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The RNA ligase RNA terminal phosphate cyclase B regulates mRNA alternative splicing and is required for mouse oocyte development and maintenance

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ABSTRACT

Recent large-scale mRNA sequencing has shown that introns are retained in 5-10% of mRNA, and these events are named intron retention (IR). IR has been recognized as a key mechanism in the regulation of gene expression. However, the role of this mechanism in female reproduction in mammals remains unclear. RNA terminal phosphate cyclase B (RTCB) is a RNA ligase; we found that RTCB conditional knockout mice have premature ovarian failure and that RTCB plays a crucial role in follicular development. RTCB regulated the splicing of transcripts related to DNA methylation and DNA damage repair. In addition, it regulated the resumption of oocyte meiosis by affecting CDK1 activation. Moreover, the loss of RTCB suppressed zygotic genome activation (ZGA) and decreased translation at the global level. In addition, Rtcb deletion resulted in the accumulation of maternal mRNAs containing unspliced introns and in a decline in the overall level of transcripts. As a result, the Rtcb^{-/-} females were sterile. Our study highlights the important role of RTCB-regulated noncanonical alternative splicing in female reproduction.

KEY WORDS: RNA processing, Transcriptome, Female fertility, Premature ovarian failure, Folliculogenesis, Oogenesis, Meiosis

INTRODUCTION

During the development of growing oocytes into fully grown oocytes, an increase in oocyte volume is coupled with active transcription events (Pangas and Rajkovic, 2006). Pre-mRNA is processed to remove introns and splice exons via alternative splicing (AS) (Darnell, 2013; Ding et al., 2014). Through AS, multiple transcripts and protein variants can be transcribed and translated from a single gene, which significantly enriches the diversity of transcriptomes and proteomes (Braunschweig et al., 2013; Stamm et al., 2005). Common AS events include intron retention (IR),

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Handling Editor: Swathi Arur
 Received 5 January 2022; Accepted 7 September 2022

skipped exons (SE), mutually exclusive exons (MXE), and alternative 5' and 3' splice sites (A5SS and A3SS). Recent studies have shown that intron retention (IR) occurs in many mRNAs and is a frequent event in plants and humans (Ner-Gaon et al., 2004; Reddy et al., 2013). IR is recognized as a key mechanism in the regulation of gene expression. Specific transcripts with IR are expressed in specific cell types or situations, and therefore control developmental transitions or cellular stress responses (Boothby et al., 2013; Braunschweig et al., 2014; Shalgi et al., 2014; Wong et al., 2013).

During mRNA retained intron splicing, two half-molecules 87 with 2'3'-cyclic phosphate and 5'-OH ends are considered 88 unconventional products (Englert et al., 2012). There are two 89 known pathways for rectifying these two ends. Classically, ligation 90 occurs via a multienzyme pathway. Classical ligases catalyze 91 the ligation of 5'-P and 3'-OH ends with the help of ATP and Mg²⁺ 92 (Desai et al., 2013). Before sealing, 2',3'-cyclic phosphate is 93 94 hydrolyzed to 3'-OH by phosphodiesterase, and 5'-OH is phosphorylated by polynucleotide kinase to generate 5'-phosphate 95 (5'-P). The second pathway depends on RTCB. The role of RTCB 96 as an RNA ligase was recently confirmed. It directly ligates the 2'3'-97 cyclic phosphate and 5'-OH terminal half molecules (Desai et al., 98 2013; Popow et al., 2011; Tanaka et al., 2011a,b; Tanaka and 99 Shuman, 2011). With the help of GTP and Mn²⁺ ions, RTCB can 100 treat 2'3'-cyclic phosphate and 5'-OH ends in one step to form a 101 phosphodiester bond (Chakravarty and Shuman, 2012; Chakravarty 102 et al., 2012; Desai and Raines, 2012; Englert et al., 2012). RTCB 103 plays a key role in tRNA maturation and splicing of Xbp1 mRNA 104 (Tanaka et al., 2011b), and can catalyze the reconnection of short 105 16S rRNA produced by stress (Maughan and Shuman, 2016). *Xbp1* A@2 is the only confirmed RTCB target mRNA (Lu et al., 2014). RTCB 107 regulates the unfolded protein response (UPR) through the 108 unconventional splicing of Xbp1 mRNA in the cytoplasm (Shinya 109 et al., 2011). RTCB also plays a key role in regulating tRNA splicing 110 after intron removal (Tanaka et al., 2011b). According to a previous 111 study, ~5% of tRNA genes in mice contain introns (Wu and 112 Hopper, 2014). tRNA introns are positioned downstream of the 113 anticodon and are smaller than mRNA introns. After intron removal, 114 these pre-tRNAs require ligation to form mature tRNAs. RTCB is 115 responsible for catalyzing the ligation of these tRNAs. 116

Immature oocytes of mice are arrested at the germinal vesicle 117 (GV) stage in growing ovarian follicles unless they are stimulated by 118 a gonadotrophin signal or released from an inhibitory follicular 119 environment. At this stage, GV oocytes can be classified based on 120 their chromatin configuration in non-surrounded nucleolus (NSN) 121 oocytes and surrounded nucleolus (SN) oocytes. NSN oocytes 122 have low developmental competence and their chromatin does not 123 form a Hoechst-positive rim around the nucleolus. In contrast, the 124

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chromatin of SN oocytes forms a Hoechst-positive rim around the
nucleolus, and transcription in SN oocytes is silenced (Tan et al.,
2009). However, many of the synthesized mRNAs are stored in the
ooplasm. The dormant mRNAs drive meiosis and early embryonic
development. The key events of meiosis mainly include germinal
vesicle breakdown (GVBD) and the following two consecutive M
phases (MI and MII) (Jones, 2011).

Splicing factors in oocytes, such as SRSF3 and BCAS2, have 132 been shown to play an important role in oocyte maturation and the 133 maintenance of transcriptome integrity (Do et al., 2018; Zhang 134 et al., 2022). Without RTCB, the lifespan of Caenorhabditis 135 elegans decreases, germ cells fail to mature into oocytes, and the 136 animals are rendered completely sterile (Kosmaczewski et al., 137 138 2014). We hypothesized that this new mechanism of rapid gene 139 regulation via RTCB may also exist in mouse oocytes. To test this hypothesis, we specifically knocked out the Rtcb gene in oocytes as 140 early as the primordial follicle stage using Gdf9-Cre. Our results 141 142 show that RTCB is responsible for regulating mRNA splicing after intron removal during oocyte meiosis. The absence of RTCB 143 144 resulted in the accumulation of a series of intron-containing 145 transcripts, mainly including DNA methylation and DNA damage 146 repair-related transcripts. As a result, the levels of these maternal 147 proteins decreased. Follicular development, meiosis recovery and early embryonic development were impeded. In summary, our 148 results confirm that RTCB-regulated alternative splicing plays an 149 150 important role in female reproduction.

