



Published in final edited form as:

*Theriogenology*. 2019 April 15; 129: 168–177. doi:10.1016/j.theriogenology.2019.02.018.

## Activin Promotes Growth and Antral Cavity Expansion in the Dog Ovarian Follicle

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### Abstract

Understanding regulators of folliculogenesis remains limited in the domestic dog (*Canis familiaris*), which challenges our ability to develop *in vitro* follicle culture systems for canid genome rescue efforts. Here, we investigated the influence of activin on dog follicle development and survival, oocyte quality, and FSH receptor expression in culture. Preantral (150 – 230  $\mu\text{m}$  diameter), early antral (231 – 330  $\mu\text{m}$ ), and antral (>330 – 550  $\mu\text{m}$ ) stage follicles were encapsulated in a fibrin-alginate hydrogel with 0, 100, or 200 ng/ml rhActivin plus 0, 0.1, 1, or 10  $\mu\text{g/ml}$  FSH for 12 or 21 d of *in vitro* culture. All follicle groups increased in diameter ( $P < 0.05$ ) with activin acting synergistically with FSH to improve ( $P < 0.05$ ) growth and antral cavity expansion (to >630  $\mu\text{m}$ ) in early antral and antral cohorts. This complementary effect was not linked to changes in FSHR mRNA expression ( $P > 0.05$ ). Although not influencing ( $P > 0.05$ ) follicle survival or transzonal projection (TZP) density in shorter term 12 d culture, activin in the presence of 1 ng/ml FSH maintained TZP density from the 12 to 21 d interval. Activin also increased oocyte diameter and improved nuclear integrity compared to un-supplemented controls. These results indicate that activin acts synergistically with FSH to promote growth and antral cavity expansion of the dog follicle *in vitro*, information useful to formulating an effective culture microenvironment for this species.

### Keywords

Activin; domestic dog; FSH; ovarian follicle; trans-zonal projection

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### Conflicts of Interest

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

## 1. Introduction

The International Union for Conservation of Nature has designated five of the 35 extant, wild canid species as endangered or critically endangered [1]. As the closest, domesticated relative, the dog is an excellent model for understanding the intricacies of canid reproductive biology, including developing knowledge and tools to foster species survival and improved conservation management. The unique biology of canids poses challenges to controlling reproductive function, including technologies common in humans as well as some laboratory and livestock species. Firstly, it is known from longitudinal measures of ovarian steroid hormone activity [2–4] and ultrasonography [5] that the ovaries of the dog and other canids generally are quiescent, lacking mature follicular development and yielding sporadic estrus only once or twice annually [6]. An explicit reasoning for this protracted, suppressed ovarian activity is not well understood. Secondly, when ovulation does occur in the dog, oocytes are immature and resume meiosis to achieve metaphase II (MII) only after intraoviductal residence for 48 to 72 hours [7, 8]. Because of this reproductive complexity, the dog offers unique opportunities for understanding regulation of canid follicle and oocyte maturation and viability.

Domestic dog oocytes from antral follicles >2 mm in diameter are more likely to resume meiosis and achieve Metaphase II *in vitro* (as high as 79.5% in our laboratory [9]) than those recovered from <0.05 mm counterparts (16.9% [9]), although in general achievement of MII dog oocytes *in vitro* ranges from 20–45% [10–12]. Thus, our objective is to develop an improved *in vitro* culture system that consistently produces antral stage follicles and, thereby, maturation-competent oocytes. Previously we explored an alginate culture system developed originally for the mouse [13, 14] to maintain the three-dimensional (3D) architecture of isolated dog follicles. Although preantral (compared with early antral) follicles produced the more robust percent growth *in vitro*, we also observed that follicles: 1) rarely grew beyond ~500  $\mu\text{m}$  diameter (compared with 5–7 mm preovulatory follicle diameter [15]); 2) were unable to maintain an antral cavity; and 3) contained oocytes that became increasingly pale over a 20 d culture period, suggesting loss of intracellular lipid (the dog is known for its dark colored, lipid-filled oocytes [16, 17]). These morphological anomalies may have resulted from poor communication between the oocyte and cumulus cells via cumulus cell trans-zonal projections (TZPs), or extensions of the plasma membrane through the zona pellucida which form junctions with the oocyte. This communication is known to direct antral cavity development [18–20], and modulate oocyte intracellular lipids [21, 22]. In sum, there is a need to explore factors which may act to promote TZP function and antrum formation for canine *in vitro* folliculogenesis.

Recent work in *in vitro* follicle culture has demonstrated a role of the protein hormone activin in promoting antrum cavitation in rat, cow, and human follicles [23–25]. Activin also has been suggested to improve cumulus-oocyte communications in cultured bovine follicles by maintaining TZPs [26] to, in turn, promote antrum formation and oocyte viability. Activin is a glycoprotein member of the transforming growth factor  $\beta$  (TGF $\beta$ ) superfamily, produced by granulosa cells of the ovarian follicle [27–32], and is believed to act both as an endocrine factor signaling pituitary FSH release, and as a paracrine/autocrine factor in the ovary [33]. Supplementing culture medium with activin has been shown to promote murine

follicle growth [34] and survival [35] as well as oocyte maturation in humans and zebrafish [30, 36]. Previous investigations into activin mechanisms have focused mainly on its role in increasing follicle sensitivity to FSH by increasing transcription of FSH receptor mRNA [37, 38]. The purpose of the present study was to understand the role of activin as a regulator of antral cavity development and cumulus cell-oocyte communication in the domestic dog.

Here, we took advantage of the dynamic Fibrin-Alginate Interpenetrating Network (FA-IPN) [39], a 3D hydrogel system which confers structural support through its alginate constituent and flexibility via its fibrin component that can be degraded by the follicle, thereby allowing further growth, including antrum expansion. FA-IPN has increased rates of successful antrum formation and oocyte maturation in murine follicle culture [40]. Our specific objectives were to determine the influence of activin, with or without FSH, on follicle growth, survival, antral cavity expansion, FSH receptor expression, TZP density, and oocyte growth and viability. We hypothesized that activin supplementation (1) acts synergistically with FSH via up-regulation of FSH receptors to promote follicle growth, (2) supports maintenance of TZP density, and (3) promotes antrum development and oocyte viability in *in vitro* culture.

## 2. Materials and Methods

All chemicals were purchased from Sigma-Aldrich (St. Louis, MO) and media from Irvine Scientific (Irvine, CA), unless otherwise stated.

### 2.1 Follicle Isolation and Encapsulation

Ovaries from 42 dogs (age 5 mo - 9 yr, breed information in Supplemental Table 1) were obtained during routine ovariohysterectomies conducted at local veterinary clinics. Each freshly-excised reproductive tract (from cervix to bilateral ovaries) were stored on ice in Leibovitz's L15 medium containing 84.2  $\mu$ M penicillin G sodium, 41.2  $\mu$ M streptomycin sulfate, and 50 mM ascorbic acid for transport to the laboratory. Within 3 hours of excision, each ovary was visually examined to estimate reproductive stage of the donor. Virtually all females were in anestrus or diestrus based on ovarian morphology, specifically absence of visible antral follicles and/or presence of corpora lutea, respectively (Anderson & Simpson 1973).

