The impact of ovulation on fallopian tube epithelial cells: evaluating three hypotheses connecting ovulation and serous ovarian cancer

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Abstract

Ovarian cancer is the most lethal gynecological malignancy affecting American women. Current hypotheses concerning the etiology of ovarian cancer propose that a reduction in the lifetime number of ovulations decreases ovarian cancer risk. Advanced serous carcinoma shares several biomarkers with fallopian tube epithelial cells, suggesting that some forms of ovarian carcinoma may originate in the fallopian tube. Currently, the impact of ovulation on the tubal epithelium is unknown. In CD1 mice, ovulation did not increase tubal epithelial cell (TEC) proliferation as measured by bromodeoxyuridine incorporation and proliferating cell nuclear antigen staining as compared to unstimulated animals. In superovulated mice, an increase in the number of pro-inflammatory macrophages was detected in the oviduct. Ovulation also increased levels of phospho-γH2A.X in TEC, indicating that these cells were susceptible to double-strand DNA breakage following ovulation. To determine which components of ovulation contributed to DNA damage in the fallopian tube, an immortalized baboon TEC cell line and a three-dimensional organ culture system for mouse oviduct and baboon fallopian tubes were developed. TEC did not proliferate or display increased DNA damage in response to the gonadotropins or estradiol alone in vitro. Oxidative stress generated by treatment with hydrogen peroxide or macrophage-conditioned medium increased DNA damage in TEC in culture. Ovulation may impact the fallopian tube epithelium by generating DNA damage and stimulating macrophage infiltration but does not increase proliferation through gonadotropin signaling.

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Introduction

Ovarian cancer is the most lethal gynecological malignancy and the fifth leading cause of cancer death among women, due in part to a lack of early detection methods (Jemal et al. 2009). As a result, ~62% of ovarian cancers are not detected until after the tumors have metastasized, resulting in high mortality rates and few treatment options (Altekruse et al. 2010). The most common histotype of epithelial ovarian cancer is serous carcinoma, which has previously been reported to arise from precursor lesions involving the ovarian surface epithelium (OSE) or ovarian epithelial inclusion cysts (Scully 1995, Auersperg et al. 2002). Emerging evidence indicates that the fallopian tube epithelium may be an alternative site of origin for serous ovarian cancers (Bell 2005, Altekruse et al. 2010). A striking resemblance between the fallopian tube epithelium and serous ovarian cancer has been observed; this includes similar gene expression profiles, cell morphologies, and expression of E-cadherin, all of which are distinct from the OSE (Auersperg et al. 2008).
Several lines of evidence support the fallopian tube as a source of neoplastic cells for the initiation of serous cancer. The majority of women with advanced stage serous cancer exhibit preneoplastic lesions in the fallopian tube in addition to a dominant ovarian mass (Jarboe et al. 2008b). A ‘p53 signature’ has been identified, which marks precursor lesions of the distal fallopian tube that are composed of morphologically benign epithelium exhibiting elevated levels of p53 protein expression (Lee et al. 2007). Identical p53 mutations have been observed in tumors of the distal fallopian tube and in ovarian or peritoneal serous carcinomas, suggesting a common source (Jarboe et al. 2008a). Women with BRCA1 or BRCA2 mutations are more likely to exhibit expression of the p53 signature as well as primary fallopian tube carcinoma, indicating that an association may exist between the BRCA mutation phenotype and serous cancer originating from the fallopian tube (Cass et al. 2005). However, much is still unknown about the association between neoplastic lesions of the fallopian tube and development of serous pelvic or ovarian cancer, largely due to a lack of tools to model neoplasia in the fallopian tube. The laying hen has been used as an animal model for spontaneous ovarian and tubal cancer, and although these animals exhibit primarily the endometrioid histotype rather than the serous histotype of adenocarcinoma, a recent analysis depicted oviductal genes expressed at high levels in cancers as the grade increased (Hakim et al. 2009, Trevino et al. 2010). A human fallopian tube ex vivo culture system was developed, which allows for co-culture of both secretory and ciliated cells of the fallopian tube (Levanon et al. 2010). This represents significant improvements over previous tubal epithelial cell (TEC) culture systems but does not allow for analysis of TEC in the context of their physiological microenvironment in contact with stroma and extracellular matrix, or for any in vivo analysis of how ovulation might damage tubal cells.

Serous ovarian cancer is largely a spontaneous disease, although ∼10% of cases are linked to mutations in BRCA1 and BRCA2 (Crum et al. 2007a). Risk factors for spontaneous serous cancer include nulliparity, infertility, and an early onset of puberty or late menarche, while pregnancy, breastfeeding, and the use of oral contraceptives are protective against ovarian cancer (Auersperg et al. 2008). An increased incidence of non-familial, spontaneous ovarian cancer is linked to an increased number of lifetime ovulations, and several hypotheses have been proposed linking events associated with ovulation to ovarian cancer initiation and progression. The first hypothesis is Fathalla’s tear-and-repair hypothesis, which states that as a result of repetitive proliferation of the OSE to repair the ovulation-induced wound in the ovarian surface, spontaneous DNA replication errors accumulate, leading to neoplasia and ovarian cancer (Fathalla 1971). OSE have been shown to proliferate in response to ovulation and to exhibit signs of DNA damage (Murdoch et al. 2001, Burdette et al. 2006), but the effects of ovulation on TEC proliferation and DNA damage are unknown. The second hypothesis linking ovulation to ovarian cancer concerns the gonadotropins, FSH, and LH, which are thought to contribute to the initiation and progression of ovarian cancer by stimulating the growth of the OSE and/or inhibiting apoptosis of damaged cells (Konishi et al. 1999). Both the OSE and the TEC have been shown to express receptors for FSH, LH, and estradiol, though the exact role of these hormones in stimulating TEC in vivo is not known (Zheng et al. 1996, Zhang et al. 2001). The third hypothesis states that inflammation induced by ovulation may generate oxidative stress that is further exacerbated by cellular proliferation and the effects of FSH and LH, leading to DNA damage and replication errors. Ovulation generates inflammatory molecules such as bradykinin, prostaglandins, and leukotrienes through ischemia–reperfusion associated with wound repair, as well as through recruitment of leukocytes to sites of ovulation (Murdoch 2008). However, an association between ovulation and inflammation in TEC has not been examined.