RESULTS

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Rtcb is highly expressed in mouse oocytes and is indispensable for female fertility

155 First, we detected *Rtcb* expression in various tissues and organs by 156 quantitative RT-PCR. The results showed that *Rtcb* was highly expressed in female germ cells (Fig. 1A). In addition, Rtcb mRNA 157 158 levels remained relatively stable in oocytes and increased drastically after the two-cell stage (Fig. 1B). These results suggest that RTCB 159 may play an important role in oocyte development and embryo 160 preimplantation. Thereafter, we detected RTCB levels by western 161 162 blotting (WB) during oocyte meiosis. The results showed that RTCB protein levels at the GVBD stage were higher than in the 163 164 germinal vesicle (GV), MI and MII stages (Fig. 1C).

165 To study the in vivo function of RTCB in oocytes, we constructed oocyte-specific Rtcb knockout mice. By crossing Rtcb-floxed mice 166 $(Rtcb^{fl/fl})$ with Gdf9-Cre transgenic mice, we deleted the Rtcb gene 167 168 in oocytes as early as the primordial follicle stage (Fig. 1D). WB and 169 immunofluorescence (IF) results revealed that RTCB protein was 170 completely depleted in GV oocytes (Fig. 1E,G). IF results showed 171 that RTCB was distributed in the nuclei and cytoplasm of the oocytes. In contrast, there was no RTCB signal in Rtcb-deleted 172 oocytes (Fig. 1E). A subcellular fractionation assay of 293T cells 173 174 also proved the nuclear and cytoplasmic localization of RTCB 175 (Fig. 1F). This result is consistent with previous reports in HeLa cells (Jurkin et al., 2014). Thereafter, we conducted a fertility test for 176 7 months by mating $Rtcb^{fl/fl}$ female mice (hereafter referred to as 177 wild type) and Rtcb^{fl/fl};Gdf9-Cre female mice (hereafter referred 178 to as $Rtcb^{oo-/-}$) with wild-type males. The $Rtcb^{oo-/-}$ female mice 179 were completely sterile (Fig. 1H). These results demonstrate that 180 RTCB in oocytes is required for female fertility in mice. 181

183 *Rtcb* deletion causes premature ovarian failure in mice older 184 than 8 weeks

To study the effect of *Rtcb* deletion on follicle development in the ovary, we collected mouse ovaries at various stages and compared their shapes and sizes. Compared with wild-type mice, the ovaries of 4-week-old $Rtcb^{oo-/-}$ mice (not stimulated by exogenous gonadotropins) were almost similar in size. However, in $Rtcb^{oo-/-}$ mice older than 6 weeks, the ovaries were significantly smaller than those of wild-type mice (Fig. 2A,B).

Subsequently, we investigated follicle development at 192 various ages using Hematoxylin and Eosin staining, and 193 immunohistochemistry (IHC). The ovaries of 4-week-old 194 *Rtcb*^{oo-/-} mice still contained many primordial, primary, 195 secondary and antral follicles (Fig. 2C). Follicular counting 196 results showed that the ovaries of 6-week-old Rtcb^{oo-/-} mice 197 (not stimulated by exogenous gonadotropins) contained similar 198 numbers of primordial, primary and bilayer secondary follicles 199 relative to wild-type mice, but fewer secondary and antral follicles 200 with multilayer granulosa cells (Fig. 2D,H). Furthermore, fewer 201 primordial, primary and secondary follicles remained in the ovaries 202 of the 8-week-old mice (Fig. 2E and I). No follicles were found 203 in the ovaries of 10-week-old or 7-month-old knockout mice 204 (Fig. 2F,G). 205

To investigate why premature ovarian failure occurs, we examined the signal of phosphorylated H2AX (γ H2AX, an unrepaired DNA lesion marker) in oocytes in ovarian sections. We found that, in 6-week-old $Rtcb^{00-/-}$ mice, more than 50% of oocytes in secondary and antral follicles had obvious DNA damage signals (Fig. 2J,K). Moreover, TdT-mediated dUTP nick-end labeling (TUNEL) showed a higher proportion of apoptotic signals in the follicles of $Rtcb^{00-/-}$ mice than in those of wild-type mice (Fig. 2L,M). In summary, these observations suggest that RTCB is essential for ovarian follicle development and that Rtcb deletion in oocytes leads to DNA damage and induced follicle atresia.

Rtcb deletion causes the accumulation of a series of maternal transcripts containing introns

Because fully grown GV oocytes were still present in the ovaries of 6-week-old $Rtcb^{oo-/-}$ mice, we analyzed the oocytes isolated from 6-week-old wild-type and knockout mice treated with pregnant mare serum gonadotropin (PMSG). The diameters of the Rtcb-null oocytes were significantly smaller than those of the wild-type oocytes (Fig. 3A,B). In addition, numerous brown cytoplasmic granules were detected in Rtcb-null oocytes (Fig. 3A). DAPI staining revealed that more than 80% of fully grown oocytes from 6-week-old wild-type mice had a surrounded nucleolus (SN) configuration. However, the proportion of SN configurations was only ~40% in the Rtcb-null oocytes (Fig. 3C,D). These results indicate that Rtcb deletion inhibited oocyte growth and nucleolar maturation.

To explore the effect of *Rtcb* deletion on maternal transcript 233 levels, we collected fully grown GV oocytes from 6-week-old wild-234 type and knockout mice for RNA sequencing (RNA-seq) analyses. 235 Gene expression levels were evaluated by fragments per kilobase 236 of transcript per million mapped reads (FPKM). According to the 237 RNA-seq results, we detected more genes with decreased mRNA 238 copy numbers [1861 when fold change >2, false discovery rate 239 (FDR) <0.05] than those with increased mRNA copy numbers 240 (498 when fold change \geq 2, FDR<0.05) (Fig. 3E). We performed 241 gene ontology (GO) analysis on the decreased transcripts and 242 found that they were mainly enriched in protein transport and 243 DNA damage repair pathway (Fig. 3F), which may partially 244 explain the phenotype of follicular atresia observed in 6-week-old 245 *Rtcb*^{oo-/-} mice. As RTCB mainly affects mRNA splicing, we tested 246 the mRNA levels of the retained introns. We detected the level 247 of transcripts containing intron retention (IR) using IRFinder 248

