mTORC1/2 inhibition preserves ovarian function and fertility during genotoxic chemotherapy

Kara N. Goldman\textsuperscript{a}, Devon Chenette\textsuperscript{b}, Rezina Arj\textsuperscript{b}, Francesca E. Duncan\textsuperscript{c}, David L. Keefe\textsuperscript{a}, Jamie A. Grifo\textsuperscript{a}, and Robert J. Schneider\textsuperscript{b,d,1}

\textsuperscript{a}Division of Reproductive Endocrinology and Infertility, Department of Obstetrics and Gynecology, New York University School of Medicine, New York, NY 10016; \textsuperscript{b}Department of Microbiology, New York University School of Medicine, New York, NY 10016; \textsuperscript{c}Department of Obstetrics and Gynecology, Feinberg School of Medicine, Northwestern University, Chicago, IL 60611; and \textsuperscript{d}Perlmutter Cancer Center, New York University School of Medicine, New York, NY 10016

Edited by Nahum Sonenberg, McGill University, Montreal, QC, Canada, and approved February 8, 2017 (received for review October 17, 2016)

The ovary contains oocytes within immature (primordial) follicles that are fixed in number at birth. Activation of follicles within this fixed pool causes an irreversible decline in reproductive capacity, known as the ovarian reserve, until menopause. Premenopausal women undergoing commonly used genotoxic (DNA-damaging) chemotherapy experience an accelerated loss of the ovarian reserve, leading to subfertility and infertility. Therefore, there is considerable interest but little effective progress in preserving ovarian function during chemotherapy. Here we show that blocking the kinase mammalian/mechanistic target of rapamycin (mTOR) with clinically available small-molecule inhibitors preserves ovarian function and fertility during chemotherapy. Using a clinically relevant mouse model of chemotherapy-induced gonadotoxicity by cyclophosphamide, and inhibition of mTOR complex 1 (mTORC1) with the clinically approved drug everolimus (RAD001) or inhibition of mTORC1/2 with the experimental drug INK128, we show that mTOR inhibition preserves the ovarian reserve, primordial follicle counts, serum anti-Mullerian hormone levels (a rigorous measure of the ovarian reserve), and fertility. Chemo- therapy-treated animals had significantly fewer offspring compared with all other treatment groups, whereas cotreatment with mTOR inhibitors preserved normal fertility. Inhibition of mTORC1 or mTORC1/2 within ovaries was achieved during chemotherapy cotreatment, concomitant with preservation of primordial follicle counts. Importantly, our findings indicate that as little as a two- to fourfold reduction in mTOR activity preserves ovarian function and normal birth numbers. As everolimus is approved for tamoxifen-resistant or relapsing estrogen receptor-positive breast cancer, these findings represent a potentially effective and readily accessible pharmacologic approach to fertility preservation during conventional chemotherapy.

mTOR | mTORC1 | mTORC2 | ovarian follicles | primordial follicles | chemotherapy | gonadotoxicity | ovarian function | fertility

**Significance**

A major unresolved issue for premenopausal women undergoing chemotherapy is infertility due to the loss of non-renewable ovarian primordial follicles. We show that pharmacologic down-regulation of the mammalian/mechanistic target of rapamycin (mTOR) pathway during chemotherapy in a mouse model prevents activation of primordial follicles, preserves ovarian function, and maintains normal fertility using clinically available inhibitors of mTOR complex (C1) and mTORC1/2. These findings represent a feasible pharmacologic approach for preservation of ovarian function and fertility during treatment with conventional chemotherapy.
Activation of the mammalian/mechanistic target of rapamycin (mTOR) pathway is critical to primordial follicle activation (13–15). mTOR is a serine/threonine kinase and a metabolic sensor that regulates mRNA translation, cell growth, proliferation, autophagy, nutrient signaling, and survival (16). Accelerated mTOR activity in the oocyte by deletion of PTEN- or TSC1-negative regulators simultaneously activates the entire pool of primordial follicles in mice, resulting in primary ovarian insufficiency (13, 14, 17). Thus, mTOR stimulators increase the activation of primordial follicles in animal models and mTOR inhibitors block the primordial-to-primary follicle transition (18), highlighting the critical importance of the PI3K/AKT/mTOR pathway (19).

