Theca-Specific Estrogen Receptor-α Knockout Mice Lose Fertility Prematurely

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Estrogen receptor-α (Esr1) mediates estrogen action in regulating at all levels of the hypothalamic-pituitary-ovarian axis. Whereas the importance of Esr1 in hypothalamus and pituitary has been demonstrated by loss of fertility in the neuron- and pituitary-specific Esr1 knockout mice, whether Esr1 plays a critical role in the ovary remains to be determined. In the ovary, Esr1 is mainly expressed in the theca/interstitial cells and germinal epithelium and thus is believed to mediate estrogen action in these cells. In this study, we assessed the importance of Esr1 in the ovarian theca cells in regulating female reproduction. The Cre-loxP approach was used to selectively delete the Esr1 gene in the theca cells, and the reproductive consequence of the deletion was measured. Adolescent theca-specific Esr1 knockout (thEsr1KO) mice (~4 months of age) are fertile and cycling. However, they begin to display an erratic pattern of estrous cycles and become infertile before they reach the age of 6 months. The ovaries of thEsr1KO mice (~4 months) have fewer corpora lutea but more antral follicles than the age-matching wild-type mice. The numbers of 17-hydroxylase-expressing cells are largely increased in the interstitium of the thEsr1KO mouse ovary. Interestingly, whereas basal levels of serum testosterone and FSH were mildly elevated, LH level was either markedly lower or undetectable in the thEsr1KO mice. When superstimulated by exogenous gonadotropins, thEsr1KO mice released significantly fewer oocytes that wild-type littermates and developed multiple hemorrhagic cysts. Taken together, this study demonstrates that theca Esr1 plays a critical role in regulating female reproduction.

In the ovary, Esr2 is mainly expressed in the granulosa cells of growing follicles and has been considered as the predominant ovarian ER form, whereas Esr1 expression is limited to germinal epithelium and theca-interstitial cells. Thus far, three different laboratories have generated at least five different lines of Esr1 knockout (Esr1KO) mice. Interestingly, however, each of the targeted mouse lines displays a different degree of fertility defects (3–9). In contrast, whereas Esr1 has been considered to be the minor ovarian ER form (8, 10–12), Esr1 knockout (Esr1KO) mice have been consistently shown to be completely infertile (3, 13–15).

Abbreviations: Cre, cAMP response element; Cyp17, 17-hydroxylase; Cyp19, aromatase; Esr1, estrogen receptor-α; Esr2, estrogen receptor-β; Esr1KO, Esr1 knockout; hCG, human chorionic gonadotropin; H&E, hematoxylin and eosin; iCre, theca-specific improved form of recombinase; nEsr1KO, neuron-specific Esr1KO; pEsr1KO, pituitary-specific Esr1KO; PMSG, pregnant mare’s serum gonadotropin; StAR, steroidogenic acute regulatory protein; thEsr1KO, theca-specific Esr1 knockout; WT, wild type.
Most interestingly, Esr1KO mice display severe ovarian disorders such as disrupted theca/stromal development, arrest of follicular development at early antral stage, formation of hemorrhagic cysts, and lack of corpora lutea (2, 3, 13–16). Similar pathological disorders were observed in the aromatase knockout mice, demonstrating a critical role of Esr1-mediated estrogen action in regulating ovarian function (17, 18).

Estron regulates female reproduction at all levels of the hypothalamus-pituitary-ovary axis. Recently the tissue-specific gene knockout approach has successfully been applied to demonstrate the Esr1-mediated estrogen action at both hypothalamus and pituitary levels. Deletion of the Esr1 gene at either level resulted in the loss of fertility. In contrast to global Esr1KO mice, milder ovarian defects were seen in the neuron-specific Esr1KO (nEsr1KO) (19) and pituitary-specific Esr1KO (piEsr1KO) mice (20). Folliculogenesis progressed beyond early antral stage without forming hemorrhagic cyst in the ovaries of the nEsr1KO and piEsr1KO mice. Furthermore, upon exogenous gonadotropins stimulation, both nEsr1KO and piEsr1KO mouse ovaries successfully ovulated and formed corpora lutea, indicating that, at some level, the deletion of Esr1 gene in the ovary is directly responsible for the severe ovarian defects seen in the global Esr1KO. We therefore hypothesize that Esr1 plays a critical role at the ovarian level in regulating female reproduction.