Development (2022) 149, dev200497. doi:10.1242/dev.200497



Fig. 1. The expression pattern of *Rtcb* mRNA and RTCB protein, and the phenotype of *Rtcb* knockout mice. (A,B) RT-qPCR results reveal the relative expression levels of mouse *Rtcb* mRNA in germinal vesicles (GV) and MII occytes (A,B), in various tissues (A), and in preimplantation embryos (B). *n*=3 biological replicates. (C) Western blotting results show the expression levels of endogenous RTCB protein in GVs, in germinal vesicle breakdown (GVBD), and in MI and MII occytes. DDB1 was used as a loading control. (D) Gene-targeting strategy for *Rtcb*-conditional knockout mouse. In *Rtcb* conditional knockout mice, the fourth exon of the *Rtcb* gene containing 100 bp is deleted, leading to premature translation termination. (E) Immunofluorescence (IF) results showing localization of endogenous RTCB in growing GV and fully grown GV occytes from wild-type and *Rtcb^{filfl};Gdf9-Cre* female mice. Scale bar: 20 µm. (F) The nuclear and cytoplasmic separation samples of 293T cells were used for western blotting, and the results showed that RTCB protein and *NC* was used as a nuclear marker, and DDB1 was used as a loading control. (G) Western blotting results show that RTCB protein is completely eliminated in *Rtcb^{filfl};Gdf9-Cre* female mice. The black arrowhead indicates the band corresponding to the RTCB protein. DDB1 was used as a loading control. (H) Cumulative number of pups per female mouse revealing the fertility of wild-type and *Rtcb^{oo-/-}* female mice. *n*=4 females for each genotype. Data are mean±s.e.m. ***P<0.001.

(Middleton et al., 2017). The IR ratio was calculated as the ratio of unspliced exon-intron junction reads to total junction reads (unspliced exon-intron and spliced exon-exon). The IR change was calculated from the IR ratio of knockout samples minus that of wild-type samples (Fig. 3G). According to IRFinder analysis, we detected many retained introns (Fig. 3I and Table S1). When IR change>0.1 and P<0.1, 91 transcripts containing IR were detected

(Fig. 3H and Table S2). In addition, we used rMATS software to analyze other AS events (Shen et al., 2014). In addition to IR, many other AS events underwent significant changes because of *Rtcb* deletion (Fig. 3J and Table S3). GO enrichment analysis of different splicing events showed that RNA splicing was the most affected (Fig. S1). Based on these results, we conclude that RTCB plays a crucial role in RNA splicing in GV oocytes. 372

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7-month-old mice with indicated genotypes. Scale bars: 500 µm. (B) The ovary weight of wild-type and Rtcb^{oo-/-} mice, as in A. Data are mean±s.e.m. *P<0.05, **P<0.01, ***P<0.001. n.s. indicates a non-significant result. (C,D) Hematoxylin and Eosin staining results revealing ovarian histology of 4- and 6-week-old mice with indicated genotypes. Scale bars: 100 µm. (E) Immunohistochemistry staining of mouse VASA homolog (MVH, a germ cell marker) in the ovaries of 8-week-old mice with indicated genotypes. Scale bars: 100 µm. (F,G) Hematoxylin and Eosin staining results reveal ovarian histology of 10-week-old and 7-month-old mice with indicated genotypes. Scale bars: 100 µm. (H) The number of follicles at indicated stages in the ovaries of 6-week-old wild-type and Rtcb^{fil} ^{fl};Gdf9-Cre female mice. Follicles were counted on serial ovarian sections after Hematoxylin and Eosin staining. Data are mean±s.e.m. *P<0.05, n.s. indicates a non-significant result. PD, primordial follicles; Pr, primary follicles; Bi, secondary follicles with two layers of granulosa cells; Mu, secondary follicles with multiple layers of granulosa cells; Atr, antral follicles. (I) The number of follicles at indicated stages in the ovaries of 8-week-old wild-type and Rtcb^{n///};Gdf9-Cre female mice. Follicles were counted on serial ovarian sections after Hematoxylin and Eosin staining. Data are mean±s.e.m. *P<0.05, **P<0.01. n.s. indicates a non-significant result. (J) Immunohistochemistry staining of phosphorylated H2AX (yH2AX, an unrepaired DNA lesions marker) in the ovaries of 6-week-old mice with indicated genotypes. Scale bars: 100 µm. (K) Quantification of γH2AX-positive oocytes in the ovaries of 6-week-old mice with indicated genotypes. Data are mean±s.e.m. ***P<0.001. (L) Immunofluorescence (IF) staining of TUNEL in the ovaries of 6-week-old mice with indicated genotypes. Scale bars: 100 µm. (M) Quantification of TUNEL-positive follicles in the ovaries of 6-week-old mice with indicated genotypes. Data are mean±s.e.m. ***P<0.001.



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Fig. 3. Transcriptome and intron retention changes in *Rtcb*-deleted fully grown oocytes. (A) Representative images of germinal vesicle (GV) stage oocytes derived from 6-week-old wild-type and $Rtcb^{oo-/-}$ females. n=5 mice for each genotype. Scale bars: 100 µm. (B) Quantitative statistical results of oocyte diameter length derived from wild-type and $Rtcb^{oo-/-}$ females in A. Data are mean±s.e.m. ****P*<0.001. (C) DAPI staining showed the non-surrounded nucleolus (NSN) and surrounded nucleolus (SN) chromatin configurations in wild-type and $Rtcb^{oo-/-}$ oocytes. Scale bars: 10 µm. (D) The rate of SNs in GV oocytes in A. Data are mean±s.e.m. ****P*<0.001. (E) Scatter plot comparing transcripts between wild-type and $Rtcb^{oo-/-}$ fully grown GV oocytes. Transcripts increased or decreased more than twofold in $Rtcb^{oo-/-}$ oocyte samples are highlighted with blue or red, respectively. (F) Gene ontology analysis of downregulated genes in $Rtcb^{oo-/-}$ fully grown GV oocytes. (G) Schematic diagram revealing distribution of sequence reads for IR. The IR ratio is calculated from the ratio of unspliced exon-intron junction reads to total junction reads (unspliced exon-intron and spliced exon-exon). (H) Scatter plot showing the number of transcripts whose IR change increased or decreased more than 0.1 with *P*<0.1. IR change is calculated by subtracting the IR value of wild-type oocyte samples from the IR value of $Rtcb^{oo-/-}$ oocyte samples. (I) The intron number corresponding to different IR ratio changes (knockout/WT). (J) The number of transcripts corresponding to various alternative splicing (AS) events detected by rMATS. FDR<0.05.