The purpose of this study was to evaluate common hypotheses regarding ovulation and the initiation of serous epithelial cancer to determine how proliferation (tear and repair), the gonadotropins, and inflammatory molecules influence normal TEC. Following ovulation, TEC might possibly be sloughed onto the ovarian surface due to the close proximity of the distal fimbriae of the fallopian tube to the ovarian rupture site by a process similar to retrograde menstruation. Sloughed cells might form inclusion cysts in which the epithelial cells are exposed to increased levels of hormones and growth factors (Kurman & Shih Ie 2010). The process of ovulation releases growth factors, steroid hormones, and inflammatory molecules, and the fallopian tube is in close proximity to the source of these factors. The current study reports that ovulation does not increase TEC proliferation in vivo. Moreover, using a novel organ culture system to study mouse TEC and baboon fimbriae TEC, as well as a newly generated immortalized baboon TEC cell line (TEC40), FSH and LH do not induce proliferation of TEC. TEC from ovulated animals exhibit increased levels of DNA damage and macrophage infiltration into the oviduct. Oxidative stress generated by hydrogen peroxide or
macrophage-conditioned media increased DNA damage in TEC40.

Materials and methods

Animals

Day 25 female CD1 mice were used for in vivo ovulation studies. For organ culture experiments, mouse oviducts were isolated from CD1 day 16 female pups. All mice were acquired through in-house breeding, and breeders were purchased from Harlan (Indianapolis, IN, USA). Fimbriae tissue was obtained from adult female baboons (Papio anubis) housed at the University of Illinois at Chicago under the supervision of Dr Fazleabas. All animals were treated in accordance with the National Institutes of Health Guide for the Care and Use of Laboratory Animals and the established institutional animal use and care protocol at the University of Illinois at Chicago. Animals were housed in a temperature and light-controlled environment (12 h light:12 h darkness) and were provided food and water ad libitum.

Experimental design of ovulation study

Induction of ovulation was performed essentially as described previously and in Fig. 1 (Burdette et al. 2006). Control mice were injected i.p. with PBS at 0900 h on d25 and 0900 h on d27. Ovulated mice were injected i.p. with 5 IU pregnant mare serum gonadotropin (PMSG; Sigma–Aldrich) at 0900 h on d25 and 5 IU human chorionic gonadotropin (hCG; Sigma–Aldrich) at 0900 h on d27. Injections containing 100 mg/kg bromodeoxyuridine (BrdU; Sigma–Aldrich) were given i.p. either at the time of the first and second hormone or PBS injections or solely with the second injection. On first injection of BrdU, animals were given water containing 0.8 mg/ml BrdU to allow for continuous labeling. At 2100 h on d27.5, animals were killed using CO₂ asphyxiation and cervical dislocation. Total basal proliferation occurring over 60 h was assessed by injecting mice (n = 6) with PBS and BrdU at 0900 h on d25, followed by continuous labeling with BrdU until 2100 h on d27.5. Abridged basal proliferation was measured by injecting mice (n = 3) with PBS and BrdU at 0900 h on d27, with labeling until 2100 h on d27.5 for a labeling time of

Figure 1 TECs do not proliferate in response to ovulation in vivo. (A) CD1 mice were injected with buffer (PBS) as a control or hormones (PMSG and hCG) to induce ovulation. BrdU was added to control injections on either day 25 or 27 to measure cellular proliferation during a 12 h (abridged basal) or 60 h (total basal) labeling period. Total cellular proliferation (total ovulatory) or proliferation in response to hCG stimulation and the process of ovarian surface rupture (post-ovulatory) was measured by adding BrdU to the d25 and d27 injections, respectively. (B) Oviductal sections (5 μm) were processed for immunohistochemistry and stained with antibodies against CK8 to mark OSE or TEC and antibodies against BrdU or PCNA to mark proliferating cells. Asterisks mark proliferating TEC. (C) The percentage of CK8-positive TEC with BrdU incorporation was scored as a percentage of the total number of CK8-positive TEC. Significant differences (P < 0.05) are between groups labeled ‘a’ and ‘b.’ (D) The percentage of CK8-positive, PCNA-positive TEC was scored as a percentage of the total number of CK8-positive TEC. No significant difference was detected between groups.
12 h. Total ovulatory proliferation was measured by ovulating animals \((n = 6)\) with PMSG and hCG, with BrdU labeling from 0900 h on d25 to 2100 h on d27.5. At 2100 h on d27.5, ovaries and oviducts, including fat pad, bursa, and part of the uterine tube, were collected. For analysis of tissues at 16 h post-hCG, animals were injected with PBS \((n = 4)\) or PMSG \((n = 5)\) at 1700 h on d25, followed by injection of PBS or hCG at 1700 h on d27, and collection of tissues at 0900 h on d28. Tissues were fixed in 4% paraformaldehyde (PFA) for 24 h, dehydrated with ethanol, paraffin embedded, and serial sectioned at 5 \(\mu\)m.