Ovarian tissue then was manipulated under a stereomicroscope using a scalpel blade and 25 g needle to mechanically-isolate follicles. Follicles were isolated from the ovarian tissue within 10 hours of excision. Only those having at least two granulosa cell layers with a homogeneously dark, circular oocyte and intact basement membrane [41] were included in the study. Isolated follicles meeting these criteria were individually-encapsulated in a bead with 22.5 mg/ml fibrinogen (bovine, Sigma) and 0.38% (w/v) alginate (FMC BioPolymers, Philadelphia, PA) according to methods described by [39]. Briefly, follicles in fibrinogen-alginate (5  $\mu$ l) were pipetted into 50 U/ml thrombin in Tris BuPH with 40 mM CaCl<sub>2</sub>. Alginate crosslinks in the presence of a divalent cation, such as calcium [42], and fibrinogen polymerizes into fibrin when thrombin activates factor XIIIa [39]. Beads then were washed in  $\alpha$ -Minimum Essential Medium containing 3 mg/ml (~45  $\mu$ M) BSA, 4.2  $\mu$ g/ml (0.72  $\mu$ M)

insulin, 3.8 µg/ml (47.5 nM) transferrin, and 5 ng/ml (63.3 nM) selenium, hereafter called 'Growth Medium' [41].

## 2.2 Follicle Culture and Growth

Each encapsulated follicle was transferred to its own individual well of a flat-bottomed 96-well culture plate in 100 µl Growth Medium supplemented with a specific concentration of recombinant human activin (Sigma) and/or FSH (Folltropin V, BioNiche Animal Health, Athens, GA). Follicles were assigned randomly to one of 12 treatments of 0, 100 (3.8 nM), or 200 ng/ml (7.7 nM) activin and 0, 0.1, 1, or 10 µg/ml (0, 0.2, 1.7, or 17.5 IU/L) FSH (3 × 4 factorial design).

As mechanical isolation can result in a small layer of ovarian stromal cells covering the isolated follicles [43], two size evaluations were taken for each follicle on the day of culture onset (Day 0) – one for determination of follicle stage (only measuring diameter within the basal lamina) and another of full the structure (diameter of follicle plus surrounding stromal cell layer) (Fig. S1). This latter size measurement was used to assess follicle growth over the culture period. Determination of stages have been defined in our previous work [44]: preantral follicle (100 – 230 µm diameter); early antral (231 – 330 µm); and antral (331 – 550 µm). For both sizing approaches, the widest apparent diameter was measured followed by determining perpendicular width. A mean for these two values was calculated and reported as diameter [45]. We used the same technique on Day 0 to determine oocyte diameter (excluding the zona pellucida). All follicle and initial oocyte diameters were assessed using an inverted microscope (Leitz DM-IL, Research Instrument Limited, Falmouth, Cornwall, UK) with a heated stage, 10x objective, and optical micrometer.

Follicles were cultured at 38.5°C in a humidified atmosphere of 5% CO<sub>2</sub> with existing medium exchanged with fresh every 72 hours. Diameter of each follicle was also measured on Days 3, 6, 9, 12, 15, and 21 following culture onset. Percentage change in diameter from Day 0 of incubation was calculated as follicle growth. Incubation of follicles was terminated at one of two time points (Day 12 or 21) and cumulus-oocyte complexes (COCs) collected to determine oocyte diameter (for subsequent growth analysis) and to evaluate TZP integrity and nuclear status (see below).

## 2.3 Follicle Survival

Follicles that survived were defined as those sustaining a normal follicular appearance. Those that failed to survive had decreased in diameter over at least two consecutive observation time points, and/or contained a degenerate or extruded oocyte (emerging from the follicle's basement membrane), or displayed fragmented granulosa cells [46]. Follicles with these degenerating traits apparent before Day 3 were considered to have had an undetectable defect prior to, or as result of, the isolation/encapsulation process. A total of 955 follicles were included in the study (136 representing the preantral stage; 480 early antral; 339 antral; Supplemental Table 2).

## 2.4 Antral Cavity Expansion

For the antrum assessment, we focused on antral cavity expansion, rather than initiation of cavitation, for two reasons. First, ~85% of our isolated follicles already had begun the cavitation process (i.e., met early antral or antral criteria; [47]). Second, we had determined in a previous study [45] that dog follicles incubated in 0.5% hydrogel developed antral cavities that subsequently collapsed. Therefore, our aim here was to both maintain antrum integrity and expand this cavity's size. We recorded follicles as being 'expanded antral' when the diameter exceeded 630  $\mu\text{m}$  and the fluid-filled cavity was maintained throughout the culture interval. For this assessment, we fixed 205 follicles on Day 21 (10 – 16 per treatment) in Bouin's solution followed by embedding in paraffin blocks. For each block, 6  $\mu\text{m}$  thick sections were cut, mounted on slides, and stained with haematoxylin and eosin for subsequent microscopic evaluations of presence of theca cells and an antral cavity.

## 2.5 Oocyte Trans-Zonal Projections (TZPs), Growth, and Intact Nuclear Status

At the end of culture on Day 12 or 21, two 25 g needles were used to tease each follicle from its encompassing FA-IPN, followed by careful tearing of the basement membrane to release the COCs. Each COC was fixed in 4% paraformaldehyde for 12 hours and then stored in wash buffer (0.2% azide, 2% normal goat serum, 1% bovine serum albumin, 0.1 M glycine, and 0.1% Triton X-100 [48] at 4°C for subsequent analyses. COCs were grouped based on hormone treatment for subsequent assessment of oocyte size, TZIP density, and nuclear status. Actin staining of fixed COCs (n = 242) was achieved using Alexa Fluor 488 Rhodamine Phalloidin [48] (Invitrogen at 1:100 dilution in wash buffer) for at least 30 min (room temperature) with simultaneous nuclear staining using 1  $\mu\text{g}/\text{ml}$  (2.2  $\mu\text{M}$ ) Hoechst 33342 (in a 90% glycerol, 10% PBS solution). No more than four oocytes per treatment group were mounted on a slide (with GVA Aqueous Mounting Solution; Genemed Biotechnologies, San Francisco, CA). Imaging was at 100x with an LSM 510 laser scanning confocal microscope (Carl Zeiss, Germany) and a 488 nm krypton/argon laser and ultraviolet light. Z-stack images were taken of each COC. The z-stack slice imaging the oocyte at its widest diameter was used for TZIP assessment. The circumference of the oocyte was measured and TZPs were pseudo-colored using ImageJ software (NIH, Bethesda, MD). TZPs were counted and density calculated per 10  $\mu\text{m}$  oocyte circumference. Oocyte diameter (in  $\mu\text{m}$  excluding the zona pellucida) was determined by taking the mean of the widest diameter across the vitelline membrane and then a perpendicular width to the initial assessment. Presence of an intact nucleus (as opposed to fragmented chromatin, or oocytes no visible nucleus) was recorded for each COC.