555 The lack of maternal RTCB proteins causes epigenetic

556 changes in oocytes

Based on previous results, we conclude that *Rtcb* deletion leads to IR accumulation and decreases many maternal mRNA levels. To determine whether IR accumulation can lead to changes in mRNA617levels, we analyzed 91 transcripts with significant IR changes (eight618transcripts from two identical genes were merged). The results619showed that the expression levels of most transcripts decreased620

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Fig. 4. The effect of maternal RTCB deficiency on epigenetic regulation. (A) Heatmap of genes showing the change of these transcripts having a significant IR accumulation due to *Rtcb* deletion. The value in the figure represents the relative level of mRNA. More details are included in Table S2.
(B) IR analysis of several transcripts using RT-qPCR. Forward primer 1 and reverse primer 1 are designed to detect the level of the total transcripts. Forward primer 1 and reverse primer 2 are designed to detect the level of transcripts containing intron retention. Data are mean±s.e.m. ***P*<0.01, ****P*<0.001. n.s. indicates a non-significant result. (C) PCR analysis results of some transcripts in A. *Gapdh* was used as a loading control. (D) Changes of relative mRNA copy numbers of indicated transcripts in wild-type and *Rtcb^{00-/-}* fully grown oocytes. Data are mean±s.e.m. ****P*<0.001. (E) Western blotting results showing the expression levels of DNMT1, DNMT3A, DNMT3B and RAD50 proteins in wild-type and *Rtcb^{00-/-}* occytes. DDB1 was used as a loading control. Total proteins from 200 oocytes were loaded into each lane. (F) Immunofluorescent staining showing the level of 5-hydroxymethylcytosine (5hmC) and 5-methylcytosine (5mC) derived from wild-type and *Rtcb*-deleted oocytes in the non-surrounded nucleolus (NSN) and surrounded nucleolus (SN) stages. Scale bars: 20 µm. (G) Quantification of 5hmC and 5mC in C. Data are mean±s.e.m. ****P*<0.01. n.s. indicates a non-significant result.

(Fig. 4A). Thereafter, we confirmed the IR of these transcripts by RT-qPCR. MBD1 (methyl-CpG binding domain protein 1), also known as CXXC3, plays an important role in DNA methylation (Jørgensen et al., 2004). DNMT3B (DNA methyltransferase 3β) is responsible for establishing DNA methylation (Chen et al., 2004). TDRD9 is an ATP-binding RNA helicase required for spermatogenesis (Guijo et al., 2018). RAD50 is a component of the MRN complex and plays a vital role in double-strand break (DSB) repair (Iijima et al., 2008; Syed and Tainer, 2018). RT-qPCR results showed that IR occurred in these transcripts and that all three introns of Tdrd9 were retained. In addition, the levels of most transcripts decreased (Fig. 4B). Visual assessment of RT-PCR products through agarose gel electrophoresis showed that IR events occur

in these transcripts after RTCB depletion. (Fig. 4C). The RNA-seq results also showed a decrease in *Dnmt1*, *Dnmt3a* and *Dnmt3b* mRNA levels (Fig. 4D).

As *Rtcb* deletion resulted in the IR of DNA methylation-related mRNAs, such as *Dnmt3b* and *Mbd1*, we detected changes in the levels of these related proteins. The results showed that the levels of DNMT1, DNMT3A, DNMT3B and RAD50 were decreased (Fig. 4E). In addition, IF staining showed that the levels of 5-methylcytosine (5mC) rather than 5-hydroxymethylcytosine (5hmC) in *Rtcb*^{oo-/-} SN oocytes were significantly downregulated compared with those in wild-type oocytes (Fig. 4F,G).

These results demonstrate that by affecting RNA splicing, RTCB regulates the transcription of other maternal epigenetic regulators



Fig. 5. The effect of Rtcb deletion on oocyte meiosis resumption. (A) Schematic diagram of oocyte culture in vitro and calculation method of oocyte germinal vesicle breakdown (GVBD) and polar body emission (PBE) rate. (B) Comparison of germinal vesicle breakdown (GVBD) kinetics in cultured wild-type and Rtcb^{oo-/-} oocytes. (C) Kinetics of polar body emission (PBE). When oocytes undergo GVBD within 6 h, they are selected for further culture. Data are mean±s.e.m. ***P<0.01. n.s. indicates a non-significant result. (D) Immunofluorescence (IF) results show spindle assembly and chromosome alignment in wild-type and Rtcb^{oo-/-} oocytes. Scale bars: 20 µm. (E) Rates of oocytes revealing normal spindle morphology as in C. Data are mean±s.e.m. n.s. indicates a non-significant result. (F) The number of ovulated eggs in E. Data are mean±s.e.m. ***P<0.01. (G) Representative images of MII oocytes derived from 6-week-old wild-type and Rtcb^{oo-/-} females treated with hCG for 16 h. n=5 mice for each genotype. Scale bars: 100 µm. (H) The proportion of GV oocytes derived from ovulated oocytes in G. (I) IF results showing spindle assembly and chromosome alignment in ovulated MII oocytes derived from wild-type and Rtcb^{00-/-} females. Scale bars: 20 µm. (J) Rates of oocytes revealing normal spindle morphology as in H. Data are mean±s.e.m. ***P<0.01. n.s. indicates a non-significant result.

and DSB repair factors during oocyte growth, and renders
developmental competence to oocytes. When *Rtcb* is deleted,
IR occurs in many DNA methylation- and DNA damage repairrelated transcripts, and the total levels of these transcripts are
downregulated.

801 Rtcb-null oocytes fail to accomplish meiosis resumption

As *Rtcb* deletion led to the accumulation of introns among maternal transcripts, we then investigated whether these retained introns affected the normal progression of oocyte meiosis by measuring the development rate at different time points in oocytes cultured *in vitro* (Fig. 5A). *In vitro* culture results showed that *Rtcb* deletion significantly downregulated the GVBD rate (Fig. 5B). However, *Rtcb*-null oocytes that were able to resume meiosis showed polar body emission (PBE) rates comparable with those of the wild-type oocytes (Fig. 5C). IF results showed that the absence of RTCB had a minor effect on chromosome alignment and spindle organization in MII oocytes (Fig. 5D,E). We also obtained oocytes from oviducts 16 h after human chorionic gonadotropin (hCG) injection. The *Rtcb*-null oocytes could be ovulated after hCG injection; however, the number of ovulated oocytes from *Rtcb*^{oo-/-} mice was only half that in the control mice (Fig. 5F). In addition, half of the ovulated oocytes were arrested at the germinal vesicle (GV) stage (Fig. 5G,H). Similar to the *in vitro* results, these ovulated *Rtcb*-null



