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Dexrazoxane Abrogates Acute Doxorubicin Toxicity in Marmoset Ovary

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ABSTRACT
Preservation of ovarian function following chemotherapy for nonovarian cancers is a formidable challenge. For prepubescent girls, the only option to prevent chemotherapy damage to the ovary is ovarian tissue cryopreservation, an experimental procedure requiring invasive surgeries to harvest and reimplant tissue, which carries the risk of cancer reintroduction. Drugs that block the primary mechanism of chemotherapy insult, such as dexrazoxane (Dexra) in the context of anthracycline chemotherapy, provide a novel approach for ovarian protection and have the potential to overcome current limitations to oncofertility treatment. Dexra is a catalytic topoisomerase 2 inhibitor that protects the mouse ovary from acute doxorubicin (DXR) chemotherapy toxicity in vitro by preventing DXR-induced DNA damage and subsequent gammaH2AX activation. To translate acute DXR ovarian insult and Dexra protection against acute ovarian DXR toxicity from mice to marmoset monkey tissue.
stromal/thecal cells as early as 2 h postinjection, followed by DNA damage in granulosa cells and later, oocytes [20]. In the ovaries of treated mice, γH2AX activation is measurable at 6 h post-DXR, with increased apoptosis in granulosa cells of growing follicles measurable by 8–10 h post-DXR-treatment [20]. Our mouse study thus defines a narrow 2-h window of opportunity to prevent DXR toxicity with novel medicinal therapy prior to the onset of somatic cell DNA damage. The current study investigates the ability of the cardio- and skin-protectant drug Dexra to shield the marmoset ovary from acute DXR ovarian insult [34–36]. Pretreatment with Dexra, a catalytic topoisomerase II inhibitor and antioxidant, attenuates DXR-induced DNA DSBs, increases ovarian cell viability, and inhibits DXR-induced γH2AX activation in cultured mouse ovaries [34]. The ability to prevent DXR-induced DNA DSBs established Dexra as the first mechanistic ovarian shield against chemotherapy toxicity [34]. With proven efficacy to protect against DXR-induced cardiomyopathy and skin extravasation without limiting DXR toxicity within cancer cells [35–38] and the ability to shield both dividing and quiescent cells, Dexra is an ideal drug to protect the ovary, which contains mixed cell populations at varying stages of cycle and quiescence.

While we have described ovarian protection from DXR insult in mouse tissue [27, 34], such studies do not always faithfully translate to humans [39]. Nonhuman primates, including marmoset monkeys, constitute a translational model that may more closely replicate both DXR ovarian insult and protection in humans [40, 41]. Marmosets are valuable nonhuman primates in which to model ovarian toxicity and fertility preservation, with a relatively short generation time (adult by ~1.5–2 yr of age) and gestation length (5 mo) [42–44]. Comparable to humans, marmoset folliculogenesis is characterized by the development of dominant preovulatory follicles under the influence of follicle-stimulating hormone (FSH) [45]. Considering species similarities between nonhuman primate and human ovaries, it is essential to assess chemotherapy toxicity and protection in a nonhuman primate prior to pioneering Dexra for drug-based fertility preservation studies in humans [39]. The current study presents an initial study demonstrating Dexra protection against DXR toxicity using an in vitro marmoset ovarian tissue culture system.

MATERIALS AND METHODS

Chemicals and Drugs

DXR was provided as DXR hydrochloride in a 2 mg/ml aqueous solution (Teva Parenteral Medicines). Dexra was provided as powdered Dexra hydrochloride (Zincedar; Pharmacia & Upjohn Company). Dexra was solubilized as 80 mM in 0.167 M sodium lactate immediately before use and diluted to a final 20 μM concentration in the culture media.

