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## **Transcriptomic profiling of neonatal mouse granulosa cells reveals new insights into primordial follicle activation†**

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Research Article

# Transcriptomic profiling of neonatal mouse granulosa cells reveals new insights into primordial follicle activation<sup>†</sup>

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## Abstract

The dormant population of ovarian primordial follicles is determined at birth and serves as the reservoir for future female fertility. Yet our understanding of the molecular, biochemical, and cellular processes underpinning primordial follicle activation remains limited. The survival of primordial follicles relies on the correct complement and morphology of granulosa cells, which provide signaling factors essential for oocyte and follicular survival. To investigate the contribution of granulosa cells in the primordial-to-primary follicle transition, gene expression profiles of granulosa cells undergoing early differentiation were assessed in a murine model. Ovaries from C57Bl/6 mice were enzymatically dissociated at time-points spanning the initial wave of primordial follicle activation. Post-natal day (PND) 1 ovaries yielded primordial granulosa cells, and PND4 ovaries yielded a mixed population of primordial and primary granulosa cells. The comparative transcriptome of granulosa cells at these time-points was generated via Illumina NextSeq 500 system, which identified 131 significantly differentially expressed transcripts. The differential expression of eight of the transcripts was confirmed by RT-qPCR. Following biological network mapping via Ingenuity Pathway Analysis, the functional expression of the protein products of three of the differentially expressed genes, namely FRZB, POD1, and ZFX, was investigated with in-situ immunolocalization in PND4 mouse ovaries was investigated. Finally, evidence was provided that

Wnt pathway antagonist, secreted frizzled-related protein 3 (FRZB), interacts with a suppressor of primordial follicle activation WNT3A and may be involved in promoting primordial follicle activation. This study highlights the dynamic changes in gene expression of granulosa cells during primordial follicle activation and provides evidence for a renewed focus into the Wnt signaling pathway's role in primordial follicle activation.

### Summary sentence

Comparing transcriptomes of two populations of granulosa cells: those from primordial follicles and a mixed group enriched for granulosa cells from primary follicles. *Frzb* expression increased in primary granulosa cells relative to primordial, and FRZB interacts with primordial follicle activation suppressor WNT3A.

**Key words:** folliculogenesis, granulosa cells, RNAseq, infertility, premature ovarian insufficiency.

## Introduction

Primordial follicle activation is integral to the fertility of sexually reproducing females as it is necessary for the follicle development and is the committing step to ovulation and subsequent fertilization. The number of primordial follicles within the ovaries defines the age of menopause in women, and therefore the end of a woman's fertility. Premature ovarian insufficiency (POI) is the early cessation or absence of ovarian function due to a reduction in the pool of primordial follicles in women before the age of 40 [1]. POI occurs in 1–3% of women, and a common cause of POI is a rapid acceleration in the rate of activation of primordial follicles [2–4]. The process of primordial follicle activation is complex, involving massive parallel molecular, cellular, and biochemical events, with limited characterization [5, 6]. Granulosa cells communicate between neighboring granulosa cells and intracellularly with the oocyte to coordinate primordial follicle activation [7]; however, the extent of their role in this process is yet to be determined. Importantly, as primordial follicles activate, there must be a sufficient number of granulosa cells supporting the oocyte, and they must all transition correctly to the activated, cuboidal morphological form for successful development [8, 9]. Improving our understanding of how the rate of primordial follicle activation is controlled is imperative to improving fertility outcomes for women at risk of premature infertility and POI.

The research in human primordial follicle activation has been hampered by small samples sizes and limitations in tissue (reviewed in [10]). The use of non-human models overcomes a number of the limitations associated with the study of human samples and is instrumental to our current understanding of primordial follicle activation. [10]. Much of the research underpinning our understanding of primordial follicle activation has focused on protein interactions and single gene knockouts in mice (reviewed in [6]). Briefly, it has been established that phosphoinositide 3-kinase (PI3K) stimulates primordial follicle activation in oocytes, and this is assisted by mammalian target of rapamycin complex 1 (mTORC1) and Forkhead box L2 (FOXO2) in the granulosa cells [11]. Consequently, well-known factors, which maintain the quiescence of primordial follicles include anti-Mullerian hormone (AMH), and phosphatase and tensin homolog deleted on chromosome 10 (PTEN) [12]. Animal studies provide important groundwork with advantages such as functional studies via conditional knockouts, drug testing, and robust characterization through omics technologies. In this study, granulosa cells isolated from mouse neonatal ovaries were utilized both before and during the initial wave of primordial follicle activation to obtain large quantities of granulosa cells originating from

primordial and primary follicles, respectively. These two distinct populations of cells were subjected to RNA-seq to compare the gene expression profiles of primordial and activating granulosa cells and identify novel factors contributing to follicle activation. This study presents the first detailed transcriptome comparison of granulosa cells from mouse primordial and activating follicles, we were able to annotate the expression occurring within granulosa cells and observed a small but significant shift in the transcriptome between the two populations of these cells. From this comprehensive dataset, we identified the Wnt pathway inhibitor, secreted frizzled-related protein 3 (FRZB) was significantly increased in expression in the population of granulosa cells isolated during the initial wave of primordial follicle activation, compared to granulosa cells isolated before activation and representing primordial follicles. Additionally, we determined an interaction between FRZB and WNT3A these granulosa cells. Our findings represent the commencement of a renewed investigation into the role of Wnt signaling factors in primordial follicle activation.

## Methods

### Ethics

C57Bl/6 mice were supplied by the University of Newcastle Animal Services Unit under ethics approval number A-2018-803. All experiments involving the use of animals were conducted in accordance with the Institutes' Animal Care and Ethics Committee guidelines.

### Tissue and cell collection

Mice were sacrificed at post-natal days one (PND1) and four (PND4). These time points were selected to coincide with immediately prior to the initial wave of primordial follicle activation when the ovary is comprised of only primordial follicles (PND1), and during the initial wave of primordial follicle activation when there is an enriched population of follicles undergoing activation (PND4) [13, 14]. See [Supplementary Material S1](#) for follicle counts of PND1 and PND4 ovaries for proportions of primordial and primary follicles. Follicles that develop during the initial wave of primordial follicle activation do contribute to post-pubertal fertility [15] and thus are used to investigate primordial follicle activation. This approach maximizes the quantity and homogeneity of the follicle types of interest retrieved for analysis that would otherwise be confounded by later-stage, gonadotropin-dependent follicles. Other biological processes do occur in the postnatal mouse ovary (reviewed

in [16]); however, the magnitude of primordial follicle activation taking place in the PND4 ovary is unparalleled by any other point in ovary development and thus makes PND1 and PND4 ovaries an effective way to explore primordial follicle activation.

