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Survivin is essential for fertile egg production and female fertility in mice

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Survivin is the smallest member of the inhibitor of apoptosis protein (IAP) family and acts as a bifunctional protein involved in mitosis regulation and apoptosis inhibition. To identify the physiological role of Survivin in female reproduction, we selectively disrupted *Survivin* expression in oocytes and granulosa cells (GCs), two major cell types in the ovary, by two different Cre-Loxp conditional knockout systems, and found that both led to defective female fertility. *Survivin* deletion in oocytes did not affect oocyte growth, viability and ovulation, but caused tetraploid egg production and thus female infertility. Further exploration revealed that Survivin was essential for regulating proper meiotic spindle organization, spindle assembly checkpoint activity, timely metaphase-to-anaphase transition and cytokinesis. Mutant mice with Survivin depleted in GCs showed reduced ovulation and subfertility, caused by defective follicular growth, increased follicular atresia and impaired luteinization. These findings suggest that Survivin has an important role in regulating folliculogenesis and oogenesis in the adult mouse ovary. *Cell Death and Disease* (2014) **5**, e1154; doi:10.1038/cddis.2014.126; published online 27 March 2014

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In mammals, ovarian folliculogenesis is a process regulated by complex networks composed of various endocrine, paracrine and autocrine factors that interact coordinately.¹ A follicle is composed of three distinct cell types: an oocyte and two different kinds of somatic cells, granulosa cells (GCs) and theca cells. The oocyte is continuously arrested at the germinal vesicle (GV) stage during follicle development, and the fully grown oocyte in the mature follicle begins the maturation process upon stimulation by the gonadotropin surge, as indicated by a significant event: GV breakdown (GVBD), followed by spindle organization and chromosome alignment as an important process for correct chromosome segregation. Following the small first polar body extrusion (PBE) produced by unequal cytokinesis the mature eggs are now ready for fertilization.^{2,3}

Successful oogenesis and oocyte maturation rely on close cooperation and communication between the oocyte and surrounding GCs. Receptors expressed on GCs such as the FSH receptor (*Fshr*) and LH receptor (*Lhcgr*), and factors secreted from GCs such as estradiol and insulin-like growth factor (IGF) are proved to be essential for folliculogenesis.⁴ Moreover, apoptosis of GCs was observed to occur earlier than that of oocytes and theca cells in atretic follicles, suggesting their vital role in folliculogenesis and follicle atresia.^{4–6}

Survivin, the smallest member of the inhibitors of apoptosis protein (IAPs) family, is a 16.5-kDa protein and contains a single repeat of BIR domain that is essential for the caspaseinhibitory function.⁷ After years of research, Survivin has been shown to have multiple roles that are not limited to apoptosis inhibition but also to regulation of the mitotic spindle checkpoint. On one hand, Survivin is proved to be a caspase inhibitor given its BIR domain, which is common to the other IAP family members. Studies showed that Survivin, associated with the hepatitis B X-interacting protein (HBXIP) or X-linked IAP (XIAP), inhibited activation of Caspase-9.8 Survivin also acts as a downstream target protein involved in several important signaling pathways, the PI3K/Akt pathway, mTOR pathway, ERK pathway, tumour suppressors (p53, PTEN) and oncogenic (Ras, Bcl-2) signaling pathways.⁹ On the other hand, Survivin together with INCENP, Borealin and Aurora B form the chromosomal passenger complex (CPC) for regulation of cell division.^{10,11} During mitosis, Survivin locates to centromeres during prophase/metaphase, relocates to the spindle mid-zone during anaphase/telophase and disappears at the end of telophase.¹² Survivin is ubiquitously expressed during embryonic and fetal development and has been proved to have roles in both neurogenesis and hematopoiesis;^{13,14} it is also expressed in various cancers.

Keywords: folliculogenesis; apoptosis; granulosa cell; meiosis; oocyte; Survivin

Abbreviations: GCs, granulosa cells; GVBD, germinal vesicle breakdown; SAC, spindle assembly checkpoint; APC/C, anaphase-promoting complex/cyclosome; PBE, polar body extrusion; MCC, mitotic checkpoint complex; COC, cumulus cell–oocyte complex; CL, corpora lutea; CC3, cleaved caspase 3; TUNEL, terminal deoxynucleotide transferase-mediated deoxyuridine triphosphate nick end labeling

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In this study, we generated mutant mice with specific deletion of *Survivin* in oocytes and GCs to investigate the function of Survivin within ovarian follicles *in vivo*. For the first time, we report here that Survivin is expressed in both mouse oocytes and GCs and it is essential for female fertility. Oocyte-specific deletion of *Survivin* did not affect oocyte survival and ovulation, but led to female infertility due to tetraploid eggs derived from failure of cytokinesis and first polar body emission. *Survivin* mutation in oocytes also caused misorganized spindles and a dysfunctional spindle assembly checkpoint (SAC). In addition, deletion of *Survivin* in GCs caused compromised follicular development, reduced ovulation and impaired luteinization, which led to defective ovarian function and female subfertility.