Animals

No live animal work was conducted in the current study. Marmoset ovaries harvested at the Wisconsin National Primate Research Centers (WNPRC) were obtained through the WNPRC’s Nonhuman Biological Material Distribution Core. Fresh marmoset ovaries were collected under general anesthesia in conjunction with independent studies. The WNPRC coprincipal investigators of the studies that harvested the monkey ovaries (J. E. Levine and D. H. Abbott) gave permission for the authors to use those ovaries for this study. The care, housing, and harvesting of tissues from female marmoset monkeys (Callithrix jacchus) at WNPRC were fully compliant with the recommendations of the Guide for the Care and Use of Laboratory Animals and the Animal Welfare Act. The Institutional Animal Care and Use Committees and Graduate School, University of Wisconsin, Madison, approved all the procedures prior to implementation.

Marmoset ovaries from 10 multiparous/nulliparous adult female common marmosets, median age ~3.5 yr, were obtained either under general anesthesia (1%–5% isoflurane/oxygen) or following premedication with ketamine (at least 15 mg/kg injected intramuscularly) and induction of a deep plane of anesthesia with an intravenous injection of sodium pentobarbital (25–50 mg/kg) to effect stoppage of the heart, permitting necropsy. The harvested ovaries enabled development of an in vitro marmoset ovarian tissue culture system to assess DXR toxicity and Dexra protection from acute DXR insult. Ovaries were placed in 5 ml of sterile PBS, and cleansed of any attached tissue. One ovary from each animal was used for this study and was sliced with the Stadie Riggs Tissue Slicer (Thomas Scientific). Ovarian tissue slices cut along the longitudinal axis of each ovary were cut into eight wedges, and each wedge was transferred randomly to a filter basket (Millipore) in a well containing ovary culture media described previously (Ham F12/DMEM phenol red, 1 mg/ml bovine serum albumin (BSA), 1 mg/ml Albumax, 0.05 mg/ml ascorbic acid, 5 units/ml penicillin 5 μg/ml streptomycin, and 0.0275 mg/ml transferrin) [34]. Four treatment groups were used for all experiments: 1) vehicle control for DXR (0.167 M sodium lactate), 2) vehicle control for DXR (PBS, pH 7.4), 2) vehicle control for DXR + 50 nM DXR, 3) vehicle control for DXR + 500 nM DXR, and 4) 20 μM Dexra + 50 nM DXR. Although Dexra pretreatment was performed in conjunction with 50 nM DXR, this was not done in conjunction with 500 nM DXR because of limited tissue availability. Cotreatment using 50 nM DXR was chosen because that concentration was sufficient to cause a toxicity response, and 500 nM is outside the upper range of circulating DXR concentrations in patients receiving chemotherapy. Ovarian tissues from three to four animals were used for each endpoint assay: neutral comet assay (NCA), Western blot analysis, or immunofluorescence. Ovarian tissue slices were treated with Dexra or vehicle control for 1 h prior to the addition of DXR to the culture media. DXR and Dere were continuously present in the culture media for the entire culture duration. Tissues were harvested for assays at 2, 4, and/or 24 h after the addition of DXR to the culture media. Small pieces of tissue were processed for the NCA to quantify DXR-induced DNA DSBs and the remaining ovarian tissue slices were used to extract protein or fixed in 4% formalin [20, 34].

NCA

Cells were isolated from marmoset ovarian slices to create a single cell suspension at 2 and 4 h post-DXR treatment. Slices were further chopped in a solution of 0.25% collagenase/PBS, pH 7.4, and incubated for 30 min total at 37°C. Every 15 min during that incubation, tissues were passed six times through a 23-gauge needle to disperse the cells. Cells were centrifuged for 10 min at 300 × g to pellet. Collagenase-containing solution was removed, and
each pellet was resuspended in 100 μl of 1 mg/ml proteinase K in PBS and incubated 5 min at room temperature (RT). At this point, the samples were blinded, resuspended in 350 μl 1% low melting agarose/PBS, plated on slides, and processed for the NCA as previously described [20, 34]. Stromal, thecal, and granulosa cells were analyzed together without distinction (no separation of different cell types) because of the small sample size. Images were collected on an Olympus microscope using a 20x objective and SPOT Plus software. At least 100 somatic cells per time point, per treatment group, per animal were imaged. Olive moment quantification of DNA DSBs was scored using CometScore software (TriTek Corporation) [20, 34].

