Acetaminophen (APAP, paracetamol) interferes with the first trimester human fetal ovary development in an *ex vivo* model.

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ABSTRACT

Context: Acetaminophen (APAP, paracetamol) is widely used by pregnant women. Although long considered safe, growing evidence indicates that APAP is an endocrine disruptor since in utero exposure may be associated with a higher risk of male genital tract abnormalities. In rodents, fetal exposure has long-term effects on the reproductive function of female offspring. Human studies have also suggested harmful APAP exposure effects.

Objective: Given that disruption of fetal ovarian development may impact women’s reproductive health, we investigated the effects of APAP on fetal human ovaries in culture.

Design and setting: Human ovarian fragments from 284 fetuses aged 7 to 12 developmental weeks (DW) were cultivated ex vivo for 7 days in the presence of human-relevant concentrations of APAP (10⁻⁸ to 10⁻³ M) or vehicle control.

Main outcome measures: Outcomes included examination of postculture tissue morphology, cell viability, apoptosis, and quantification of hormones, APAP and APAP metabolites in conditioned culture media.

Results: APAP reduced the total cell number specifically in 10-12 DW ovaries induced cell death and decreased Ki67-positive cell density independently of fetal age. APAP targeted sub-populations of germ cells and disrupted human fetal ovarian steroidogenesis, without affecting prostaglandin or inhibin B production. Human fetal ovaries were able to metabolize APAP.
Conclusions: Our data indicate that APAP can impact first trimester human fetal ovarian development, especially during a 10-12 DW window of heightened sensitivity. Overall, APAP behaves as an endocrine disruptor in the fetal human ovary.

Key words: N-acetyl-p-aminophenol, paracetamol, human fetal ovary, hormones, metabolites.
INTRODUCTION

Acetaminophen (N-acetyl-p-aminophenol, APAP, paracetamol) is one of the most widely used over-the-counter mild-analgesics, with more than half of pregnant women reporting its consumption\textsuperscript{1,2}. Although considered safe, its innocuousness during pregnancy is now questioned regarding newborn health\textsuperscript{1–4}.

Several experimental studies have demonstrated the endocrine disrupting capacity of APAP in fetal testis\textsuperscript{5–8}. Although studies showed that elements of the steroidogenic pathway are present in the human fetal ovary (hFO)\textsuperscript{9–12}, and that it displays endocrine activity\textsuperscript{13}, the effect of APAP on fetal ovarian endocrine capability has not been reported.

During fetal life, female germ cells go through crucial morphogenetic events, starting with migration while actively proliferating, followed by sex differentiation and commitment to meiosis, culminating with primordial follicles assembly\textsuperscript{14}. Fetal germ cell development, and, more broadly, ovarian development, influences later reproductive lifespan since it establishes a dynamic but finite stockpile of follicles; the ovarian reserve. Studies in rodents demonstrate that APAP reduces both fetal ovarian germ cell numbers and total adult follicle numbers, resulting in reduced fertility\textsuperscript{15–17}. Its impact on ovarian development in offspring also suggests intergenerational effects through epigenetic germline alterations\textsuperscript{15,18}. To the best of our knowledge, only one study on the effect of APAP on hFO has been reported to date and it demonstrated that, \textit{ex vivo}, APAP may interfere with germ cell development\textsuperscript{18}. 


Metabolic processing effectively deactivates drugs while facilitating their elimination, although it can generate toxic metabolites for some drugs, including APAP\textsuperscript{19}. Although the liver is the key organ for drug metabolism, other organs have metabolizing activities\textsuperscript{20}. Fetal exposure to xenobiotics is mainly modulated by the maternal and placental metabolic capacities\textsuperscript{21}. The human fetal liver also expresses several key enzymes involved in phase I and phase II metabolism\textsuperscript{22,23} and is able to biotransform xenobiotics\textsuperscript{24–26}.

We aimed to investigate the effect of APAP on first trimester hFO development by using an established ex vivo culture model\textsuperscript{27}. We show that APAP interferes with first trimester hFO development, altering the viability of a germ cell subpopulation. We also found that APAP disrupts hFO steroidogenic capability without perturbing prostaglandin and inhibin B production, suggesting specific endocrine disruption. Finally, we show, for the first time, that the hFO can metabolize APAP into an inactivated sulfated conjugate.

**MATERIALS AND METHODS**

**Collection of human fetal ovaries**

284 first trimester human fetuses (7–12 developmental weeks (DW)) were obtained from legally induced terminations of pregnancy performed in Rennes University Hospital. Tissues were collected following women’s written consent, in accordance with legal procedures agreed by the National agency for biomedical research (recommendation #PFS09-011; Agence de la Biomédecine) and the Local ethics committee of Rennes Hospital (notification #11-48). Termination of pregnancy was induced using a standard combined mifepristone and misoprostol protocol, followed
by aspiration. The whole aspiration material was collected in a sealed sterile pouch rapidly stored at 4°C and kept cold at all times. Ovaries were isolated under a binocular microscope (Olympus SZX7, Lille, France) within 2 hours after sample collection and immediately placed into ice cold phosphate-buffered saline. Gestational age, previously determined by ultrasound, was further confirmed by measurement of foot length. Ovaries were recovered from aspiration products using a binocular microscope (Olympus SZX7, Lille, France) and immediately placed in cold phosphate-buffered saline.