Results

Oocyte-specific/GC-specific disruption of the *Survivin* **gene.** The mutant mice with oocyte-specific and GC-specific disruption of the *Survivin* gene, in which exon II of the *Survivin* gene was respectively targeted in oocytes and GCs,¹⁵ were generated by crossing *Sur^{F/F}* mice with transgenic mice expressing *Gdf9* or *Cyp19* promotermediated Cre recombinase. Survivin always expressed in oocytes and GCs in the *Sur^{F/F}* ovary (Figure 1a). Western blot analysis showed in *Sur^{F/F}*; *Gdf9-Cre* mouse oocytes and *Sur^{F/F}*; *Cyp19-Cre* mouse GCs, the Survivin protein was depleted (Figure 1b). Immunofluorescent analysis of oocytes from *Sur^{F/F}*; *Gdf9-Cre* female revealed loss of Survivin localization on kinetochores (Figure 1c).

Survivin is essential for female fertility. Breeding assays showed that female $Sur^{F/F}$; *Gdf9-Cre* mice were completely infertile, while the $Sur^{F/F}$; *Cyp19-Cre* mice were evidently subfertile and gave birth to about 70% fewer pups than control mice (Figure 1d). The decreased number of pups born in $Sur^{F/F}$; *Cyp19-Cre* mice appeared to be related to decreased ovulation as seen in the result of the super-ovulation assay (Figure 1e).

Depletion of Survivin causes spindle defects, chromosome misalignment, premature anaphase entry and first PBE failure of oocytes. To investigate the primary defect leading to female infertility after oocyte-specific Survivin depletion, we first performed superovulation. We found that oocvtes were ovulated normally, while no polar body was extruded in SurF/F; Gdf9-Cre mice (Figure 2a). To further understand the defects in oocytes caused by Survivin deletion, we employed immunofluorescent staining to observe spindles and chromosomes in the oocytes. It was observed that a number of misaligned chromosomes existed in superovulated Sur^{F/F}; Gdf9-Cre oocytes (see white arrows in Figure 2b), and the rate of oocytes with misaligned chromosomes was significantly higher than that in SurF/F eggs (92.3 ± 5.1% versus 13.3 ± 8.9%) (Figure 2c). Another interesting phenotype of SurF/F; Gdf9-Cre oocytes is that they displayed defective spindle morphologies (see red arrows in Figure 2b). In many abnormal spindles, bundles of microtubules were seen scattered with chromosomes attached to them (see arrows in Figure 2b'). In addition, we

performed chromosome spreads to address chromosome morphology. As expected, $Sur^{F/F}$; Gdf9-Cre oocytes displayed 40 univalent chromosomes that were twice in number compared with those in $Sur^{F/F}$ oocytes (Figure 2d), indicating that oocytes yielded from female $Sur^{F/F}$; Gdf9-Cre mice all displayed tetraploidy. Ovaries of indicated genotypes were fixed after injection of PMSG (48 h) and hCG (8 h). After H&E staining, we found that oocytes in follicles remained at the MI stage in the $Sur^{F/F}$ group, whereas chromosomes were segregating (Figure 2e, white arrows), or an abnormal polar body was extruding (Figure 2e, black arrow) in $Sur^{F/F}$; Gdf9-Cre oocytes, indicating premature anaphase entry.

The above phenotypes, especially the precocious anaphase entry, were further confirmed by incubating Sur^{F/F}; Gdf9-Cre GV oocytes in M2 medium in vitro. After culture for 2 h, the oocytes that had undergone GVBD were counted and the results showed no obvious difference in GVBD rates between $Sur^{F/F}$ and $Sur^{F/F}$; Gdf9-Cre oocytes (88.4 ± 2.0%) versus 83.3 ± 0.9%) (Supplementary Figure S1A). As was the case in *in vivo* experiments, we observed a significantly lower PBE rate in Sur^{F/F}; Gdf9-Cre oocytes (2.9 ± 3.9% versus 75.9 ± 0.7%) (Supplementary Figure S1B) after culture for 12 h. It was observed that multipolar spindles and misaligned chromosomes dominantly existed in Sur^{F/F}; Gdf9-Cre oocytes after culture (Supplementary Figure S1C). There was a significant increase in frequency of abnormal bipolar spindles $(82.1 \pm 6.2\% versus 14.2 \pm 7.2\%, Supplementary Figure S1D)$ and misaligned chromosomes $(97.4 \pm 3.4\%)$ versus 20.1 ± 3.2%, Supplementary Figure S1E). In SurF/F; Gdf9-Cre oocvtes, abnormal spindles contained more than two poles and scattered poles with different orientations, which was confirmed by staining of y-tubulin (see arrows in Supplementary Figure S1F). To address whether homologous chromosome segregation prematurely took place in Sur^{F/F}; Gdf9-Cre oocytes, we performed chromosome spreading and found prematurely segregated homologs after culture for 8h (Supplementary Figure S1G, left). Univalent chromosomes were found in oocytes from both mutant genotypes after culture for 12 h (Supplementary Figure S1G, right), suggesting that homologous chromosomes were precociously segregated in Sur^{F/F}; Gdf9-Cre oocytes.