**Protein Lysate Preparation and Western Blot Analysis**

Whole-cell protein lysates were prepared from cultured marmoset ovarian tissue homogenized in RIPA buffer that included phosphatase inhibitors and complete protease inhibitors (Roche/Life Technologies). Protein quantification was determined using the Bio-Rad DC Protein Assay per the manufacturer’s instructions (Bio-Rad). Protein lysates were heated for 5 min at 75°C in 1X Laemmli sample buffer prior to loading 10 or 20 μg total protein per lane, and the samples were separated by size on 4%-20% SDS-PAGE gradient gels (Bio-Rad) under reducing conditions. Proteins were transferred to polyvinylidine fluoride-Fl (Millipore), and the membranes were preblocked in TBS-T (20 mM Tris Base, 137 mM NaCl, 1 M HCl) + 5% BSA (for phospho-specific antibodies) or + 5% milk for 1 h at RT. Western blots were probed with rabbit anti-S139-phosphorylated γH2AX (1:500; Abcam), rabbit anti-caspase-9 (1:1000; Cell Signaling), rabbit anti-PTEN (phosphatase and tensin homolog) (1:1000; Cell Signaling), or mouse anti-β-actin (1:10000; Sigma) in TBS-T + 5% BSA or milk overnight at 4°C. Blots were washed with TBS-T then probed simultaneously with secondary antibodies, donkey anti-rabbit Alexa 680 (Molecular Probes) and donkey anti-mouse IRdye 800 (LiCor), both at 1:15 000 in TBS-T, dried, and scanned using the LiCor Odyssey System (University of Wisconsin-Small Molecule Screening Facility). Density measurements were taken using the Odyssey software.

**Phospho-γH2AX Immunostaining**

Phospho-γH2AX protein expression was detected in 5 μm ovarian tissue sections as previously described [20]. Tissue sections were deparaffinized, hydrated through graded ethanol, and rinsed for 5 min in distilled-deionized H2O. Sections were blocked for 1 h at RT using 10% goat serum in PBS followed by a PBS wash. Sections were then incubated with rabbit anti-phospho-γH2AX (Cell Signaling) at a 1:480 dilution in PBS with 1% goat serum overnight at 4°C. Slides were rinsed three times for 5 min in PBS. Secondary staining was performed with goat anti-rabbit Alexa Fluor 488 (Invitrogen) at a 1:400 dilution in PBS for 30 min at RT in the dark. Slides were rinsed three times for 5 min each in PBS. Slides were mounted with ProLong Gold antifade reagent containing 4',6-diamidino-2-phenylindole (DAPI) (Invitrogen) to counterstain nuclei, and the coverslips were placed. Slides were imaged on the Nikon A1 confocal laser microscope (three ovaries/treatment type).
Fluorescence Microscopy for Detection of Phospho-γH2AX Staining

Images for p-γH2AX-stained slides were collected using a Nikon A1 laser-scanning confocal microscope (WNPRC for Biological Imaging). Alexa Fluor 488 fluorochrome-conjugated Alexa Fluor 488 was detected at an excitation wavelength of 488 nm and emission wavelength of 525 nm. To detect DAPI, the excitation was at 403.6 nm and the emission was collected at 450–465 nm. Laser settings were determined by imaging control samples and setting laser intensities to minimize autofluorescence. Laser settings were then held constant for all the experimental samples. Each image was taken at the Z plane providing maximal signal in the given section. Merged and individual channels were collected under simultaneous excitation. Follicle types were identified using standard morphology and size ranges from parallel-stained hematoxylin and eosin slides [46]. Phospho-γH2AX-positive granulosa or stroma cells had green nuclear staining. A predefined cutoff level to label an entire follicle as p-γH2AX positive was used based on previously reported TdT-mediated dUTP nick end labeling assay and follicle atresia studies [20, 46]; all the follicles were considered positive if they had at least four p-γH2AX-positive granulosa cells except for primordial follicles, which were considered positive with greater than or equal to one p-γH2AX-positive granulosa cell. The mean p-γH2AX-positivity index was calculated as the (number of p-γH2AX-positive follicles)/(total follicle count) for each follicle type based on 2–3 sections/ovary/treatment type.