**Ex vivo culture**

Recovered ovaries were processed as previously described. Briefly, to ensure the viability of the tissue in terms of nutrients accessibility all along the experimental procedure, each ovary was cut into 1 mm³ size explants. Ovaries from fetuses younger than 10 DW were cultured in two separate wells, one ovary under control (vehicle) condition and the other exposed to APAP. Ovaries from 10 DW old fetuses onwards were cut into 1 mm³ pieces allowing more than 2 conditions. The same number of ovarian explants (3 to 4 depending on the endpoint) was used for each condition. One mm³ ovary pieces were placed into cell culture inserts (0.4 μM pores, Millipore #PICM01250) in wells filled with 400 μl of phenol red-free Medium 199 (Invitrogen Life Technologies, Cergy-Pontoise, France) supplemented with 50 μg/ml gentamycin, 2.5 μg/ml fungizone (Sigma Aldrich Chemicals, Saint-Quentin Fallavier, France) and 1 g/l insulin, 0.55 g/l transferrin and 0.67 mg/l sodium selenite (Corning, Fisher Scientific). Cultures were incubated at 37°C for 7 days with 5% CO₂ and humidity. Explants were immediately exposed to either vehicle at a final concentration of 0.1% v/v (dimethyl sulfoxide, DMSO) or APAP (Sigma Aldrich...
Chemicals, Cas #103-90-2). Commonly used therapeutic doses of APAP lead to 100 to 200 µM APAP in the plasma\textsuperscript{28,29}, and APAP freely crosses the placenta\textsuperscript{30}. APAP concentrations used in this study ranged from 0.01 to 100 µM to assess potential dose-response effects and concentration of 1,000 µM was used to explore toxic effects. Upon replacement, culture media were snap-frozen and stored at −80°C for further analyses.

**Single cell dissociation and flow cytometry**

Single cell suspensions from explants were obtained with a 2-step sequential enzymatic digestion procedure\textsuperscript{27}. Live cells were counted on a Malassez hematocytometer using Trypan Blue. When ovaries were cultivated in a single condition, the total number of cells resulted from the live cell counts. Alternately, the total number of cells per condition was calculated from the number of cells counted after dissociation and the ratio of fragments per condition to the initial number of fragments per ovary. Apoptotic cells were labeled with FITC Annexin-V (BD pharmlingen #556419) following the manufacturer’s recommendations, coupled to necrotic nucleus staining with 3 µg/ml 7-Aminoactinomycin D (7-AAD; FluoProbes, Interchim #132303)\textsuperscript{27}. A subpopulation of germ cells was identified through plasma membrane M2A labelling (Table 1). Cell death and germ cell dynamics were evaluated by acquiring a minimum of twenty thousand events using a FACSCalibur (BD Biosciences). Data were analyzed using CellQuest software. The number of M2A-positive germ cells was calculated from the percentage of positive cells and the total cell number.
Immunohistochemistry, stereology, and in situ hybridization

Immunohistochemistry was performed as previously described$^{27}$ on Bouin’s fixed, paraffin-embedded ovaries using antibodies as detailed in table 1. Stained sections were examined and photographed with a light microscope (Olympus BX51). A mean of 6.15 +/- 0.14 photos at 40x magnification were taken per condition in areas containing germ cells (at least 2 photos per explant), with a mean area of 191303 +/- 4366 µm2. Same photos were used for proliferation and germ cell counts. ImageJ software (US National Institutes of Health, Bethesda, MD, USA) was used to perform the cell counting and surface measurement. Proliferating cells were reported as the percentage of KI67-positive cells in a population of KI67-negative cells. Germ cells were identified based on their morphology and size (either < 9 µm, or ≥ 9 µm)$^{31,32}$ calculated with ImageJ. Density of germ cells was the number of germ cells per µm2.

In situ hybridization was performed using RNAscope 2.5 (Advanced Cell Diagnostics) on paraformaldehyde-fixed, paraffin-embedded ovaries according to the manufacturer’s instructions with a minor modification, i.e. the fifth amplification step was increased from 30 to 60 min$^{33}$. Probes are detailed in Table 2.

Prostaglandin measurements

Prostaglandin E2 (PGE2) and prostaglandin D2 (PGD2) levels were measured in culture media of ovarian explants after 1 day of exposure using an ELISA assay (PGE2 EIA Kit—Monoclonal, #514010 and PGD2 – MOX ELISA Kit, #512011, Cayman Chemical Company Ann Arbor, MI, USA) according to the manufacturer’s instructions. Each sample was diluted 1:2 in sample EIA diluent solution prior to reactions and assayed in duplicate. Intra- and inter-assay CV were 3.7–30.4% and 6.4–35% for PGE2 and PGD2, respectively.
Measurements of APAP and its main metabolites

Acetaminophen (APAP), APAP-β-D-glucuronide (APAP-GLU) and APAP-sulfate (APAP-SUL), within culture media collected 24 h and 72 h after culture onset, were simultaneously quantified by liquid chromatography coupled - high resolution mass spectrometer (LC-HRMS). Briefly, an eight calibration points range of APAP, APAP-GLU and APAP-SUL (Sigma Aldrich) (0-5 mg/L) was prepared using culture medium (i.e. M199 with ITS) diluted with methanol (Fisher Scientific UK, Loughborough, Leicestershire, UK). Then, 25 µl of medium diluted 5:18 in methanol and standard solutions were enriched with 100 ng of APAP-D4 (Sigma Aldrich) as an internal standard. Following homogenization, samples were incubated at +4°C for 15 min prior to centrifugation (10 min, +4°C, 3,000g). Target compounds were ionized by using an HESI-II ion source for the electrospray ionization. Analytes were separated by using an Accela Pump (Thermo Scientific) equipped with a Thermo Fisher Hypersil Gold C18 column (3.0 mm, 2.1, 100 mm) as previously described with minor modifications (available upon request). Saline water / twice charcoal dextran stripped aged female rabbit serum (1/1, v/v) was used as the matrix for calibrators.

Steroid measurement

Dihydrotestosterone (DHT), testosterone (T), estradiol (E2), Δ4-androstenedione (4-dione), progesterone (Prog), 17 hydroxy-progesterone (17OHP), and pregnenolone (Preg) were assayed simultaneously by GC/MS as previously described with minor modifications (available upon request). Saline water / twice charcoal dextran stripped aged female rabbit serum (1/1, v/v) was used as the matrix for calibrators.
and quality control (QC) standards. Accuracy, target ions, corresponding deuterated internal control, range of detection, low LOQ (LLOQ), and intra & inter assay CVs of the quality control are reported in Table 3.