Mice were housed under a controlled lighting regime (12 h light: 12 h dark) at 21–22°C and supplied with food and water ad libitum. Neonatal mice were euthanized by asphyxiation with carbon dioxide, followed by decapitation. Ovaries were collected immediately after euthanasia and either snap-frozen, fixed in 10% neutral buffered formalin for  $\leq 10$  h, or placed into 37°C Leibovitz L-15 media supplemented with 1% fetal bovine serum (FBS) for immediate granulosa cell collection. For a detailed description of the methodology and rationale behind time point selection and granulosa cell collection, see [17].

Briefly, the outer tissue and bursa were removed from the ovary, ovaries then transferred into Dulbecco Modified Eagle/F-12 medium (DMEM/F12), containing 0.02% collagenase, and 0.02% DNase I. Replicates contained between 4 and 8 ovaries, though some replicates contained pooled samples of up to 15 ovaries, as specified where appropriate. After incubation for 45–60 minutes at 37°C in 5% CO<sub>2</sub> with gentle pipetting every 15 minutes, cells were retrieved by centrifugation (1500 g, 3 min), supernatant removed, and incubated with 0.1% trypsin/EDTA at 37°C in 5% CO<sub>2</sub> for 15–20 minutes. Cells were recovered by centrifugation (1500 g, 3 min), and resuspended in ovary culture media (containing DMEM/F12 with 0.1% bovine serum albumen, 4% penicillin–streptomycin, 5% fetal bovine serum, 0.5% ITS-G and 3% L-glutamine). An aliquot of cells was removed for cell quantification, and the remaining cells incubated in ovary culture media with 0.5% ascorbic acid 18–24 hours at 37°C for either collection or fixing. When collecting granulosa cells, cells were incubated with 0.1% trypsin/EDTA for five minutes, collected (800 g, 15 min) and washed 3  $\times$  5 minutes at 1200 g in Hanks' Balanced Salt Solution, counted, and stored at –80°C until use. Cell populations were enriched for the cell target of interest, granulosa cells, rather than the other major cell type in neonatal ovaries, the oocytes, and stromal cells, as determined by oocyte and stromal counts using immunofluorescent targeting of DDX4 and Vimentin, respectively. Counts of 300 cells across four biological replicates indicate an average abundance of 0.4% oocytes and 0.8% stromal cells present in isolated granulosa cell samples. Granulosa cell viability prior to freezing and/or downstream experiments was determined via Trypan blue (Invitrogen, Carlsbad, USA; cat no. T102872) and accepted at  $\geq 80\%$  viability.

### RNA extraction and RT-qPCR

Granulosa cell RNA was extracted using the RNeasy Plus Micro Kit (Qiagen, Hilden, Germany; cat no. 74034) as per the manufacturer's instructions. Post-extraction, cDNA was synthesized by reverse transcription with the SuperScript IV VIL0 system (Invitrogen, Carlsbad, USA; cat no. 11766050). Quantitative real-time PCR (RT-qPCR) was performed in triplicate on cDNA with a reaction equivalent to 20 ng of total RNA. Predesigned and validated gene-specific TaqMan Gene Expression Assays (Life Technologies, Carlsbad, USA) were used for RT-qPCR. Each TaqMan gene expression assay contained gene specific, exon spanning forward and reverse primers for each of the genes of interest (Avpr1a (Mm00444092\_m1), Cdkn2b (Mm00483241\_m1), Ddr2 (Mm00445615\_m1), Fam171a1 (Mm01332727\_m1), Frzb (Mm00441378\_m1), Ifitm1 (Mm00850040\_g1), Rpl19 (Mm02601633\_g1), Tcf21 (Mm00448961\_m1), Zfx (Mm03053842\_s1)) and

fluorogenic minor groove binder probes consisting of a target-specific oligonucleotide labelled with a fluorescent dye FAM (6-Carboxyfluorescein) or VIC (2'-chloro-7-phenyl-1,4-dichloro-6-carboxy-fluorescein), and a non-fluorescent quencher. Data were normalized to the expression of the transcript encoding 60S ribosomal protein L19 (Rpl19). Triplicate expression values of each gene was set relative to the reference gene via the  $\Delta\Delta$ CT method [18] and is presented as the mean  $\pm$  SEM with statistical analysis determined by unpaired Student *t*-test.

### RNA sequencing and mapping transcripts

Eight, libraries were sequenced on the Illumina NextSeq 2  $\times$  75bp high output via Auckland Genomics at the University of Auckland. The RNA samples consisted of four replicates each of two groups of mouse granulosa cells (isolated from PND1 and PND4), each replicate contained pooled samples of cells isolated from 4–15 ovaries. The bioinformatics team at Auckland Genomics Facility performed the following analyses on the RNA sequencing output. The overall quality of the data was visualized using the program FastQC [19]. TrimGalore [20] was used to check and remove adapters from the sequences. Sequences with a Phred score less than 30 were trimmed. Reads with a length shorter than 25 base pairs were discarded. Hisat2 [21, 22] was used to map the cleaned reads to the mouse transcriptome (grcm38). Output sam files from hisat2 were then sorted by genomic position and converted to bam format using samtools sort [23]. To generate expression estimates for each sample, the mapped reads were then assembled into transcripts using StringTie. Differential expression analysis was performed using the R software package Ballgown [24]. Low abundance transcripts with a variance less than one across the samples were removed. To view the similarity between the samples, the Pearson correlation and distance between the samples were calculated. Differential gene expression between the two treatments was calculated using FPKM (fragments per kilobase million) values. The data were then sorted by their adjusted *p*-value (false discovery rate) and absolute log<sub>2</sub> fold change values.

### In silico analysis of gene expression

The analysis of genes expressed within PND1 and PND4 neonatal mouse granulosa cells was undertaken in silico using a number of techniques. Briefly, transcript abundance data were assessed via volcano plots to visualize trends associated with differentially accumulating genes in the primordial granulosa cells compared to activating granulosa cells (PND1 versus PND4). The threshold for significant differentially expressed transcripts was defined as having a two-fold change of FPKM in either direction (log<sub>2</sub> fold change  $\pm 1$ ), and a significance level of  $P \leq 0.05$ . Consistency of gene expression among biological replicates for transcripts above the determined threshold was visualized via a heat map of expression (FPKM) for each gene of each replicate normalized to the PND1 average FPKM for the respective gene. Differential expression that did not meet the threshold of statistical significance, but relevant to demonstrate the cell type of interest being captured are presented in Table 2.