Fig. 6. *Rtcb*-null oocytes with inhibitory CDK1 lack phosphorylated lamin A/C and fail to GVBD. (A) Western blotting results showing the level of indicated proteins in GV oocytes derived from wild-type and $Rtcb^{oo-/-}$ females. DDB1 was used as a loading control. Total proteins from 200 oocytes were loaded in each lane. (B,C) Immunofluorescent (IF) staining showing the level of p^{T14/Y15}CDK1 and p^{S22}lamin A/C in oocytes at GV and GV+2 h stages derived from wild-type and Rtcb-deleted mice. Scale bars: 20 µm. (D,E) Quantification of p^{T14/Y15}CDK1 and p^{S22}lamin A/C in B and C. Data are mean±s.e.m. **P*<0.05, ****P*<0.001. n.s. indicates a non-significant result. (F) Western blotting (WB) results show the level of Flag-CDK1^{T14A,Y15F} in GV oocytes derived from wild-type and $Rtcb^{oo-/-}$ females. The FLAG lines were microinjected with the mRNA products of an empty vector with a FLAG tag. DDB1 was used as a loading control. Total proteins from 100 oocytes were loaded in each lane. (G) GVBD rates of cultured wild-type and $Rtcb^{oo-/-}$ oocytes with or without CDK1^{T14A,Y15F} expression in M16 medium. Data are mean±s.e.m. ****P*<0.001. n.s. indicates a non-significant result.

MII oocytes showed normal chromosomal alignment and spindle
organization (Fig. 5I,J). In summary, these results suggest that a lack
of RTCB mainly affects meiosis resumption and has a minor effect
on the subsequent PBE process.

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917CDK1 activity is reduced in *Rtcb*-null oocytes and lamin A/C918cannot be phosphorylated

919 We examined the effects of *Rtcb* deletion on the levels of meiotic 920 resumption-related proteins. Cyclin B1 is a key cell cycle-regulating protein that promotes G2/M transition (Adhikari and Liu, 2014). 921 922 CDC20 is a positive regulator of the anaphase-promoting complex (APC) (Sudakin et al., 2001). However, WB results revealed that the 923 924 levels of cyclin B1 and CDC20 proteins increased because of *Rtcb* deletion. Resumption of meiosis requires activation of cyclin-925 926 dependent kinase 1(CDK1), and phosphatase cell division cycle 25 927 B (CDC25B) is an activator of cyclin-dependent kinases (Lincoln 928 et al., 2002). The dephosphorylation of threonine 14 and tyrosine 15 929 of CDK1 by CDC25 phosphatases is a key step in the activation of 930 CDK1-cyclin B protein kinase (Morgan, 1995). We found that the levels of CDC25B and p-T161-CDK1 proteins were lower than those in the wild-type oocytes (Fig. 6A).

Phosphorylated lamin A/C (p^{S22} lamin A/C) is a direct substrate of active phosphorylated CDK1 (Heald and McKeon, 1990). Therefore, we examined changes in the inhibitory phosphorylation of CDK1 ($p^{T14/Y15}$ CDK1) and p^{S22} lamin A/C. In wild-type GV oocytes, $p^{T14/Y15}$ CDK1 levels decreased and p^{S22} lamin A/C levels increased when GVBD was about to occur (GV oocytes cultured for 2 h). However, in the *Rtcb*-null oocytes cultured in the same manner, the $p^{T14/Y15}$ CDK1 levels remained high and lamin A/C was not successfully phosphorylated (Fig. 6B-E). This observation indicates that the decline in CDC25B leads to the persistent phosphorylation of CDK1 on threonine 14 and tyrosine 15, and the failure of CDK1 phosphorylation on threonine 161. Consequently, lamin phosphorylation and GVBD failed to occur.

To further test our hypothesis, we overexpressed continuously activated CDK1 (CDK1^{T14A,Y15F}) in wild-type and *Rtcb*-null oocytes at the germinal vesicle (GV) stage (Fig. 6F). The GVBD rate of *Rtcb*-null oocytes overexpressing CDK1^{T14A,Y15F} was ~70%

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Fig. 7. The lack of maternal RTCB protein causes defects in early embryonic development. (A) Representative images of one-cell stage embryos derived from wild-type and Rtcb^{oo-/-} females. n=5 mice for each genotype. Scale bars: 100 µm. (B) Immunofluorescence (IF) results showing the pronucleus formation of Rtcb^{Q+/d+} and Rtcb^{Q-/d+} zygotes. Scale bar: 20 µm. (C) The rate of one cell having normal pronuclei in A. (D) Representative images of two-cell stage embryos and a blastocyst derived from wild-type and Rtcb^{oo-/-} females. n=5 mice for each genotype. Scale bar: 100 µm. (E) Developmental rates of maternal Rtcb-deleted embryos at the time when wild-type embryos reached corresponding stages. Data are mean±s.e.m. ***P<0.001. (F) L-homopropargylglycine (HPG) staining results show protein synthesis activity at the one-cell and two-cell stages in Rtcb^{Q+/3+} and Rtcb^{Q-/3+} embryos. Before staining, these embryos were incubated in medium containing 50 µM HPG for 1 h. Scale bars: 20 µm. (G) Quantification of HPG signal intensity in F. Data are mean±s.e.m. ***P<0.001. n.s. indicates a non-significant result. (H) IF of 5-ethynyl uridine (EU) fluorescence showing RNA transcription in Rtcb^{φ+/δ+} and Rtcb^{φ-/δ+} two-cell stage embryos. Scale bar: 20 μm. (I) Quantification of EU signal intensity in H. Data are mean±s.e.m. *P<0.05. (J) RT-qPCR results showing the expression level of zygotic genome activation (ZGA)-related genes at the one- and two-cell stages in Rtcb^{2+/d+} and Rtcb^{2-/d+} embryos Data are mean±s.e.m. ***P<0.001. n.s. indicates a non-significant result.

(Fig. 6G). This result indicates that RTCB plays an important role in maintaining the levels of activated CDK1 proteins.

Maternal RTCB protein is essential for zygotic genome activation and early embryonic development

Thereafter, we superovulated 6-week-old wild-type and Rtcb^{oo-/-} females and mated them with wild-type males. The diameters of the zygotes obtained from Rtcb^{oo-/-} mice were significantly smaller than those of the wild-type mice, and brown cytoplasmic granules aggregated in maternal *Rtcb*-deleted (hereafter *Rtcb*^{$Q-/d^+})$) zygotes</sup> (Fig. 7A). IF results showed many abnormal or multiple pronuclei in $Rtcb^{\varphi-/\delta^+}$ zygotes (Fig. 7B). The proportion of $Rtcb^{\varphi-/\delta^+}$ zygotes containing normal pronuclei was significantly lower than that of $Rtcb^{\varphi+/\delta^+}$ zygotes (Fig. 7C). Subsequently, zygotes were cultured in



¹¹⁴³ splicing is essential for oocyte growth and embryo development (wild type, top). *Rtcb* deletion in oocytes (bottom) resulted in intron retention (IR) of many ¹¹⁴⁴ transcripts such as *Mbd1*, *Rad50* and *Dnmt3b*. IR accumulation led to the downregulation of corresponding protein expression and insufficient oocyte growth. ¹¹⁴⁵ In addition, by influencing the protein level of p-CDK1 and p-Lamin A/C, the meiosis recovery of most oocytes was blocked. Even if a few oocytes could ¹¹⁴⁶ complete the meiosis recovery, the previous insufficient growth and abnormal pronucleus formed during the fertilization process led to the failure of ZGA. As a result, most zygotes were arrested at the one-cell stage.