**Organ culture**

Oviducts from d16 prepubertal mice were dissected from the ovary and uterus and gently uncoiled with forceps in dissecting medium composed of Leibovitz (L15) medium with 2 mM l-glutamine, 100 U penicillin, and 100 \(\mu\)g streptomycin (Invitrogen). Each oviduct was cut in four pieces, termed organoids. Fallopian tube fimbriae from adult baboons were collected into dissecting medium and the distal ends of the baboon fimbriae were isolated using a scalpel (baboon organoids). Each organoid was placed into a 0.5% alginate (w/v in PBS) droplet formed on a mesh fiber as described previously (Jackson et al. 2009). The alginate-encapsulated organoid was placed into 50 mM calcium chloride for 2 min to cross-link the alginate into a gel. The encapsulated organoid was then placed into growth media for 7 days. Growth media was composed of alpha-MEM (Invitrogen), 100 U penicillin, and 100 \(\mu\)g streptomycin, and 3 mg/ml BSA (MP Biomedicals, Solon, OH, USA), 10% v/v FBS (Invitrogen), 10 or 100 mlU/ml FSH (Sigma), 10 or 100 mlU/ml LH (Sigma), or 10 \(n\)M estradiol (Sigma). BrdU was added to the media at a final concentration of 10 \(\mu\)M 24 h prior to fixation.

**Cell culture**

Fallopian tube fimbriae from adult baboons were rinsed briefly in 70% ethanol and placed into Leibovitz (L15) medium with 1000 U collagenase (Sigma) and 2× trypsin–EDTA (Invitrogen) for 1 h at 37°C. Tissue was vortexed briefly and the media was collected into a new microcentrifuge tube and centrifuged at 800 \(g\) for 3 min to pellet cells. Cells were maintained in MEM with 10% FBS, 2 mM l-glutamine, 100 \(\mu\)M non-essential amino acids, 1 mM sodium pyruvate, 100 U/ml penicillin, and 100 \(\mu\)g/ml streptomycin (Invitrogen).

To immortalize the baboon TEC, cells were transfected with pW2T+ t+ to express the SV40 large and small T antigens (Dr Kathy Rundell, Northwestern University, Chicago, IL, USA) and Lipofectamine 2000 according to the manufacturer’s instructions (Invitrogen). Following transfection and subsequent passaging, untransfected cells died, leaving immortalized cells expressing SV40. Cells were cultured for several months and were verified for expression of SV40, secretory, and ciliated markers at passage number 16. The immortalized baboon TEC cell line was named TEC40, for TEC immortalized with SV40.

For proliferation assays, TEC40 cells were seeded into 96-well plates at 5×10^4 cells/ml. The next day, complete media containing 10% FBS and 10 or 100 mlU/ml FSH or LH was added to plates and the cells were allowed to grow for 4 days. Proliferation was measured with CellTiter 96 Aqueous One Solution (Promega) according to the manufacturer’s directions. Spectrophotometric analysis at 490 nm was completed using a Biotek EL312e microplate reader (Fisher Biotek, Pittsburgh, PA, USA).

To prepare macrophage-conditioned medium (MCM), RAW264.7 cells were maintained in DMEM with 10% FBS, 100 U penicillin, and 100 \(\mu\)g streptomycin. Conditioned medium was prepared as previously described (An et al. 2009). Cells were plated at 5×10^5 cells/ml in a 6-well plate 24 h prior to treatment with Escherichia coli lipopolysaccharide serotype O55:B5 (Fisher Scientific, Pittsburgh, PA, USA) in serum-free media. Cells were treated with serum-free media. Conditioned media containing 5 \(\mu\)g/ml lipopolysaccharide for 24 h before conditioned media was collected.

For analysis of phospho-γH2A.X foci, TEC40 cells were serum starved for 18 h prior to treatment with hormones, hydrogen peroxide, or MCM. FSH (10 or 100 mlU/ml), LH (10 or 100 mlU/ml), estradiol (10 \(n\)M), hydrogen peroxide (Sigma–Aldrich; 1 mM), or MCM was added to fresh serum-free medium. TEC40 were treated for 20 min, followed by fixation, and immunofluorescent staining as described below.

**Luciferase assays**

TEC40 cells were plated in triplicate in phenol red-free DMEM with 10% charcoal-stripped FBS (Gibco), 100 U/ml penicillin, and 100 \(\mu\)g/ml streptomycin into 12-well plates 24 h prior to transfection. Cells were co-transfected with 0.25 \(\mu\)g pCMV Sport-βgal plasmid and 2.5 \(\mu\)g of equal amounts of the ERE-luc reporter plasmid and ERx or pGL3-Basic plasmids using the calcium phosphate precipitation method (Sambrook et al. 1989). The ERE-luc reporter plasmid and ERx expression plasmids were obtained from
Dr Benita Katzenellenbogen, University of Illinois, Champaign, IL, USA. After cells were incubated with plasmids for 4 h, cells were washed, glycerol shocked, and incubated in media with or without 50 nM estradiol for 20 h. Cells were harvested in reporter lysis buffer and luciferase activity measured relative to β-galactosidase activity as described previously (Jaffe et al. 2003).