Depletion of Survivin led to failure of spindle checkpoint protein localization to kinetochores, and premature activation of anaphase-promoting complex/cyclosome. Considering the phenotypes of chromosome misalignment and precocious chromosome segregation, we next asked whether the SAC is under proper control in *Sur^{F/F}*; *Gdf9-Cre* oocytes. In SurF/F oocytes, there was clear localization of Bub3 and BubR1 (two SAC proteins)^{16,17} at kinetochores. In sharp contrast, there was no visual centromeric localization of both SAC proteins in SurF/F; Gdf9-Cre oocytes (Figures 3a and b), suggesting that Survivin was required for the localization of SAC proteins to kinetochores. Bub3, BubR1, Mad2 and Cdc20 form the mitotic checkpoint complex (MCC) that inhibits the ability of Cdc20 to activate the anaphasepromoting complex/cyclosome (APC/C) in mitosis¹⁸ and a similar mechanism was determined for meiosis.19 For the evaluation of premature activation of APC/Ccdc20, we used time-lapse live imaging to observe the degradation of Cyclin



Figure 1 Survivin expression in oocytes and granulosa cells is essential for female fertility. (a) Expression of Survivin protein in ovaries of WT D15 and D21 mice. (b) Verification of Survivin deletion in oocytes and granulosa cells from mice with indicated genotypes. GV oocytes were collected from minced ovaries and 200 oocytes were used for each sample. GAPDH was used as a loading control. (c) Localization of Survivin (green) in $Sur^{F/F}$; *Gdf9-Cre* and $Sur^{F/F}$ oocytes. Oocytes at 8 h of culture were fixed for immunofluorescent staining. Red, DNA. Bar = 20 μ m. (d) Infertility of the female $Sur^{F/F}$; *Gdf9-Cre* mice and subfertility of the female $Sur^{F/F}$; *Cyp19-Cre* mice. Continuous breeding assessment showed the cumulative number of progeny per female mouse. (e) Lack of or defective ovulation in $Sur^{F/F}$; *Cyp19-Cre* mice. Oocytes were collected at 13 and 16 h of hCG treatment, respectively (n = 6 for each genotype in each group). *P < 0.05

B1 which is one of the key substrates of APC/Ccdc20. As shown in the movies, we found that Cyclin B1 started to be degraded in *Sur^{F/F}*; *Gdf9-Cre* oocytes at about 2 h ahead of that in *Sur^{F/F}* oocytes (Figures 3c and d, also see Supplementary Movie 1A and B in Supplementary Material), which was confirmed by the quantitation of total fluorescence of Cyclin B1-GFP in both the genotypes (Figure 3e). This result validated the finding that deletion of *Survivin* resulted in failure of SAC assembly to kinetochores for inhibition of Cdc20, which triggers precocious anaphase I onset during the first meiosis.

Depletion of *Survivin* caused cytokinetic abscission failure. To explore the underlying reason for PBE failure, we performed live oocyte imaging to analyze spindle dynamics (Figure 4) and F-actin (Figure 5) during oocyte maturation. In *Sur^{F/F}*; *Gdf9-Cre* oocytes, it was shown that the midbody microtubules were absent at telophase I (red arrow, Figure 4b, also see Supplementary Movie 2B in Supplementary Material) when cytokinesis was initiated, which led to unequal chromosome segregation (white arrows, Figures 4b and 5b, also see Supplementary Movies 2A and 3B in Supplementary Material). In addition, the rate of transient PBE (membrane protrusion for accuracy) at indicated time points showed that the transient PBE rate of Survivin-deficient oocytes reached the peak at 8 h of culture and all the extruding polar bodies were abnormal, being smaller than the normal size (red and white arrows. Figure 4c), and further retracted within 2h (Figure 4d, 10 h). In SurF/F oocytes, the contractile ring was present throughout the PBE process (red arrows, Figure 5a, also see Supplementary Movie 3A in Supplementary Material). Whereas in Sur^{F/F}; Gdf9-Cre oocytes, the assembly of F-actin at the midbody (contractile ring) was not disturbed but disappeared soon along with the retraction of the polar body (red arrows, Figure 5b, also see Supplementary Movie 3B in Supplementary Material), suggesting abscission failure. Immunofluorescent staining showed complete absence of the midbody spindle at telophase in SurF/F; Gdf9-Cre oocytes (Figure 5c). In meiosis, it has been reported that Aurora C instead of Aurora B has pivotal roles in oocyte maturation^{20,21} and Aurora B is responsible for npg