In this study, we assessed the reproductive consequence of the ovary-specific deletion of Esr1 gene as a measure of determining the importance of the intraovarian role of Esr1-mediated estrogen action. The floxed Esr1 mouse line that was used for generating piEsr1KO mice and a theca-specific improved form of recombinase (iCre)-expressing mouse line (Cyp17Cre) that we generated recently (21) were used to produce theca-specific E\(\alpha\) knockout (thEsr1KO) mice. Here we report a comprehensive analysis on the reproductive phenotypes of the thEsr1KO.

Materials and Methods

Materials

Pregnant mare’s serum gonadotropin (PMSG) and human chorionic gonadotropin (hCG) were purchased from Sigma (St. Louis, MO). Cell strainers were supplied by Becton Dickinson Falcon (Billerica, MA). Media for experiments were obtained from Life Technologies, Inc. (Grand Island, NY). Molecular reagents were purchased from Invitrogen (Carlsbad, CA).

Generation of thEsr1KO

Two mutant mouse lines were used as founders. The floxed Esr1 (Esr1\textsuperscript{flox/flox}) mouse line, which carries two loxP sites in the introns flanking exon 3 of the Esr1 gene, was used as a target of Esr1 gene excision. This mouse was previously used for generating conventional Esr1KO and piEsr1KO mice (20). The transgenic mouse line, Cyp17Cre, in which an enhanced form of Cre recombinase (iCre)-gene expression is driven by a promoter of 17β-hydroxylase a1, was used to selectively delete Esr1 in the theca cells as was previously described (21). Briefly, a female Esr1\textsuperscript{flox/flox} mouse was first bred with a male Cyp17Cre mouse, which resulted in the production of F1 heterozygotes, Esr1\textsuperscript{flox/+} Cyp17Cre. Then the male F1 Esr1\textsuperscript{flox/+} Cyp17Cre mice were backcrossed with the female Esr1\textsuperscript{flox/flox} mouse, which produced four different genotypes including Esr1\textsuperscript{flox/flox} Cyp17Cre, Esr1\textsuperscript{flox/+} Cyp17Cre, Esr1\textsuperscript{flox/wt} Cyp17Cre, and Esr1\textsuperscript{flox/+}. The Esr1\textsuperscript{flox/+} Cyp17Cre mouse was named thEsr1KO. The genotypes of individual mice were determined by PCR using ear-biopsy DNA. The following primers were used for genotyping the mice: Esr1-P1 (5’-tct ccc gat aac aat aat aac at-3’), Esr1-P2F (5’-ggt tca gaa gag gac aat-3’), Esr1-P3 (5’-ggc att acc tcc ctt ggg agt c-3’), Cre-P1 (5’-ggg ctt ggc cag gga tgt ctc gca ggc g-3’), and Cre-P85 (5’-ggt aaa cag cat tgc tgt cac ttc-3’). Primer combinations of Esr1-P1 + Esr1-P3 and Esr1-P2F + Esr1-P3 were used to determine the presence or absence of loxP sequence (flox or wild type) or deletion of exon 3 (Esr1\textsuperscript{-}). Presence of Cre recombinase was determined using primers Cre-P1 and Cre-P85.

Superovulation assay

Two- and 6-month-old mice were injected with 5 IU PMSG and, 48 h later, the mice were additionally injected with 5 IU hCG. One group of mice was euthanized by CO\(_2\) overdose 48 h after PMSG injection, and the ovaries were collected for histology. The other group of mice was euthanized 22 h after hCG injection, oocyte-cumulus complexes were retrieved from the ampulla of the oviduct, and the number of ova were counted microscopically using a CKX41 inverted microscope (Olympus, Tokyo, Japan) equipped with a digital camera SN H1045062-H (Olympus). The ovaries were immediately fixed in 10% buffered formalin for later histology. The oocyte-cumulus complexes were further examined before and after cumulus cells were removed by using hyaluronidase (80 U/ml) treatment. The quality of oocytes was analyzed according to the standard morphological criteria (22, 23).