**RNA isolation and RT-PCR**

Total RNA was extracted from 1 mg of adult mouse ovary, d16 mouse oviduct, baboon fimbriae, or 1 x 10⁶ TEC40 cells using a Qiagen QiaShredder column and Qiagen RNeasy kit according to manufacturer’s directions (Qiagen, Inc.). cDNA was prepared by reverse transcribing 1 μg total RNA using MMLV-RT (Fermentas, Inc., Glen Burnie, MD, USA). cDNA was used for PCR amplification using the following primers: mouse FSHR forward (CCTTGCTCC-TGGTGCTCTTGTG), reverse (CTCGGTACCCCTTCATCTTGT); mouse LHR forward (CGCCGACTATCTCTCACCTA), reverse (GACAGATTTGAAGAGGTGTTGCTG); mouse ERz forward (CAGATTAGAGTGCATTG), reverse (GGCCGACCAATCAT); and mouse GAPDH forward (AGGCCGTGGTGAACGGATTTG), reverse (TGTA GACCATGTAGTGGAGTCA). For PCR amplification of baboon cDNA, the following human primers were used: human FSHR forward (GTATCATCCTG-GATCTGT CACT), reverse (CA TCCCTCAGGA-GGTCAGAAG); human LHR forward (GGAGGCGACGGTTGCTCTAC C), reverse (CAGTTCCACTCT-CAGCAAGCAT); human ERz forward (TCTCTACGACCCCTTCAGT), reverse (GGTACAA-TCCACAAAGCC TG); and human GAPDH forward (ATGGGGTAAGTGAAGGTCTCG), reverse (GGGGT-GGTCATAGAGTGGCAAACA). For cycling conditions for reactions were 94°C 2 min; 34 cycles of 94°C for 30 s, 55°C for 30 s, 68°C for 30 s; 68°C for 5 min. Reactions were run on a 1% (w/v) agarose gel and visualized using an Alpha Innotech gel documentation system (Santa Clara, CA, USA).

**Immunohistochemistry**

Organoids were removed from culture media, recross-linked in 50 mM calcium chloride for 2 min, and fixed in 2% PFA with 50 mM sodium cacodylate and 10 mM calcium chloride for 16–24 h. All reagents were obtained from Vector Laboratories, Inc. (Burlingame, CA, USA) unless otherwise indicated. Xylene and ethanol were obtained from Surgipath Medical Ind., Inc. (Richmond, IL, USA). Antigen retrieval was performed using 10 mM sodium citrate, pH 6.0. Slides were washed in Tris-buffered saline (TBS) with Tween 20 (20 mM Tris, 500 mM NaCl, and 0.1% Tween 20, pH 7.4). When staining for BrdU, tissues were treated with 4 M hydrochloric acid for 10 min, followed by 0.1 M sodium tetraborate for 10 min. Tissues were blocked for 15 min in 3% hydrogen peroxide (Fisher Scientific) followed by avidin and biotin blocking according to manufacturer’s instructions. Slides were incubated in TBS-3%BSA-10% serum of the secondary antibody host for 1 h at room temperature. After blocking, slides were incubated overnight at 4°C in primary antibody in 3% BSA-TBS-10% serum. Control slides received serum block instead of the primary antibody. The primary antibodies against BrdU (rat, 1:200 dilution; Abcam, Cambridge, MA, USA), cytokeratin 8 (CK8) (CK8 TROMA-1 antibody, rat, 1:200; Developmental Studies Hybridoma Bank, Iowa City, IA, USA), E-cadherin (rabbit, 1:50; Cell Signaling Technology, Denver, MA, USA), F4/80 (rat, 1:100; Abcam), acetylated tubulin (mouse, 1:500; Sigma–Aldrich), oviductal glycoprotein 1 (OVGP1, rabbit, 1:250; Abcam), Pax8 (rabbit, 1:250 Proteintech Group, Chicago, IL, USA), phospho-γH2A.X (rabbit, 1:100; Cell Signaling Technology), or SV40 T Ag (rabbit 1:50; Santa Cruz Biotechnology, Santa Cruz, CA, USA) were incubated on tissue sections overnight at 4°C. Slides were rinsed three times for 5 min in TBS-Tween and then incubated at room temperature for 30 min in biotinylated secondary antibody in 3% BSA-TBS. After washing slides in TBS-Tween, avidin/biotin complex (ABC) reagent was added and incubated for 30 min at room temperature. Slides were washed in TBS and antigen–antibody–HRP complex was visualized using diaminobenzidine reagent for 1 min. Slides were counterstained with hematoxylin.

**Immunofluorescence**

TEC40 cells were plated in an 8-well chamber slide at a density of 5 x 10⁶ cells per well. Cells were fixed in 4% PFA for 5 min, washed with PBS, and permeabilized with 0.2% Triton-X100 in PBS for 10 min. Cells were then washed twice with PBS and blocked with 10% goat serum in PBS for 10 min. Cells were incubated in primary antibody diluted in 10% serum in PBS for 1 h at room temperature. Cells were washed and incubated with anti-rat AlexaFluor-594-conjugated secondary antibodies (Invitrogen) for 30 min to detect CK8 staining. For double immunofluorescent staining of CK8 and phospho-γH2A.X, cells were incubated with anti-rat AlexaFluor 594 and anti-rabbit AlexaFluor 488 for 30 min. For all other double

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immunofluorescent staining, cells were incubated with both AlexaFluor594 and a biotinylated anti-rabbit secondary antibody for 30 min. Cells were washed with PBS and incubated with ABC reagent for 30 min. After washing with PBS, cells were fluorescently labeled with AlexaFluor488 using a streptavidin–HRP tyramide signal amplification kit (TSA kit #22, Invitrogen) for 10 min at room temperature. Cells were washed in distilled water and slides were mounted with Vectashield Mounting Medium with DAPI (Vector Laboratories). The proliferating cell nuclear antigen antibody (PCNA; rat) was used according to the manufacturer’s protocol (Zymed, San Francisco, CA, USA).