Results

CY Chemotherapy Reduces Primordial Follicle Counts in a Dose-Dependent Manner and Activates the Transition to Growing Follicles. To establish the effects of chemotherapy regimens on primordial follicles, we conducted a dose-finding pilot study of CY exposure. Mice received weekly i.p. injections of vehicle (normal saline), 75 mg/kg CY, or 150 mg/kg CY over 3 wk followed by sacrifice 1 wk following the final dose. No gross abnormalities in the animals were identified at necropsy and by pathohistologic examination. No significant differences in weight at sacrifice were found between treatment groups (Fig. S2A). Because growing follicles may be overrepresented when serial ovarian sections are analyzed and estimations of follicle counts have been shown to be equivalent when whole ovaries are serially sectioned compared with interval sectioning (20), ovaries were sectioned in five 100-μm intervals (interval sectioning). The mean number of primordial follicles per ovarian surface area (mm²), a standard measure of gonadotoxicity, was inversely related to treatment dose, with the untreated control group having the highest average number of primordial follicles per mm² (3.5 ± 0.5) compared with a conventional dose of 75 mg/kg CY (1.4 ± 0.2) and a sterilizing dose of 150 mg/kg CY (0.2 ± 0.1) (Fig. S2B).

To determine the acute impact of CY exposure on acute ovarian follicle dynamics, we treated mice with one high dose of CY (150 mg/kg) followed by characterization at 24 h postexposure. There were no differences in untreated and treated mean weight, mean ovarian weight, or ovarian surface area. However, CY-treated mice had significantly fewer (60% reduction) primordial follicles than control mice (Fig. 1A). CY treatment also induced rapid follicle activation, demonstrating 2.5 times more growing follicles per mm² than untreated controls (Fig. 1B and C).

mTOR Inhibitors Down-Regulate the PI3K/AKT/mTOR Pathway in Ovaries and Preserve Primordial Follicles When Administered CY. Mice were treated with CY, with or without mTOR inhibitors, as shown in the schema for treatment, and tissues were harvested (Fig. S3). CY treatment moderately increased AKT/mTOR pathway activation, shown by phosphorylation of S473 AKT and S65 4E-BP1 to P-AKT and P-4E-BP1, respectively, compared with untreated controls, measured within 2 h of the last daily mTOR inhibitor treatment (Fig. 2A). Inhibition of mTOR complex 1 (mTORC1) with everolimus (RAD001) or mTORC1/2 with INK reduced phosphorylation of 4E-BP1 and S6 kinase (S6K), with and without CY. Inhibition of AKT phosphorylation was greater in ovaries exposed to INK than RAD. mTORC1/2 inhibition by INK was also more effective in blocking phosphorylation of all downstream targets, consistent with the weaker activity of allosteric RAD (21). Phosphorylation of AKT and S6K was effectively blocked in ovaries exposed to CY with RAD or INK, more so than with mTOR inhibitors alone, suggesting possible synergism (Fig. S4).

Following immunoblotting of whole-ovary lysates, immunohistochemistry (IHC) was used to assess the activity of mTORC1 and mTORC1/2 within primordial follicles using 4E-BP1 and S6K phosphorylation within 2 h of the last inhibitor dose. Following treatment with CY and mTORC1/2 (INK) inhibition, primordial follicles averaged fourfold reduced mTOR activity compared with approximately twofold reduction for mTORC1-only inhibition (RAD) (Fig. 2B–E). Overall, INK was a more effective mTORC1 inhibitor than RAD, even in non–CY-treated controls. RAD efficacy was tested by examining S6K phosphorylation and, in contrast to 4E-BP1 phosphorylation, inhibition of mTORC1-only (RAD) reduced S6K phosphorylation by twofold in follicles of CY-treated animals compared with fourfold for mTORC1/2 inhibition (Fig. 2D and E).

