millicell-CM, 0.4-μm pore size; Millipore Corp., Billerica, MA). Approximately 400 μL of culture medium was added to the compartment below the membrane insert, such that ovaries on the membrane were covered with a thin film of medium. Up to six ovaries were placed in each well. The ovaries were incubated at 37°C, 5% CO2, for 4 days. Every other day, 150 μL of media was replaced with fresh culture media.

Histologic Analysis and Follicle Classifications

Ovaries from 8- and 12-day-old mice were fixed overnight in a 4% paraformaldehyde solution at 4°C and then dehydrated in an ethanol series and embedded in paraffin wax. Sections (5 μm) were stained with hematoxylin and eosin (H&E). The number of follicles at each developmental stage was counted and averaged in three serial sections from the largest cross-sections through the center of the ovary (17, 18). Only follicles that contained an oocyte nucleus were counted. Follicles were classified as primordial (stage 0), primary (stage 1), and secondary (stage 2) as previously described (19). Follicle counting results were calculated as percentages to account for differences between preculture and postculture ovaries.

Alginate Hydrogel and FA Gel Preparation

Alginate hydrogel was prepared as described in previous reports (9, 14). FA gel was prepared as described in (16). Tissue fibrin sealant kits (Baxter, Deerfield, IL) were used according to the kit instructions. Fibrinogen and thrombin were reconstituted with aprotinin (3000 KIU/mL) solution and calcium chloride (40 mM) separately. Appropriate concentrations of both solutions were attained by dilution in tris-buffered saline solution. The FA gel was prepared by mixing 50 mg/mL fibrinogen solution with 0.5% alginate solution at 1:1 and then adding the same volume of 50 IU/mL thrombin solution to the mixture.

Isolation, Encapsulation and Culture In Vitro of Preantral Follicles

After 4 days of ovary tissue culture, secondary follicles were mechanically isolated using insulin-gauge needles and placed into L15 media (Invitrogen, Carlsbad, CA) with 1% fetal calf serum (FCS), then transferred into αMEM supplemented with 1% FCS and incubated at 37°C, 5% CO2, for 2 hours. Follicles with centrally located oocytes and at least two layers of granulosa cells were encapsulated into alginate beads (0.25% w/v) or FA beads (0.125% alginate, 12.5 mg/mL fibrinogen) using the method reported by Xu et al. (15). Encapsulation in FA beads was performed as described by Shikanov et al. (16). Alginate and FA beads containing follicles were washed twice in culture media. One bead was placed in each well of a 96-well plate, in 100 μL culture media and incubated at 37°C, 5% CO2, for 12 days. Every other day, 50 μL of the media was replaced by fresh culture media, and follicle survival and diameter were assessed as described previously (15). At the end of the culture period, the media was replaced by L15 medium (100 μL) containing alginate lyase (10 units/mL; Sigma-Aldrich), and the beads were incubated for 30 minutes at 37°C. Follicles were then removed from the degraded alginate bead by mechanical isolation (14, 15).

In Vitro Maturation and Fertilization of Oocytes

Cumulus-enclosed oocytes (CEOs) were collected from antral follicles released from alginate or FA beads. The CEOs were placed in αMEM, 10% FCS, 1.5 IU/mL hCG, and 5 ng/mL epidermal growth factor (EGF; Sigma-Aldrich) for 18 hours at 37°C, 5% CO2 (28). Sperm was collected from the cauda epididymis of proven CD-1 male breeder mice using Percoll gradient centrifugation as described previously (14). The sperm was capacitated for 30 minutes in IVF media (KSO; Specialty Media, Phillipsburg, NJ) containing 3 mg/mL bovine serum albumin and 5.36 mM D-glucose. Approximately 5–10 metaphase II (MII)-stage oocytes were placed in a 100-μL droplet of IVF medium containing sperm, placed under mineral oil, and incubated for 7–8 hours at 37°C, 5% CO2. Fertilized oocytes were washed three times in fresh KSO to remove all sperm and then transferred into a 50-μL fresh KSO microdrop under mineral oil overnight. Embryos that cleaved to the two-cell stage were recorded as fertilized (8, 15).