**Statistical analysis**

Distributions of cell numbers, percentages and densities in response to different APAP concentrations were compared using mixed regression models to account for the testing of multiple conditions for a same fetus. The fetus was considered as a random effect and the treatment group (DMSO, APAP at different concentrations) as a fixed effect. The global effect of the treatment group was tested with a Fisher’s test and comparison of each treatment effect against the DMSO condition used a Student's t test. Linear models were directly applied or after log-transformation when necessary. A negative binomial regression model was used for the number of germ cells. Fetal age (<10 vs ≥10 DW) and its interaction with the treatment group were included as fixed effects in the models. As interactions appeared significant, we stratified all the analyses according to the age group. Data analyses were computed using the SAS software (SAS/STAT version 9.4; SAS Institute Inc). For hormone measurements, APAP-exposed values were normalized by the corresponding DMSO ones. Global analysis of the effects on ratios was analyzed thanks to a non-parametric Kruskall-Wallis test, followed by Mann-Whitney rank sum test for comparisons between control and APAP-treated samples with the GraphPad Prism 5 software package (GraphPad Software, Inc., La Jolla, CA). Statistically significant post-hoc tests are indicated by (*) if the P-value is below .05, (**) below .01, and (***) below .001.
RESULTS

APAP interferes with ovarian development.

Overall, exposure of 7-12 DW ovarian explants to APAP at various concentrations for 7 days affected the total ovarian cell number differently depending on the age of the ovaries (p<0.0001, Figure 1A). With the exception of $10^{-8}$M, all APAP concentrations significantly decreased overall cell numbers in 10-12 DW explants (p<0.01, Figure 1B), with a dramatic decline of 67% (p<0.0001) at $10^{-3}$M APAP (Figure 1B). In similar exposure conditions, we scored KI67-positive and -negative cells on immunostained sections of 8-12 DW ovarian explants. All tested APAP concentrations significantly decreased the percentage of proliferating KI67-positive cells, except $10^{-5}$M in 10-12 DW explants (Figure 1C). The percentage of dead cells according to sample age and culture condition was then determined by Annexin-V-FITC/7AAD labelling and flow cytometry (Figure 1D). In 7-9 DW ovarian explants, only exposure to $10^{-4}$M APAP induced a significant increase of the dead cell ratio, whereas in 10-12 DW explants, all APAP concentrations except $10^{-4}$M, increased (p<0.05) both apoptosis and necrosis (Table 4), with again an especially pronounced effect of $10^{-3}$M APAP (p<0.001).

APAP targets a subpopulation of germ cells.

The percentage of M2A-positive germ cells reported to the total cell number remained stable with all the APAP concentrations tested, apart from $10^{-3}$ M, which on 10-12 DW explants had a massive impact (p<0.001) (Figure 2A). However, the number of M2A-positive cells significantly decreased in 10-12 DW ovarian explants exposed to APAP concentrations as low as $10^{-7}$M (p<0.05). In 7 DW explants, a loss of M2A-positive cell number was observed with $10^{-4}$M APAP exposure (Figure 2B).
Immunostaining revealed a progressive loss of immature LIN28-positive cells with increasing concentrations of APAP (drastic depletion at $10^{-3}$M), but only a slight reduction in pre-meiotic SYCP3-positive cell numbers (Figure 3A-B). We further quantified germ cell numbers based on their diameter (< 9 µm, ≥ 9 µm) and chromatin compaction on histological sections. Compared to 10-12 DW explants, where the density of < 9 µm germ cells was noticeably reduced by APAP at concentrations as low as $10^{-8}$M, a weak but statistically significant loss of the smaller germ cells was detectable after exposure to $10^{-6}$ and $10^{-4}$M APAP in 8-9 DW explants. Regarding the larger germ cells (≥ 9 µm) counts, APAP had little impact, except for the $10^{-3}$M concentration in 10-12 DW explants (Figure 3C). The percentage of KI67-positive germ cells, whether < 9 or ≥ 9 µm, remained steady regardless of the APAP exposure concentration or fetal age (Figure 3D).

**APAP dysregulates sex steroid production without altering PGE2 and PGD2 production by human fetal ovary.**

Prostaglandin D2 (PGD2), E2 (PGE2), and inhibin B (Figure 4), as well as sex steroid (Figure 5-6) levels were measured in media after either 24h or 7 days culture. Neither prostaglandins nor inhibin B production by hFO were significantly altered by APAP except after exposure of 10-12 DW ovarian explants to the toxic concentration of $10^{-3}$M APAP (PGE2 p<0.001; inhibin B p<0.01). Dihydrotestosterone, testosterone and Δ4-androstenedione levels were below the limit of quantification, while progesterone, pregnenolone, 17OHP and E2 were measurable in all samples, consistent with the expression of steroidogenesis pathway enzymes (Figure 5). Only exposure to $10^{-3}$M APAP drastically decreased levels of these steroids in 8-9 DW
samples while in 10-12 DW samples, both $10^{-4}$ and $10^{-3}$M APAP exposures significantly decreased progesterone and E2 levels and at concentrations as low as $10^{-5}$M, lowered 17-OHP levels (Figure 6). To evaluate whether E2 may have a local impact, we analyzed the expression of estrogen receptors using *in situ* hybridization within first trimester human fetal ovaries. While all 3 receptors were expressed, the level of *ESR2* was consistently higher than that of *ESR1* and *GPER* (Figure 6C-E).