Using conditions established above, mTOR inhibitors were treated with and without CY to assess the effects of cotreatment on follicle counts (schema, Fig. S3). Female mice were randomized into six groups (n = 5 per group) and treated with 75 mg/kg CY in three weekly doses with or without RAD or INK administered by daily oral gavage, followed by sacrifice 1 wk after the final dose of chemotherapy. Markers of toxicity were compared, including mouse weights (g) pre- and posttreatment, ovarian surface area (mm²), and ovarian weight (μg) (Fig. S5). There were no differences between groups when comparing ovarian surface area at sacrifice or ovarian weight at sacrifice (Fig. S5A and B). Mouse weight was similar at baseline between all groups with the exception of the RAD+CY group, which had an average weight of 7.9% less than control at baseline (16.4 ± 0.1 vs. 17.7 ± 0.5, P = 0.03) (Fig. S5C). All mice treated with mTOR inhibitors weighed significantly more at sacrifice than they did at baseline, a phenomenon not seen in mice treated with CY or in untreated controls (Fig. S5C). Mice treated with 75 mg/kg CY alone had a 64% reduction in primordial follicles per mm².
were fewer primary follicles per mm² with mTORC1 or mTORC1/2 inhibition when cotreated with CY, compared with mTORC1 inhibition alone, supporting a synergistic effect of cotreatment on prevention of follicle activation (Fig. 3C). Secondary follicle counts and antral follicle counts were not statistically different among all treatment groups, despite a trend toward fewer antral follicles in the 75 mg/kg CY group (Fig. 3D and E). Of note, total follicle counts were lower in all CY-treated mice compared with control, RAD alone, and INK alone, suggesting that mTOR inhibitors incompletely prevented primordial follicle activation and loss. This is likely secondary to the timing of administration (2 d before CY) and time needed to achieve steady-state concentrations, which is ∼7 d (22, 23) (Fig. S6). Strikingly, ovaries of CY-treated mice demonstrated a ratio of growing to primordial follicles more than twice that of ovaries from mice cotreated with CY+RAD or CY+INK (Fig. 3F), supporting our finding that mTOR inhibitors maintain ovarian quiescence and preserve the primordial follicle pool. The histological follicle counts also substantiate the molecular data, indicating that more than twofold, but no more than fourfold, down-regulation of mTORC1 and/or coinhibition of mTORC2 is all that is required to maintain ovarian follicle quiescence.

**mTOR Inhibition Prevents Chemotherapy-Mediated Reduction in Serum Anti-Mullerian Hormone in a Dose-Dependent Manner.** Anti-Mullerian hormone (AMH) is produced by the granulosa cells of preantral and small antral follicles, correlates with histological primordial follicle numbers, and is one of the most important measures of ovarian reserve used clinically (24, 25). To investigate the impact of CY treatment on serum AMH, 8-wk-old mice were administered 75 mg/kg CY, 150 mg/kg CY, or vehicle (control) weekly for 3 wk and killed 1 wk following the final treatment. Untreated mice had significantly higher levels of serum AMH compared with 75 mg/kg CY-treated animals, which declined further at 150 mg/kg CY (Fig. 4A), indicative of chemotherapy-driven loss of follicles. Using conditions established above, mTOR inhibitors were treated with and without CY, and effects of cotreatment on serum AMH were assessed. There was no statistically significant difference in circulating AMH levels between untreated controls and RAD- or INK-treated groups in the absence of CY treatment (Fig. 4B). However, mice cotreated for 3 wk with weekly 75 mg/kg CY and daily RAD, and killed at 11 wk of age, showed significantly higher AMH levels compared with mice treated with CY alone (Fig. 4C). Importantly, even a small degree in AMH levels reflects a clinical reduction in ovarian reserve (26). The same positive trend in AMH levels existed with INK+CY treatment but did not reach statistical significance.