Statistics

All the experiments were performed in triplicate for each treatment group. Graphs and statistical analyses were generated using OriginLab. One-way ANOVAs were conducted with Bonferroni tests for means comparisons (mean ± SEM). Data were normalized to the controls to allow pooling across experiments and were presented as fractional change compared to the controls. Differences were considered significant at *P*, 0.05. For follicle counts, ovarian tissue slices from the same animal were treated at the same time, and each tissue slice was analyzed as an independent sample.

RESULTS

Dexra Protects Marmoset Ovarian Tissue from DXR-Induced DNA Damage

To test the hypothesis that in vitro administration of Dexra protects marmoset ovarian tissue from DXR-induced DNA damage, cultured marmoset ovarian slices were pretreated with vehicle control for 1 h, followed by the addition of 50 or 500 nM DXR, or with 20 μM Dexra 1 h prior to the addition of 50 nM DXR. NCA showed no increase in DNA DSBs in treated cells at 2 h post-DXR addition (data not shown). By 4 h post-DXR, however, both 50 nM and 500 nM DXR, which span the 100–400 nM range of DXR serum blood concentration in humans [47], increased DNA DSBs by 36% and 37%, respectively, compared to the control (n = 3, *P* < 0.05; Figure 1). Dexra pretreatment prevented DNA damage induced by 50 nM DXR, maintaining DNA DSBs at control values (Fig. 1). These data demonstrate that Dexra shielded cultured marmoset ovarian tissue from early DXR-induced DNA damage.

Dexra Attenuates DXR-Induced Increase in γH2AX Phosphorylation in Marmoset Ovarian Tissue

Phosphorylation of γH2AX was assessed in lysates of marmoset ovarian tissue culture from four marmoset animals at

Fig. 3B. Dexra pretreatment decreases the number of follicles containing phospho-γH2AX foci-positive granulosa cells. Marmoset ovarian tissue culture slices were treated with vehicle control or DXR ± Dexra and fixed at 24 h post-DXR treatment. Ovarian sections were stained for p-γH2AX (green) and nuclei (blue, DAPI). A) Representative fluorescent images (A–H) demonstrate formation of p-γH2AX foci in marmoset ovarian sections for each treatment condition from the same animal (A and B = control, C and D = 50 nM DXR, E and F = Dexra + 50 nM DXR, G and H = 500 nM DXR). Bar = 500 μm for all the images. B) Summary bar graphs of p-γH2AX-positive antral follicles at 24 h after 50 and 500 nM DXR treatment ± Dexra. A growing follicle is considered positive if it contains at least four p-γH2AX-positive granulosa cells per follicle; *P* < 0.05 and **P* < 0.001, one-way ANOVA, Bonferroni means comparison (n = 3 animals).
FIG. 4. DXR-induced formation of phospho-γH2AX foci in granulosa cells of antral follicles of marmoset ovarian tissue. Ovarian sections were stained for p-γH2AX (green) and nuclei (blue, DAPI). A) Representative fluorescent images (A–L) of marmoset ovarian sections from three different animals treated with 500 nM DXR are shown (A–F illustrate primordial, primary, and secondary follicles, G–L illustrate antral follicles). The arrows in C and F point to p-γH2AX-positive oocytes from primordial follicles. B) Digitally zoomed images show p-γH2AX-positive oocytes from primordial follicles (white arrows). Bar = 100 μm for all the images.
4 h post-DXR addition to the culture media. Both 50 and 500 nM DXR increased the density of the band corresponding to phosphorylated γH2AX in two of the four animals at this time point. Dexra pretreatment protected ovarian tissue in the two animals that exhibited a DXR response at 4 h post-DXR treatment as demonstrated by phospho-γH2AX levels that were not different from the control on quantified Western blots. A representative Western blot probed with anti-phospho-γH2AX antibody at 4 h post-DXR treatment and individual bar graphs from the two responsive animals are shown in Figure 2.