## 2.6 Granulosa Cell FSHR Expression

Additional short-term (3 d) cultures were carried out to evaluate the influence of activin supplementation on dog FSH receptor expression. This evaluation required collecting the ovaries from another 14 donors (age, 6 months – 2.5 years) and then the recovery of 639 additional follicles representing each of the three target stages; Supplemental Table 3). After isolation, these follicles were cultured in FA-IPN (as above), including in 0, 100, or 200 ng/ml activin and 0 or 10  $\mu\text{g}/\text{ml}$  FSH (3  $\times$  2 design). On the third day after culture onset, granulosa cells were collected in RNeasy and stored at -20°C on the basis of the three

follicle stages by six hormone treatments (4 – 5 cultures per pool; 18 total groups). After later thawing, total RNA was isolated using an Agilent Absolutely RNA Nanoprep kit with on-column DNase digestion (manufacturer's instructions). Synthesis of cDNA was performed with a Transcriptor High Fidelity cDNA Synthesis Kit (Roche) with 30 ng total RNA. A reverse-transcription polymerase chain reaction (RT-PCR) for FSH receptor was performed with FastStart Essential DNA Green Master (Roche) using  $\beta$ -actin as an endogenous control (primer details, Supplemental Table 4) on a LightCycler 96 (Roche).

## 2.7 Statistical Analyses

Follicle growth data were log-transformed to achieve normality followed by model selection using an Akaike information criterion (AIC) [49] in RStudio (Version 1.0.143) with the following factors: activin concentration; FSH concentration; follicle stage at isolation; and all related interactions, as well as age of follicle donor. Culture replicate (i.e. cultured follicles from an individual donor dog for all but nine experiments, in which follicles isolated from two dogs were pooled: See Supplemental Table 1) was included as a random variable in the model. All factors and interactions were included in the final model that relied on a standard least squares model and post-hoc Tukey-Kramer HSD test for significance via JMP 9.0 software (SAS, Cary, NC).

Follicle survival was evaluated using a proportional hazards model with donor dog age, follicle stage at isolation, activin concentration, FSH concentration, and the interactions of the latter three as model effects. Donor dog age was treated as a categorical variable on the basis of <8 mo of age (prepubertal, n = 4 culture replicates, Supplemental Table 1), 8 mo to <2 y (peripubertal, n = 15 replicates), 2 to 4 y (adult, n = 5 replicates), and >8 y (aged, n = 2 replicates). Due to the random nature of collections from local, commercial spay clinics, precise animal age was not always available; furthermore, there were no donors 5 to 8 y of age. Follicles that survived through the 12 or 21 d culture end points were censored. We used a Chi square likelihood ratio test to examine a specific type of follicle mortality associated with oocyte extrusion; our earlier data have suggested that this subcategory of follicle loss may be associated with a breakdown in oocyte and surrounding cumulus cell connectivity [45].

Proportions of follicles forming an expanded antrum were compared among groups using a Chi square likelihood ratio test. TZP density, proportion of oocytes with intact nuclei, and oocyte diameter differences among treatment groups and time points (Days 0, 12, and 21) were evaluated using a nonparametric Wilcoxon test. RT-PCR data were analyzed via the Pfaffl method, accounting for primer efficiency [50]. Results are expressed as the mean  $\pm$  SEM, and significant differences set at  $P < 0.05$ .

## 3. Results

### 3.1 Follicle Growth

Ovarian follicles, regardless of stage or treatment group, experienced growth over the 21 d culture. Overall, this growth tended to be rapid in the first 3–6 days of culture, and level off to a more steady, sustained growth in the latter half of the incubation period. Our statistical



model demonstrated that follicle stage at isolation, activin concentration, FSH concentration, and interaction of follicle stage x activin, activin x FSH, and follicle stage x activin x FSH influenced ( $P < 0.05$ ) subsequent follicular growth *in vitro*. Early antral follicles tended to increase in size over 21 day culture the most, on average  $40.7 \pm 3.4\%$  compared to  $37.5 \pm 7.9$  and  $32.4 \pm 2.9$  for preantral and antral counterparts, respectively. Providing activin during incubation (in the absence of FSH) increased ( $P < 0.05$ ) follicle size compared to the controls as early as Day 3 (Fig. 1). For example, in the absence of FSH and activin, early antral follicles grew only  $11.8 \pm 3.6\%$  by Day 3 compared to the same stage follicles exposed to 100 ng/ml activin and 0  $\mu\text{g/ml}$  FSH, which increased  $22.7 \pm 3.3\%$  in diameter.

Figure 1 illustrates the interaction between follicle stage and hormonal supplementations. Preantral follicles experienced no benefit ( $P > 0.05$ ) of supplemental activin. Conversely, presence of 100 or 200 ng/ml activin enhanced ( $P < 0.05$ ) follicle growth for the early antral and antral cohorts, but only in the presence of FSH. For example, early antral follicles exposed to 0.1  $\mu\text{g/ml}$  or 1  $\mu\text{g/ml}$  FSH plus 100 ng/ml activin increased ( $P < 0.05$ ) in size  $52.6 \pm 6.7\%$  and  $63.8 \pm 14.1\%$ , respectively, by Day 21 compared to only  $17.9 \pm 6.5\%$  and  $23.2 \pm 14.8\%$  in these FSH dosages alone. There was no influence ( $P > 0.05$ ) of increased activin concentration (100 versus 200 ng/ml) on *in vitro* growth, regardless of follicle stage.

There was an effect of donor dog age on the ability of ovarian follicles to grow *in vitro*. Specifically, follicles isolated from adult / prime breeding age individuals (2 – 4 y old) increased in size the most ( $P < 0.05$ ) compared to the same stage follicles recovered from prepubertal (<8 mo) or peripubertal (8 mo - <2 y) bitches (Fig. S2A).

### 3.2 Follicle Survival

The parametric survival model demonstrated that the only significant factors were FSH dosage and donor age. Original follicle stage (Fig. S3A), activin supplementation (Fig. 2A) or their interactions had no influence ( $P < 0.05$ ) on follicular survival. Follicles had improved ( $P < 0.05$ ) survival when cultured with 0 or 10  $\mu\text{g/ml}$  FSH compared to the lowest rate after treatment with 0.1  $\mu\text{g/ml}$  (Fig. S3B). Follicles recovered from <2 y old donors also underperformed with higher mortality ( $P < 0.05$ ) compared to all older groups (Fig. S2B).

When evaluating follicle loss specifically due to oocyte extrusion, there was an impact ( $P < 0.05$ ) of follicle stage and FSH presence. The greatest frequency (13%) of this anomaly occurred in preantral follicles exposed to exogenous FSH (Fig. 2B). Although oocyte extrusion was not attenuated by activin presence, age of donor bitch increased the likelihood ( $P < 0.05$ ) of this phenomenon occurring (circa 53% and 57% in bitches < 8 mo and 8 mo - 1.5 yr of age compared to 75% and 86% for donors 2 – 4 yr and > 8 y of age, respectively).