**mTOR Inhibitors Preserve Fertility in Chemotherapy-Treated Mice.** We next investigated whether mTOR inhibition also preserves fertility during chemotherapy. Eight-week-old female mice were treated with a nonsterilizing dose of 75 mg/kg CY in four weekly doses, with or without RAD or INK cotreatment. One additional week of treatment was used to further impact on subfertility, providing even more stringent criteria for the assessment of fertility than used earlier. To ensure two full cycles of primordial follicle activation following treatment, breeding commenced 8 wk following the final treatment. Mice were harem-bred with proven male breeders and given 8 wk to breed (Fig. S3). Studies have shown that rodents are capable of normal mating behavior as early as 14 d following CY exposure (27).

There was no difference in mouse weight following 4 wk of treatment (P > 0.05) (Fig. S7). After an 8-wk delay, all mice had incrementally gained weight and there were no differences in mouse weights between groups, supporting no differences in systemic toxicity between groups (P > 0.05) (Fig. S7). Two out of five female CY-only treated mice were infertile, compared with no infertile females in the RAD+CY and INK+CY groups. The CY-treated animals also had significantly fewer pups per litter compared with controls (3.4 ± 1.7 vs. 8.8 ± 0.5, P < 0.005) (Fig.
Mice treated with 75 mg/kg CY i.p. in three weekly doses, ±RAD or INK daily, and killed 1 wk after the final dose of CY. CY-treated mice were 64% reduced in primordial follicles compared with untreated controls. Mice treated with 75 mg/kg CY and RAD had 43% more primordial follicles compared with CY alone (P < 0.05). Mice treated with CY and INK were 54% increased in primordial follicles compared with CY alone (**P < 0.005). Representative images are shown. (C) Mice cotreated with RAD+CY or INK+CY tended toward fewer primary follicles compared with CY alone (n.s., not significant; P > 0.05). (D and E) Secondary follicle and antral follicle counts were not statistically different among treatment groups despite trending toward fewer antral follicles in the 75 mg/kg CY group (n.s., P > 0.05). (F) Ovaries of CY-treated mice had twice the ratio of growing to primordial follicles compared with all other treatment groups (**∗∗∗P < 0.0001). Ovaries of mice cotreated with CY+RAD or CY+INK had ratios of growing to primordial follicles matching untreated controls. Results are derived from five mice per treatment group with SEM shown.

5.4). Importantly, RAD+CY (7.4 ± 1.2) and INK+CY (7.4 ± 0.9) treatment groups produced significantly more pups per litter compared with CY alone (3.4 ± 1.7, P < 0.05). There were no differences in litter size between RAD alone, INK alone, and untreated controls. There were no differences in the percentage of pups born live, pup weight, or pup anomalies among treatment groups (Fig. 5 B and C). The time from the start of breeding (days) to birth was similar between groups, arguing against altered endocrine effects (Fig. 5D). There was a trend toward increased time from male introduction to birth in the CY group compared with untreated controls, which did not reach significance.