KSOM medium. Although most $Rtcb^{\varphi+/\sigma^+}$ zygotes developed 1149 to the two-cell stage, $\sim 70\%$ of $Rtcb^{\varphi-/\delta+}$ zygotes underwent cell 1150 cycle arrest at the one-cell stage. However, fewer than 20% of 1151 $Rtcb^{\varphi-/\delta^+}$ zygotes developed into blastocysts (Fig. 7D,E). These 1152 results indicate that maternal RTCB is important for early 1153 1154 embryonic development. However, as Rtcb-null oocytes suffer from maturation obstacles, subsequent defects in early embryonic 1155 1156 development may be secondary effects caused by defective oocytes.

1157 To detect the global translation level, we incubated wild-type and $Rtcb^{\varphi-/\delta^+}$ embryos at the one- and two-cell stages with 1158 L-homopropargylglycine (HPG), an analog of methionine that can 1159 1160 be examined using the Click-iT cell reaction kit (Life Technologies) 1161 (Zhang et al., 2019). At the one-cell stage, the level of the HPG signal was not significantly different between the wild-type and 1162 $Rtcb^{\varphi-/\delta^+}$ zygotes. However, $Rtcb^{\varphi-/\delta^+}$ two-cell embryos showed 1163 weaker HPG signals than the wild-type embryos (Fig. 7F,G). 1164

To evaluate whether zygotic genome activation (ZGA) was 1165 suppressed in arrested $Rtcb^{\varphi-/\delta^+}$ embryos, we labeled newly 1166 1167 synthesized RNAs with 5-ethynyl uridine (EU) added to the culture medium 2 h before fixation of the one-cell stage. EU signals were 1168 lower in $Rtcb^{Q^{-/d^+}}$ embryos than in $Rtcb^{Q^{+/d^+}}$ (Fig. 7H,I). To confirm 1169 that ZGA was suppressed in the arrested $Rtcb^{Q^{-/O^{+}}}$ embryos, we 1170 examined the expression of six ZGA-related transcripts [MuERV-L 1171 1**A7O3** (Erv4), Npl, Gucula (xxxxx), Zscan4c, Eif1a and Zscan4b] using quantitative RT-PCR at the one- and two-cell stages. The levels of 1173 1174 these transcripts should increase during ZGA in $Rtcb^{Q+/d^+}$ embryos; 1175 however, their transcriptional activation was significantly suppressed in $Rtcb^{Q-/O^+}$ embryos (Fig. 7J). 1176

In summary, because of the lack of RTCB protein in oocytes,ZGA is suppressed and the global protein translation level is

downregulated, leading to embryonic development arrest. These results indicate that maternal RTCB proteins are important for ZGA and early embryonic development.

DISCUSSION

Among the various AS events, intron retention has been frequently reported in various species. mRNA isoforms that contain introns are not useless. They are temporarily preserved in the cytoplasm and then spliced to synthesize functional proteins during developmental transitions or the cellular stress response. The ligation of these RNAs after intron removal is regulated by the RNA ligase RTCB. According to previous studies, IR occurs in many physiological processes, including the unfolded protein response, tRNA and 16S rRNA splicing (Maughan and Shuman, 2016; Shinya et al., 2011; Tanaka et al., 2011b). However, the role of IR in oocyte meiosis is still vague.

Many maternal mRNAs and proteins stored in the ooplasm of GV stage-arrested oocytes drive oocyte meiosis and early embryonic development. Our study revealed a previously unreported function for RTCB in mouse oocyte maturation. RTCB conditional knockout mice exhibited a phenotype of female infertility, suggesting the important role of RTCB-mediated mRNA splicing in female fertility. The female infertility phenotype can be attributed to multiple factors.

In contrast, the lack of RTCB caused premature ovarian failure in 1235 female mice after 6 weeks of age, and we detected the apoptosis 1236 signal of granulosa cells on ovarian sections at this stage; however, 1237 the ovaries of 4-week-old $Rtcb^{00-/-}$ mice still contained many primordial, primary, secondary and antral follicles. However, 1239 regardless of *in vivo* superovulation or *in vitro* culture, most GV 1240

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1241 oocytes from 6-week-old RTCB knockout mice have difficulty 1242 resuming meiosis. This result was not unexpected because Rtcb 1243 deletion resulted in the accumulation of many IR-containing 1244 mRNAs in the GV stage, including DNA methylation (Mbd1, 1245 Dnmt3b) and DNA damage repair-related mRNA (Rad50). These results imply that RTCB may be involved in regulating follicle 1246 activation, except for the first wave. Combined with changes in the 1247 1248 oocyte transcriptome, premature ovarian failure may be a secondary effect caused by the decline in transcripts enriched in protein 1249 1250 transport and DNA damage repair pathways.

1251 IR accumulation of these transcripts leads to changes in protein levels. RAD50 is crucial for DNA damage repair, and decreased 1252 RAD50 protein levels may lead to GVBD failure (Inagaki et al., 1253 1254 2016). MBD1 and DNMT3B are crucial for regulating DNA 1255 methylation (Hendrich and Bird, 1998; Nowialis et al., 2019). The 1256 decreased levels of related proteins result in a series of epigenetic 1257 changes that impair the developmental potential of the oocyte and 1258 embryo. Although the oocytes that resumed meiosis were able to extrude the polar body and had normal chromosome alignment and 1259 spindle assembly, most of the MII oocytes from Rtcb^{oo-/-} mice had 1260 1261 difficulty with fertilization and ZGA was suppressed. Defects occur 1262 during subsequent fertilization and early embryonic development. 1263 This result illustrates that RTCB may be dispensable for oocyte maturation after the resumption of meiosis. However, epigenetic 1264 1265 changes could affect the developmental potential of oocytes, and the 1266 defects become evident at the zygote stage; therefore, the process of ZGA cannot be completed. GVBD failure, premature ovarian failure 1267 1268 and ZGA defects caused by epigenetic changes in GV stage oocytes 1269 lead to infertility in female mice. These phenotypes also illustrate 1270 that RTCB-regulated IR splicing is important for female 1271 reproduction.