### 3.3 Antral Cavity Expansion

Antral cavity expansion (Fig. 3A) occurred in early antral and antral stage follicles cultured in the presence of both activin and FSH (Fig. 3B). Only a single follicle categorized as 'preantral' at the onset of culture developed a large antral cavity (in presence of 100 ng/ml activin and 10  $\mu\text{g/ml}$  FSH). No follicle experienced expanded cavitation in the absence of activin or FSH. The combined supplementation of 100 or 200 ng/ml of activin and 0.1, 1, or 10  $\mu\text{g/ml}$  FSH produced 36 total follicles with cavity expansion (3.4% of total cultured),

with most of this cavitation (88.9%) occurring Days 6 to 15 from incubation onset (i.e., midculture). Histology at 21 d of incubation revealed an absence of cells with theca-like morphology compared to freshly-isolated, sized-matched control follicles (Fig. 3C). Presumptive small antral cavities also were observed in ~40% of follicles; these cavities stained pink in the presence of eosin (Fig. 3C), a trait not observed in cavities of fresh controls. Donor dog age had no influence ( $P > 0.05$ ) on antral cavity expansion.

### 3.4 Oocyte Trans-Zonal Projections (TZPs), Growth, and Intact Chromatin

Confocal microscopic images of TZPs in freshly-isolated oocytes and cultured follicles on Days 12 and 21 after FSH and activin exposure (and respective oocytes on Day 21) are depicted in Figure 4A–C). Regardless of treatment, vesiculation was notable along the cortical edge of oocytes at the junction with TZPs in 66% and 58% of Day 12 and 21 oocytes with TZPs, respectively (Fig. 4B,C). Average TZP density for COCs from freshly-isolated preantral versus antral follicles was  $2.4 \pm 0.4$  and  $3.5 \pm 0.1$  TZPs/10  $\mu\text{m}$  oocyte perimeter. Both these values were higher ( $P < 0.05$ ) than for COCs from cultured follicles (average across all treatments,  $0.51 \pm 0.10$  TZPs/10  $\mu\text{m}$ ). Neither hormone influenced ( $P > 0.05$ ) incidence of vesiculation. TZP density was reduced ( $P < 0.05$ ) in follicles exposed to the highest (200 ng/ml) activin concentration over the 12 d incubation (Fig. 4E). For example, oocytes from follicles cultured for this interval in 0 or 100 ng/ml activin and 1  $\mu\text{g/ml}$  FSH averaged  $0.72 \pm 0.20$  and  $0.54 \pm 0.27$  TZPs/10  $\mu\text{m}$ , respectively. By contrast, those cultured in the same FSH concentration, but with 200 ng/ml activin averaged only  $0.37 \pm 0.25$  TZPs/10  $\mu\text{m}$  ( $P < 0.05$ ). Follicles cultured in 200 ng/ml activin consistently experienced reduced TZP density on Day 12 compared to 0 or 100 ng/ml activin counterparts (regardless of FSH dosage). However, at the Day 21 evaluation and in the presence of 1  $\mu\text{g/ml}$  FSH, it was clear that activin was required to maintain TZP density. At this time, the average number of TZPs/10  $\mu\text{m}$  was less ( $P < 0.05$ ) in absence of activin ( $0.09 \pm 0.5$ ) compared to 100 ng/ml ( $0.89 \pm 0.40$ ), or 200 ng/ml ( $0.46 \pm 0.31$ ) activin, respectively. These latter values were similar to the highest average TZP density measured in oocytes at the abbreviated (12 d) incubation time point.

For oocytes from surviving follicles, growth occurred from Day 0 to 12 followed by a plateau through to Day 21 (Fig. 5A). Supplementation with 100 ng/ml activin increased ( $P < 0.05$ ) oocyte diameter ( $87.5 \pm 2.9$   $\mu\text{m}$ ) at Day 21 compared to non-supplemented controls ( $77.2 \pm 3.1$   $\mu\text{m}$ ) (Fig. 5A). FSH concentration had no influence ( $P > 0.05$ ) on oocyte size at Day 12 or 21. Presence of activin at 100 or 200 ng/ml, compared to activin absence, produced a higher proportion of oocytes with intact chromatin, with significance reached on Day 12 for 0.1  $\mu\text{g/ml}$  FSH and Day 21 for 10  $\mu\text{g/ml}$  FSH (Fig. 5B). FSH concentration had no effect ( $P > 0.05$ ) on intact nuclear status of recovered oocytes on Days 12 or 21 (Fig. 5B).

### 3.5 Granulosa Cell FSHR Expression

There was no statistical inference ( $P > 0.05$ ) to be made from an activin impact on FSHR expression, largely because of variation among replicates (Fig. 6). Of all the comparisons, activin may have been stimulative only in the preantral cohort in the absence of FSH compared to the fresh, non-cultured control. FSH and activin supplementation alone or in



combination had no influence ( $P > 0.05$ ) on level of the FSHR transcript in incubated early antral and antral follicles.

#### 4. Discussion

Until now, all studies examining the role of activin have focused on traditional livestock, laboratory rodent, and non-human primate models. Here we were interested in exploring the influence of this hormone and its mechanisms of action in the reproductively-complex dog that only displays ovarian follicular activity only 1–2 times yearly. Previous investigations have revealed that FSH can stimulate growth of cultured dog follicles [41], but additional LH supplementation fails to promote antral cavity formation [45]. Sequentially increasing FSH concentrations [51], supplementing with both FSH and high concentrations of insulin [52] or growth hormone [53] *in vitro* results in antrum development. Yet, in each of these previous studies, follicle growth and antral cavity expansion have been limited to  $<500 \mu\text{m}$  in diameter (only ~25% of the size needed to produce an oocyte capable of achieving MII [9]). The goal of our present study was to explore, for the first time, the potential role of activin on follicle/oocyte growth and survival, including in regulating two crucial functions, antral cavity expansion and cumulus cell-oocyte communication.

From the adaptation of a 3D culture system (already known useful in other models) [39, 54], we made four significant discoveries useful to the dog while extending the informational base on activin's role in mammals at large. First, the influence of activin was specific to follicle stage, with virtually all effects exerted at the level of the early antral or antral cohorts. The preantral stage likely lacks an active mechanism or receptor to respond to activin stimulation. Second, activin's role required presence of FSH to achieve a synergistic effect on follicle growth and antral cavitation in both the early antral and antral stage follicle. In the absence of FSH, there was no influence of activin on follicle growth or cavity expansion. Third, although having a negligible effect on follicle survival *in vitro*, activin supplementation promoted cumulus cell-oocyte communication by partially sustaining TZP density in long-term culture. Finally, although having no growth promotion influence on the resident oocyte, it was clear that activin facilitated preservation of egg nuclear integrity and normality. There was no evidence that these impacts of activin were mediated via changes in FSHR mRNA expression.