Figure 2 Deletion of *Survivin* disturbed oocyte maturation *in vivo*. (a) Percentages of eggs with first polar body extrusion from superovulated $Su^{F/F}$ and $Su^{F/F}$; *Gdf9-Cre* females, respectively, after injection of PMSG (48 h) and hCG (12–14 h). Red arrows, first polar body. Bar = 100 μ m. **P*<0.0001. (b and b') Immunofluorescent staining of superovulated eggs. White arrows showed misaligned chromosomes and red arrows showed abnormal spindle poles in (b). Yellow arrows showed bundles of microtubules in (b'). Red, DNA; Green, α -tubulin; PB, polar body. Scale bars: 20 μ m in (b) (Merge); 10 μ m in (b) (DNA and Tubulin) and (b'). (c) Rates of oocytes with misaligned chromosomes in superovulated $Su^{F/F}$; *Gdf9-Cre* oocytes, respectively. **P*<0.0001. (d) Chromosome spreads of superovulated oocytes from the indicated genotypes. Polar body (PB) and the numbers of chromosomes were indicated. Bar = 20 μ m. (e) Hematoxylin and eosin-stained ovary sections from $Su^{F/F}$; *Gdf9-Cre* females. Ovaries of the indicated genotypes were fixed after injection of PMSG (48 h) and hCG (8 h). Only oocytes and cumulus cells of the sections were shown. Yellow arrow showed membrane protrusion and white arrows showed the segregating chromosomes

abscission in mitosis.²² Here, we found that the assembly of Aurora C (Myc-conjugated) in the contractile ring was disturbed in *Sur^{F/F}*; *Gdf9-Cre* oocytes (Figure 5d). Taken together, as shown in Figure 5e, oocyte-specific deletion of *Survivin* impaired the assembly and maintenance of midbody microtubules that ensures proper chromosome segregation, and caused cytokinetic abscission failure, followed by retraction of the extruding polar body.

Disruption of *Survivin* in GCs causes defects in folliculogenesis, ovulation and luteinization. Since the *Sur* ^{*F*/*F*}; *Cyp19-Cre* mice could barely ovulate oocytes in the superovulation assay, we then examined whether folliculogenesis was defective. At 24 h of PMSG treatment, the secondary follicles were growing fast, filled with rapidly increasing GCs in control mice; in contrast, unhealthy follicles with sparsely distributed GCs were observed in *Sur*^{*F*/*F*}; *Cyp19-Cre* mice (Supplementary Figure S2A). This tendency became more obvious after 48 h of PMSG

treatment (Figure 6a). The granulosa layers of *Sur^{F/F}*; *Cyp19-Cre* mice were much thinner than the control. To quantify this discrepancy, the diameter of antral follicles was measured and a 20% decrease in mutant mice was found compared with control mice (Supplementary Figure S2B, right panel). This result was also supported by the smaller size and decreased weight of ovaries from *Sur^{F/F}*; *Cyp19-Cre* mice (Supplementary Figure S2B, left panel).

Typically in mutant mice, there were only 4–5 layers of membrana GCs (black arrow) and only one layer of cumulus cells (yellow arrow); furthermore, many of these GCs displayed pyknotic nuclei that indicated morphological apoptosis (Figure 6a, 20X). To test whether the dominant antral follicles could maintain ovulation, we administered hCG at 48 h of PMSG to simulate ovulation. At 8 h of hCG priming, the cumulus cell–oocyte complex (COC) had undergone expansion in control ovaries as expected; however, there were too few cumulus cells (Supplementary Figure S2C, yellow arrow) to expand and even thinner membrana granulosa



Figure 3 Precocious anaphase initiation during $Sut^{F/F}$; *Gdf9-Cre* oocyte maturation. (**a** and **b**) Immunofluorescent staining showing the localization of spindle assembly checkpoint (SAC): proteins Bub3 (**a**) and BubR1 (**b**) in oocytes at the Pro-MI stage of the indicated genotypes. Green: Bub3 in (**a**), BubR1 in (**b**); Red, DNA. Scale bars: 20 μ m in (**a**) (Bub3), (**b**) (BubR1); 10 μ m in (**a**) and (**b**) (Merge). (**c** and **d**) Dynamics of Cyclin B1-GFP fluorescence during $Sut^{F/F}$ and $Sut^{F/F}$; *Gdf9-Cre* oocyte maturation. Oocytes microinjected with Cyclin B1-GFP mRNA were transferred to the live imaging station after 2 h of incubation. Time points indicated the time lapse from the start of corresponding Supplementary Movie 1A and B. White arrows showed segregated chromosomes in (**a**) and (**b**). Yellow arrow showed polar body in (**a**) and membrane protrusion in (**b**). Bar = 25 μ m. (**e**) Cyclin B1-GFP fluorescence. Arrows indicated the time points of polar body extrusion in $Sut^{F/F}$; *Gdf9-Cre* oocytes

(Supplementary Figure S2C, black arrow) in the preovulatory follicles of mutant ovaries. As the process continued, most COCs of mutant mice failed to ovulate and these follicles were left in the ovaries and turned into atretic follicles (Figure 6b, asterisks).