Fertility assay

For fertility assays, cycling 45- to 50-d-old Esr1\textsuperscript{flox/flox}, Esr1\textsuperscript{flox/flox} Cyp17Cre, and their heterozygote Esr1\textsuperscript{flox/+} Cyp17Cre were individually housed with a proven Esr1\textsuperscript{flox/-} male. Each cage held one or two females and one male, and their fertility was evaluated by counting the number of litters and litter size per breeding pair. Additionally, variable age-matching female Esr1\textsuperscript{flox/flox} Cyp17Cre and male Esr1\textsuperscript{flox/+} Cyp17Cre mice were bred for a long-term (6–8 months) fertility assay. At the end of the assay, the mice were killed and the tissues were fixed in 10% buffered formalin. Animal handling procedures were carried out in accordance with the University of Kentucky Animal Care and Use Committee. All mice were maintained on a 14-h-light, 10-h-dark cycle and given a continuous supply of food and water.

Determination of estrous cyclicity

Using vaginal lavage techniques, the pattern of estrous cycles was determined in three different age groups of mice (2, 4, and 6 months old) for more than 21 d. Vaginal lavage was performed daily, in the mornings at the same time each day, by flushing the vagina with 0.9% sodium chloride. The cell samples were then examined microscopically and scored. Estrus was determined by the presence of cornified cells. Metestrus was scored by the presence of large round cells with an irregular border. A high density of leukocytes indicated the stage of diestrus, whereas small nucleated cells indicated proestrus (24). The cycling data are expressed as either estrus or nonestrus.

Tissue collection, histology, and immunohistochemistry

For all the female mice used in this study, stage of the estrous cycle was determined and recorded before the animals were killed. Animals were deeply anesthetized and perfused intracardially with PBS, followed by 4% paraformaldehyde. Tissues were then embedded in paraffin blocks, serially sectioned at 4 μm, and stained with hematoxylin and eosin (H&E). Embedding and staining procedures were performed by standard protocols. The histology was examined by BX51 microscope (Olympus) equipped with a digital camera and photography was performed using the MagnaFire-SP imaging system (Olympus). Immunohistochemical staining was performed on deparaffinized sections by the linked streptavidin–biotin complex technique using a linked streptavidin–biotin detection kit (Dako, Carpenteria, CA) with 3-amin-9-ethylcarbazole (Dako) as a chromogen. Mouse monoclonal Esr1 antibody (6F11; Novoceastra, Newcastle, UK) at a dilution of 1:40, antihuman 17-hydroxylase (Cyp17) antibody (kindly provided by Dr. Alan Conley, Uni-
mRNA measurement

Total RNA was isolated from 2- and 6-month-old mice ovaries, which were collected 22 h after hCG injection. Trizol (Invitrogen) and RNeasy kit (QIAGEN Inc., Valencia, CA) were used for purifying the total RNA following standard procedure following manufacturer’s recommendations. Briefly, Moloney murine leukemia virus reverse transcriptase (Invitrogen; 28205-013) was used for cDNA synthesis. Specific CYP17i, aromatase (Cyp19), and steroidogenic acute regulatory protein (StAR) Taqman probe (Applied Biosystems, Foster City, CA) and Taqman gene expression kit was used for quantitative analysis of mRNA, and the PCR was carried out using iQ5 real-time PCR detection system (Bio-Rad, Hercules, CA). As an internal control, glyceraldehyde-3-phosphate dehydrogenase Taqman probe was used. The PCRs were performed using the following conditions: 5 min at 95 °C followed by 40 cycles of denaturation (1.5 sec at 95 °C), annealing, and extension (1 min at 60 °C) with FAM signal reading after extension. For all time points, triplicates of samples were used for the assay.

Statistical analysis

Statistical significances were assessed using Fisher’s exact test for fertility assay and Student’s two-tailed t test for estrous cycle pattern analysis and serum hormone assay and ΔΔct test for analysis of real-time PCR assay. For all statistical analyses, P < 0.05 was considered significant, if not specifically mentioned.