**Discussion**

Inhibitors of mTORC1 and mTORC1/2 have a growing role in cancer treatment as part of multiagent chemotherapeutic regimens and are being explored for the treatment of a growing list of malignant as well as nonmalignant conditions. Here we further elucidated the critical role of the mTOR pathway in primordial follicle activation and demonstrated that mTOR inhibitors can have a second significant function in promoting follicular quiescence when administered as pretreatment and cotreatment with conventional gonadotoxic chemotherapy. We show that daily administration of mTOR inhibitors that achieve only a two- to fourfold attenuation of mTOR activity can maintain ovarian follicles in their primordial state during chemotherapy treatment, maintain normal serum AMH levels, and preserve normal fertility. Although estrus cyclicity was not assessed, our data do not support an interpretation involving an effect of treatments on estrus cyclicity. The impact of CY on fertility was therefore related to ovarian reserve rather than ovulation. In the mouse, the cycle of follicle maturation requires ~10 to 12 d from primordial follicle to secondary follicle, and an additional 6 to 12 d for development to the antral follicle (28) (see Fig. S8 for follicle classifications). We obtained tissue and serum 1 wk after the third and final treatment dose, so the difference observed in AMH level likely reflects the impact of the first 1 to 2 wk of treatment on the primordial follicle pool. Primordial follicle counts were significantly greater in the RAD+CY group and INK+CY group compared with CY alone, and AMH levels were significantly greater in RAD+CY compared with CY. The actual AMH values were identical between RAD+CY and INK+CY groups, but AMH did not reach significance between the INK+CY and CY groups. This may be attributed to variance but invites further investigation. Whereas primordial follicle counts were significantly greater in mice cotreated with mTOR inhibitors and CY compared with CY alone, total follicle counts were lower among all CY-treated mice compared with mice not exposed to CY. Follicular burnout is likely one of many processes occurring in the ovary during chemotherapy exposure, and growing follicles, vasculature, and ovarian stroma are also susceptible to chemotherapy effects. Importantly, although oral RAD001 is rapidly absorbed, steady-state concentrations are not reached for ~7 d (22). In our study, mTOR inhibitors were administered for 2 d before the first exposure to chemotherapy, likely insufficient to prevent early recruitment and subsequent loss of primordial follicles during the first CY administration. Future studies will be needed to determine the optimal timing of mTOR inhibitor administration before chemotherapy.

Our data suggest that mTOR inhibitors may represent a fertility-sparing pharmacologic therapy that can be administered alongside gonadotoxic chemotherapy. Oocyte or embryo cryopreservation are proven methods of fertility preservation for peri- and postmenarchal females but are time-consuming, costly, may be medically contraindicated, and preserve only a limited number of gametes or embryos with no maintenance of ovarian function. Ovarian tissue cryopreservation is still an experimental technique and requires an initial surgery to remove ovarian tissue followed by subsequent surgery for autotransplantation, and may risk reintroducing malignant cells (29).

**Fig. 3.** Cotreatment with mTORC1 and mTORC1/2 inhibitors protects the primordial follicle pool in CY-exposed ovaries. (A and B) Eight-week-old C57BL/6 mice were treated with 75 mg/kg CY i.p. in three weekly doses, ±RAD or INK daily, and killed 1 wk after the final dose of CY. CY-treated mice were 64% reduced in primordial follicles compared with untreated controls. Mice treated with 75 mg/kg CY and RAD had 43% more primordial follicles compared with CY alone (P < 0.05). Mice treated with CY and INK were 54% increased in primordial follicles compared with CY alone (**P < 0.005). Representative images are shown. (C) Mice cotreated with RAD+CY or INK+CY tended toward fewer primary follicles compared with CY alone (n.s., not significant; P > 0.05). (D and E) Secondary follicle and antral follicle counts were not statistically different among treatment groups despite trending toward fewer antral follicles in the 75 mg/kg CY group (n.s., P > 0.05). (F) Ovaries of CY-treated mice had twice the ratio of growing to primordial follicles compared with all other treatment groups (**∗∗∗P < 0.0001). Ovaries of mice cotreated with CY+RAD or CY+INK had ratios of growing to primordial follicles matching untreated controls. Results are derived from five mice per treatment group with SEM shown.