1272 In addition, we investigated the differences between AS types after *Rtcb* deletion in oocytes. Surprisingly, many AS events were 1273 1274 observed, including SEs. However, no relevant studies have 1275 examined the correlation between RTCB and exon skipping. GO enrichment analysis showed that the relevant transcripts were 1276 mainly enriched in the category of RNA processing, which differed 1277 1278 from the transcripts found in IRFinder. These results indicate that 1279 RTCB may act through other mechanisms in oocytes.

1280 Our study found that *Rtcb* deletion caused IR accumulation and 1281 significantly downregulated the global levels of these IR-containing transcripts. In addition, this has not yet been reported. Different 1282 1283 mRNA degradation pathway may cause decreased levels of IR-1284 containing transcripts, which may be because they cannot 1285 synthesize proteins and are degraded by nonsense-mediated 1286 mRNA decay, as the intron retention of these transcripts affects 1287 their open reading frame. However, the decrease in the levels of 1288 other IR-free transcripts remains unclear.

1289 Knocking out the Rtcb gene in mature mouse B cells leads to 1290 structural defects within the endoplasmic reticulum and insufficient 1291 antibody secretion (Jurkin et al., 2014). However, our study found 1292 that *Rtcb* deletion in oocytes led to obstacles in the meiotic process, 1293 premature ovarian failure and loss of fertility. Xbp1 mRNA was the 1294 only confirmed target mRNA of RTCB. The different phenotypes 1295 in germ cells show that *Rtcb* affects tRNA and Xbp1 mRNA, 1296 potentially has other substrates, such as *Rad50* and *Dnmt3b*, and has 1297 an impact on fertility.

1298 Our study examined the role of *Rtcb* across the life of the oocyte 1299 and transition to embryo. IR-containing mRNAs are spliced 1300 appropriately through RTCB to synthesize proteins and perform 1301 their functions. This mechanism can regulate oocyte meiosis and the 1302 subsequent early embryonic development.

MATERIALS AND METHODS

Mice

 $Rtcb^{fl/fl}$ mice were engineered as described previously (Li et al., 2021) and provided by Prof. Nai-Zheng Ding from the College of Life Science, Shandong Normal University, Jinan City, China. To construct oocyte-specific knockout *Rtcb* mice, *Rtcb^{fl/fl}* mice were bred with the *Gdf9-Cre* mouse line to excise loxP-flanked exon 4, thus generating *Rtcb*-conditional-knockout mice.

All mice were bred under specific pathogen-free (SPF) conditions in a stable environment, including 50-70% humidity, 20-22°C, a 12/12 h light/ dark cycle, and food and water provided *ad libitum*. All the mice had a C57BL/6J genetic background. The experimental procedures were approved by the Zhejiang University Institutional Animal Care and Research Committee (approval ZJU20210252 to H.-Y.F.) and mouse care and use were conducted following the relevant guidelines and regulations of Zhejiang University. The primers used for genotyping are listed in Table S4. The age, sex and size of each sample are shown in the related figure legends.

Oocyte culture

Three-week-old female mice were intraperitoneally primed with 5 IU of pregnant mare serum gonadotropin (PMSG) and fully grown oocytes were obtained in M2 medium (M7167; Sigma-Aldrich) 44 h later. These oocytes were cultured in mini-drops of M16 medium (M7292; Sigma-Aldrich) covered with mineral oil (M5310; Sigma-Aldrich) and then incubated at 37° C in a 5% CO₂ atmosphere.

Cell culture

HEK 293T cells were obtained from ATCC and were recently authenticated and tested for contamination. Cells were cultured in DMEM plus 10% fetal bovine serum and 1% penicillin-streptomycin solution at 37°C in a humidified 5% CO_2 incubator.

Superovulation and fertilization

Three-week-old female mice were injected with 5 IU PMSG. These mice were injected with 5 IU of human chorionic gonadotropin (hCG) 44 h later. After an additional 16 h, oocyte-cumulus complexes were separated from the oviducts. Individual oocytes were obtained via digestion with hyaluronidase (Sigma-Aldrich). To acquire early embryos, 12-week-old wild-type males were mated with female mice. Vaginal plugs indicate successful mating. Embryos were obtained from oviducts at the indicated time points after hCG injection.

Plasmid construction and in vitro transcription

Continuously activated CDK1 was constructed by mutating T14 and Y15 of human CDK1 to alanine and phenylalanine, respectively. The corresponding cDNA was then subcloned into a FLAG-tagged expression vector (pDEST). The expression vector was linearized with HindIII and the linearized RNA was transcribed *in vitro* using the SP6 mMESSAGE mMACHINE kit (Invitrogen, AM1450). Phenol/chloroform extraction and ethanol precipitation were performed for RNA extraction and purification, respectively. The transcribed RNAs were *in vitro* polyadenylated using a poly (A) tailing kit (AM1350; Invitrogen).

Microinjection of mRNAs

All microinjections were performed using an Eppendorf TransferMan NK2 micromanipulator. Fully grown oocytes were cultured in M2 medium containing 2 μ M milrinone to suppress spontaneous GVBD. Approximately 5-10 pl of 500 μ g/ml mRNA was microinjected into the cytoplasm of the oocytes. As a negative control, 5-10 pl of 500 μ g/ml transcribed RNA from the empty vector (pDEST-FLAG) was microinjected into the control oocytes. The oocytes were cultured in M16 medium containing 2 μ M milrinone at 37°C in a 5% CO₂ atmosphere.

Quantitative RT-PCR

Total RNA was extracted from oocytes using an RNeasy Mini Kit1362(QIAGEN) according to the manufacturer's instructions. Reverse1363transcription was performed using a Superscript RT kit (Bio-Rad).1364

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Quantitative RT-PCR was conducted using Power SYBR Green PCR 1365 Master Mix (Applied Biosystems, Life Technologies) with an ABI 7500 1366 Real-Time PCR system (Applied Biosystems). The primers used are listed in 1367 Table S4. 1368

1369 Immunofluorescence 1370

Oocytes were fixed with 4% paraformaldehyde (PFA) for 30 min and 1371 permeabilized for 20 min with 0.3% Triton X-100 in PBS. Antibody 1372 staining was performed according to previously described protocols (Zhang 1373 et al., 2020). The antibodies used in these experiments are listed in Table S5. 1374 Imaging was performed using a Zeiss LSM710 confocal microscope. 1375 Quantitative analysis of the fluorescence signals was performed using the 1376 ImageJ software.