**Human fetal ovary metabolizes APAP mainly into APAP-SUL**

Using LC-HRMS, we measured APAP, as well as its main glucuronide (APAP-GLU) and sulfate (APAP-SUL) metabolites in culture media collected 24h and 72h after culture onset (Figure 7). No APAP was detected in vehicle (DMSO)-exposed culture media (data not shown). A consistent linear relationship between measured and expected (theoretical) APAP levels was evident in both 24h and 72h culture media, except for the $10^{-8}$M concentration that was below the limit of quantification (Figure 7A-B). Regardless of the APAP concentration, the exposure time and fetal age, the only metabolite found within culture media was APAP-SUL, APAP-GLU being below the limit of detection. The amount of APAP-SUL produced was similar for ovaries from 8-10 or 10-12 DW fetuses and increased linearly with the concentration of APAP (data not shown). The sulfation rates (*i.e.* the mean ratios between measured levels of APAP-SUL and APAP) ranged from 0.5% to 2% depending on the concentration and the exposure time (Figure 7C). This rate after 2 consecutive days of culture (72h) was usually twice as high as after 24h (Figure 7D). Our bulk RNAseq data indicates that several sulfotransferases, including SULT1E1, SULT1A1, SULT4A1 and SULT1C4 are expressed in hFO (Figure 7E). Immunohistochemistry for SULT1E1 revealed immunoreactivity in endothelial cells as well as in currently
unidentified structures within hFO, strong immunoreactivity in the mesonephros, as well as in human fetal liver which were used as positive controls (Figure 7F).

DISCUSSION

Given the wide use of APAP by pregnant women and inter-species variability with rodent models, we took advantage of a previously described ex vivo model of hFO culture\textsuperscript{27} to investigate whether APAP interferes with development of the first trimester hFO. Our study shows that concentrations equivalent to, or below, peak therapeutic APAP plasma levels\textsuperscript{28,29} interfere with both development and steroidogenesis of the hFO, despite the ovary’s innate metabolizing capability.

Ovarian development relies on growth and differentiation, the former sustained by proliferation of both the somatic and germ cell lineages\textsuperscript{39}, until oogonia commit to meiosis. In line with a previous study from our group\textsuperscript{27}, we report that the organotypic culture system allows for ovarian development. Under these conditions, APAP did not globally alter the total number of cells in 7-9 DW ovaries but reduced the percentage of proliferative KI67-positive cells while moderately inducing apoptosis at 10^{-4}M. This apparent discrepancy could be attributed to marginal effects observed for these parameters. In contrast, in 10-12 DW ovaries, the loss of total cells was consistently accompanied by a reduction of the percentage of KI67-positive cells and an increased percentage of dying cells. The hindering effect of APAP observed on cell proliferation is consistent with previous reports\textsuperscript{40,41}. Our data defines a possible window of sensitivity of human fetal ovaries for APAP between 10 and 12 DW.
Maternal and fetal blood APAP concentrations, as well as pharmacodynamics were reported to be similar according to several experimental strategies and PBPK modeling\textsuperscript{30,42,43}. To the best of our knowledge, there are unfortunately no available findings regarding APAP distribution in the human fetal ovary. Hence, we cannot exclude that the exposure of the ovary can be different \textit{in vivo} via blood flow and \textit{ex vivo} via exchanges with culture medium. However, the range of APAP concentrations used in this study was chosen in order to cover peak plasma concentrations after therapeutic doses and subsequent concentrations corresponding to biotransformation and clearing. We also included a typical concentration corresponding to APAP surdosage and classically used as a cytotoxicity-inducing concentration in several cellular models\textsuperscript{41,44-46}. Of note, inter-individual variability was high for APAP concentrations below $10^{-4}$M, while almost absent at the toxic concentration of $10^{-3}$M. Therefore, $10^{-3}$M APAP-induced cytotoxicity in our \textit{ex vivo} human fetal ovary culture model is consistent with cell line models.

Since germ cells commit to meiosis from 11 DW onwards, a biologically consistent hypothesis to explain the difference in response of 7-9 DW ovaries compared to that of 10-12 DW could be that APAP differentially affects germ cells according to their differentiation stage. The percentage of M2A-positive cells was not affected by APAP regardless of the drug concentration and the developmental stage considered, while the total number of M2A-positive cells was significantly decreased, in particular in 10-12 DW ovaries explants exposed to APAP at $10^{-7}$M and above. In addition, the density of germ cells, quantified on histological sections according to their size and chromatin compaction, indicated an impact of APAP on small ($< 9 \, \mu m$) germ cells.
regardless of age, but not on larger germ cells at sub-toxic APAP concentrations. This suggests that APAP selectively impacts small germ cells, in agreement with previous observations showing a decrease in the density of immature TFAP2C-positive germ cells after exposure to $10^{-5}$M APAP for 7 days\textsuperscript{18}. The developmental stages of exposure in rodent studies encompassed two crucial consecutive, morphogenetic events occurring in the germ cell lineage, \textit{i.e.} mitosis and meiosis, and therefore did not allow for the identification of a specific window of sensitivity\textsuperscript{47}. Our data does not support the hypothesis that APAP induces a shift of oogonia towards meiosis\textsuperscript{15} in first trimester hFO as we did not observe any increase in the density of larger germ cells. By contrast, proliferating hepatic HepaRG cells were reported to be more sensitive to APAP than differentiating ones\textsuperscript{48}, and APAP is known to interfere with mitotic checkpoints and DNA synthesis\textsuperscript{40,41,45,46,49}. Since small germ cells are actually actively proliferating oogonia, a working hypothesis that will require further investigations is a preferential effect of APAP on cycling germ cells.

In the fetal rat ovary, APAP-induced germ cell depletion was associated with a slight decrease in PGE2 content\textsuperscript{15}. In the NTera2 germ cell line, cell number was reduced following exposure to either APAP or PGE2 receptor antagonists, whereas PGE2 agonists prevented the APAP-induced loss in cell number\textsuperscript{18}. In humans, since PGE2 is considered as an autocrine regulator of germ cell development\textsuperscript{50}, it was hypothesized that APAP-induced dysregulation of prostaglandin could also explain deleterious impacts on germ cells\textsuperscript{18}. However, we show here that exposure levels equivalent to or lower than therapeutic doses of APAP had no effect on PGE2 nor PGD2 levels. This is in contrast to ibuprofen\textsuperscript{27}, which is consistent with the weaker
inhibitory effect of APAP on prostaglandin production\textsuperscript{51,52}. As our results suggest that APAP impacts the hFO independently of prostaglandin inhibition, but its mechanisms of action remain to be elucidated.