We further investigated whether the luteinization process was affected at 24 and 48 h of hCG administration when the corpora lutea (CL) of the control mice were well developed (Supplementary Figure S2D; Figure 6c, left panel), and found that luteinization was slower in the mutant mice. Many atretic follicles were not filled with luteinized GCs at 24 h of hCG treatment (Supplementary Figure S2D, right panel, asterisks), and the development of CLs did not complete even after 48 h of hCG administration (Figure 6c, right panel, asterisks). CL counting also confirmed that the numbers of developed CLs displayed significant differences between the Sur^{F/F}; Cyp19-Cre mice and control mice (Supplementary Figure S2E, right panel). The results from comparison of ovary size and weight between the two groups (Supplementary Figure S2E, left panel) may be further supporting evidence for the conclusion that luteinization of mutant mice was impaired. Analysis of serum estradiol and progesterone showed a marked decrease in both hormone levels in comparison with the control mice (Figure 6d).

Survivin deletion led to GCs apoptosis and impaired ovarian function. To characterize the observed changes in Survivin knockout ovaries at the cellular level, we first examined the proliferative rate of GCs in the developing follicles. On the basis of immunostaining for the mitosis marker phosphohistone H3 (pHH3) (Supplementary Figure S3A), it appeared that there was no significant difference in GC proliferation especially in normal secondary follicles between the two groups. To test the apoptosis level in mutant ovaries, we next employed the terminal deoxynucleotide transferase-mediated deoxyuridine triphosphate nick end labeling (TUNEL) assay and immunostaining for cleaved caspase 3 (CC3). The numbers of apoptotic follicles (Figure 7a) and CC3-positive GCs (Figure 7b) were significantly increased in the PMSG-primed Survivin mutant ovaries. When the Survivin knockout GCs were isolated and cultured in vitro for immunofluorescent staining, the number of CC3-positive GCs increased even further (Figure 7c). Western blot analysis showed that the levels of pro-caspase 9 and pro-caspase 3 were reduced in the Survivin mutant GCs, indicating more cleavage activity of these caspases (Supplementary Figure S3B). As shown in Figure 7d, the mutant GCs showed an evident decrease in mRNA expression levels of Fshr and Lhgcr. Moreover, we examined the expression of aromatase (Cyp19a1) and found that it was



Figure 4 Impaired midbody in $Sur^{F/F}$; *Gdf9-Cre* occytes at late meiosis I. (**a** and **b**) Dynamics of spindle and chromosomes during maturation in $Sur^{F/F}$ and $Sur^{F/F}$; *Gdf9-Cre* occytes. β -Tubulin-GFP mRNAs were microinjected into GV occytes of the indicated genotypes and occytes were transferred to the live imaging station after 2 h of incubation when most occytes had undergone GVBD. Time points indicated the time elapsed from the start of corresponding Supplementary Movie 2A and B. Red arrow indicated the absence of spindle midbody microtubules in (**b**); White arrows indicated segregated chromosomes in (**a**) and (**b**); yellow arrow showed the polar body in (**a**) and membrane protrusions in (**b**). Bar = 25 μ m. (**c**) Polar bodies extruded by $Sur^{F/F}$ occytes at 12 h of culture (upper row, white arrows) and membrane protrusions in $Sur^{F/F}$; *Gdf9-Cre* occytes at 8 h of culture (lower row, red arrows). (**d**) Cumulative rates of polar body extrusion and membrane protrusion during maturation of $Sur^{F/F}$ and $Sur^{F/F}$; *Gdf9-Cre* occytes. The numbers of analyzed occytes were indicated (*n*)

dramatically downregulated in *Sur^{F/F}*; *Cyp19-Cre* mice, suggesting that the differentiation of GCs to the preovulatory stage was impaired.

Discussion

We found that Survivin in both oocyte and surrounding GCs is essential for female fertility. During follicle development, primordial follicles with Survivin depletion in the oocyte could develop into antral follicles with normal granulosa and cumulus cells, and release eggs. However, deletion of *Survivin* in oocytes caused impaired oocyte maturation and infertility. Our findings also provided evidence that specific deletion of *Survivin* in GCs led to abnormal ovarian function and female subfertility. Survivin is required for spindle assembly, SAC protein kinetochore localization and the accurate regulation of anaphase onset during oocyte meiotic maturation. In mitosis, cells lacking Survivin are unable to align their chromosomes, fail to recruit Aurora B to kinetochores and become polyploid at a very high frequency.²³ Survivindepleted cells displayed delayed cell-cycle progression and accumulated in prometaphase with misaligned chromosomes.²⁴ In human cells, Survivin also regulates assembly and stability of microtubules during late mitosis.²⁵ We recently showed that Survivin knockdown in vitro led to precocious PBE but did not affect the bipolar spindle formation.²⁶ In our previous studies, we showed that BubR1 knockdown induced acceleration of mouse oocyte meiotic progression,¹⁷ while Bub3 deletion by RNAi did not disturb the meiotic process but abrogated oocyte meiotic