**Western blotting**

TEC40 cells were serum-starved overnight prior to treatment with FSH (10 or 100 mIU/ml) or LH (10 or 100 mIU/ml) in serum-free media for 24 h. Cells were lysed in modified RIPA buffer (25 mM Tris–HCl, pH 7.6, 150 mM NaCl, 1% (v/v) Triton-X, 1% (v/v) sodium deoxycholate, 0.1% SDS, 1× Complete Mini Protease Inhibitor Cocktail tablets (Roche), and 1× Phosphatase Inhibitor Cocktail III (Sigma)). Protein concentration was determined using the bicinchoninic acid assay (Pierce, Rockford, IL, USA). Cell lysate (30 μg) was run on a 10% SDS–PAGE gel under reducing conditions and transferred to polyvinylidene fluoride membrane. Membranes were blocked for 1 h at RT in 5% BSA (for pERK1/2) or 5% nonfat dry milk (for actin) in TBS-T. Primary antibodies against phospho-p44/42 MAPK (Erk1/2) (Thr 202/Tyr204) (rabbit, 1:1000 dilution; Cell Signaling Technology) or actin (rabbit, 1:1000 dilution; Sigma) were incubated on blots overnight at 4 °C. After washing, membranes were incubated with goat anti-rabbit HRP secondary antibody (Cell Signaling) at 1:100 in blocking buffer for 1 h. Membranes were washed and incubated with SuperSignal West Femto substrate (Thermo Scientific, Rockford, IL, USA) before imaging on an Alpha Innotech gel documentation system.

**Imaging and counts**

Immunohistochemistry or immunofluorescent images were acquired on a Nikon E600 microscope using a DXM1200 digital camera and NIS Elements software (Nikon Instruments, Melville, NY, USA). Immunofluorescent images were also acquired on a Zeiss LSM510 laser scanning confocal microscope (Carl Zeiss MicroImaging, LLC, Thornwood, NY, USA). Using ImageJ software (National Institutes of Health, Bethesda, MD, USA), the number of CK8-positive tubal cells that were also positive for BrdU, PCNA, or phospho-γH2A.X were counted and expressed as a percentage of total number of CK8-positive tubal cells. At least three random fields from at least two independent experiments were counted.

**Statistical analysis**

All values are expressed as the mean±s.e.m. Tukey’s multiple comparison tests were used to assess differences between experimental groups and control groups. For comparison between two groups, a Student’s t-test was used. P<0.05 was considered statistically significant.

**Results**

**Ovulation does not drive proliferation of TEC in vivo**

In order to investigate whether ovulation increases proliferation of TEC, incorporation of BrdU was quantified in control CD1 mice or mice that were hormonally induced to ovulate (Fig. 1A). Day 25 immature mice were used to ensure that no previous ovulations had occurred, leaving the ovaries and oviducts undamaged by previous ovulation-related inflammation, wound repair, or the influence of hormones. Mice were injected with either PBS as a control or a combination of 5 IU PMSG and hCG to induce ovulation (n=6) (Burdette et al. 2006). Ovulation was verified to have occurred by the presence of corpora lutea within the ovary. Proliferation in response to both PMSG and hCG (BrdU treatment on d25, 60 h labeling period) and post-ovulatory proliferation in response to hCG injection (BrdU treatment on d27, 12 h labeling period) were determined and compared to proliferation in PBS-treated animals. To quantify the percentage of proliferating TEC, serial sections of oviduct were stained with antibodies against CK8 to mark epithelial cells, or BrdU to mark proliferating cells (Fig. 1B). Tubal cells positive for both CK8 and BrdU were scored as a percentage of total CK8-positive tubal cells (Fig. 1B and C). For TEC, animals in the abridged basal group showed 3.5% proliferating cells compared with 6.3% proliferating cells in the post-ovulatory group, over the course of a 12 h labeling period. The total basal animals exhibited 18.5% proliferating epithelial cells compared with 17.7% in the total ovulatory animals, both of which underwent labeling for 60 h. Significant differences were not detected when comparing ovulated animals.
to unstimulated animals and controlling for labeling time in vivo.

To verify TEC proliferation using an endogenous marker of proliferation, serial sections were stained for CK8 or PCNA and counted as described above (Fig. 1B). Basal animals included both total basal and abridged basal animals, as the only difference between these groups was the time of BrdU incorporation, which was not necessary for measuring PCNA (n=9). Ovulated animals included both total ovulatory and post-ovulatory animals (n=12). In accordance with the measurement of BrdU labeling as an indicator of cell proliferation, basal animals showed 7.0% PCNA-positive cells, compared with 9.0% for ovulated, which was not significantly different (Fig. 1D). Therefore, ovulation does not increase TEC proliferation in vivo.