Although steroidogenic pathway members are reportedly expressed as early as 11 DW\textsuperscript{9–13}, only two studies have reported conversion of pregnenolone and androgens into androgens and estrogens, respectively, within the hFO\textsuperscript{53,54}. We show here that APAP can interfere with progesterone, 17OHP and E2 production. During pregnancy, circulating estrogens are mostly conjugated to serum proteins, therefore reducing the levels of bioactive-free estrogens\textsuperscript{55}. The roles for a local production of estrogens at this developmental stage, as well as the consequences of its alterations remain to be determined in humans. The alteration of the oogonia pool with decreased local estrogens could be consistent with the phenotype reported in CYP19A1-deficient rabbits\textsuperscript{56}. However, to the best of our knowledge, this is the first report of an \textit{ex vivo} specific endocrine disruption of the steroidogenesis in the hFO.

In the adult liver, APAP is mainly detoxified by glucuronidation and sulfation while a small proportion is oxidized to form the reactive metabolite N-acetyl-\textit{p}-benzoquinone imine (NAPQI)\textsuperscript{57,58}. Our recent bulk RNAseq data\textsuperscript{38} indicates that the ovaries do not express the main cytochromes P450 (CYPs), metabolizing APAP into NAPQI, namely CYP2E1, CYP3A4 and CYP1A2 (data not shown). However, previous reports showed that APAP can be toxic \textit{via} a CYP-independent pathway, which can eventually lead to mitochondrial dysfunction\textsuperscript{59,60}. Hence, further investigations are required to determine whether this CYP-independent mechanism of APAP toxicity might explain some of our results including impaired production of sex steroids. In
contrast, we show that fetal ovaries as young as 8 DW and at least up to 12 DW metabolize APAP mainly through sulfation, as observed in the fetal liver\textsuperscript{57}. The stability of sulfation rates despite increasing APAP concentrations suggests that, at least under our experimental conditions, the underlying detoxification system was neither preferentially-induced nor saturated. The rapid production of sulfo-conjugates (\textit{i.e.} during the first 24h) also suggests that sulfo-transferases are already expressed in non-exposed ovaries, rather than being induced by the exposure to APAP. Although we cannot rule out the involvement of several SULT enzymes, of those detected at the mRNA level, only SULT1E1 was detected \textit{in situ}. This is consistent with the reports that SULT1E1, one of the main sulfotransferase enzymes known to metabolize APAP, is not only expressed but active in human fetal liver and in extra-hepatic tissues\textsuperscript{61--64}. In 7 DW fetuses, we also observed a very high expression of this enzyme in the mesonephros, suggesting a possible local detoxification role \textit{in vivo}. The precise identity of the ovarian cell type expressing SULT1E1 remains an avenue of research.

In conclusion, we demonstrate that biologically relevant concentrations of APAP can interfere with hFO development \textit{ex vivo}. The underlying mechanisms are likely to differ depending on fetal age and APAP concentration, but they may be independent of the prostaglandin pathway. We also confirm that active steroidogenesis takes place within first trimester hFOs. Furthermore, we provide evidence that the hFO displays intrinsic detoxification capacities, although this might not be sufficient to provide significant protection against prolonged or acute exposure to xenobiotics. Nevertheless, we demonstrate, for the first time, that APAP disrupts human fetal ovarian endocrine activity and is therefore an endocrine disruptor in the developing human female.
Authors’ roles

L.L.L., S.L.P. and S.M.G. designed, performed and analyzed experiments (organ collection for SMG and L.L.L.; culture, histology, cell counts and flow cytometry for S.L.P and L.L.L.; culture, histology, cell counts for S.M.G.). N.C. performed the statistical analyses and contributed to the writing of the corresponding section. L.L. performed culture; C.D.L performed prostaglandins measurements; F.G. performed the steroids measurements and contributed to the writing of the corresponding section. B.F. and T.G. designed APAP and metabolites measurements; T.G. and L.L.L. performed APAP and metabolites measurements; V.L. supervised the collection of the first trimester human fetal ovarian samples. S.M.G. and A.D.R. conceptualized the project. L.L.L. and S.M.G. prepared the visualization of the data and wrote the original draft. S.L.P., A.D.R. contributed to critical discussions, reviewed and edited the manuscript.

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FIGURE LEGENDS

Figure 1. Effects of APAP on human fetal ovary growth. (A) Fetal age distribution of total ovarian cell number after 7 days of exposure to 10^{-6}, 10^{-5} and 10^{-4} M APAP normalized to control values as a function of age (developmental weeks: DW) (n = 91). (B) Quantification of total viable cell number per ovary following 7 days of exposure to either control vehicle (Ctrl, white bars) or APAP at 10^{-8}-10^{-3} M (gray to black bars) in fetuses at 7, 8-9 and 10-12 DW. Cell counts were performed with Trypan Blue exclusion staining of 7-12 DW fetal explants, and are expressed as 10^6 cells per ovary. (C) Percentage of KI67-positive cells relative to the total number of cells assessed on immunostained sections 7 days after exposure to either control vehicle (Ctrl, white bars) or APAP at 10^{-8}-10^{-3} M (gray to black bars) at 8-9 and 10-12 DW. (D) Percentage of apoptotic (solid bars) and necrotic cells (hatched bars) after 7 days of exposure to either control vehicle (Ctrl, white bars) or APAP at 10^{-8}-10^{-3} M (gray to black bars) assessed with Annexin-V versus 7-AAD labeling of fetal ovarian explants at 7-9 and 10-12 DW. Data are represented as mean +/- SEM. Mixed linear models with fetus as a random fetus effect were performed for cell numbers, densities and percentages. (*) , (**) and (***) above the bars indicate * P < .05; ** P < .01; *** P < .001 from the t-test comparisons with the control condition. The number of samples tested per condition is indicated at the bottom of each bar.