Although we identified multiple roles for activin in the dog system, it was somewhat unexpected that activin had no discernible effect on the preantral follicle, especially as activin A is known to promote growth of these same stage follicles in the cow [55]. This hormone clearly was not the primary driver for initiating antrum development in the dog. Cavity initiation in the preantral follicle most likely demands theca cells, an obligation lacking in our current microenvironment (per our histological findings). The pink eosin staining in antral cavities, indicative of an internal richness in intrafollicular proteins, may also be due to the lack of theca cells that facilitates fluid incursions into the follicle via aquaporins [56]. One possible resolution for the dog system, currently under exploration, is evaluating other promoters of theca cell development, for example, kit-ligand that is known to help fully stimulate antrum cavitation in early stage bovine follicles [57].

Once at the early antral stage, the dog follicle had developed a sensitivity to the presence of combined activin and FSH. This synergism resulted in increased growth and antral cavity expansion, which was similar to what has been observed in the cow, human, and rat [23–25]. Interestingly, supplementing primary rat follicles with activin seems to serve a priming function for a more robust response to gonadotrophin when the activin is withdrawn [58]. In this same species, activin also appears to promote FSH receptor mRNA expression in granulosa cells [37, 38, 59], a mechanism that we did not observe in the dog. As activin also has been postulated to block the stimulatory effects of FSH on early stage preantral follicles in the cow [55] and rat [58], it would be useful to examine the influence of this hormone only on this stage dog follicle before FSH exposure.

Incubation with a specific combination of activin and FSH (1 µg/ml) supported the maintenance of TZP density from culture days 12 through 21. The benefit of activin to TZP density was reduced by the highest activin supplementation (200 ng/ml) during the first 12 d of culture, possibly due to accelerated granulosa cell proliferation and follicle expansion that impeded new projections and/or normal connectivity to oocytes. This same phenomenon was also likely occurring during oocyte extrusion. Premature oocyte discharge has been described previously in the domestic dog [51] and has occurred in follicular cultures of the cow [60] goat [61], and human [62]. As none of our control follicles (0 FSH) experienced extrusion, it is likely that our preantral cohort was inadequately primed for the level of granulosa cell proliferation produced in the presence of FSH. However, our ability to recover TZP density in the presence of both activin and FSH indicated the exciting potential of achieving longer-term, successful follicular cultures. The phenomenon of vesiculations at the junction of the oocyte cortex and TZP (observed here in 57% of dog follicles) has also been described in an ultrastructural evaluation of *in vitro* and *in vivo* matured dog oocytes [63, 64], as well as in the human COC [65]. In these studies, deeply embedded TZPs with a ‘bulbous ending’, perhaps resulting from an in-folding of the oolemma, were noted, although frequency of this observation went unreported. Ultrastructural evaluations in other species have demonstrated that TZPs end in foot-processes adjoining the oocyte’s plasma membrane [66]. Therefore, these vesiculations may have originated from a normal feature, but one where the TZP simply grew atypically large. Although our findings revealed no definitive cause for this structural peculiarity, we would suggest that such formations may serve as useful markers for effectiveness of *in vitro* maturation of both the follicle and its resident oocyte. Further, as TZP density in cultured follicles, regardless of treatment, were reduced compared to fresh controls, it is evident that activin is not the sole factor promoting this interaction. Future studies should explore the function of growth differentiation factor 9 in simulating TZP formation, as demonstrated in the mouse [67], in dog follicles.

We determined that activin played a beneficial role in ensuring both oocyte form and integrity. It is known from previous *in vivo* studies that the dog oocyte reaches a maximal size of ~113 µm (range, 86 to 132; [8], typically near the early antral follicular stage [44]). It was encouraging that our described microenvironment was capable of producing oocytes of this desired diameter from incubated follicles by 12 d post-culture onset. However, there was as much as 26% reduction in oocyte size during the second phase of culture (through 21 d) in the presence of 100 ng/ml activin and 10 µg/ml FSH. A more detailed analysis revealed that follicles exposed to 200 ng/ml activin in the presence of 0, 0.1, or 10 µg/ml FSH

maintained oocyte diameter throughout the 3 wk incubation (growing 0.4, 6.1, and 5.8%, respectively). Resident oocytes from follicles supplemented with 100 ng/ml activin and 1 µg/ml FSH (the group also producing the highest TZP densities at Day 21) increased in size a modest 2% over the latter half of culture. This information plus finding that activin supplemented follicles produced oocytes with more intact nuclei at both mid- and late incubation clearly demonstrated that this hormone promoted oocyte growth, survival, and integrity.

Lastly, although not a major focus of the present study, the influence of ovarian donor age on follicle viability *in vitro* was important. We identified clear advantages for developing these microenvironments using ovarian materials recovered from adult bitches. Prepubertal donors presented more challenges to achieving successful follicle growth and survival in culture but a lower prevalence of premature oocyte extrusions, probably at least in part due to naivete and lack of earlier gonadotropin priming. Therefore, our observations on the donor-age dependent performance of follicles should influence the source of ovaries for future studies.

In summary, we identified a probable role for activin in domestic dog folliculogenesis, promoting growth of early antral and antral stage follicles, largely through antrum cavity expansion and in cooperation with FSH. This improved ability to reach later developmental stages *in vitro* appears due to activin maintaining TZP density and, therefore, oocyte viability for a 21 d culture interval. Using the permissible FA-IPN incubation system, the follicle growth cap of 500 µm diameter achieved in our recent previous investigation [45] was surpassed 1.8-fold (to 800 to 900 µm), largely by the current protocol expanding the antral cavity. As a large antral follicle is critical to producing a meiotically-competent oocyte, our findings here edge us closer to the ultimate goal of developing an *in vitro* folliculogenesis system for the domestic dog.

## Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

## Acknowledgements

The authors thank: Dr. Ariella Shikanov (University of Michigan) and the laboratories of Drs. Lonnie Shea and Teresa Woodruff (Northwestern University) for instruction in fibrinogen preparation; veterinary clinics in Front Royal, Stephen's City, and Harrisonburg, VA for donating domestic dog ovaries; Dr. Megan Brown (SCBI) for assisting with model selection statistical analyses; and Dr. Lara Moutham and two anonymous contributors for thoughtful reviews of the manuscript.

### Funding

This research was supported by a Smithsonian Institution Doctoral Fellowship (to J.N.), the Smithsonian Scholarly Studies Program, and by a grant from the National Institutes of Health KO1RR020564 (to N.S.).