Three-dimensional oviduct culture system retains markers of secretory and ciliated cells

To determine the effects of individual components of ovulation, such as the gonadotropins and inflammation, on TEC proliferation and DNA damage, a three-dimensional organ culture system was developed to study normal primary TEC in their microenvironment. Alginate, a widely used biomaterial in tissue engineering, was used as described previously (Jackson et al. 2009) to maintain the physiological architecture of the oviduct, with the TEC lining the lumen of the tube. Sexually immature d16 CD1 mice were used in order to study oviductal epithelium that had not been affected by ovulations. Each oviduct was separated into four pieces with a scalpel and placed into an alginate droplet (Fig. 2) and cultured for 7 days in basal media (BSA or FBS) or serum-free media supplemented with FSH, LH, or estradiol at cycling or menopausal concentrations (King et al. 2011). Oviducts were analyzed by immunohistochemistry for normal endogenous markers of TEC and compared to uncultured mouse oviducts (Fig. 3A and Supplementary Figure 1, see section on supplementary data given at the end of this article). The oviduct culture system retained cytoplasmic markers of normal oviductal epithelium, CK8 and E-cadherin. As assayed by PCNA staining, cultured and uncultured oviducts exhibited similar TEC proliferation rates (Supplementary Figure 1, see section on supplementary data given at the end of this article). Cultured oviducts continued to express oviductin (OVGP1), a secreted glycoprotein characteristic of the oviduct (Umezu et al. 2003). A subpopulation of TEC from cultured and uncultured oviducts stained positively for nuclear expression of Pax8, a marker of secretory cells in the oviduct, while a separate subpopulation of TEC stained positively for acetylated tubulin, a marker of ciliated cells.

In primates, the TEC of the distal fimbriae of the fallopian tube are comparable to the TEC of the mouse...
oviduct (Critoph & Dennis 1977, Komatsu & Fujita 1978). However, TECs are found on the outer surface of the fimbriae and are in contact with the ovary and face the peritoneal cavity, rather than facing the lumen of the oviduct as in mice. To account for these anatomical differences between mice and primates, a three-dimensional culture system for distal baboon fimbriae was developed using the alginate hydrogel method described above (Jackson et al. 2009). Adult female baboons (n=3) were killed and individual fimbria was placed into alginate hydrogel droplets and cultured for 7 days. Cultured and uncultured fimbriae were evaluated by immunohistochemistry for markers of secretory and ciliated cells (Fig. 3B and Supplementary Figure 1, see section on supplementary data given at the end of this article). Fimbria cultures retained markers of TEC (CK8, E-cadherin), oviductin expression, and similar levels of cell proliferation (PCNA). A subpopulation of baboon fimbriae TEC was positive for expression of acetylated tubulin, while other TECs were positive for expression of Pax8, a marker of secretory cells. Therefore, the three-dimensional organ culture system for mouse oviduct and baboon fallopian tube fimbriae retains markers of the cell types normally present in these tissues.

**Immortalization of baboon fimbriae TEC allows co-culture of secretory and ciliated cells**

An immortalized baboon fimbriae TEC cell line (TEC40; TEC immortalized with SV40) was generated to observe the effects of individual aspects of ovulation on cultured TEC. Development of a two-dimensional immortalized TEC cell line allows for comparison of immortalized TEC to immortalized OSE cells (IOSE cells) and to previous studies using traditional two-dimensional culture on plastic of normal and cancer cell lines. TEC40 cells retained an epithelial morphology and were analyzed for retention of ciliated and secretory cell markers by immunofluorescence at passage number 16, after several weeks in culture (Fig. 4A–F). As demonstrated in the baboon fimbriae organ culture system, TEC40 retained the epithelial cell markers CK8, E-cadherin, and expression of the secreted glycoprotein OVP1. A subset of cells was positive for Pax8, while another subset was positive for acetylated tubulin. TEC40 exhibited immunofluorescence staining for SV40 T/t, indicating that the cultured cells were immortalized by expression of the viral oncogenes that disrupt p53 and retinoblastoma.

To determine whether TEC40 were responsive to FSH and LH signaling, cells were treated with cycling or menopausal levels of hormones and analyzed for expression of phospho-MAPK (pERK1/2), which is downstream of both FSH and LH signaling in OSE and granulosa cells (Choi et al. 2002, 2009). Cells treated with FSH or LH showed a dose-dependent accumulation of phospho-MAPK (pERK1/2), indicating that FSH and LH signaling is present in TEC40 cells (Fig. 4G). Additionally, transcripts for FSHR and LHR were detected in TEC40 cells (Supplementary Figure 2, see section on supplementary data given at the end of this article).

To determine whether TEC40 cells were responsive to estradiol, cells were co-transfected with pCMV Sport-β-gal and ERE-luciferase reporter plasmids, with or without co-expression of a plasmid expressing ERα. TEC40 exhibited no significant change in activation of the ERE-luciferase reporter relative to the β-galactosidase reporter when cells were treated with 50 nM estradiol. This corresponds to a lack of expression of ERα in TEC40 cells as assayed by RT-PCR (Supplementary Figure 2, see section on supplementary data given at the end of this article). When ERα was overexpressed, TEC40 cells exhibited approximately fourfold induction of the ERE-luciferase reporter (Fig. 4H).
TECs do not proliferate in response to FSH, LH, or estradiol

The mouse oviduct and baboon fimbriae organ culture systems were used to evaluate the effects of FSH, LH, and estradiol on TEC proliferation in vitro. For mouse oviduct cultures, no significant increase in proliferation was observed in response to FSH, LH, or estradiol compared with BSA or FBS controls. Cycling levels of FSH or estradiol suppressed TEC cell proliferation (Fig. 5A). For baboon fimbriae cultures, no significant difference in proliferation was observed for tissues cultured with FSH, LH, or estradiol (Fig. 5B). To verify the presence of FSHR, LHR, and ERα in these tissues, RNA was extracted from adult mouse ovary as a positive control, d16 mouse oviduct, and adult baboon fimbriae and subjected to RT-PCR. Transcripts for FSHR, LHR, and ERα were detected in the mouse ovary and oviduct and the baboon fimbriae, though FSHR shows weak expression in mouse oviduct (Supplementary Figure 2, see section on supplementary data given at the end of this article).