Figure 2. Effects of APAP on germ cells. Percentage (A) and number (B) of M2A-positive germ cells was assessed by flow cytometry following 7 days of exposure to either vehicle (-, white bar) or APAP 10^{-8}-10^{-3} M (gray to black bars). Total M2A-germ cells number per ovary is expressed as 10^3 germ cells per ovary at 7, 8-9 and 10-12
DW. Data are represented as mean +/- SEM. Mixed linear models with fetus as a random effect were performed for cell numbers, densities and percentages. (*), (**), and (***) above the bars indicate * P< .05; **P < .01; ***P < .001 from the t-test comparisons with the control condition. The number of samples tested per condition is indicated at the bottom of each bar.

Figure 3. Differential effects of APAP on germ cell populations. (A-B) Representative immunostaining of immature oogonia (A) and pre-meiotic germ cells (B) with antibodies directed against LIN28 and SYCP3, respectively, after 7 day of exposure of 11.6 DW ovarian explants to control vehicle (DMSO) or different concentrations of APAP (10^{-6}-10^{-3}M) at 8-9 and 10-12 DW. Immunoreactive germ cells appear brown (3,3-diaminobenzidine tetrachloride staining), and sections were counterstained with hematoxylin. Scale bar= 100 µm. (C) Density of germ cells (number of cells per µm^2) in 8-9 DW and 10-12 ovarian explants exposed to 10^{-8}-10^{-3}M APAP for 7 days. Cells were quantified as a function of their size (<9 µm and ≥ 9µm). (D) The percentage of KI67-positive germ cells for each category of germ cells was quantified on immunostained sections as a function of KI67 immunoreactivity and cell size (<9 µm and ≥ 9 µm). Data are represented as mean +/- SEM. Mixed linear and negative binomial models with fetus as a random effect were performed. (*) and (***) above the bars indicates P < .05 and P < .001 from the t-test comparisons with the control condition, respectively. The number of samples tested per condition is indicated at the bottom of each bar.
Figure 4. Effects of APAP on the production of prostaglandin and inhibin B by human fetal ovaries. (A-B) Measurement of PGE2 (A) and PGD2 levels (B) normalized with control after 24h of exposure of 8-9 and 10-12 DW samples to either vehicle (-, white bars) or APAP at 10^{-8}-10^{-3} M (gray to black bars). (C) Measurement of inhibin B levels normalized with control after 24h of exposure to either vehicle (-, white bars) or APAP at 10^{-8}-10^{-3} M (gray to black bars) produced by 10-12 DW ovarian explants. Data are represented as mean +/- SEM. The P-value of the Kruskall–Wallis test is indicated on the top, ** above the bar indicates that the Wilcoxon post-hoc test compared to control has a P-value <0.01. The number of samples per condition is indicated at the bottom of each bar.

Figure 5. Steroidogenesis machinery and production in human fetal ovaries. (A) Steroid levels measured by mass spectrometry as a function of developmental weeks (DW). Each dot represents the measurement of steroid levels in the medium of human fetal explants from one donor cultivated for 7 days in control conditions. (B) Expression of transcripts encoding enzymes of the steroidogenic pathway, namely the hydroxysteroid dehydrogenase HSD3B2, and the cytochrome P450 CYP11A1, CYP17A1, and CYP19A1 (aromatase) were evaluated in human fetal ovaries by RNAsequencing\textsuperscript{38}. Data are expressed as means +/- SEM of fragments per kilobase of exon model per million reads mapped (fpkm). (C) Representative low magnification (upper panel) and high magnification (lower panels) of in situ localization of CYP11A1 and CYP17A1 proteins, and CYP19A1 mRNA in a 11.8 DW-old ovary, using immunohistochemistry and RNAscope, respectively. Scale bar = 100 µm.
Figure 6. Effects of APAP on human fetal ovarian sex steroids production. (A-B) Amounts of progesterone, pregnenolone, 17 hydroxyprogesterone (17OHP) and estradiol (E2) produced by APAP-exposed explants, normalized with amounts produced by control explants, after 7 days of exposure. Data from 8-9 DW (A) and 10-12 DW ovarian explants (B) are shown for normalized values of APAP at $10^{-6}$-$10^{-3}$M (gray to black bars). Data are represented as mean +/- SEM. (**) above the bar indicates the Wilcoxon post-hoc test compared to control has P-values <0.01. The number of individuals per condition is indicated in the exposure key. (C-E) Representative in situ hybridization of nuclear estrogen receptors ESR1 (C), ESR2 (D), and the membrane estrogen receptor GPER mRNAs (E) in a 13.6 DW human fetal ovary. Lower panel photos show higher magnifications. Scale bar: 100 µm.

Figure 7. Metabolization of APAP by the human fetal ovary. (A-B) APAP availability in culture media. Relationship between theoretical and measured APAP levels (A) and ratios between measured and theoretical APAP levels (B) were determined for each APAP concentration after 24h of culture as well as after an additional 48h of culture (72h). (C) APAP sulfation rates (measured [APAP-SUL]/[APAP]) in 8-10 DW and 10-12 DW ovaries for each initial theoretical APAP concentration after 24h of culture as well as during the following 48h (72h). The number of samples per condition is indicated in parentheses below the x-axis. (D) Ratios between measured [APAP-SUL] levels produced by 8-10 DW and 10-12 DW ovaries after 24h and the subsequent 48h (72h) of exposure were calculated for each initial APAP concentration. The number of samples per condition is the same as in C. Data are represented as mean +/- SEM. (E) SUL1E1, 1A1, 2A1, 4A1 1C4 transcript levels in human fetal ovary as a function of developmental age. Data are
expressed as fragments per kilobase of exon model per million reads mapped (fpkm). Note that the Y-axis scale for \textit{SULT1E1} is 6x higher than for the other \textit{SULTs}. (F) Immunostaining of SULT1E1 using the age-matched fetal liver as a positive control, and in uncultured ovaries of 11.8, 16 and 26 DW. SULT1E1-positive cells appear brown (3,3-diaminobenzidine tetrachloride staining), and sections were counterstained with hematoxylin. Ov: ovary; Meso: mesonephros; MD: Müllerian duct. Scale bar= 100 µm.
TABLE LEGENDS

Table 1. Antibodies used for immunohistochemistry (IHC) and flow cytometry (FC) staining. 10 mM Tris, 1 mM EDTA buffer, pH 9 (TE) or 10 mM citrate buffer, pH 6 (citrate). ABC: Avidin-biotin complex (VECTASTAIN® Elite ABC-HRP Kit, Vector Labs); RTU: ready to use.