Figure 5 Cytokinesis failure in $Sur^{F/F}$; *Gdf9-Cre* oocytes. (**a** and **b**) Dynamics of F-actin during $Sur^{F/F}$ and $Sur^{F/F}$; *Gdf9-Cre* oocyte maturation. Oocytes were microinjected with Alexa 488-phalloidin and transferred to the live imaging station after 1 h of incubation. Time points indicated the time elapsed from the start of corresponding Supplementary Movie 3A and B. Red arrows showed actin ring in (**a**) and (**b**); white arrows indicated segregated chromosomes in (**a**) and (**b**); yellow arrows showed the polar body in (**a**) and membrane protrusion in (**b**). Bar = 25 μ m. (**c** and **d**) Localization of midbody spindle and Myc-Aurora-C at telophase in $Sur^{F/F}$; *Gdf9-Cre* oocytes. Oocytes used for staining were fixed after 8 h of incubation in the $Sur^{F/F}$; *Gdf9-Cre* group and 12 h in the $Sur^{F/F}$ group, respectively. Green: α -tubulin in (**c**); Myc-Aurora C in (**d**); Red: DNA. Bar = 25 μ m. (**e**) Scheme represents the process of polar body emission in oocytes of indicated genotypes

arrest induced by nocodazole.¹⁶ In the present study, indeed the absence of kinetochore localization of BubR1 and Bub3 in prometaphase I was observed in *Survivin*-deleted oocytes. Furthermore, microinjection of the kinase-deficient Aurora C (AurC-KD) mRNA into mouse oocytes also disrupted Bub1 and BubR1 targeting to kinetochores and caused premature chromosome segregation.²¹ Interestingly, our data also showed that the kinetochore localization of Aurora C was disturbed in Survivin-depleted oocytes (data not shown).

Survivin is essential for the completion of cytokinesis in mouse oocyte meiotic maturation. In meiosis I, the course of PBE mainly includes three processes: the establishment of polarity, membrane protrusion and successful abscission.^{27,28} *Survivin*-deleted oocytes showed membrane protrusion but failed to extrude the polar body, suggesting that these defective oocytes completed the first two steps of PBE but failed to undergo the third step, that is, abscission. In *Drosophila*, Survivin facilitates the localization of Polo kinase and Rho at the equatorial cortex in spermatocytes,²⁹ and these kinases may modify substrates like CHO1/MKLP that is responsible for assembly of the midbody microtubule complex.³⁰ In human retinal pigment epithelial (RPE) cells, Survivin depletion disrupts the association of the inner centromere protein (INCENP),

Aurora-B and MKLP-1 with the spindle midzone and the midbody.²⁵ In our study, we found that Survivin depletion disrupted the midbody microtubules in telophase I but did not affect the dynamic movement of F-actin to form a contractile ring. We also found that the kinase protein Aurora C, which carries out functions in oocyte meiosis rather than Aurora B,³¹ was also disturbed in telophase in Survivin-depleted oocytes.

Survivin in GCs has fundamental roles in folliculogenesis. In a recent study, a known splice variant of Survivin. Survivin-2B, has been reported to regulate cell death.³² Here, our results showed that the Survivin depletion in GCs led to abnormal follicle growth, ovulation failure (follicular atresia) and impaired luteinization, which were caused by massive GCs apoptosis. In the absence of healthy GCs, ovarian follicles are destined to undergo atresia because of loss of the main supporting signals. It is known that FSH controls follicular development by regulating GC proliferation and differentiation;^{33,34} and FSH responsiveness is essential for the formation of the antrum and growth to the preovulatory stage.³⁵ In our study, GC viability and differentiation to the preovulatory stage was impaired in the Sur^{F/F}; Cyp19-Cre mice. The lower expression of the Cyp19a1 in the mutant GCs suggested a weakened FSH responsiveness. Then, the downregulation of the Fshr in mutant GCs



Figure 6 Impaired folliculogenesis, ovulation and CL development in $Sur^{F/F}$; *Cyp19-Cre* mice. (a) Representative H&E staining of ovaries 48 h after PMSG priming of ~ d23 female mice. Bar = 500 μ m. The two panels below were at a \times 20 magnification (black arrow, the membrana GCs; yellow arrow, the cumulus cells). (b) Representative H&E staining of ovaries from mice primed with PMSG for 48 h at ~ d23, followed by hCG for 16 h. Bar = 500 μ m; asterisk, follicles that failed to ovulate oocytes. (c) Representative H&E staining of ovaries from mice primed with PMSG for 48 h at ~ d23, followed by hCG for 48 h. Bar = 500 μ m, asterisk showed undeveloped CLs. (d) Levels of both serum estradiol and progesterone were decreased in *Sur* ^{F/F}; *Cyp19-Cre* mice. **P* < 0.05