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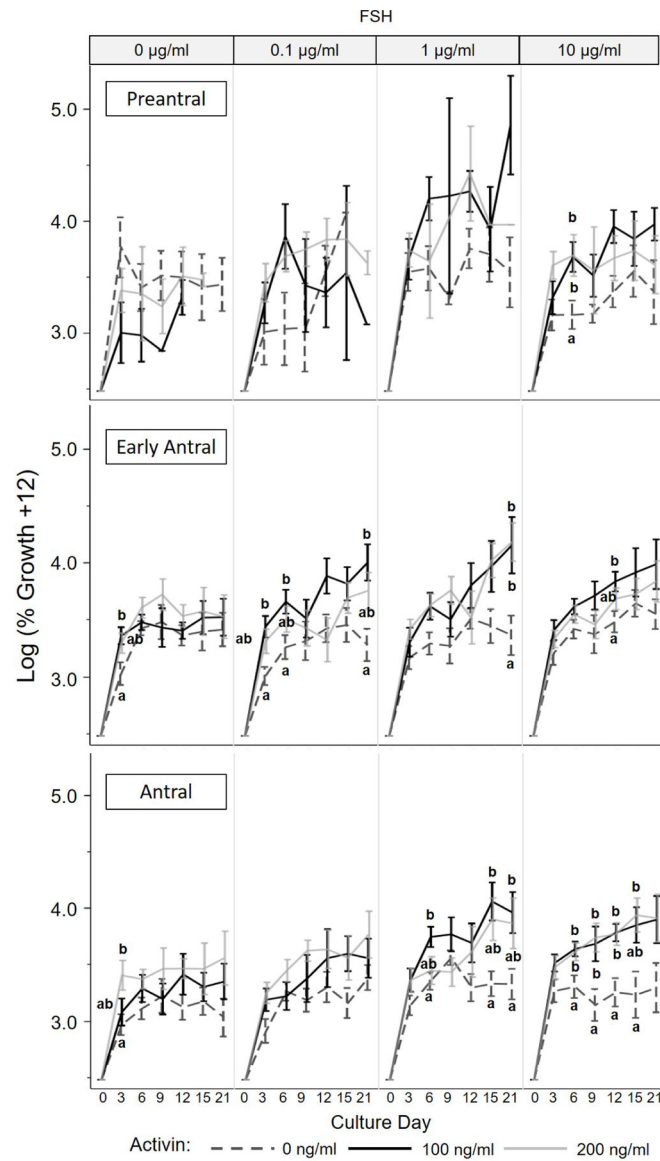
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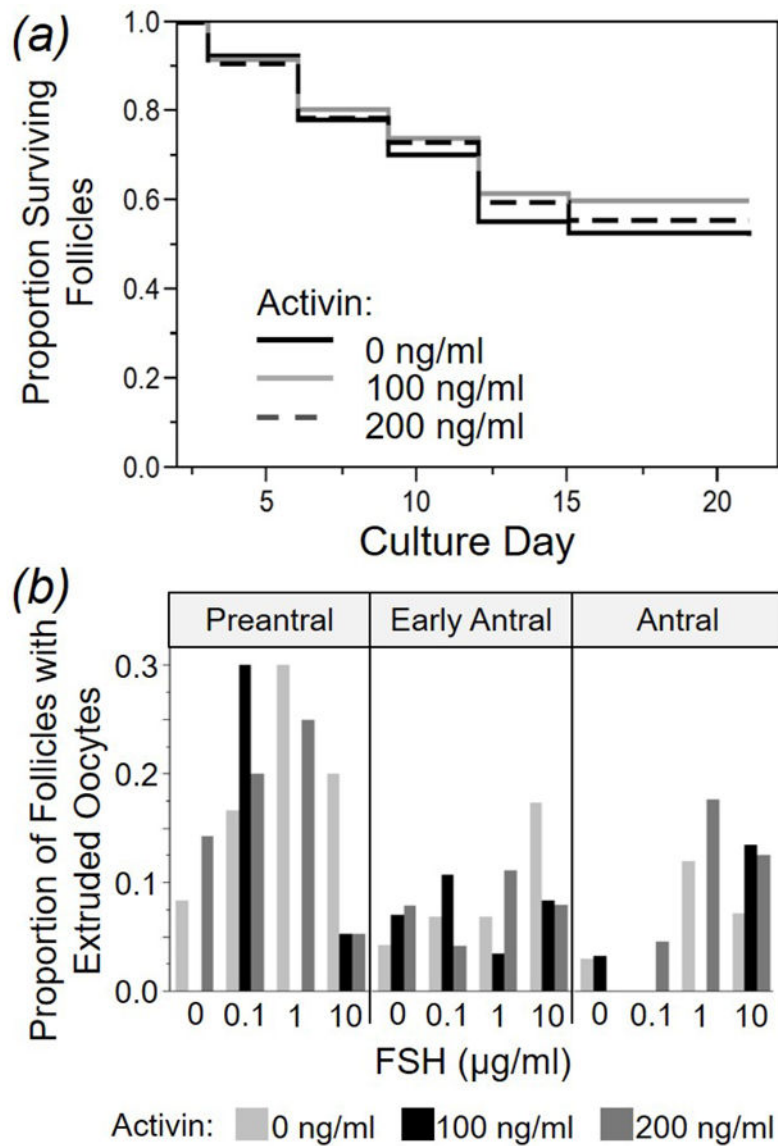
### Highlights

- Domestic dog ovarian follicles have a stage-specific response to activin supplementation, with early antral and antral but not preantral stage follicles responding to activin in culture.
- Activin and FSH have a synergistic effect on growth and antral cavitation in both early antral and antral stage domestic dog follicles that is not mediated via changes in FSHR mRNA expression.
- Activin facilitated improved preservation of dog oocyte nuclear integrity and trans-zonal projection density, but not follicle survival, during *in vitro* culture.