Table 2. Probes used for RNAscope.

Table 3. GC-MS fetal ovarian cell supernatant analytical control validation. LLOQ : low limit of quantification; QC : quality control.

Table 4. Comparison of cell death, apoptosis and necrosis occurrence under different exposure concentrations against control condition, stratified by age: P-values derived from t-test pairwise comparisons of each exposure concentration against the control condition are indicated. The number of samples per condition (n) is indicated in parenthesis in the concentration row.

Data Availability

All data generated or analyzed during this study are included in this published article.
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Table 1. Antibodies used for immunohistochemistry (IHC) and flow cytometry (FC) staining. 10 mM Tris, 1 mM EDTA buffer, pH 9 (TE) or 10 mM citrate buffer, pH 6 (citrate). ABC: Avidin-biotin complex (VECTASTAIN® Elite ABC-HRP Kit, Vector Labs); RTU: ready to use.

| Target protein | Primary antibody reference and RRID | Application | Antigen retrieval | Primary antibody dilution | Secondary antibody reference | Secondary antibody dilution | Signal enhancement |
|----------------|-----------------------------------|-------------|------------------|---------------------------|------------------------------|-----------------------------|---------------------|
| Cleaved caspase 3 | Cell Signaling Tech. #9661 (AB_2341188) | IHC | TE, 40 min, 80°C | 1:150 | Goat anti-Rabbit Biotin conjugated Dako #E0432 | 1:500 | ABC |
| CYP11A | Sigma Aldrich #HPA016436 (AB_1847423) | IHC | citrate, 40 min, 80°C | 1:150 | Dual Link Dako #K4063 | RTU | |
| CYP17A | Abcam #ab48019 (AB_869326) | IHC | citrate, 40 min, 80°C | 1:200 | Rabbit anti-goat Biotin conjugated Dako #E0466 | 1:500 | ABC |
| KI67 | Dako #M7240 (AB_2142367) | IHC | citrate, 40 min, 80°C | 1:100 | Dual Link Dako #K4063 | RTU | |
| Protein | Source | Catalogue Number | Format | Dilution | Enzyme/Reagent | Incubation Conditions | Antibody Conjugate | Sensitivity |
|---------|--------|------------------|--------|----------|----------------|-----------------------|--------------------|-------------|
| LIN28   | Abcam  | #ab46020 (AB_776033) | IHC    | 1:300    | Dual Link Dako #K4063 | RTU                  |                    |             |
| M2A     | Abcam  | #ab77854 (AB_1566117) | FC     | 1:200    | R-Phycoerythrin coupled anti-Mouse IgG (H+L) Jackson ImmunoResearch #715-116-150 | RTU                  |                    | 1:200       |
| SYCP3   | Sigma Aldrich | #HPA039635 (AB_2676607) | IHC    | 1:250    | Dual Link Dako #K4063 | RTU                  |                    |             |
| SULT1E 1 | Sigma | #HPA028728 (AB_10601894) | IHC    | 1:250    | Goat anti-Rabbit BIOTIN conjugated Dako #E0432 | ABC                  |                    | 1:500       |
Table 2. Probes used for RNAscope.

| Target genes | RNAscope probes                  | Cat No |
|--------------|----------------------------------|--------|
| ESR1         | RNAscope® Probe- Hs-ESR1         | 310301 |
| ESR2         | RNAscope® Probe- Hs-ESR2         | 470151 |
| GPER         | RNAscope® Probe- Hs-GPER         | 553361 |
| CYP19A1      | RNAscope® Probe- Hs-CYP19A1      | 489941 |
Table 3. GC-MS fetal ovarian cell supernatant analytical control validation. *LLOQ:* low limit of quantification; *QC:* quality control

| Accuracy (%) | Analytes | Target (GC-MS) or Precursor ion analyte (GC-MS/MS) / IS (m/z) | Range (pg/ml) | Mean (23 runs; pg/ml) | Intra- & Inter assay CVs (%) |
|--------------|----------|---------------------------------------------------------------|---------------|-----------------------|-----------------------------|
|              |          |                                                               |               |                       | LLOQ Mean Intra- & Inter assay CVs | Low QC Mean Intra- & Inter assay CVs | Middle QC Mean Intra- & Inter assay CVs | High QC Mean Intra- & Inter assay CVs |
| 95 - 109     | T        | 482 / 485                                                     | 3.9 - 944     | 3.6                   | 153.7                       | 310.4                       | 621.5                       |
|              |          |                                                               |               |                       | 15.1 - 18.8            | 3.7 - 5.6                   | 3.6 - 4.9                   | 3.4 - 5.1                   |
| 94 - 108     | DHT      | 484 / 487                                                     | 1.4 - 333     | 1.7                   | 30.8                       | 62.5                       | 124.4                       |
|              |          |                                                               |               |                       | 15.7 - 19.9             | 4.0 - 5.8                   | 3.8 - 5.6                   | 3.3 - 5.2                   |
| 94 - 107     | E2       | 660 / 664                                                     | 0.2 - 56.0    | 0.22                  | 2.87                       | 6.07                       | 12.88                       |
|              |          |                                                               |               |                       | 17.3 - 19.8             | 4.3 - 5.7                   | 3.6 - 5.2                   | 3.3 - 5.4                   |
| 89 - 113     | Preg     | 298 / 302                                                     | 25 - 6000     | 5.8                   | 14.4                       | 77.9                       | 198.9                       |
|              |          |                                                               |               |                       | 16.1 - 19.4             | 4.3 - 6.7                   | 4.1 - 5.3                   | 4.6 - 5.8                   |
| 93 - 110     | 4-dione  | 482 / 488                                                     | 11.7 -        | 1.6                   | 9.8                        | 48.1                       | 198.5                       |
|              |          |                                                               |               |                       |                            |                            |                            |                            |
| Range  | Test   | Value 1 | Value 2 |
|--------|--------|---------|---------|
| 94 - 105 | Prog  | 510 / 518 | 16.5 - 4000 |
|        |       | 16.7 | 63.4 |
|        |       | 15.9 - 19.5 | 3.3 - 5.3 |
|        |       | 15.2 - 19.5 | 3.2 - 5.3 |
|        |       | 30.0 - 5.5 |
| 88 - 116 | 17OHP | 465 / 469 | 33 - 8000 |
|        |       | 36.3 | 64.6 |
|        |       | 15.8 - 19.7 | 4.7 - 8.5 |
|        |       | 4.1 - 6.1 |
|        |       | 4.4 - 7.2 |
Table 4. Comparison of cell death, apoptosis and necrosis occurrence under different exposure concentrations against control condition, stratified by age: P-values derived from t-test pairwise comparisons of each exposure concentration against the control condition are indicated. The number of samples per condition (n) is indicated in parenthesis in the concentration row.