Given the heterogeneous response at 4 h post-DXR, γH2AX phosphorylation was further assessed using immunofluorescence in marmoset ovarian tissue slice wedges that were harvested 24 h after treatment with DXR ± Dexra or vehicle control (n = 3 animals, 6–10 ovarian tissue slices per treatment). In contrast to the split phospho-γH2AX response at 4 h posttreatment, immunostaining at 24 h post-DXR revealed DXR increased the percentage of antral follicles containing p-γH2AX-positive granulosa cells in tissue from all animals when compared to vehicle control. In vehicle-treated tissue, 18.9% ± 4.4% of antral follicles were scored as phospho-γH2AX positive, with 50 and 500 nM DXR-treated tissue exhibiting 61.7% ± 10.3% (P < 0.001) and 69.9% ± 16.6% (P < 0.001) positive antral follicles, respectively (Fig. 3 and Supplemental Fig. S1). Dextra pretreatment prevented the observed increase in γH2AX phosphorylation that occurred with 50 nM DXR treatment, maintaining the percentage of p-γH2AX-positive antral follicles at 19.2% ± 6.2% (P < 0.05), comparable to vehicle control (Fig. 3 and Supplemental Fig. S1). These data suggest that Dexra protects DXR-vulnerable antral follicles from chemotherapy insult, potentially delaying or diminishing loss of the growing follicle population. Contrary to antral follicles, γH2AX phosphorylation was observed with low frequency in granulosa cells of other preovulatory follicle types (primordial, primary, and secondary), often below the threshold of 1–4 positive cells/follicle at 24 h post-DXR treatment (Fig. 4). Interestingly, a handful of oocytes of primordial follicles exhibited γH2AX phosphorylation (~5%, Fig. 4) with 500 nM DXR treatment. Nuclear foci of phosphorylated γH2AX were noted in the highly vascularized corpora lutea (Fig. 5) [48], demonstrating the marked heterogeneity of γH2AX phosphorylation of different follicle types in cultured marmoset ovarian tissue.

**Dextra Attenuates DXR-Induced Caspase-9 Cleavage in Marmoset Ovarian Tissue**

To evaluate whether Dexra prevents acute DXR-induced apoptotic signaling, we quantified procaspase-9 cleavage as a marker for apoptosis. Quantification of the density of bands in Western blots corresponding to pro- and cleaved (activated) caspase-9 revealed that treatment with 50 or 500 nM DXR appeared to increase the ratio of cleaved to procaspase-9 at 24 h post-DXR treatment by 29% and 39%, (P > 0.05) over control, respectively (n = 3, Fig. 6). The cleaved to procaspase-9 ratio in tissue pretreated with Dextra prior to 50 nM DXR was not different from control-treated samples (<5% change), suggesting that Dexra attenuated the apparent trend in DXR-induced procaspase-9 cleavage at 50 nM, suggestive of a reduction in apoptosis.

**Dextra Ameliorates DXR-Induced Changes in AKT1 Phosphorylation in Marmoset Ovarian Tissue**

To investigate whether DXR insult affects mediators of cell survival and primordial follicle activation, we quantified phosphorylated AKT1 (v-akt murine thymoma viral oncogene homolog 1) and total PTEN protein expression at 2, 4, and 24 h post-DXR treatment. There were no changes in the density of the phosphorylated AKT1 band at 2 and 4 h post-DXR treatment (50 or 500 nM) on Western blots probed with anti-phospho AKT1 antibodies (not shown). There were trends toward a 25% (P > 0.05) increase in the density of phosphorylated pAKT1 on Western blots at 24 h after the addition of 50 nM DXR and ~45% increase (P < 0.05) following treatment with 500 nM DXR (n = 4; Fig. 7, A and B). Dexra prevented the upward trend in phosphorylated AKT1 following 50 nM DXR, producing levels similar to vehicle control (n = 4, P > 0.05; Fig. 7, A and B). These data suggest that Dexra may guard against the noted upward trend in DXR-induced perturbations in pAKT1, a cell survival and primordial follicle activation protein. PTEN protein expression was not affected by DXR because the corresponding band densities did not change with DXR treatment in marmoset ovarian tissue at
any tested time point (n = 4; Fig. 7, C and D, show 4 h post-DXR treatment while 2 and 24 h data not shown).