Ingenuity pathway analysis (IPA) was used to explore biological networks for differentially expressed genes with significance  $P \leq 0.05$ . IPA queries a large database of experimental observations between molecules and uses this information to construct biological networks that represent cause-effect relationships between mammalian genes, proteins, and their functions with probability

**Table 1.** Antibody details.

Antibody target	Species raised in	Dilution	Company	Catalogue number
FOXL2	Goat	1:100	Abcam	ab5096
FRZB	Goat	1:50	ThermoFisher	PA5-47793
GATA4	Rabbit	1:100	Abcam	ab84593
POD1	Rabbit	1:100	Bios USA	BS-8688R
WNT3A	Rabbit	1:100	Abcam	ab28472
ZFX	Rabbit	1:200	ThermoFisher	PA5-34376

calculations [25]. The granulosa cell dataset of significantly differentially expressed genes ([Supplementary Material S2](#)) was interrogated for enrichment of functional pathways using bioinformatic enrichment tools available via the Database for Annotation, Visualization and Integrated Discovery (DAVID; v6.8) [26, 27] then, separate gene lists according to the clusters identified in the heatmap were assessed with the same tools. DAVID Gene Ontology (GO) annotation tools were utilized for exploration of biological process [28] in differentially expressed genes, GO identifiers identified within the dataset were also subject to enrichment analysis to determine their representation [29], for full list of DAVID GO terms and identifiers returned for this dataset, see [Supplementary Material S3](#).

### Immunolocalization

For immunofluorescence of mouse ovaries, tissues were washed with and embedded in Optimal Cutting Temperature (OCT) compound (ProSciTech, IA018), then snap-frozen in dry ice. Blocks were serially sectioned (5  $\mu$ m thick) with a cryostat (Leica Biosystems) onto Superfrost Plus slides (ThermoFisher; cat no. 4951PLUS4) three ovaries per section for biological triplicate. Before use, slides were dried to room temperature for 5 min and rehydrated in PBS. Heat mediated antigen retrieval was performed by warming slides for 30 min at 65°C in 10 mM sodium citrate buffer (pH 6). To prevent non-specific antibody binding, sections were blocked in phosphate-buffered saline solution with 5% donkey serum and 1% bovine serum albumen for 1.5 h at room temperature. Sections were probed with antibodies in Table 1 (antibody of interest, colocalized with a granulosa cell marker, GATA4 or FOXL2) and incubated overnight at 4°C. After washing, sections were incubated with secondary antibodies Alexa Fluor 488 donkey anti-rabbit IgG (ThermoFisher; cat no. A21206), Alexa Fluor 555 donkey anti-goat IgG (Abcam, cat no. A21432) at a dilution of 1:200 for 1 h at room temperature. Slides were counterstained with 4'-6-Diamidino-2-phenylindole (DAPI) and mounted in anti-fade reagent Mowiol (13% w/v Mowiol4-88, 33% w/v glycerol, 66 mM Tris (pH 8.5), 2.5% w/v 1, 4 diazobicyclo-[2.2.2] octane). Images were taken on an Olympus Fluoview 1000-IX81 confocal microscope (Olympus, Center Valley, USA) under fluorescent optics.

For immunocytochemistry of fixed granulosa cells, wells were permeabilized in phospho-buffered saline (PBS) with 0.01% Triton-X then blocked for 2 h at room temperature in PBS containing 10% donkey serum and 3% bovine serum albumen. Cells were probed overnight at 4°C with anti-FRZB antibody (ThermoFisher; cat no. PA5-47793), and anti-WNT3A antibody (Abcam, Cambridge, UK, cat no. ab28472), each at a dilution of 1:200. After washing, sections were incubated with secondary antibodies: Alexa Fluor 488 donkey anti-rabbit IgG (ThermoFisher; cat no. A21206), or Alexa Fluor 555 donkey anti-goat IgG (Abcam, cat no. A21432) at a dilution of 1:200

for 1 h at room temperature. Cells were counterstained with 4'-6-Diamidino-2-phenylindole (DAPI) and a layer of PBS was added to prevent drying out. Images were taken on an EVOS FL (AMF4300; ThermoFisher).

### Duolink proximity ligation assay

Protein interactions were detected via Duolink proximity ligation assay kit (Merck, DUO92105) using an anti-rabbit plus probe and an anti-goat minus probe as according to the manufacturer's instructions. Briefly, cells isolated by dissociation were fixed for immunocytochemistry as above, then blocked with the kit blocking solution for 1 h at room temperature. Antibodies were applied at a ratio of 1:100 with kit antibody diluent solution and incubated overnight at 4°C. Following this, Duolink probes were added to cells at a ratio of 1:5 with kit diluent solution and incubated for 1 h at 37°C in a humidified chamber and excess was washed in kit wash buffer. Cells were incubated in a humidified chamber heated to 37°C in kit solutions for ligation of probes (30 min), and amplification of the signal (100 min), then cells were counterstained with 4'-6-Diamidino-2-phenylindole (DAPI). The plus and minus probes were bound to the anti-WNT3A antibody, and antibody anti-FRZB, respectively. Where the distance between the two bound probes is <40 nm a red signal is generated, which indicates an interaction of proteins of interest [30].