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Figure 7 Survivin deletion led to GC apoptosis and downregulation of key ovarian genes. (**a** and **b**) TUNEL assay and immunostaining of CC3 in ovaries from control and $Sur^{F/F}$; *Cyp19-Cre* mice treated with PMSG for 48 h showing GC apoptosis. (**c**) Immunofluorescence for CC3 in cultured GCs showed more apoptotic cells in Survivin-depleted GCs. Green, CC3; Red, DNA. Bar = 20 μ m. (**d**) Relative expression of key ovarian genes (*Fshr, Lhgcr* and *Cyp19a1*) in control and $Sur^{F/F}$; *Cyp19-Cre* mice after PMSG treatment. **P* < 0.05

provided a molecular mechanism that might explain the poor follicle growth in *Sur^{F/F}*; *Cyp19-Cre* mice. On the other hand, LH initiates ovulation, terminates GC proliferation and regulates the transition of GCs to lutea cells (LCs).³⁶ Here, we demonstrated a dramatic decrease of *Lhgcr* expression in the mutant GCs, which might account for the reduced ovulation and impaired CL formation observed in *Sur^{F/F}*; *Cyp19-Cre* mice.

A recent report showed that human oocytes/embryos not only expressed but also secreted surviving, suggesting that Survivin may have an important role in human oogenesis and embryogenesis.³⁷ Thus, our results may provide new insights for understanding functions of Survivin in human oogenesis, which will aid in understanding pathology of human infertility or subfertility.

Materials and Methods

Mice. The *Survivin^{flox/flox}* (referred to as *Sur^{F/F}*) mice in a C57BL/6J genomic background were provided by Tak W Mak.¹⁵ Transgenic mouse lines carrying *growth differentiation factor 9* (*Gdf-9*) and *cytochrome P450* (*Cyp-19*) promotermediated Cre recombinase were provided by Dr. Heng-Yu Fan.^{38,39} Both transgenic mouse lines have C57BL/6J genomic background. The *Sur^{F/F}* mice were crossed with *Gdf9-Cre* mice and *Cyp19-Cre* mice, respectively. After multiple rounds of crossing, we had generated mutant female mice lacking *Survivin* in ocytes (referred to as *Sur^{F/F}*; *Gdf9-Cre*) and GCs (referred to as *Sur^{F/F}*; *Cyp19-Cre*). The mice were housed under controlled environmental conditions with free access to water and food. Illumination was provided between 0800 and 2000 h.

Oocyte collection and culture. Animal care and handling were conducted according to the guidelines of the Animal Research Committee of the Institute of Zoology, Chinese Academy of Sciences. GV stage oocytes were isolated from ovaries of 6- to 9-week-old female mice and cultured in M2 medium (Sigma, St. Louis, MO, USA) under paraffin oil at 37°C, 5% CO₂ in air. They were collected

at different times of culture for immunofluorescent staining, western blotting, microinjection and chromosome spreads.

Superovulation. To induce synchronized follicular growth and ovulation, each female mouse at \sim d23 was injected with 7.5 IU of PMSG followed by 7.5 IU of hCG 48 h to promote ovulation. Mice were killed at 12–14 h or 16 h of hCG treatment and COCs were recovered from each oviduct. After a 5-min treatment with hyaluronidase (1 mg/ml) in M2 medium (Sigma), oocytes were collected and counted.

Antibodies, immunofluorescence and confocal microscopy. Antibodies used in the experiments were purchased from the following companies: rabbit monoclonal anti-Survivin (Cell Signaling Technology, Beverly, MA, USA); mouse monoclonal anti-z-tubulin, anti-y-tubulin (Sigma), anti-myc-FITC, anti-GAPDH (Invitrogen, Carlsbad, CA, USA); sheep polyclonal anti-BubR1 (Abcam, Cambridge, MA, USA); rabbit polyclonal anti-Bub3 (Santa Cruz, Dallas, TX, USA). Secondary antibodies were purchased from ZhongShan Golden Bridge Biotechnology Co., Ltd (Beijing, China). Oocytes for immunofluorescent staining were fixed in 4% paraformaldehyde in PBS for 30 min at room temperature. Then, they were transferred to membrane permeabilization solution (0.5% Triton X-100) for 20 min and blocking buffer (1% BSA-supplemented PBS) for 1 h. At last, oocytes were incubated overnight at 4°C with antibodies described above in appropriate dilutions. Then, the oocytes were mounted on glass slides and examined with a laser scanning confocal microscope (Zeiss LSM 710 META, Oberkochen, Germany).

cDNA cloning, mRNA synthesis and microinjection. Mouse Aurora C cDNA was obtained by RT-PCR using total RNA from 150 mouse oocytes. The following two nested primers were used: F1: ACTTCAACTTGCCCTCAC, R1: ACAGAGCCTGGAGACCTT; F2: TCAGGCCGGCCGATGGAGACCCCAGCACCTCA ACCA, R2: GTTGGCGCGCCCTAGCAAGGAAGCACCCTTCGA. The full Aurora C cDNA was then cloned to pSC2 +-Myc vector. The Myc-Aurora C-pCS2 + plasmid was linearized by *Sal* and purified by the gel extract kit (Promega, Madison, WI, USA). SP6 message machine (Ambion, Austin, TX, USA) was used for producing capped mRNA which was then purified by the RNeasy cleanup kit (Qiagen, Hilden, Germany). Microinjection was performed using an Eppendorf microinjector (Hamburg, Germany) and completed within 30 min.