**DISCUSSION**

Girls who survive cancer are at increased risk for premature ovarian insufficiency due to chemotherapy toxicity. DXR chemotherapy induces ovarian damage leading to increased follicular demise and decreased fertility, with the extent of toxicity depending on the age of the patient and dose used [26, 49, 50]. We previously characterized DNA damage, subsequent γH2AX activation, and apoptosis in the mouse ovary over the acute insult period following DXR treatment [20]. Furthermore, we recently demonstrated that pretreatment with Dexra prevents DXR-induced ovarian toxicity in cultured mouse ovaries [34]. Prior to translation of drug-based ovarian protection to humans, it is important to validate efficacy and safety of putative ovoprotective drugs in a nonhuman primate model. The present study translates DXR acute insult and Dexra protection from mouse to a marmoset ovarian tissue model.

Dexra pretreatment prior to DXR prevented DXR-induced DNA damage in cultured marmoset ovarian tissue as measured by NCA at 4 h post-DXR, consistent with our previous in vitro mouse study [34]. To further examine the effect of Dexra on cellular responses to DNA damage in marmoset tissue, we quantified the levels of phosphorylated (activated) H2AX. Phosphorylation of H2AX is the earliest and most sensitive downstream cellular response to DNA damage [51] and typically parallels DNA DSBs [52]. The time line for γH2AX phosphorylation is species and tissue specific, however, and may vary depending on the type and dose of the offending agent [53, 54]. Only two out of four marmosets showed increased p-γH2AX band density in response to 4 h DXR treatment despite the fact that tissues from all the animals exhibited DXR-induced DNA damage at this time point. Dexra prevented the increase in γH2AX phosphorylation in the two responsive animals. Given that 4 h was the earliest time to detect DNA damage, γH2AX activation may have been more consistently detected if assayed at later time points. To corroborate the Western blot results, we assessed γH2AX phosphorylation by immunofluorescence at 24 h post-DXR treatment. Because there is no precedent for classifying entire follicles as p-γH2AX-positive nor is the impact of γH2AX activation on follicle survival known, a scoring system was adopted from previous publications that used the TdT-mediated dUTP nick end labeling assay to score follicles as apoptotic [20, 46]. DXR treatment for 24 h increased p-γH2AX positivity in granulosa cells of antral follicles in all animals, suggesting that Western blots of tissue harvested at 4 h posttreatment may have contained samples that were collected at too premature a time to consistently detect γH2AX phosphorylation. Consistent with the Western blots of responsive animals, Dexra abrogated DXR-induced p-γH2AX phosphorylation in all animals at 24 h posttreatment. The immunofluorescence data importantly revealed striking heterogeneity of p-γH2AX immunostaining that was dependent upon follicle type and ovarian structure, with preferential γH2AX phosphorylation in granulosa cells of antral follicles and in the corpus luteum. The follicle- and corpus luteum-dependent activation of γH2AX would have contributed significantly to the mixed results of the Western blots because of the heterogeneity of the marmoset ovarian tissue slices. Future in vivo marmoset studies will determine the timing, pattern, and consequence of γH2AX activation in the follicular, stromal, and oocyte compartments of nonhuman primate ovaries.