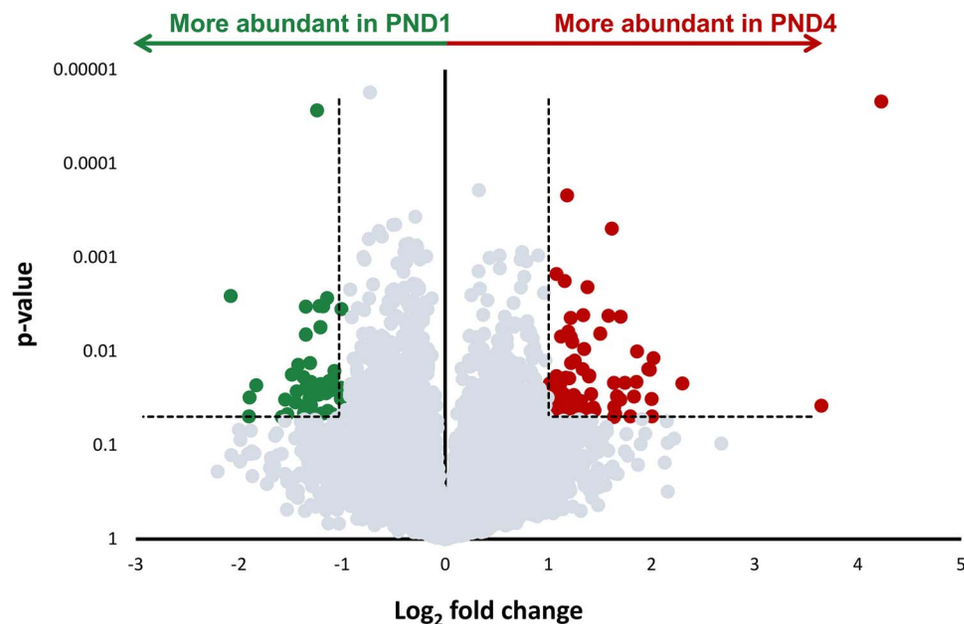
## Results

### Subtle changes in granulosa cell gene expression during primordial follicle activation

To investigate granulosa cell transcriptional changes associated with primordial follicle activation, RNA-seq analysis was performed on large quantities of granulosa cells isolated from primordial (PND1) ovaries or activating and primary follicles (PND4). A comparison between the PND1 and PND4 date sets did not reveal any transcripts with a false discovery rate  $\leq 0.1$ . A total of 11 784 mapped genes were identified from RNA transcripts during bioinformatics processing, including genes characteristic of granulosa cells and primordial follicle activation (Table 2). There was a total of 131 transcripts differentially expressed ( $\log_2$  fold change  $\pm 1$ ;  $P \leq 0.05$ ) between PND1 and PND4 granulosa cells (Figure 1). By comparing the transcripts in the PND1 granulosa cells with those found in the PND4 granulosa cells, 82 RNA transcripts were identified as significantly more abundant whilst 49 transcripts were identified as significantly more abundant in PND4 granulosa cells compared to PND1 (Figure 1); for a complete list of differentially expressed genes, see [Supplementary Material S2](#). This represents a very small (1.1%) change in transcripts from the total number identified between these two fundamental developmental stages.

**Table 2.** Average number of fragments per kilobase million (FPKM) for notable genes detected via RNAseq from granulosa cells isolated at PND1 and PND4.

Gene symbol	Gene name	Relevance	Average reads PND1 (FPKM)	Average reads PND4 (FPKM)
<i>Inha</i>	Inhibin alpha	Marker of granulosa cells	44.4	58.9
<i>Gata4</i>	GATA binding protein 4	Marker of granulosa cells	71	64.4
<i>Amb</i>	Anti-Mullerian hormone	Marker of mature granulosa cells	0.06	3.4
<i>Akt1</i>	AKT serine/threonine kinase 1	Promotes primordial follicle activation	130.1	138.8
<i>Bmp4</i>	Bone morphogenetic protein 4	Promotes primordial follicle activation	12.6	13.4
<i>Lbx8</i>	LIM Homeobox 8	Suppresses primordial follicle activation	2	0.9

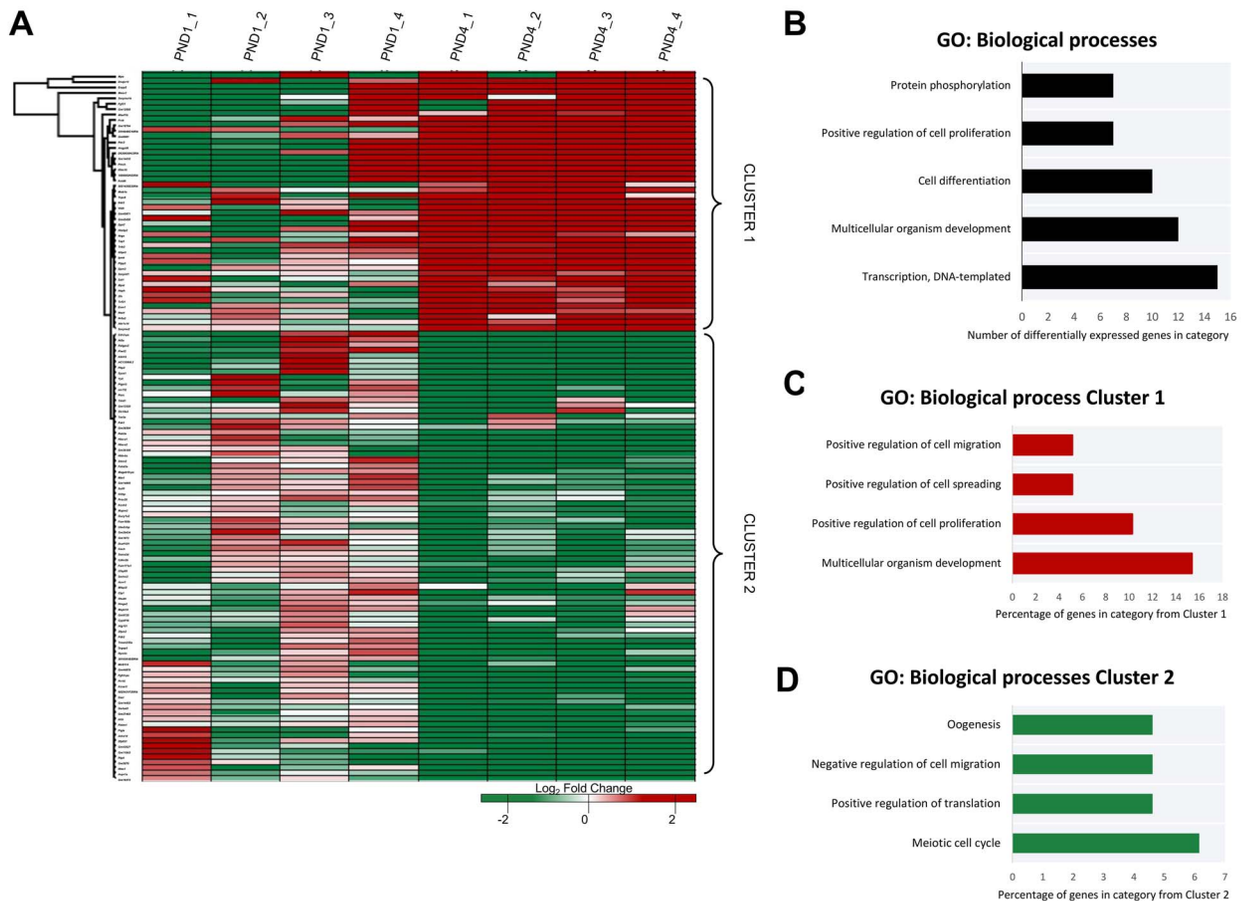
**Figure 1.** Differential gene expression profiling of neonatal mouse granulosa cells. The volcano plot indicates the differential expression between RNA transcripts from postnatal day 1 (PND1), and postnatal day 4 (PND4) granulosa cells ( $n=4$  replicates from each group). Mapped genes expressed in higher abundance in PND1 granulosa cell RNA transcripts compared to PND4 granulosa cells are colored green. Genes expressed in higher abundance in PND4 granulosa cells compared to PND1 are colored red. Dashed line indicates the threshold for differential expression; genes with high statistical significance ( $P \leq 0.05$ ), and log<sub>2</sub> fold change of greater than  $\pm 1$ . Points that fall below these thresholds are colored grey.