Since TEC40 cells were responsive to FSH and LH, but not estradiol (Fig. 4G and H), TEC40 cells were analyzed for proliferation in response to cycling and menopausal levels of FSH and LH. After 4 days in culture, cells cultured in cycling (10 mIU/ml) levels of FSH showed a slight but statistically significant increase in proliferation (Fig. 5C). No difference in proliferation was observed at higher concentrations of FSH or LH.

Ovulation induces inflammation and oxidative stress in vivo

As ovulation did not stimulate TEC proliferation in vitro through a tear-and-repair process (Fig. 1), and menopausal levels of gonadotropins did not induce proliferation in mouse oviduct culture, baboon fimbriae organ culture, or immortalized TEC40 cells (Fig. 5), ovulation-induced inflammation in TEC was evaluated. Tissues from control mice or ovulated mice were analyzed for nuclear phospho-H2A.X foci 12 or 16 h after hCG injection (Fig. 6A). Phospho-γH2A.X is a histone protein incorporated at sites of double-strand DNA breaks, which are frequently formed due to exposure to oxidative damage (Bartkova et al., 2005). In control animals, 6.5% of TEC exhibited phospho-γH2A.X foci compared with 20.0% of TEC from ovulated animals at 12 h post-hCG and 16.3% of TEC from ovulated animals at 16 h post-hCG (Fig. 6B). No difference in proliferation rates were observed in TEC from
ovulated animals at 12 h compared with 16 h post-hCG (data not shown).

It is possible that the DNA damage observed in vivo in response to ovulation may be due to the inflammatory reaction associated with ovulation. Macrophage infiltration into human fallopian tube mucosa has previously been demonstrated in response to ovulation, and macrophages are known to secrete pro-inflammatory mediators (Gaytan et al. 2007). Therefore, oviducts from control or ovulated mice were analyzed for expression of F4/80 antigen, a surface glycoprotein that is expressed by macrophages during maturation and activation in response to inflammation (Austyn & Gordon 1981, Robinson-Smith et al. 2007). Control animals showed no macrophage infiltration, while macrophages were found at the periphery of oviducts of ovulated animals at 12 h after hCG injection (Fig. 6C). At 16 h after hCG injection, macrophages were found at the periphery of oviducts, adjacent to cumulus–oocyte complexes, and associated with the tubal epithelium. Therefore, macrophage infiltration and phospho-γH2A.X foci are detected in the oviduct after ovulation in vivo.

To determine which aspect of ovulation correlates with increased DNA damage, TEC40 cells were treated with cycling or menopausal levels of FSH or LH or 10 nM estradiol to examine the impact of hormonal components of ovulation on DNA damage. To analyze the effects of oxidative stress associated with inflammation, 1 mM hydrogen peroxide, macrophage-conditioned media from RAW264.7 cells (MCM), or MCM from RAW264.7 cells stimulated with E. coli lipopolysaccharide (MCM + LPS) was added to TEC40 cells. Cells were analyzed for the presence of phospho-γH2A.X foci after treatment with basal media, hormone, hydrogen peroxide, MCM, or MCM + LPS (Fig. 7). In response to hydrogen peroxide or MCM + LPS, a significant increase in DNA damage was observed.

**Discussion**

In addition to a family history of breast and ovarian cancer, a primary risk factor for serous ovarian cancer is an increased number of lifetime ovulations (Auersperg et al. 1997). While the OSE is a commonly accepted site of origin for many histotypes of ovarian cancer, emerging evidence suggests that the TEC may be an alternative source for high-grade serous cancer (Crum et al. 2007b). The goal of this study was to examine the impact of ovulation on TEC by evaluating three commonly accepted hypotheses regarding the origin of ovarian cancer – the tear-and-repair
hypothesis, the gonadotropin hypothesis, and the inflammation hypothesis. Despite well-documented increases in OSE proliferation in the CD1 mouse strain 12 h post-hCG injection (Burdette et al. 2006), TEC in this study did not proliferate in response to ovulation in vivo, at either 12 or 16 h post-hCG injection. However, it is possible that TEC proliferation may occur at a more distant time point following ovulation. Although no increase in TEC proliferation was observed at these early time points following ovulation, heightened DNA damage and macrophage infiltration were detected.