Live imaging of oocytes and chromosome spread. For live oocyte imaging, oocytes were respectively microinjected with β 5-tubulin-GFP mRNAs, Cyclin B1-GFP mRNAs or Alexa 488-phalloidin (Invitrogen). The oocytes were cultured in M2 medium with Hoechst 33342 (5 ng/ml) for 2 h and then transferred to the Perkin Elmer precisely Ultra VIEW VOX Confocal Imaging System (Perkin-Elmer, Waltham, MA, USA) equipped with 37°C incubator and 5% CO₂ supply. The DNA in oocytes was labeled in blue using Hoechst 33342, and changed into red in the figures and movies.

For chromosome spreads, oocytes were left in hypotonic solution (1% Na Citrate) for 20 min at room temperature and fixed with methanol/glacial acetic acid (3:1). The oocytes were then stained with PI for 5 min.

Histological analysis, immunohistochemistry and TUNEL assay. Ovaries used for histological analysis were collected from female mice after a PMSG 48-h treatment. Then, they were fixed in 4% paraformaldehyde (pH 7.5) overnight at 4°C, dehydrated, and embedded in paraffin. Paraffin-embedded ovaries were sectioned at a thickness of 8 μ m for hematoxylin and eosin (H&E) staining. One or both ovaries from more than three mice of each genotype were used for each time point.

Immunohistochemistry was performed on 8- μ m sections using the Vectastain ABC kit (Vector Laboratories, Burlingame, CA, USA). Rabbit monoclonal anti-Survivin was used to detect the expression of Survivin in mouse ovary. Rabbit anti-phosphohistone H3 (Upstate Laboratories, Lake Placid, NY, USA) and rabbit anti-cleaved caspase-3 (Cell Signaling Technology) were used to evaluate cell proliferation and apoptosis in follicles. For direct comparison, ovary sections from four individual females of each genotype were processed together.

Analysis of apoptosis in ovarian follicles was carried out by TUNEL assay using the ApopTag Plus *in situ* apoptosis detection kit (Chemicon International, Temecula, CA, USA). At least four different specimens from each genotype were analyzed in parallel.

GCs isolation, culture and immunofluorescence. Immature female mice (\sim d23) were injected with 7.5 IU PMSG, and ovaries were harvested 48 h later. Ovaries were dissected and placed in DMEM/F12. Large antral follicles were

punctured to extrude GCs, which were then strained and spun down. GCs were harvested and cultured in DMEM/ F12 medium on cover glasses in culture dishes. After 24 h of culture, cells were fixed in 4% paraformaldehyde and immunofluorescent staining was performed as previously described⁴⁰ with primary anti-cleaved caspase-3 antibody and secondary FITC-conjugated goat anti-rabbit IgG antibody (ZhongShan Golden Bridge Biotechnology Co.).

Western blot analysis. A total of 250 mouse oocytes per sample or GC extracts containing 30 μ g protein were mixed with SDS sample buffer and boiled for 5 min at 100°C for SDS-PAGE. Western blotting was performed as described previously⁴⁰ using the antibody dilution anti-Survivin: 1 : 1000; anti-GAPDH: 1 : 2000.

RNA isolation and quantitative real-time PCR analysis. Total RNA from primary mouse GCs was extracted using the RNeasy micro purification kit (Qiagen). Single-stranded cDNAs were generated with the cDNA synthesis kit (Invitrogen). The resultant cDNAs were used for real-time PCR. Real-time PCR was conducted by using SYBR Premix Ex Taq kit (TaKaRa Biotechnology (Dalian) Co., Ltd., Japan) in ABI prism 7500 Sequence Detection System (Applied Biosystems, Foster City, CA, USA). Analysis of relative gene expression was measured by real-time quantitative PCR and the $2^{-\Delta \Delta CT}$ method.

Serum analysis. Mice were anesthetized after PMSG 48 h or PMSG 48 h-hCG 48 h treatment and blood was collected by eyeball removal. Estradiol and progesterone measurements were made at a commercial laboratory (Chemclin Co., Beijing, China). At least 10 mice were anesthetized for each group.

Statistical analysis. All experiments were repeated at least three times. Statistical analysis was performed using SPSS (SPSS China, Shanghai, China). Data were expressed as mean \pm S.E.M. and P < 0.05 was considered as statistically significant.

Conflict of Interest

The authors declare no conflict of interest.

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