Hormone Assays

E2 and P were measured in conditioned media collected on follicle culture days 2, 6, and 12. Conditioned media from each time point were pooled together, and the average concentration at each time point was determined from three independent experiments. All measurements were performed by electrochemiluminescent assay using an Immulite 2000 Analyzer (Roche, Indianapolis, IN) in the Endocrine Services Lab at the Oregon National Primate Research Center (Oregon Health and Science University, Portland, OR). Interassay variations were 6.1% for E2 and 5.4% for P, and the limits of sensitivity were 5 pg/mL for E2 and 0.03 ng/mL for P.

Statistical Analysis

All experiments were performed at least three times. Values are given as mean ± SEM, and statistical analysis was done using Student’s t test. Differences were considered significant at \( P < 0.05 \).

RESULTS

Follicle Development in Organ Culture

Ovaries from 8-day-old mice contained mostly primordial follicles (84.8 ± 3.2%), with a few primary (8.8 ± 2.5%) and secondary follicles (6.4 ± 5.2%; Fig. 1A, 1C, 1F). After 4 days of organ culture in
vitro (Fig. 1B, 1D), primordial follicles represented a smaller percentage of the total follicle pool (65.7 ± 0.5%), similar to the follicle distribution seen in ovaries from 12-day-old mice (65 ± 10.6%; Fig. 1E, 1F). The proportion of secondary follicles increased significantly during the 4-day culture, from 6.4 ± 5.2% to 24.5 ± 3.3% (P < 0.05; Fig. 1A vs. 1B; Fig. 1C vs 1D). The ratio of activated follicles in the cultured ovaries was similar to that of 12-day-old ovaries (25.8 ± 10%). There were no differences in the proportion of primary follicles in the 8-day-old ovaries before or after culture and in the 12-day old ovaries (Fig. 1F).

**Secondary Follicle Growth in the Alginate and FA Culture Systems**

A total of 430 secondary follicles, with a diameter range of 111–137 μm, were isolated from the cultured ovaries (Fig. 2A), embedded in FA beads or alginate beads, and cultured for 12 days (Fig. 2B). At day 12, the majority of follicles had survived the culture period in either FA beads (74.8 ± 4.6%) or alginate beads (68.6 ± 5.5%; Table 1). Antrum formation and the appearance of a laminar-like theca cell layer were seen more frequently in follicles cultured in the FA system compared with follicles cultured in alginate (antrum, 72.0 ± 3.9% vs. 59.7 ± 5.6% [P < 0.05]; theca layer, 72.3 ± 3.2% vs. 64.7 ± 4.6% [P < 0.05]; Fig. 2C; Table 1). Follicle diameter increased significantly, from 124 ± 2.2 μm at day 0 to 362.4 ± 10.1 μm in alginate and 371.6 ± 8.8 μm in FA (Fig. 2D). Follicle-enclosed oocytes in both groups also increased in size during the 12-day culture (Fig. 2E); however, final oocyte diameter was larger in the FA-cultured follicles compared with alginate-cultured follicles (73 ± 0.6 μm vs. 69.3 ± 0.7 μm; P < 0.05). By comparison, the average oocyte diameter in secondary follicles from 24-day-old mice was 73 ± 0.6 μm.

FIGURE 1

Representative photomicrographs of H&E-stained paraffin sections of whole ovaries before and after culture. (A) Control, uncultured 8-day-old mouse ovary. (B) Eight-day-old mouse ovary after 4 days of organ culture. (C) H&E staining of uncultured 8-day-old mouse ovary, which contains mainly primordial follicles with a few primary and secondary follicles. (D) H&E staining of 8-day-old mouse ovary after 4 days of organ culture. More secondary follicles were observed. (E) H&E staining of uncultured 12-day-old mouse ovary. (F) Follicle distribution in mouse ovaries before and after 4-day organ culture. 0, primordial follicle; 1, primary follicle; 2, secondary follicle; scale bars in A and B = 150 μM; other scale bars = 200 μM; letters indicate a statistically significant difference between groups (P < 0.05).
Development and differentiation of representative secondary follicles cultured in vitro. (A) Secondary follicles with centrally located immature oocytes isolated from cultured ovarian tissues. (B, C) Follicles maintained their 3D structure with proliferation of granulosa cells, antrum formation (white arrowhead), and development of theca cell layers (black arrowhead) after 12 days of culture in 0.25% alginate or FA. (D) Follicle diameter in both culture systems increased significantly during the culture period. (E) Oocyte size increased significantly over the culture period. Statistically significant differences were observed between groups as indicated with different letters (P < 0.05). Scale bar = 100 μM. (F, G) Average values of E₂ (F) and P (G) secretion were measured in conditioned culture media from secondary follicle cultures.