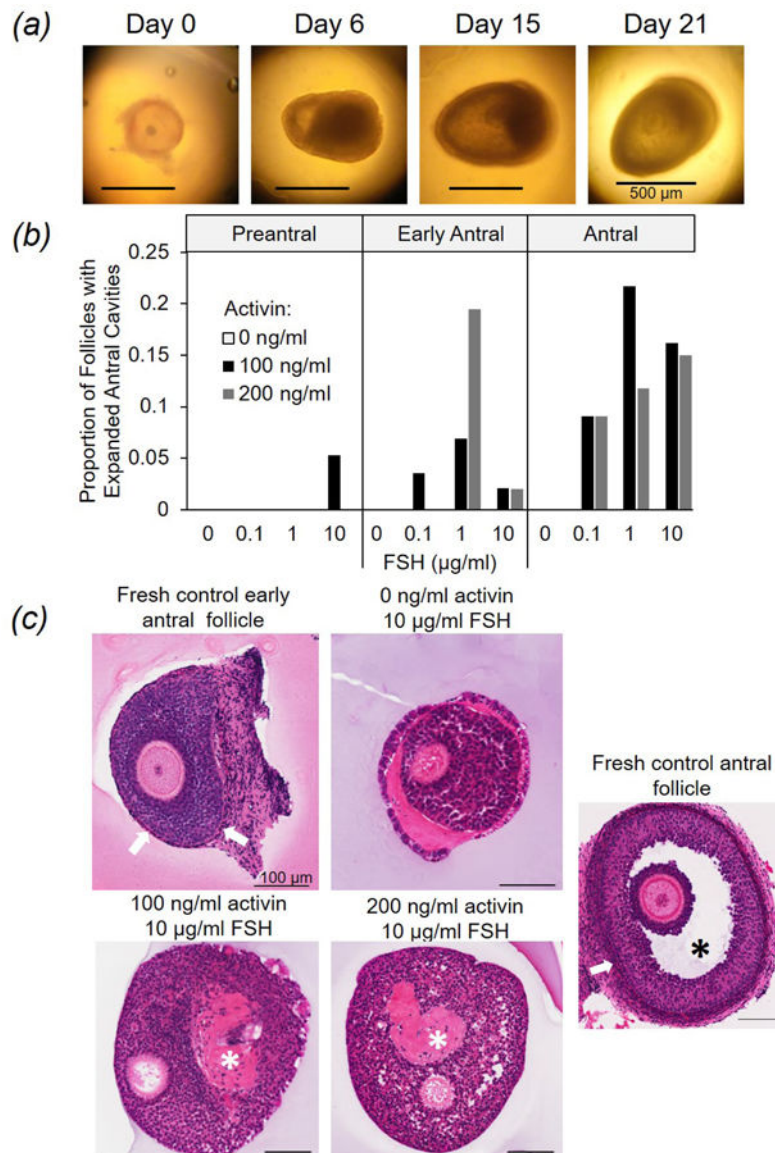


**Figure 1.**

Growth (log transformation of percent diameter increase compared to Day 0 defined as onset of culture)  $\pm$  SEM of preantral, early antral, and antral stage dog ovarian follicles cultured *in vitro* with various activin (0, 100, or 200 ng/ml) and FSH (0, 0.1, 1, or 10  $\mu$ g/ml) concentrations. Values with differing letters within the same follicle stage, FSH treatment, and culture day represent differences ( $P < 0.05$ ) among activin concentration groups.

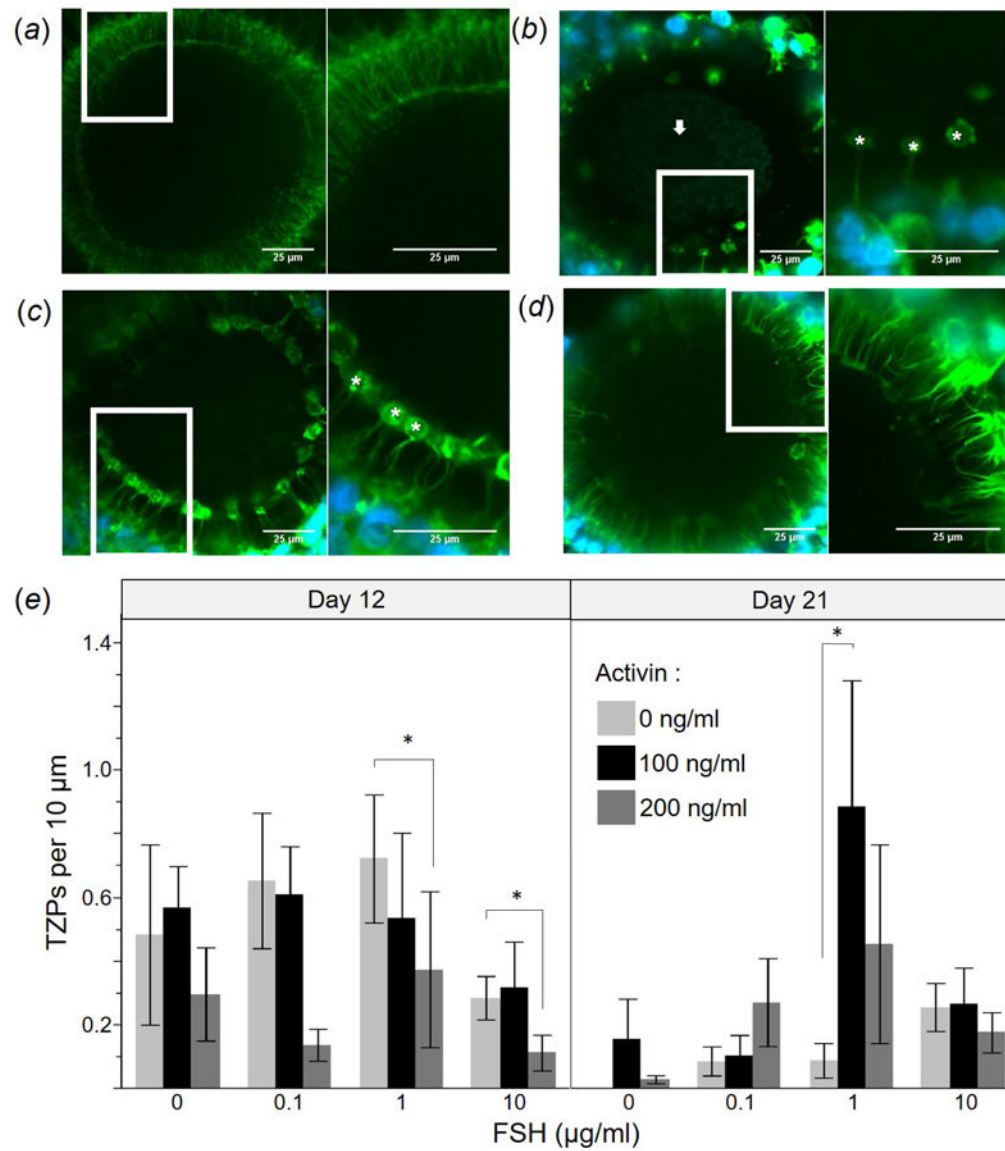


**Figure 2:** Proportions of dog ovarian follicles (A) surviving in 0, 100, or 200 ng/ml activating (combining all follicle stages and FSH treatment groups) over a 21 d culture, and (B) experiencing oocyte extrusion based on original stage at isolation and hormone concentration *in vitro* (Activin, 0, 100, or 200 ng/ml; FSH, 0, 0.1, 1, or 10 µg/ml).