Results

Generation of theca-specific Esr1KO mice

A Cyp17iCre transgenic mouse line that selectively expresses iCre in the theca cells was successively crossed with the floxed Esr1 mouse line that we previously used for generating pEsr1KO mice (20). The genotypes of the resulting offspring were determined by PCR, and the female mice with the genotype of Esr1f/flox/Cyp17iCre were named thEsr1KO (Fig. 1). The deletion of the Esr1 gene in the theca cells of the thEsr1KO mouse was demonstrated by the absence of Esr1-positive staining in the thecainterstitial cells, whereas a normal level of Esr1 expression was seen elsewhere including the oviduct (Fig. 1).

thEsr1KO mice are fertile but lose fertility prematurely

A prolonged preliminary observation on the reproductive activity of the thEsr1KO mice indicated that whereas no noticeable fertility defect was seen in male thEsr1KO mice up to the ages of 6 months, female thEsr1KO showed signs of an age-dependent

![Image](https://academic.oup.com/endo/article-abstract/150/8/3855/2456512)

**TABLE 1. Effect of theca-specific Esr1 deletion on fertility**

<table>
<thead>
<tr>
<th>Genotypes</th>
<th>Age (months)</th>
<th>No. of mice used (n)</th>
<th>No. of delivered litters (%)</th>
<th>No. of weaned litters (%)</th>
<th>No. of pups per weaned littera</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild typeb</td>
<td>2</td>
<td>23</td>
<td>18 (78.2)</td>
<td>12 (52.2)</td>
<td>6.5 ± 0.69</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>24</td>
<td>19 (79.2)</td>
<td>15 (62.5)</td>
<td>7.7 ± 0.63</td>
</tr>
<tr>
<td></td>
<td>6</td>
<td>11</td>
<td>10 (90.9)</td>
<td>7 (63.6)</td>
<td>9.1 ± 1.38</td>
</tr>
<tr>
<td>thEsr1KO</td>
<td>2</td>
<td>14</td>
<td>7 (50.0)</td>
<td>3 (21.4)</td>
<td>2.3 ± 1.20</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>10</td>
<td>3 (30.0)</td>
<td>1 (10.0)</td>
<td>4.0 ± 0.00</td>
</tr>
<tr>
<td></td>
<td>6</td>
<td>10</td>
<td>1 (10.0)</td>
<td>1 (10.0)</td>
<td>2.0 ± 0.00</td>
</tr>
</tbody>
</table>

a Number of pups per litter was calculated by the total numbers of weaned pups divided by the numbers of weaned litters; some litters were lost (mostly by cage by young mother mice) before they reached weaning ages, and therefore the numbers of pups of the litters could not be counted. Fertility of the thEsr1KO mice was significantly affected in the 4- and 6-month (P = 0.015 and 0.0003, respectively) but not in the 2-month-old (P = 0.145) thEsr1KO mice compared with control mice.

b Some heterozygotes (Esr1flox/Cyp17iCre) were included in this group; no fertility difference between wild type and heteros were seen.
fertility loss. We therefore compared reproductive activity in different age groups of thEsr1KO mice by pairing them with proven wild-type (WT) males. Whereas no significant difference was found in adolescent (aged 2–3 months) groups, a dramatic decline of fertility was seen in the 4- and 6-month-old thEsr1KO mice compared with wild-type or heterozygote littermates (Table 1). The fertility defect in these mice was reflected in the aberrant pattern of estrous cycles. No thEsr1KO mouse showed a normal pattern of cycles when they reached the age of 4 months or older but displayed increased days of diestrus (Fig. 2, A and B). This is in contrast to the cycling pattern of age-matching WT mice, which showed a normal pattern of estrous cycles and were fertile, indicating that the thEsr1KO mice experience a significant change in their reproductive function prematurely.