A heat map was generated to illustrate the expression (normalized to the average FPKM of the PND1 samples) of each gene within the threshold of significantly differentially expressed genes across each of the sample replicates (Figure 2A). The top biological process occurring in differentially expressed genes were identified using gene ontology (GO) tools (Figure 2B). These processes are reflective of the dynamic events occurring before and during a mass wave of primordial follicle activation, namely transcription (GO:006351), cell differentiation/proliferation (GO:0030154/GO:0008284), phosphorylation (GO:0006468), and genes relating to early development (GO:0007275). Two clusters of differentially abundant genes are apparent from the heatmap; Cluster 1 comprising of transcripts that are more abundant in the PND4 granulosa cells compared to the PND1, and Cluster 2 of transcripts with a decreased abundance in PND4 relative to PND1 granulosa cells. Within Cluster 1, among the top GO annotation of biological processes in terms of the percentage of genes represented within the dataset (Figure 2C) are those associated with development and growth; multicellular organism development (GO:0007275), positive regulation of cell proliferation (GO:0008284), spreading (GO:1900026), and migration (GO:0010634). These processes signify activation occurring,

which is governed by the upregulation of signaling pathways encompassing both the activation and repression of genes and proteins, while granulosa cells differentiate from their squamous to cuboidal structure [6, 11], as is reflective of ovarian events at this time. Cluster 2 (Figure 2D) contains notable biological processes: meiotic cell cycle (GO:0051321), negative regulation of cell migration (GO:0030336), oogenesis (GO:0048477), and positive regulation of translation (GO:0045727). These biological processes are indicative of primordial follicles, known to be regulated by genes associated with development of the ovary [16] that establish the ovarian reserve after germ cell nest breakdown, and are the exclusive follicle type in the ovaries at PND1 [13, 31].

### Validation of differentially expressed genes

To verify the differential gene expression between primordial granulosa cells and granulosa cells undergoing primordial follicle activation in light of the false discovery rate, eight candidate genes were selected for validation of RNA-seq data using RT-qPCR. Genes above the threshold of significance for differential expression were chosen, while also selecting genes with varying degrees of differential



**Figure 2.** Gene expression clustering of neonatal mouse granulosa cells. (A) A heatmap differential expression within the defined threshold ( $P \leq 0.05$ ,  $\log_2$  fold change of  $\pm \geq 1$ ,  $n = 131$ ) depicts the consistency of expression of mapped transcripts identified as having significant between biological replicates. Different genes are represented in different rows, and different replicates in different columns. Expression values (fragments per kilobase million) are represented as a color scale from green for lower expressions to red for higher expressions and normalized to the average FPKM for the PND1 group. Differentially expressed genes were annotated via gene ontology (GO) analysis in DAVID (v6.8). (B) GO information of the biological process categories with highest count of genes from the differentially expressed genes. Clusters of differential expression were identified from the heat map and annotated via gene ontology (GO) analysis in DAVID (v6.8). (C) GO information of four biological process categories from Cluster 1 are presented as the percentage of genes from within that cluster belonging to a given category, and similarly (D) the GO information of biological process categories from Cluster 2 for four notable categories with the greatest percentage. For full list of DAVID GO terms and identifiers returned for this dataset, see [Supplementary Material S3](#).

expression within the threshold. Four genes decreased abundance in PND1 (primordial) granulosa cells compared to PND4 (activating) granulosa cells (frizzled related protein [*Frzb*], transcription factor 21 [*Tcf21*], discoidin domain receptor 2 [*Ddr2*], and zinc finger protein x-linked [*Zfx*]), and conversely, four genes exhibiting increased abundance in PND1 granulosa cells relative to PND4 (Interferon-induced transmembrane protein 1 [*Ifitm1*], Family with sequence similarity 171 member A1 [*Fam171a1*], Cyclin-dependent kinase inhibitor 2B [*Cdkn2b*], arginine vasopressin receptor 1A [*Avpr1a*]), see Table 3. Experiments were performed on at least six pooled biological replicates ( $n = 4-15$  animals per sample). The transcript encoding the 60S ribosomal protein L19 (*Rpl19*) was used as an endogenous control in all RT-qPCR analyses and confirmed the differential expression of all eight candidate genes while following a parallel trend to the RNA-seq data (Figure 3). Six out of the eight genes validated remained significant ( $P \leq 0.05$ ) following Bonferroni correction for multiple comparisons. Collectively, these results validate the accuracy our RNA-seq dataset. Importantly this outcome