**Fig. 4.** Serum AMH decreases after CY treatment whereas mTOR inhibitor cotreatment maintains AMH concentration. (A) AMH serum concentrations were measured in untreated 8-wk-old C57BL/6 mice, mice exposed to three weekly doses of 75 mg/kg CY, and mice exposed to three weekly doses of 150 mg/kg CY. Mice treated with 75 mg/kg CY had significantly lower serum AMH levels compared with control (**P < 0.005), as did mice treated with 150 mg/kg CY (P < 0.05). (B and C) AMH serum concentrations were measured in 8-wk-old mice that underwent long-term treatment (3 wk) with polysilanylglycrolidone (PVG) alone daily, 75 mg/kg CY weekly for 3 wk, RAD daily, RAD daily plus 75 mg/kg CY weekly, INK daily, or INK plus 75 mg/kg CY weekly. There were no differences in AMH concentrations in mice treated with RAD or INK alone compared with untreated controls. Mean ± SEM. Mice cotreated for 3 wk with weekly 75 mg/kg CY and daily RAD and killed 1 wk after the final CY treatment had a significantly higher AMH compared with CY alone (12.5 ∼ vs. 11.1, P > 0.05). Mice cotreated for 3 wk with weekly 75 mg/kg CY and daily INK and killed 1 wk after the final CY treatment had a higher AMH level compared with CY alone but this did not reach significance (12.6 vs. 11.1, n.s., P > 0.05). Results are derived from five mice per treatment group with SEM shown.
Intriguingly, in endometriosis, which contributes to diminished ovarian reserve and an accelerated decline in fertility (1), enhanced follicular recruitment and burnout are observed in the ovarian cortex of ovaries with endometriomas (30). The insulin growth factor system, which regulates signaling through pathways including AKT/mTOR, is altered in endometriotic tissues, suggesting a similar relationship between mTOR activation and follicular burnout in endometriosis (31). Data implicate mTOR up-regulation in the poor reproductive outcomes of overweight and obese patients, who are more likely to be infertile, have higher miscarriage rates, and poorer obstetric outcomes compared with normal-weight peers (32–34). Obese rats demonstrate accelerated ovarian follicle development and follicle loss and activation of the mTOR pathway (35), whereas SIRT1 activators, which suppress mTOR, improve the ovarian reserve (36). Caloric restriction in rats also down-regulates mTOR activity and preserves the ovarian reserve (37). mTOR inhibitors may be attractive for prevention of iatrogenic and noniatrogenic depletion of ovarian reserve (31).

Beyond the important scope of fertility, primary ovarian insufficiency has devastating emotional, psychosocial, and physical consequences, contributing to depression, cognitive dysfunction, bone loss, sexual dysfunction, and even cardiovascular mortality, prevention of which all require maintenance of normal ovarian function (2, 38–40). Moreover, with the continuing rise in average age at first birth in the United States and the reciprocally reduced fertility rate (41, 42), the possibility of extending reproductive potential is attractive. Investigating the relationship between mTOR and the ovarian reserve may provide opportunities for therapeutic intervention. Because the PI3K/AKT/mTOR pathway plays a critical role in primordial follicle activation (13, 43), this relationship has been investigated as a means to promote follicle growth in patients with POI. In murine models, inhibitors of PTEN (a negative regulator of PI3K) and TSC1 and TSC2 (negative regulators of mTOR) have been shown to activate the mTOR pathway and primordial follicles (13, 14, 17). Similarly, AKT stimulators (both PI3K activators and PTEN inhibitors) have been used in both preclinical and experimental clinical models to activate in vitro cultured ovarian cortical strips, retransplant the activated cortical strips, and stimulate follicle growth in vivo (44). In a prospective study of 37 women with primary ovarian insufficiency who underwent in vitro activation of ovarian cortical tissue with an AKT stimulator followed by in vitro fertilization, 24 oocytes were retrieved from six patients, ultimately leading to two successful deliveries (44, 45). Moreover, Rictor/mTORC2 was recently implicated as a key regulator of folliculogenesis, with inactivation of this pathway ubiquitously leading to premature ovarian failure (15).

Studies are needed to assess the long-term reproductive impact of mTOR inhibitors, and specifically the impact on future pregnancies and offspring. Although few large studies exist regarding the impact of mTOR inhibitors on long-term fertility and pregnancy outcomes, case series and observational studies describe fertility outcomes in patients who have been treated with rapalogs following solid-organ transplant. Following kidney transplant, women treated with long-term sirolimus (13 mo or longer) developed amenorrhea, which resolved with its discontinuation, and men who were azoospermic developed normal semen parameters and fathered children following discontinuation (46). Case reports describe temporary azoosperma in men treated with sirolimus, which resolved after its discontinuation or switch to an mTOR inhibitor (47, 48). A rodent study further demonstrated the reversible impact of sirolimus on male fertility (47, 48).