**thEsr1KO mice have decreased ovulatory capacity**

As a measure of assessing the defects in fertility, ovarian phenotypes of 4-month-old WT (Esr1flox/flox), thEsr1KO, and global Esr1KO (Esr1−/−) were examined. Externally, no significant difference was seen between WT and thEsr1KO ovaries, whereas multiple hemorrhagic cysts were apparent in the Esr1KO ovaries (Fig. 2C). Internally, the thEsr1KO ovary showed signs of defects that are intermediate between WT and Esr1KO: reduced numbers of corpora lutea, increased numbers of late preantral and early antral stage follicles, interstitial hypertrophy, and signs of the formation of hemorrhagic cysts but much fewer compared with Esr1KO ovary (Fig. 2C). The ovarian defects of thEsr1KO mice resulted in decreased ovulatory capacity and oocyte quality as was determined by superovulation induction followed by oocyte examination (Fig. 3, A and B). Two- and 6-month-old WT and thEsr1KO mice were injected with PMSG followed by hCG. Twenty-two hours after hCG injection, oocytes were retrieved from oviducts and examined. Whereas the numbers of retrieved oocytes in the 2-month-old thEsr1KO mice were marginally smaller than age-matching WT mice (27 vs. 20 oocytes), significantly fewer numbers of oocytes were released in the 6-month-old thEsr1KO mice than age-matching WT mice (22 vs. six oocytes) (Fig. 3A). Furthermore, whereas oocytes from WT mice were healthy and had well-defined germinal vesicles, the majority of oocytes from thEsr1KO mice were unhealthy and had well-defined germinal vesicles, the majority of oocytes from thEsr1KO mice displayed signs of degeneration (Fig. 3B). Furthermore, the gonadotropin injection (PMSG alone or PMSG + hCG) resulted in the formation of hemorrhagic follicle in the thEsr1KO ovaries, whereas hemorrhage was rarely seen in the WT ovaries (Fig. 3, C and D).

**thEsr1KO mice have higher serum testosterone and lower LH levels**

Previous studies (25, 3) suggested that ovarian androgen synthesis is negatively regulated by the paracrine action of estrogen, in which estradiol produced by granulosa cells directly acts on Esr1 in the theca cells to suppress Cyp17 transcription and therefore testosterone production. The deletion of Esr1 specifically in the Cyp17-expressing cells of the thEsr1KO mice offers an outstanding experimental condition...
to test the hypothesis. As an initial step, serum levels of sex steroids and gonadotropins were measured in the 2- and 6-month-old WT and thEsr1KO female mice (Fig. 4, A–C). Whereas no significant difference in estradiol level was seen (5–16 pg/ml range in WT and 5–26 pg/ml range in thEsr1KO mice), testosterone level in the thEsr1KO mice was moderately increased numbers of gonadotrophs (5.5 LH

Interestingly, pituitary of thEsr1KO mice had undetectable (6 month groups) in the thEsr1KO mice (Fig. 4, A–C). It has long been speculated that estrogen plays critical roles in the ovary. However, due to the complexity of the mechanism involved, the exact role of estrogen in the ovary remains unclear. Recent studies have shown that estrogen has a direct effect on folliculogenesis and ovulation. However, the mechanism by which estrogen affects these processes is not fully understood.

Discussion

It has long been speculated that estrogen plays critical roles in the ovary. However, due to the complexity of the mechanism involved, the exact role of estrogen in the ovary remains unclear. Recent studies have shown that estrogen has a direct effect on folliculogenesis and ovulation. However, the mechanism by which estrogen affects these processes is not fully understood.

FIG. 3. Superovulation in thEsr1KO mice. Two- and 6-month-old WT and thEsr1KO mice were superovulated using PMSG (5 IU/mouse) and hCG injections (5 IU/mouse), and the numbers of oocytes were retrieved from ampulla at hCG 22 h and examined. A, Number of retrieved oocytes (n, numbers of mice used for each group). *, P < 0.05 vs. age-matching WT mice. B, Oocyte-cumulus complex (left panels) retrieved from the oviducts and cumulus-free oocytes (right panels) after hyaluronidase treatment. Note the well-expanded cumulus mass from WT but dense clumps of cumulus mass from thEsr1KO mice ovaries. C, External morphology of ovaries of 2- and 6-month-old WT and thEsr1KO mice. D, H&E staining of 6-month-old WT and thEsr1KO mice ovaries. Note that whereas multiple numbers of corpora lutea are seen in the WT and thEsr1KO mice ovaries, blood filled-cystic follicles are seen only in the thEsr1KO mice. Arrows indicate the hemorrhagic follicles.