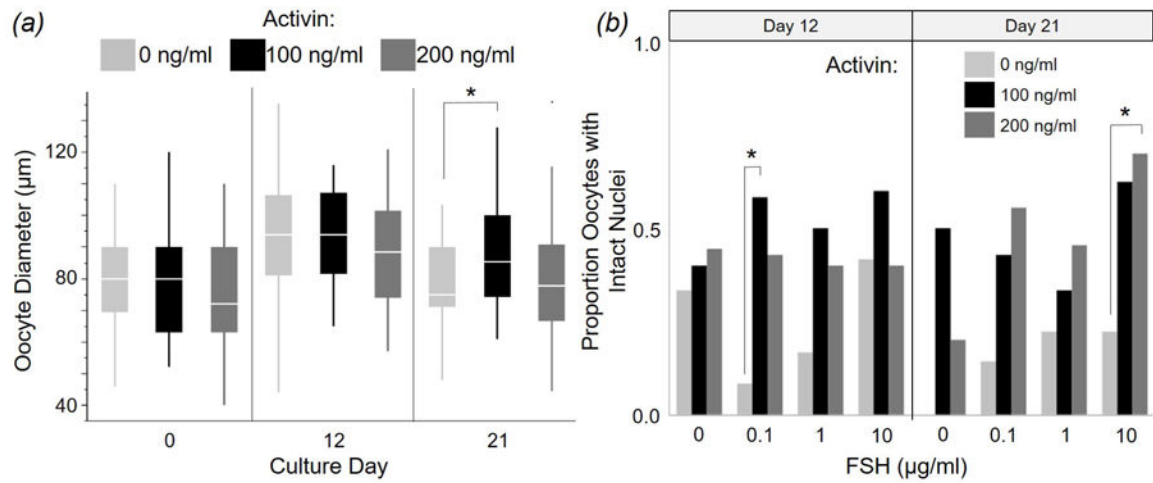
**Figure 3:**

Antral cavity formation in cultured dog ovarian follicles. (A) Images of a single, early antral follicle exposed to 100 ng/ml activin and 1  $\mu\text{g/ml}$  FSH developing an antral cavity over 21 d (Day 0 = culture onset). (B) Proportion of follicles developing an expanded antral cavity based on original stage at isolation and hormone concentration (activin, 0, 100, or 200 ng/ml; FSH, 0, 0.1, 1, or 10  $\mu\text{g/ml}$  FSH). (C) Hematoxylin and eosin staining of freshly-isolated early antral and antral follicles (controls) and counterparts cultured 21 d (activin, 0, 100, or 200 ng/ml and FSH, 10  $\mu\text{g/ml}$ ). White arrows indicate theca cell layers, and asterisks (\*) mark antral cavities in fresh (black) versus cultured (white) follicles.



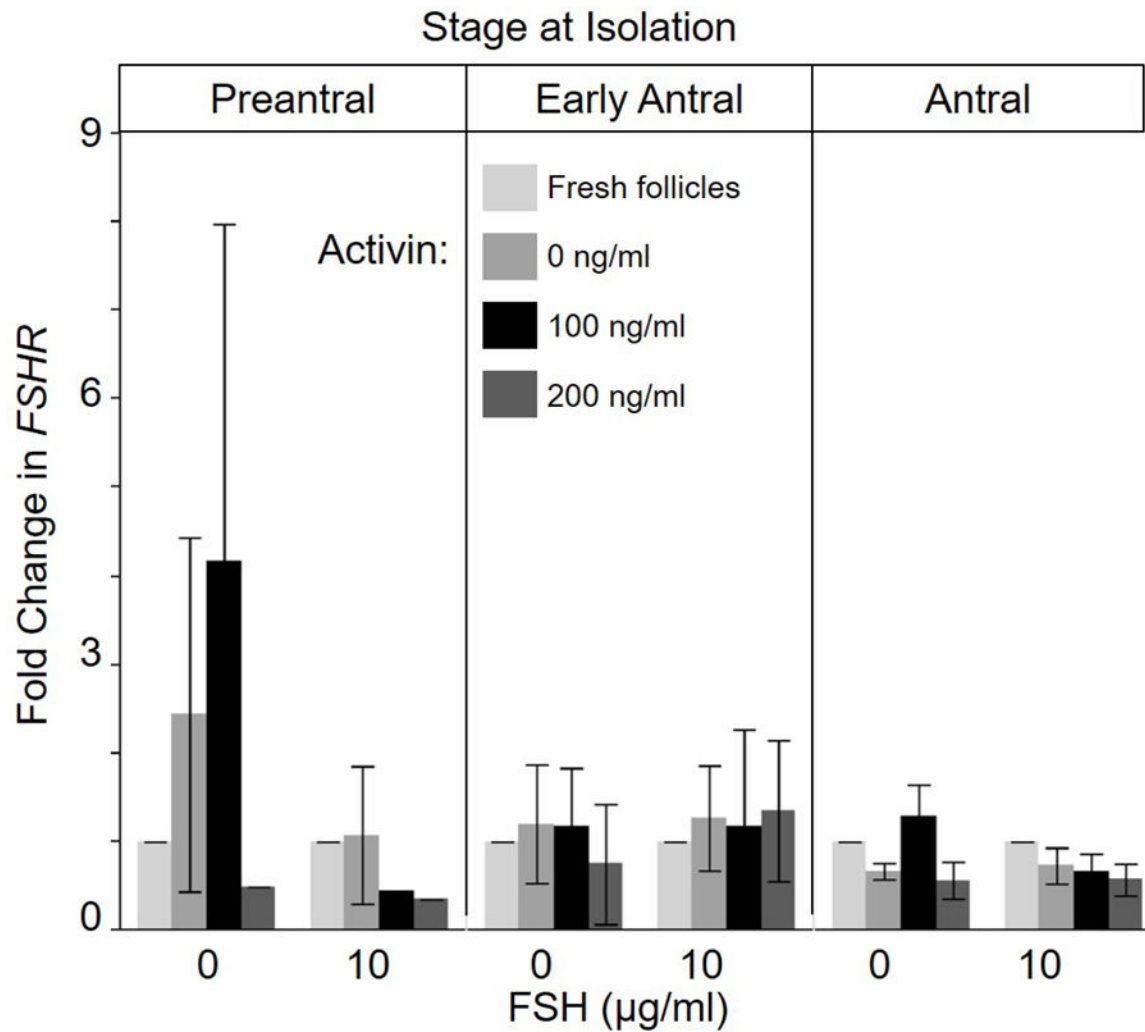
**Figure 4:** Confocal microscopy images depicting trans-zonal projections (TZPs) in oocytes of dog ovarian follicles that were (A) freshly-isolated (control) or cultured for 21 d in 1 μg/ml FSH and (B) 0, (C) 100, or (D) 200 ng/ml activin. Inserts within panels indicate magnified (2x) areas depicted in each right panel; asterisk (\*) indicates vesicularisation, and arrow denotes nucleus. (E) TZP density (mean ± SEM) after 12 and 21 d culture (activin, 0, 100, or 200 ng/ml; FSH, 0, 0.1, 1, or 10 μg/ml). Asterisks (\*) indicate differing ( $P < 0.05$ ) values among activin concentrations within a given FSH dosage and culture day.





**Figure 5:**

(A) Oocyte growth based on diameter changes during culture of dog follicles for 12 or 21 d while exposed to 0, 100, or 200 ng/ml activin (combining all FSH treatment groups). (B) Proportion of chromatin-intact oocytes from donor follicles supplemented with activin (0, 100, or 200 ng/ml), and FSH (0, 0.1, 1, or 10 µg/ml) *in vitro*. All values are means ± SEM. Asterisks (\*) indicate differing ( $P < 0.05$ ) values among activin concentrations within a time point.



**Figure 6:**

Fold change (mean  $\pm$  SEM) in *FSHR* expression on Day 3 of cultured dog ovarian follicles (Day 0 = incubation onset) using *B actin* as reference gene. Four to five independent cultures were pooled by treatment group (0, 100, or 200 ng/ml activin, and 0 or 10 µg/ml FSH) and initial follicle stage (preantral, early antral, or antral). No differences ( $P > 0.05$ ) were found.