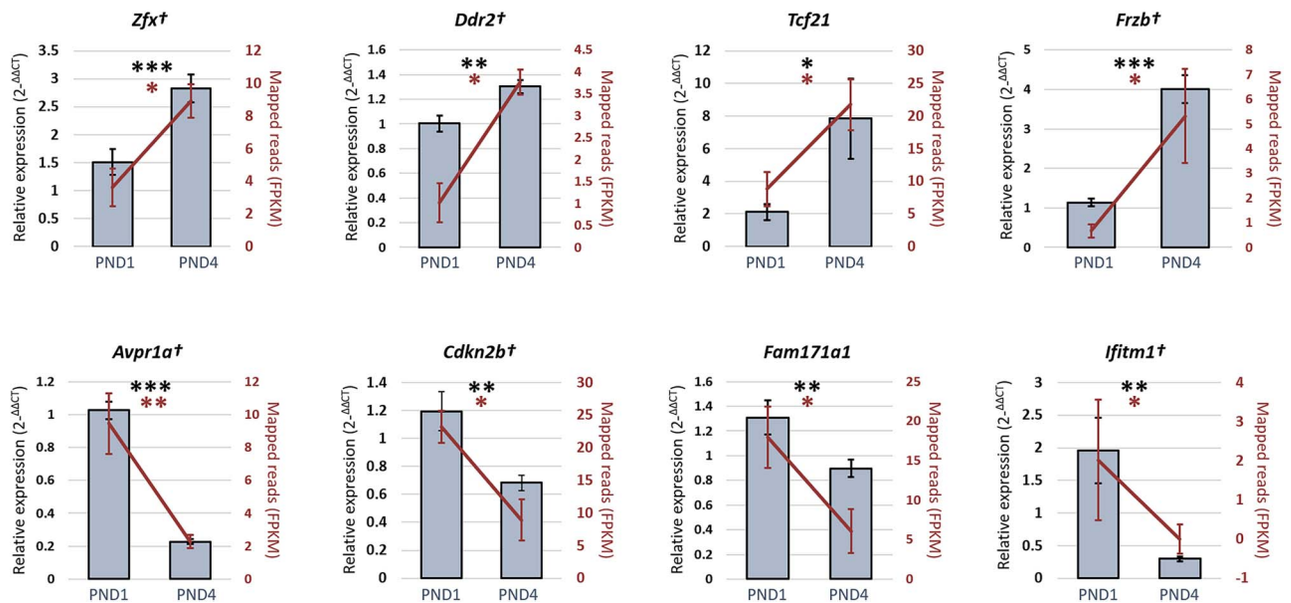
strengthens our justification for the use of this valuable dataset to explore the differential temporal gene expression between mouse neonatal granulosa cells, representative of primordial and activating subtypes.

### Validating protein expression in the mouse ovary

Following gene expression validation, the functional protein expression of notable genes from the RNA-seq dataset was investigated. Through utilizing the Ingenuity Pathway Analysis (IPA) network analysis feature, a number of genes from the dataset were identified to encode proteins that interact with molecules demonstrated to be involved in primordial follicle activation. A proportion of the IPA-generated network map is presented in Figure 4 and identifies six molecules from the dataset interacting through binding or regulation and includes two of those validated in via RT-qPCR (FRZB, and ZFX). Primordial follicle activation component TGFBR from the dataset was also linked within the network. To explore a potential

**Table 3.** Targets selected for validation of RNAseq.

Gene symbol	Absolute fold change	Abundance in PND4 relative to PND1	P-value	Gene name
<i>Frzb</i>	3.56	Increased	0.023	Frizzle-related protein
<i>Tcf21</i>	2.7	Increased	0.014	Transcription factor 21
<i>Ddr2</i>	2.59	Increased	0.019	Discoidin domain receptor 2
<i>Zfx</i>	2.5	Increased	0.027	Zinc finger protein x-linked
<i>Ifitm1</i>	4.05	Decreased	0.012	Interferon-induced transmembrane protein1
<i>Fam171a1</i>	3.95	Decreased	0.016	Family with sequence similarity 171 member A1
<i>Cdkn2b</i>	3.91	Decreased	0.016	Cyclin-dependent kinase inhibitor 2B
<i>Avpr1a</i>	2.99	Decreased	0.004	Arginine vasopressin receptor 1A



**Figure 3.** The RT-qPCR validation of differentially expressed genes within neonatal mouse granulosa cells. In verifying the RNA-seq data (represented on the secondary y-axis by the fragment per kilobase of exon per million, FPKM, for each mapped read), eight genes that displayed significantly different levels of expression in PND1 granulosa cells compare to PND4 granulosa cells (see Table 3) were selected for orthogonal validation using RT-qPCR. Validation experiments were performed in triplicate using  $\geq$  six distinct pools of biological samples ( $n=4-15$  animals per sample), statistical significance was investigated via a student *t*-test. The 60S ribosomal protein L19 gene was employed as an endogenous control to normalize the expression levels of target genes. Six out of 8 genes remained significant after Bonferroni correction for multiple comparison and are indicated by †. Data are presented as mean  $\pm$  SEM. \* $P < 0.05$ , \*\* $P < 0.01$ , \*\*\* $P < 0.001$ .

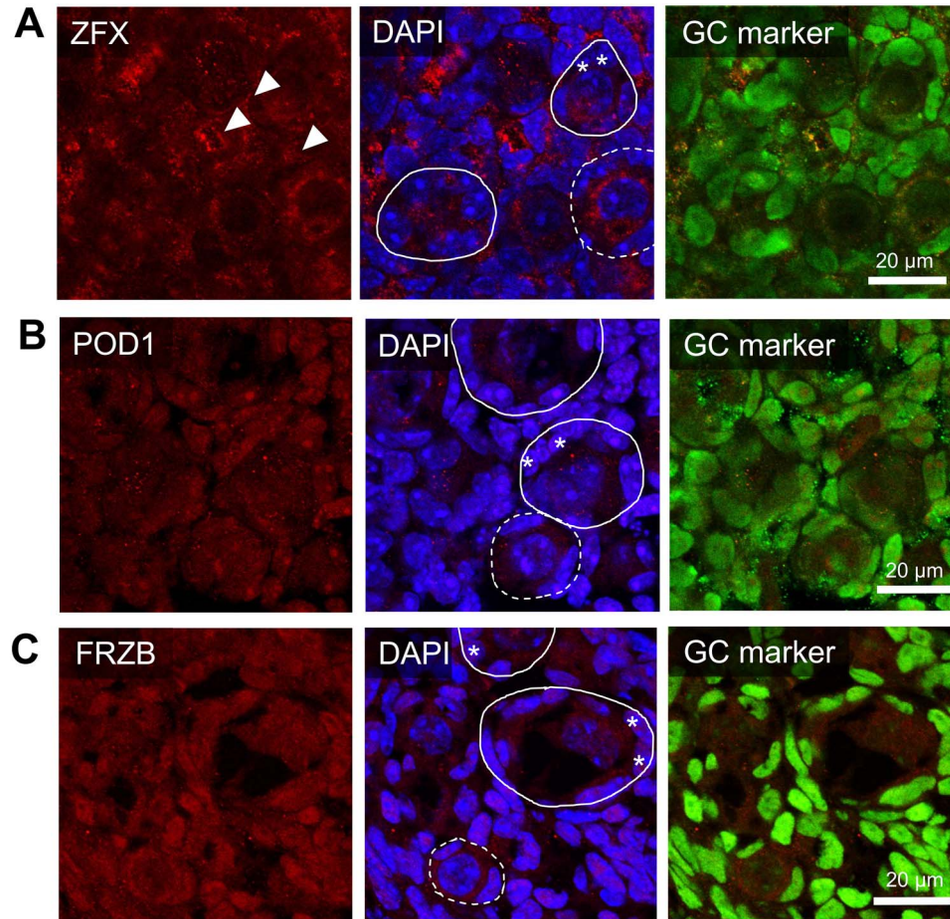
relationship with primordial follicle activation, the two molecules from the pathway analysis network that had been validated at the gene expression level were selected for *in situ* immunolocalization in neonatal mouse ovaries (Figure 5), granulosa cells staged according to morphology, described elsewhere [32]. Podocyte-expressed 1 (POD1) did not feature in the IPA-generated network, but was also selected for further investigation, as the expression of its gene (*Tcf21*) increased 3.7-fold ( $P < 0.05$ ) in PND4 granulosa cells compared to PND1 granulosa cells as determined by RT-qPCR. The relationship of POD1 to primordial follicle activation that made it of interest is depicted in addition to the network map in Figure 4. ZFX protein was localized to the oocyte cytoplasm of both primordial and primary follicles and was also observed in the extracellular space proximal to the granulosa cells of follicles (Figure 5A). There was some expression of POD1 in all granulosa cells, but POD1 protein was more intensely localized in the nuclei of some granulosa cells (Figure 5B). Oocytes were also observed to have punctate expression