Apoptosis occurs when cells fail to repair DNA damage despite recruitment of DNA repair proteins. When cell damage is severe and DNA repair insufficient, apoptosis-inducing downstream signaling effectors, including caspase-9, are activated, leading to cellular demise [55]. A marker of mitochondrial stress signaling and early apoptosis, caspase-9 cleavage was quantified to determine whether Dexra...
inhibits DXR-induced follicle apoptosis signaling in marmoset ovarian tissue. Cleaved caspase-9 activates procaspase-3, leading to the formation of the apoptotic bodies [56]. Our data indicating a trend toward DXR-induced caspase 9 cleavage are consistent with previous studies demonstrating DXR activates caspase-3 in granulosa cells (which occurs downstream of caspase-9) and caspase-2 and -12 in oocytes in mice [57, 58]. Dextra-mediated attenuation of this DXR-induced trend in caspase-9 cleavage parallels our previous study in which pretreatment with Dextra prevented DXR-induced demise of immortalized murine granulosa cells assayed by cell cytotoxicity [34].
One proposed model for chemotherapy-induced ovarian insult has been termed the burn-out model [59]. The model describes enhanced primordial follicle activation due to depletion of growing follicles and subsequent primordial follicle growth and attrition, thus depleting the ovarian reserve [59]. Primordial follicle activation is initiated by oocyte growth and transition of granulosa cells from flat to cuboidal cells. Members of the PI3K signaling pathway, including AKT1 and PTEN, initiate primordial follicle activation [60]. To test the hypothesis that DXR treatment alters PI3K signaling, we examined AKT1 phosphorylation and PTEN expression. DXR appeared to activate AKT1 in marmoset ovarian slices, while Dexra maintained phosphorylated AKT1 at baseline control values. PTEN protein levels on the other hand did not change with DXR treatment in marmoset ovarian tissue culture at the time points tested; hence, restoration of PTEN by Dexra administration could not be tested in our model. PTEN is expressed specifically in oocytes, whereas AKT1 is expressed in granulosa cells. Depletion of PTEN from oocytes results in hyperactivation of AKT1, which in turn leads to rapid depletion of the primordial follicle pool [60]. The trend of DXR-induced AKT1 activation observed in the present study appeared independent of PTEN depletion, but is consistent with chemotherapy-induced premature ovarian failure. Future studies may assess the relationship between these cell division regulator proteins and both primordial follicle activation and ovarian cell survival following DXR insult.

Recently, there have been some concerns about the use of high-dose Dexra (Dexra:DXR 10:1 mg ratio) therapy for cardiac protection because of the possible association between high-dose Dexra and increased secondary blood cancers in children [61] and the suggestion that high-dose Dexra might diminish the anticancer efficacy of DXR in breast cancer patients [62]. The potential risk concerns raised by Tebbi et al. [61] did not reach statistical significance, and the slowed remission rate or patient survival. The majority of studies point toward an absence of pathogenic concern regarding Dexra, including a recent Cochrane review and other studies [63–65]. Nevertheless, the use of Dexra to provide skin and cardiac protection has been greatly restricted by the FDA to cancer patients receiving high doses of DXR (>300 mg/m²) [66]. To mitigate the risk for complications following Dexra treatment, the current study utilized Dexra in a 1:1 mg ratio with DXR, rather than the 10:1 mg ratio currently utilized clinically [34, 67]. Administering Dexra at a dose tenfold lower than that used for cardioprotection appears to successfully protect somatic ovarian cells from acute DXR damage. Future studies are needed to confirm the long-term efficacy and safety profile of low-dose Dexra therapy in vivo prior to clinical use.

Taken together, the results from this study suggest that Dexra protects against acute DXR toxicity in marmoset ovarian tissue, preventing DNA damage, possibly through inhibition of apoptotic signaling. Our findings are limited by the low numbers of the animals tested because of the limited availability of marmoset ovarian tissue. These findings nevertheless support the use of the marmoset as a nonhuman primate for modeling ovarian toxicity and fertility preservation studies. This monkey is particularly suited for fertility studies given its relatively short generation time [40, 68]. These results are also encouraging for future translation of ovoprotection to an in vivo marmoset model to allow systematic assessment of the whole ovary for DXR toxicity and to assess the promise of low-dose Dexra protection.

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