Although our study is limited by the use of a murine model, the mouse and human ovary maintain many functional similarities in that the ovarian reserve in both is maintained in primordial follicles that localize to the ovarian cortex, and follicular development occurs through the same stages (49, 50). It is important to note that the mTOR inhibitor regimens used in clinical oncology practice are based on maximum tolerated doses and are associated with stomatitis, fatigue, gastrointestinal distress, headache, and rash (51). Dosages used in our study were extrapolated from these regimens to mice, and may thus be greater than necessary for fertility preservation. Future studies will assess the effects of mTOR inhibitors and alkylating chemotherapy on tumor dynamics, and determine optimal and minimal dose levels and regimens.

Materials and Methods

In Vivo Murine Model. Studies were approved by the New York University School of Medicine Institutional Animal Care and Use Committee and conducted in accordance with their guidelines. Female C57BL/6 mice aged 8 wk (Taconic Biosciences) were housed under pathogen-free conditions in autoclaved individually ventilated cages (IVCs) that included autoclaved food and water. Female C57BL/6 mice were intraperitoneally injected at the time of estrus with 0.1 mg of 17β-estradiol (Sigma-Aldrich; E2113) to induce the estrus cycle. Female C57BL/6 mice aged 8 wk were housed under pathogen-free conditions in autoclaved individually ventilated cages (IVCs) that included autoclaved food and water. Female C57BL/6 mice were intraperitoneally injected at the time of estrus with 0.1 mg of 17β-estradiol (Sigma-Aldrich; E2113) to induce the estrus cycle. In vivo activation of ovarian cortical tissue was therefore performed using a combination of sirolimus (31 mg/kg, i.p.) and doxorubicin (1 mg/kg, i.p.) at the time of estrus. Female C57BL/6 mice aged 8 wk were housed under pathogen-free conditions in autoclaved individually ventilated cages (IVCs) that included autoclaved food and water. Female C57BL/6 mice were intraperitoneally injected at the time of estrus with 0.1 mg of 17β-estradiol (Sigma-Aldrich; E2113) to induce the estrus cycle. In vivo activation of ovarian cortical tissue was therefore performed using a combination of sirolimus (31 mg/kg, i.p.) and doxorubicin (1 mg/kg, i.p.) at the time of estrus. Female C57BL/6 mice aged 8 wk were housed under pathogen-free conditions in autoclaved individually ventilated cages (IVCs) that included autoclaved food and water. Female C57BL/6 mice were intraperitoneally injected at the time of estrus with 0.1 mg of 17β-estradiol (Sigma-Aldrich; E2113) to induce the estrus cycle. In vivo activation of ovarian cortical tissue was therefore performed using a combination of sirolimus (31 mg/kg, i.p.) and doxorubicin (1 mg/kg, i.p.) at the time of estrus. Female C57BL/6 mice aged 8 wk were housed under pathogen-free conditions in autoclaved individually ventilated cages (IVCs) that included autoclaved food and water. Female C57BL/6 mice were intraperitoneally injected at the time of estrus with 0.1 mg of 17β-estradiol (Sigma-Aldrich; E2113) to induce the estrus cycle. In vivo activation of ovarian cortical tissue was therefore performed using a combination of sirolimus (31 mg/kg, i.p.) and doxorubicin (1 mg/kg, i.p.) at the time of estrus.

**Statistical Analysis.** One-way analysis of variance, Student’s t test, chi-square test, and Fisher’s exact test were used where appropriate. Data are presented as mean ± SEM with significance set at *P* < 0.05. The ratio of growing to primordial follicles was calculated by dividing the total number of growing follicles (primary, secondary) in each treatment group by the total number of primordial follicles in each treatment group.

Additional Methods and Information. Additional information can be found in SI Materials and Methods.