1378 **EU** incorporation assay

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Embryos were cultured in KSOM medium with 100 µM 5-ethynyl uridine 1379 (EU) for 2 h. Fixation, permeabilization and staining were performed using 1380 the Click-iT RNA Alexa Fluor 488 Imaging Kit (ThermoFisher, 48 1381 C10329) according to the manufacturer's protocol. The embryos were 1382 imaged using a Zeiss LSM710 confocal microscope. 1383

1384 **Detection of protein synthesis**

1385 Oocytes were incubated in M2 medium primed with 100 mM 1386 L-homopropargylglycine (HPG) for 1 h. These oocytes were then fixed 1387 for 30 min in 4% PFA. HPG signals were examined using a Click-iT HPG 1388 Alexa Fluor Protein Synthesis Assay Kit (Life Technologies). The mean 1389 HPG signal was measured and quantified using ImageJ software.

1391 **Histological analysis**

Ovaries were harvested and fixed in formalin overnight, processed and 1392 embedded in paraffin wax, according to standard protocols. The ovaries 1393 were serially sectioned at 5 µm and stained with Hematoxylin and Eosin. 1394 IHC was performed according to standard protocols. All the antibodies and 1395 their related dilution factors are listed in Table S5. 1396

1397 Western blot analysis

1398 Oocytes were lysed in a loading buffer containing β-mercaptoethanol and 1399 then heated at 95°C for 10 min. Immunoblotting and SDS-polyacrylamide 1400 gel electrophoresis were performed following standard procedures using the 1401 Mini-PROTEAN Tetra Cell System (BioRad). All the antibodies and their 1402 related dilution factors are listed in Table S5.

RNA-seq library preparation and data analysis 1404

1405 Fully grown GV oocytes were collected from 6-week-old wild-type and Rtcbfl/fl;Gdf9-Cre female mice after PMSG (10 fully grown GV oocytes per 1406 140Q4 sample). Ten oocytes with equal amounts of ERCC were subjected to each RNA-seq library construction. Total RNA was extracted using the RNeasy 1408 micro kit (Qiagen) following the manufacturer's instructions after picking 1409 the cells into 350 µl of lysis buffer supplied in the RNeasy micro kit 1410 (Qiagen). Libraries were prepared using a NEBNext Ultra RNA Library 1411 Prep Kit (Illumina). The libraries were sequenced on an Illumina platform 1412 with 150 bp paired-end reads. We used a script to filter out low-quality 1413 reads, and the clean reads were mapped to the mouse genome mm10 with 1414 TopHat (version 2.0.6). Gene expression levels were calculated and normalized to FPKM using cuffquant and cuffnorm. In RNA-seq data 1415 normalization, sequencing reads were mapped to an ERCC reference to 1416 obtain the percentage of ERCC reads in total reads. Gene expression levels 1417 were then normalized by multiplying the raw FPKM values by a 1418 normalization factor (normalization factor=percentage of ERCC in wild-1419 type oocytes/percentage of ERCC in the sample). FPKM and mRNA copy 1420 numbers represent gene expression levels. Only the expressed genes 1421 (FPKM>1 in at least one sample) were considered in all analyses. A two-1422 tailed Student's t-test was used to generate statistically significant values. 1423

1424 **IR change analysis**

1425 Intron retention (IR) was analyzed using IRFinder software. The IR ratio was calculated as the ratio of unspliced exon-intron junction reads to total 1426

junction reads (unspliced exon-intron and spliced exon-exon). The IR 1427 change was calculated from the IR ratio of knockout samples minus that of 1428 wild-type samples. According to IRFinder, only these transcripts (IR change>0.1) exhibited obvious IR accumulation in all the analyses. Changes in IR in genes are shown in Table S1, and the IR number of the filter is shown in Table S2.

Analysis of various AS events

Using replicate multivariate analysis of transcript splicing (rMATS) (Shen et al., 2014), we detected five different AS events, including SE, A5SS, A3SS, MXE and retained intron (RI) (Shen et al., 2014). Various AS events were considered only when FDR<0.05. AS events are listed in Table S3.

Statistical analysis

Results are presented as mean±standard error of the mean (s.e.m.). Most experiments comprised at least three independent samples and were repeated at least three times. The results for the two experimental groups were compared using two-tailed unpaired Student's t-tests. *P<0.05, **P<0.01, ***P<0.001. At P<0.05, results were considered statistically significant. n.s. indicates a non-significant result.

Acknowledgements

We thank the Life Sciences Institute core facilities at Zhejiang University for their technical assistance.

Competing interests

The authors declare no competing or financial interests.

Author contributions

Conceptualization: H.-Y.F.; Methodology: J.-C.J., Y.-W.W., Y.-S.Y., H.-Y.F.; Software: H.Z., Y.-W.W., Y.-S.Y.; Validation: H.Z., J.-C.J.; Formal analysis: J.-C.J., Y.-W.W.; Investigation: H.Z., J.-C.J.; Resources: N.-Z.D.; Data curation: Y.-W.W.; Writing - original draft: H.Z., J.-C.J., N.-Z.D.; Writing - review & editing: H.Z., H.-Y.F.; Visualization: H.Z., J.-C.J., Y.-S.Y., H.-N.W., N.-Z.D., H.-Y.F.; Supervision: H.-N.W., N.-Z.D., H.-Y.F.; Project administration: H.-Y.F.; Funding acquisition: H.-N.W., H.-Y.F.

Funding

This work was supported by the National Key Research and Development Program of China (2021YFC2700100), the National Natural Science Foundation of China (31930031, 31890781 and 31771659; 32072939 to H.-N.W.), the National Ten Thousand Talent Program, the Natural Science Foundation of Zhejiang Province (D22C68649) and the Key Research and Development Program of Zhejiang Province (2021C03098 and 2021C03100).

Data availability

RNA-seq data have been deposited in GEO under accession number GSE190681.

Peer review history

The peer review history is available online at https://iournals.biologists.com/dev/ lookup/doi/10.1242/dev.200497.reviewer-comments.pdf

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1613 Summary: The important role of RNA terminal phosphate cyclase B-regulated non-canonical splicing in female reproduction and theAQ11 mechanisms involved.

S.No.	Funder name	Funder ID	Grant ID
1	National Key Research and Development Program of China		2021YFC2700100
2	National Natural Science Foundation of China	http://dx.doi.org/10.13039/501100001809	31930031 32072939 31771659 31890781
3	National Ten Thousand Talent Program		
4	Natural Science Foundation of Zhejiang Province	http://dx.doi.org/10.13039/501100004731	D22C68649
5	Key Research and Development Program of Zhejiang Province		2021C03098 2021C03100