As shown in Fig. 2F and 2G, the secretion patterns of E2 and P were consistent with observed changes in follicle morphology and cell differentiation in the cultured follicles. During the first 6 days of culture, both E2 and P levels rose more slowly than during the last 6 days of culture. There was no significant difference in steroid secretion between follicles cultured in alginate or FA.

**Oocyte Meiosis and Fertilization Competence**

CEOs (n = 96 from alginate-cultured follicles [Fig. 3A], and n = 50 from FA-cultured follicles [Fig. 3E]) were stimulated with hCG and EGF for 18 hours. After treatment, significant cumulus cell expansion was observed in both groups (Fig. 3B, 3F). Most of the oocytes in both groups resumed meiosis, underwent germinal vesicle breakdown (GVBD), and matured to MII with extrusion of a first polar body (Fig. 3C, 3G; Table 1). In the FA-cultured group, 86 ± 0.9% of the oocytes could be fertilized and cleaved to two-cell embryos, whereas in the FA-cultured group, 54 ± 4% of MII oocytes formed two-cell embryos (P < 0.05; Fig. 3D vs. 3H; Table 1).

**DISCUSSION**

In addition to providing key insights into early follicle growth and its effect on later oocyte maturation, these studies bring us one step closer to improving current in vitro follicle culture methods, which may ultimately have a clinical application in the preservation of fertility (15).

Individual early-stage follicle culture in vitro is not yet feasible, because the complete set of factors that drive progression of primordial or primary follicles toward secondary stages has not been identified. However, organ culture maintains the in vivo condition of oocytes that reached MII was higher in the FA-cultured group than in the alginate-cultured group (88 ± 8.7% vs. 61.3 ± 2.4%; P < 0.05; Table 1). In the alginate-cultured group, 33 ± 1.7% of the MII oocytes could be fertilized and cleaved to two-cell embryos (P < 0.05; Fig. 3D vs. 3H; Table 1).

**FIGURE 3**

Meiotic and fertilization competence of oocytes from follicles cultured for 12 days in alginate (A–D) or FA (E–H) were assessed by in vitro maturation (IVM) and in vitro fertilization (IVF). CEOs isolated from antral follicles retrieved from alginate (A) or FA (E) culture systems were induced with hCG for 18 hours in vitro. In both environments, cumulus cells around the oocytes expanded. (C, G) Oocytes resumed meiosis and extruded the first polar body (arrowhead). (D, H) Two-cell embryos were obtained by IVF of MII oocytes. Scale bar = 50 μM.

![Image](image-url)
microenvironment of the follicles, including the surrounding stromal cells and their intercommunication with early-stage follicles, and the connectivity between cellular compartments within the follicle. The growth of primordial and primary follicles in vitro has been accomplished using organ culture (3-6), but the efficiency has been low (20, 21), and it is generally accepted that organ culture alone is not able to support complete growth and development of follicles and oocytes competent for fertilization. For this reason, we chose to develop a two-step culture system that combined early follicle growth within the intact ovary with a hydrogel-based follicle culture system to support the further growth and development of secondary follicles.

Previous reports have described the growth of preantral follicles using a nonspherical, 2D in vitro culture system. Ola et al. (22) achieved a 48% follicle survival rate and 38% antrum formation, and Oktem et al. (23) reported a survival rate of 47% in a standard 2D culture system. Haidari et al. (24) showed a follicle survival rate of 68% and an antrum formation rate of 54%. In this study, we achieved a follicle survival rate of 68% and an antrum formation rate of 59% using an alginate-based, 3D follicle culture system. We achieved higher rates of follicle survival (74%) and antrum formation (72%) using an FA culture system. Gomes et al. (25) showed that a flat, 2D, adhesive environment leads to a distortion of follicle morphology, marked extracellular matrix modifications, and high rates of spontaneous follicle disruption. In contrast, 3D gel environments are able to maintain follicular structure with an in vivo-like basal lamina architecture that minimizes spontaneous disruption. Oktem et al. (23) demonstrated that 3D culture with extracellular matrix provides a better milieu for in vitro growth and survival of immature mouse preantral follicles, compared with conventional 2D culture.