of POD1. FRZB was expressed in the oocyte cytoplasm (Figure 5C) and exhibited differential localization in granulosa cells.

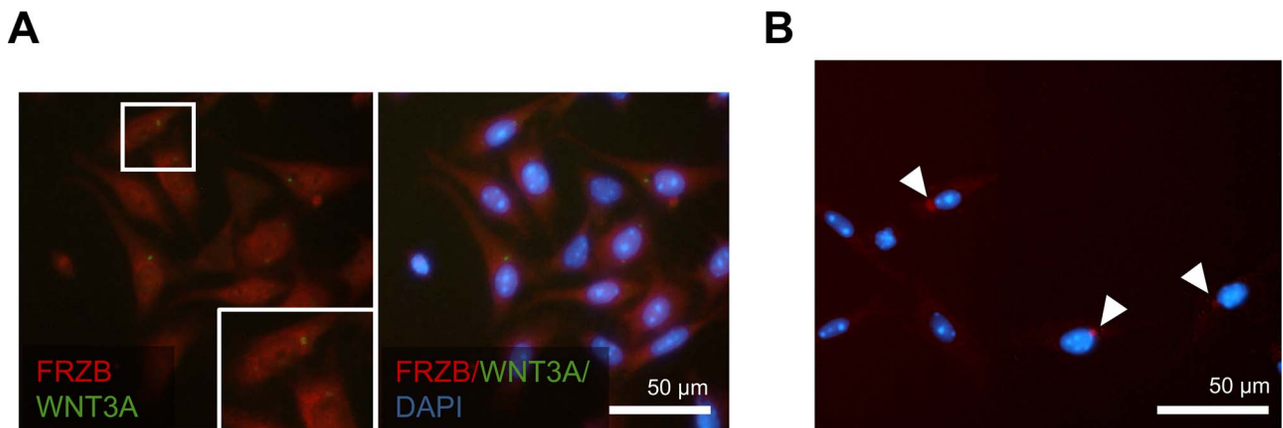
The expression and localization of FRZB was further investigated in neonatal mouse ovaries and granulosa cells to identify possible links to the process of primordial follicle activation considering it was placed among the top known genes in differential expression due to its 3.5-fold increase ( $P < 0.001$ ) in gene expression in PND4 granulosa cells compared to PND1, and its protein localization to granulosa cells (Figure 5C). Additionally, FRZB was of interest due to its known interaction with 35% of the molecules within the IPA-generated network (Figure 4), and its canonical role as a Wnt-pathway inhibitor.

The expression of FRZB was investigated in granulosa cells and to validate its role in the Wnt pathway via WNT3A, a known suppressor of primordial follicle activation [33] was also investigated. WNT3A protein localization has not previously been reported in neonatal ovaries, only its gene expression [34].





**Figure 5.** Expression and localization of proteins of interest within the PND4 ovary. The *in situ* immunofluorescent expression of three proteins of interest (A-C) were explored in the neonatal mouse ovaries and were co-localized with a nuclear marker (DAPI, blue), a granulosa cell marker (GATA4 or FOXL2, green), and either (A) ZFX (B) POD1 or (C) FRZB in red. Representative images from PND4 selected as they include populations of primordial, activating and primary follicles. Representative images are indicative of  $n=4-6$  biological replicates of both PND1 and PND4 performed in triplicate. Images taken at 60 $\times$  magnification, scale bars represent 20  $\mu\text{m}$  with dotted circles outlining primordial follicles, solid lines outlining activating or primary follicles. Arrows indicate extracellular staining regions; asterisks indicate activating granulosa cells.



**Figure 6.** FRZB interacts with WNT3A in granulosa cells. (A) Immunocytochemistry of granulosa cells isolated from a post-natal day (PND) 4 mouse ovary. Frzb (red) co-localized with Wnt pathway activator WNT3A. Insets show zoomed image of boxed area. (B) Duolink proximity ligation assay of granulosa cells isolated from PND4 mouse ovary, red dot indicates proteins are interacting; representative images are indicative of  $n=3$  biological replicates performed in triplicate. Images taken at 40 $\times$  magnification. Scale bar represents 50  $\mu\text{m}$ .

of the current dataset, we interpret our data with caution and validated changes biologically, with RT-qPCR experiments mirroring expression changes of the RNA-seq. The advantage of this study over previous studies characterizing gene expression characterization in young stage follicles is the targeted focus on the granulosa cell, and the use of relatively large populations of cells to more accurately reflect the heterogeneity within these cells. Where granulosa cell transcriptome studies have been performed on human samples, they have been small in sample size and/or utilize microarray assays [44–47], both of which have limitations in identifying unique genes and accounting for sample heterogeneity. Furthermore, this study presents a granulosa cell-focused investigation of primordial follicle activation in mice. Previous mouse model studies using bulk RNA-seq to capture primordial follicle activation or early ovarian time points analyze a compendium of cell types and do not focus specifically on the granulosa cell to granulosa cell transition [48, 49]. Single cell sequencing has, so far, been underutilized in the study of primordial follicle activation and lacking depth in the analysis of granulosa cells during this important physiological event [39].