No dramatic difference of steroidogenic enzyme gene expressions but highly populated theca cells in the interstitium of thEsr1KO ovary

To further investigate the cause of the elevation of testosterone level in the thEsr1KO mice, the expression patterns of steroidogenic enzymes involved in the androgen synthesis were analyzed. To overcome any existing physiological differences that may affect steroidogenic gene expression at the time of assay, a superovulation regimen was used for the assay. Two- and 6-month-old thEsr1KO mice and their littersmates were induced superovulation by PMSG and hCG injections. Twenty-two hours after hCG injection, mRNA expression levels of Cyp17, aromatase (Cyp19), and StAR were measured by quantitative PCR using total RNA extracted from whole ovarian homogenates (Fig. 5). Whereas not dramatically different, Cyp17 mRNA expression levels were higher in thEsr1KO mice in both age groups. No significant difference in Cyp19 mRNA expression was seen between the two genotypes, whereas the expression levels were elevated in the aged (6 month) groups. StAR mRNA levels were not different in the 2-month group, but the level was significantly lower in the thEsr1KO mice of the aged group. After finding out that there was not a dramatic difference of the key steroidogenic gene mRNA expressions, we tested whether there was a change in the number of testosterone-producing cells (Cyp17 positive cells). An immunohistochemical examination on the superovulated 6-month group of mouse ovaries revealed that thEsr1KO ovaries were heavily populated with Cyp17-positive cells (Fig. 6A). Interestingly, the majority of the theca cells in the thEsr1KO ovaries were found in interstitial areas in contrast to the WT ovary in which Cyp17-positive cells are mostly localized in the theca interna of preovulatory follicles. To determine whether the increase of theca cells is a general phenotype of thEsr1KO mice, we then examined the ovaries of younger groups (4 months old) and older groups (10 months old) on the day of diestrus when the Cyp17 expression was supposed to be minimal. As was in the gonadotropin-injected mouse ovaries, more cyp17-positive cells were seen in the ovaries of thEsr1KO than WT mice (Fig. 6B).
of estrogen action and the near-ubiquitous expression of estrogen receptors, it has been challenging to determine the intracellular role of estrogen. In this study, we determined the physiological significance of Esr1 expression in the theca cells by using a tissue-specific gene knockout approach. By cross-breeding of floxed Esr1 mice and a newly generated Cyp17iCre mouse line, theca cell-specific deletion of Esr1 gene was achieved. Reproductive outcome, pattern of estrous cycles, ovarian morphology, and ovariocytic response to gonadotropin stimulation of the thEsr1KO mice in comparison with WT mice were used as a measure of the significance of the Esr1 expression in the theca cell.

The Cyp17-iCre driven deletion of Esr1 gene lead to sex-dependent fertility outcome: whereas the majority of female thEsr1KO mice lose fertility before they reach the ages of 6 months, male counterpart (Leydig cell-specific Esr1KO; LeyEsr1KO) mice do not show any apparent sign of fertility loss up to the ages of 6 months (data not shown). This sex-dependent reproductive performance indicates that Esr1 plays a role critically related to fertility regulation in the ovary but not the testis. This finding is not surprising because it was shown the fertility defect in the global Esr1KO male mice were not caused by testicular defects but epididymal malfunction (15). However, considering the fact that Esr1 is expressed in the Leydig cells of the testis (26–28) and aging global Esr1KO mice displayed testicular hypotrophy (15), future analysis on LeyEsr1KO mouse testis is warranted.