The development of a culture system that supports follicle growth and oocyte maturation beginning at the early follicle stage might make it possible to access a significantly greater number of follicles for in vitro maturation and IVF. O'Brien et al. (3) performed 8-day organ culture of ovaries isolated from newborn mice followed by 2D culture of COCs. In their optimized protocol, a GVBD rate of 62% and an MII rate of 44% were achieved. In this study, we describe a two-step protocol that combines traditional organ culture and a novel hydrogel-based 3D follicle culture technique. Whole ovaries from 8-day-old mice, which contained primarily primordial follicles with a few primary and secondary follicles, were cultured to support early-stage follicle growth and development into the secondary follicle stage. We believe that the cultured ovary acts as an incubator, where important stromal-cell and cell-cell interactions remain intact, and the presence of local paracrine and autocrine factors support primordial and primary follicle growth. With 4-day culture of 8-day-old ovaries, we were able to achieve a similar degree of early-stage follicle development and transition to secondary follicles as in 12-day-old ovaries. In the second step, secondary follicles were isolated from the cultured ovaries and grown in alginate beads for 12 days to support further follicle development, as described previously (12-16). During this time, follicles significantly increased in mean diameter, with formation of an antral cavity and proliferation and differentiation of granulosa cells and theca cells. The mean diameter of oocytes also increased and cumulus cells expanded significantly in response to hCG. The majority of oocytes resumed meiosis and were competent to undergo GVBD and polar body extrusion, and fertilized oocytes developed to two-cell embryos. With this novel protocol, we demonstrated the ability to produce embryos starting from early-stage follicles from 8-day-old mice.

Furthermore, we found that FA hydrogel was superior to alginate in regard to follicle growth and differentiation, thus producing a larger percentage of oocytes competent for fertilization and a greater number of two-cell embryos than alginate alone. Studies have shown that the efficiency of producing fertilizable oocytes in vitro is influenced by many factors (26, 27). Fibrin is naturally derived protein, and commercial fibrin consists of thrombin and fibrinogen that is cryoprecipitated from blood plasma, as well as small amounts of fibronectin, transforming growth factor-I, basic fibroblast growth factor, epidermal growth factor, vascular endothelial growth factor, and other biomolecules (28). Some of these factors play an important role in follicle development (29), and fibrin itself supports a number of cellular processes, including growth, proliferation, and differentiation (28). The FA hydrogel also has unique dynamic mechanical properties, as cell-secreted proteases degrade the fibrin in the surrounding bead and remodel the local environment. Alginate is produced by brown algae and permits diffusion of hormones and other molecules from the surrounding environment (30). Thus, the combination of alginate and fibrin maintains the 3D architecture of follicles and provides an environment that supports follicle growth.

In conclusion, the present study introduces a novel, robust, two-step culture strategy for in vitro growth of early-stage follicles. Organ culture provided follicles with an in situ growth environment, and the hydrogel-based 3D culture scaffold promoted further development to terminally differentiated follicles. The maintenance of the follicle architecture supports the critical cellular interactions between adjacent somatic cells and between somatic and germ cells, which we believe facilitated coordinated growth and differentiation of granulosa and theca cells and the oocyte in culture. This two-step in vitro culture system creates new possibilities for preserving the fertility of women who must undergo life-saving but potentially fertility-threatening treatments. In particular, the ability to successfully culture follicles to produce mature oocytes in vitro starting at the primordial follicle stage provides prepubertal female cancer patients with a new fertility-sparing option. For adult patients, this technique eliminates the need to wait for the menstrual cycle or to expose patients with hormone-sensitive cancers to exogenous hormones for IVF, and thus has the potential for preserving fertility in a much greater number of female cancer patients.

Acknowledgments: The authors thank Tyler Wellington for sectioning all tissue analyzed in this study.

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