In the PND4 mouse ovary, POD1 protein was expressed in the nuclei of granulosa cells, consistent with previous reports of POD1 expression increasing postnatally in the mouse [50]. This study provides further evidence, demonstrating that the expression of *Tcf21* also increases by a factor of 3.7 in granulosa cells from PND4 ovaries. This group also identified POD1 as a negative regulator of steroidogenic factor-1 (SF-1), an important regulator of granulosa cell regulated ovary development [51], suggesting a possible role in the differentiation of granulosa cells. Further studies focusing on whether there is a link between POD1-induced repression of SF-1 and activation of primordial follicles would be valuable, as the SF-1 gene *Nr5a1* was also detected in the granulosa cells of our RNAseq experiment (for entire dataset, see GEO accession No. GSE162927).

ZFX is a transcription factor previously speculated to be involved in POI as it occurs in the POI-critical region of the X chromosome [52], and early research in mice showed a reduced ovarian reserve when *Zfx* was deleted, but the mechanism by which this occurs is not known [53]. The extracellular expression of ZFX in the ovaries was notable considering that in other somatic cells, ZFX is associated with promoting cell proliferation through cell cycle control [54, 55], but was not in the nucleus of activating granulosa cells, which are known to proliferate. Our results demonstrate that *Zfx* expression is upregulated in granulosa cells from PND4 mice during primordial follicle activation. Our protein localization studies indicate that ZFX is predominately expressed in oocyte and extracellular space suggesting the subcellular reorganization of the transcription factor. The bidirectional communication between oocytes and granulosa cells is a critical feature of folliculogenesis and follicular survival [7, 56]. These findings provide a renewed advocacy for investigating the role of ZFX to gain insight into primordial follicle activation and POI in the context of ZFX and oocyte-granulosa cell–cell communications.

An important finding in this study was the 3.5-fold increase in mRNA expression of *Frzb* in PND4 granulosa cells indicative of activating granulosa cells. Importantly, FRZB directly and indirectly has association with or influences cellular function of molecules known to be involved in primordial follicle activation including Wnt ligands like WNT3A, in addition to ERK1/2, and the TGF $\beta$  superfamily [5, 33, 57], providing further evidence to investigate its role in primordial follicle activation. This study also provides compelling evidence for an interaction between FRZB and WNT3A

in granulosa cells. This is consistent with previous findings identifying two sites on the FRZB protein on which WNT3A may bind [58]. WNT3A has been implicated in the maintenance of primordial follicle quiescence through FOXO3 [33]. When *Wntless* (gene required for secretion of all WNT ligands) was deleted in granulosa cells with an *Sf1-cre* mouse line, the transition from squamous to cuboidal granulosa cell morphology, a hallmark of primordial follicle activation, was impaired and coupled with the lack of translocation of FOXO3 [59]. This study provides novel evidence linking FRZB to the potential Wnt signaling cascade involved in primordial follicle activation through its ability to bind to WNT3A in the granulosa cells. A recent human granulosa cell transcriptome study observed a downregulation of Wnt family signaling in granulosa cells from primordial follicles compared to primary follicle granulosa cells [38], indicating that WNT3A, and by extension FRZB, may have a conserved role in the human ovary. Taken together with our findings, this emphasizes the need for further studies to determine the role of FRZB in primordial follicle activation, including a detailed investigation into a potential FRZB-WNT3A-FOXO3a relationship.

One limitation of the method of granulosa cell isolation to model processes occurring during primordial follicle activation is evident in the functions related to development (multicellular organism development), and genes observed in PND1 samples (such as *Dazl*). This is likely to be due to the remnants of naive pre-granulosa cells adjacent to germ cell nests, the structural precursors of primordial follicles. Germ cell nests begin to break down in the hours post-birth in mice, and contain pre-granulosa cells surrounding clusters of multiple germ cells, eventually giving rise to the single-oocyte encapsulated primordial follicle [31], thus ovaries from a PND1 (~24 h post birth) mice may contain residual germ cell nests.

This study has utilized a method for the isolation of mouse granulosa cells for the study of primordial follicle activation by taking advantage of the initial wave of primordial follicle activation in neonatal ovaries. The transcriptome of primordial and activating granulosa cells presented in this study has identified a range of factors previously associated with primordial follicle activation, which signify that future research could be well-spent redoubling efforts into extricating the roles of these factors, such as SF-1, ZFX, and WNT3A, and their up- and down-stream effectors. Co-immunoprecipitation assays for example, would be valuable to validate the binding between WNT3A and FRZB, in addition to rigorous assessment of the proteins via quantifying localization or via proteomics and phosphoproteomics. The findings in this study are enhanced by the validation of transcriptome data via RT-qPCR, confirming the dramatic changes in gene expression of granulosa cells during primordial follicle activation. Additionally, novel evidence is presented for Wnt antagonist *Frzb* as a potential regulator among the genetic network controlling primordial follicle activation through its protein interaction with murine primordial follicle suppressor WNT3A in granulosa cells. Understanding the factors, which dictate the maintenance of primordial follicle quiescence or their activation into primary follicles will inform our knowledge of early follicle development and may lead to future diagnostic methods and treatment regimens for women with infertility and premature ovarian insufficiency.

## Supplementary material

Supplementary material is available at *BIOLRE* online.

## Author contributions

EAF collected the samples, performed all laboratory experiments (excluding RNA-seq, library prep and mapping), and in silico data analysis. EAF contributed to study design, wrote the manuscript, analyzed the data and created the figures. ERF contributed to sample preparation and collection, and prepared samples for immunolocalization experiments. SDR, EAM, and ELB contributed to interpretation of the data, the preparation, and critical feedback of the manuscript. EAM funded the RNA-seq and bioinformatics. JMS conceived and designed the study. JMS funded the remainder of the study, contributed to data analysis and manuscript preparation and had the final decision on the manuscript.

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## Conflict of interest

The authors declare that there is no conflict of interest.

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