| Age (DW) | Cell death type | $10^9$M (n=19) | $10^7$M (n=27) | $10^5$M (n=29) | $10^3$M (0) |
|----------|----------------|----------------|----------------|----------------|-------------|
| < 10     | Cell death     |                | 0.3614         | 0.2662         | 0.0081      |
|          | Apoptosis      |                | 0.7866         | 0.6258         | 0.0154      |
|          | Necrosis       |                | 0.9383         | 0.4202         | 0.0178      |
| ≥ 10     | Cell death     | 0.0200         | <.0001         | 0.0001         | 0.0003      | 0.1         | <.0001 |
|          | Apoptosis      | 0.6102         | 0.0045         | 0.0019         | 0.0139      | 0.4103      | <.0001 |
|          | Necrosis       | 0.3147         | 0.0019         | 0.0007         | 0.0058      | 0.1482      | <.0001 |
Figure 1

A. Age distribution of total ovarian cell number

B. Total ovarian cell number

C. Proportion of proliferating cells

D. Proportion of dying cells
Figure 2

A. Proportion of germ cells

7 DW

| Cell Type | Ctrl | 10^6 | 10^7 | 10^8 | 10^9 |
|-----------|------|------|------|------|------|
| APAP      |      |      |      |      |      |
| M2A+ cell % | 23   | 5    | 7    | 11   |      |

8-9 DW

| Cell Type | Ctrl | 10^6 | 10^7 | 10^8 | 10^9 |
|-----------|------|------|------|------|------|
| APAP      |      |      |      |      |      |
| M2A+ cell % | 22   | 8    | 15   | 14   |      |

10-12 DW

| Cell Type | Ctrl | 10^6 | 10^7 | 10^8 | 10^9 |
|-----------|------|------|------|------|------|
| APAP      |      |      |      |      |      |
| M2A+ cell % | 33   | 12   | 16   | 21   | 10   |

p<0.0001

B. Number of germ cells

| Cell Type | Ctrl | 10^6 | 10^7 | 10^8 | 10^9 |
|-----------|------|------|------|------|------|
| APAP      |      |      |      |      |      |
| M2A+ cell number \(10^3\) | 23   | 5    | 7    | 11   |      |

| Cell Type | Ctrl | 10^6 | 10^7 | 10^8 | 10^9 |
|-----------|------|------|------|------|------|
| APAP      |      |      |      |      |      |
| M2A+ cell number \(10^3\) | 20   | 4    | 10   | 11   |      |

| Cell Type | Ctrl | 10^6 | 10^7 | 10^8 | 10^9 |
|-----------|------|------|------|------|------|
| APAP      |      |      |      |      |      |
| M2A+ cell number \(10^3\) | 30   | 11   | 16   | 18   | 22   |

p<0.0001

-34%

-28%

-32%

-25%

-20%

-19%

-96%

***
Figure 3

C. Germ cell density

D. Proportion of proliferating germ cells
Figure 4

A. PGE2 8-9 DW

B. PGD2

C. Inhibin B 10-12 DW
Figure 5

A. Levels of steroids

| Steroid | 0 | 2 | 4 | 6 | 8 | 10 | 12 DW |
|---------|---|---|---|---|---|-----|-------|
| Progesterone | 0 | 10 | 30 | 50 | 70 | 90 | 18 |
| Pregnenolone | 0 | 10 | 30 | 50 | 70 | 90 | 18 |
| 17-OHP | 0 | 10 | 30 | 50 | 70 | 90 | 18 |
| E2 | 0 | 10 | 30 | 50 | 70 | 90 | 18 |

B. Expression of enzymes of the steroidogenic pathway

| mRNA | 0 | 0.5 | 1 | 1.5 | 2 | 2.5 | 3 | 3.5 | 4 |
|------|---|-----|---|-----|---|-----|---|-----|---|
| HDS3B2 | 0 | 0.5 | 1 | 1.5 | 2 | 2.5 | 3 | 3.5 | 4 |
| CYP11A1 | 0 | 0.5 | 1 | 1.5 | 2 | 2.5 | 3 | 3.5 | 4 |
| CYP17A1 | 0 | 0.5 | 1 | 1.5 | 2 | 2.5 | 3 | 3.5 | 4 |
| CYP19A1 | 0 | 0.5 | 1 | 1.5 | 2 | 2.5 | 3 | 3.5 | 4 |

C. Immunohistochemistry images of CYP11A1, CYP17A1, and CYP19A1 mRNAs.