In the case of females, thEsr1KO mice are born fertile but gradually lose fertility leading to near complete infertility by the age of 6 months, when the age-matching WT females maintain their fertility at the highest level (Table 1). The loss of fertility in the thEsr1KO female mice is accompanied by the loss of estrous cyclicity (Fig. 2, A and B), a physiological presentation of cyclic changes of circulating estrogen levels (29). It is presumed, therefore, that the steroidogenic activity is altered in the thEsr1KO mice. In fact, the main functional role of ovarian theca cells is to synthesize androgens (testosterone and androstenedione) that in turn serve as the substrate of Cyp19 aromatase for estrogen production in the granulosa cells. Whereas comparison of steroid levels at different stages of estrous cycles in cycling WT and thEsr1KO mice would be a desirable way to determine the role of Esr1 in theca cell steroidogenesis, this approach was not applicable due to the acyclic nature of the thEsr1KO mice (Fig. 2, A and B). Therefore, alternatively, the serum levels of steroids on the mornings of diestrus were measured along with gonadotropin levels. Concurrent with the previous reports (2, 25), moderately elevated estradiol, testosterone (Fig. 4A), and FSH (Fig. 4B) levels were seen. Surprisingly, however, LH level was significantly lower in the thEsr1KO or undetectable (Fig. 4C). Whereas the present study did not find the direct cause of the gradual loss of fertility in the thEsr1KO female mice, both the aberrant levels of steroids and gonadotropins are believed to be responsible. In regard to the lower basal level of circulating LH, we found, unexpectedly, increased numbers of gonadotrophs with approximately similar level of LH content in each cell in the pituitary in the thEsr1KO pituitary as determined by immunohistochemistry using anti-LHβ antibody (Fig. 4, D and E), indicating that secretion of LH is compromised by the Esr1 deletion in the theca cells but not the hormone synthesis. The potential cause of defect in LH secretion is currently under investigation.

Unexpectedly, ovaries of the gonadotropin-injected thEsr1KO mice developed hem-
Positive cells were seen in the ovaries of thEsr1KO than WT mice. A, Antral thEsr1KO mice were examined on the day of diestrus. Large numbers of Cyp17-internal of preovulatory follicles. B, Ovaries of 4- and 10-month-old WT and contrast to the WT ovary in which Cyp17-positive cells are mostly in the theca of the theca cells in the thEsr1KO ovaries were found in interstitial areas in

Population of Cyp17-positive cells after gonadotropin stimulation. A, Ovaries of 6-month-old WT and thEsr1KO mice were collected 48 h after PMSG and 22 h after hCG injection, formalin fixed, paraffin embedded, and stained with anti-Cyp17 antibody. Note that the Cyp17-positive cells (dark brown color) are heavily populated in the ovaries of 6-month-old thEsr1KO mice. The majority of the theca cells in the thEsr1KO ovaries were found in interstitial areas in contrast to the WT ovary in which Cyp17-positive cells are mostly in the theca interna of preovulatory follicles. B, Ovaries of 4- and 10-month-old WT and thEsr1KO mice were examined on the day of diestrus. Large numbers of Cyp17-positive cells were seen in the ovaries of thEsr1KO than WT mice. af, Antral follicle; CL, corpus luteum; cf, cystic follicle; pf, preantral follicle.

FIG. 6. Population of Cyp17-positive cells after gonadotropin stimulation. A, Ovaries of 6-month-old WT and thEsr1KO mice were collected 48 h after PMSG and 22 h after hCG injection, formalin fixed, paraffin embedded, and stained with anti-Cyp17 antibody. Note that the Cyp17-positive cells (dark brown color) are heavily populated in the ovaries of 6-month-old thEsr1KO mice. The majority of the theca cells in the thEsr1KO ovaries were found in interstitial areas in contrast to the WT ovary in which Cyp17-positive cells are mostly in the theca interna of preovulatory follicles. B, Ovaries of 4- and 10-month-old WT and thEsr1KO mice were examined on the day of diestrus. Large numbers of Cyp17-positive cells were seen in the ovaries of thEsr1KO than WT mice. af, Antral follicle; CL, corpus luteum; cf, cystic follicle; pf, preantral follicle.

orrhagic follicles, whereas such response was not induced in the WT ovaries (Fig. 3, C and D). Whereas hemorrhagic cysts were seen in not all but some unstimulated thEsr1KO ovaries, all of the gonadotropin-injected mouse ovaries had the hemorrhagic follicles, demonstrating that the gonadotropin is the direct cause of the hemorrhagic response in the thEsr1KO mouse ovary. Whereas a cohort of previous studies showed that either the global deletion of aromatase or Esr1 resulted in the formation of hemorrhagic cysts, whereas less number of oocytes was released from thEsr1KO ovary, indicating that the Esr1 may play a critical role in antral stage folliculogenesis. The possibility of Esr1 in the theca cell proliferation and differentiation is being sought. Taken together, the present study demonstrates that the theca Esr1 plays a critical role in regulating female reproduction.

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