**Figure 6** Ovulation induces double-strand DNA breaks in vivo and increases oviduct-associated macrophages. (A) Oviducts from basal or ovulated mice (12 or 16 h post-hCG) were sectioned and analyzed by immunohistochemistry for CK8 (red), phospho-\(\gamma\)H2A.X (green), and DAPI (blue). Calibration bars, 10 \(\mu\)m. Lower panels show CK8 staining (brown), which indicates the location of TEC cells. Upper panels are higher magnification of boxed area in lower panel. L, lumen of oviduct; P, periphery of oviduct. Yellow arrows point to damaged nuclei. (B) Positive cells were scored as CK8-positive TEC cells with nuclear phospho-\(\gamma\)H2A.X nuclear foci, relative to the total number of CK8-positive TEC cells. Data represents average fold change \(\pm\) S.E.M. compared with control. Significant differences \((P<0.05)\) are between groups labeled a and b. (C) Oviducts from basal or ovulated mice (12 or 16 h post-hCG) were sectioned and analyzed by immunohistochemistry for F4/80 macrophage glycoprotein (brown staining indicates antigen). COC, cumulus oocyte complex. Top panel, total magnification 400×. Bottom panel, total magnification 200×. Calibration bars, 20 \(\mu\)m.
In order to examine the components of ovulation that may contribute to fallopian tube neoplasia, such as the influence of gonadotropins and inflammatory factors on TEC proliferation and DNA damage, novel tools were developed. Using an alginate hydrogel matrix, sections of mouse oviduct or baboon distal fimbriae were cultured in a three-dimensional organ culture system, allowing for precise control of hormones or growth factors in contact with TEC. This system offers certain advantages over traditional cell culture, including retention of the underlying stromal tissue, and culture of non-transformed, non-immortalized cells. However, luminal structures are particularly sensitive to constriction within an alginate matrix, as has been observed for follicular antrum formation (Xu et al. 2006, West-Farrell et al. 2009). Compared with uncultured mouse oviducts, those cultured in alginate for 7 days showed a reduction in the thickness of the epithelial layer lining the interior of the oviduct, which likely reflects compression of the oviductal lumen due to growth within the alginate gel. Despite this compression, the cultured mouse oviducts showed similar rates of proliferation of TEC as observed in uncultured oviducts and retained both secretory and ciliated epithelium, with the highest proportion of ciliated cells found in sections of the oviduct closest to the ovary (infundibulum and ampulla) and decreasing toward the uterus (isthmus). This study also used baboon fallopian tube epithelium, which is structurally similar to human tissue with the TEC on the outer surface of the tissue. As a corollary to the organ culture system, an immortalized baboon fallopian tube cell line was generated that retained both secretory and ciliated TEC, allowing for comparative analysis of the effects of the gonadotropins on cell growth on two-dimensional plastic versus three-dimensional organ culture, as well as a renewable source of cells for studies on downstream signaling pathways and neoplastic transformation in TEC. The TEC40 cells were immortalized with SV40, which allows direct comparison of the IOSE cell lines that were generated in a similar manner (Auersperg et al. 1994, Karst et al. 2011). Similar to the IOSE cells and immortalized human TEC, the TEC40 cells do not form colonies in soft agar (data not shown), indicating that they could be used to model early transformative events (Karst et al. 2011).

Using three-dimensional organ culture to evaluate the gonadotropin hypothesis of ovarian cancer demonstrated that although FSH and LH at the tonically high levels observed during menopause induce proliferation of OSE, no increase in proliferation or DNA damage was detected in TEC, despite a common embryonic origin for these cells in the coelomic epithelium (Auersperg et al. 2008, Auersperg 2011). Transcripts for FSHR, LHR, and ERα were detected in the mouse oviduct and baboon fimbriae, indicating that signaling components responsive to FSH, LH, and estradiol are present in the tissues. Interestingly, in the mouse oviduct cultures, the gonadotropins suppressed proliferation of TEC, which may be due to a lack of growth factors in the media permitting analysis of the isolated effects of each hormone, whereas in vivo the cells are exposed to a variety of cytokines, growth factors, and hormones, resulting in a slightly higher basal rate of proliferation. At cycling levels, FSH caused a significant increase in TEC40 cell proliferation. Although these cells are morphologically normal, expression of SV40 T/t antigen disrupts p53 and retinoblastoma, leading to genetic changes in the cells that may allow for potential alterations in FSH-mediated signaling. The TEC40 cells exhibited little or no expression of ERα, which may also be reflective of altered signaling related to immortalization of the cells with SV40 or loss of receptor in long-term culture.
Oviduct TEC expressed increased levels of phospho-\(\gamma\)H2A.X, a marker of double-strand DNA breaks, after ovulation at 12 and 16 h after administration of hCG. Using the oviduct and fimbriae organ culture systems to determine whether FSH or LH caused DNA damage revealed no increase in phospho-\(\gamma\)H2A.X foci relative to controls as expected, as FSH and LH have not been shown to generate reactive oxygen species. Metabolites of estradiol have been shown to generate reactive oxygen species, increase inflammation, and play a role in progression of hormone-responsive cancers (Straub 2007). Postmenopausal estrogen therapy has been correlated with an increased risk of ovarian cancer, but the effect of estrogens on TEC is unknown (Ness et al. 2002). However, treatment of TEC with estradiol did not increase proliferation or DNA damage in culture after 7 days. Hydrogen peroxide has been frequently used to model oxidative stress generated by ovulation (Murdoch et al. 2001, Symonds et al. 2008). TEC40 cells treated with hydrogen peroxide or MCM+LPS exhibited increased levels of phospho-\(\gamma\)H2A.X foci, providing a common lesion in cell lines exposed to oxidative stress and in TEC in vivo after ovulation. Induction of H2A.X foci in response to oxidative stress was not likely a result of the altered p53 and Rb status of the TEC40 cells, as OVCA420 cells, which express wild-type p53 and Rb, exhibited similar levels of DNA damage in response to hydrogen peroxide or MCM+LPS (data not shown).

Ovulation caused an increase in activated macrophages associated with the oviduct and suggests one potential mechanism for how cells proximal to the site of ovulation may be susceptible to DNA damage. At 12 h post-hCG administration, activated macrophages were found primarily at the periphery of oviducts, likely associated with blood vessels, where they may be available for extravasation into the tubal epithelium. At 16 h post-hCG, examination of oviducts showed the presence of cumulus–oocyte complexes within the oviductal lumen, and activated macrophages were observed adjacent to and in contact with tubal epithelium near sites of ovulated oocytes. Macrophages associate with cumulus–oocyte complexes (Akkoyunlu et al. 2003), which together with the expelled follicular fluid from the site of ovulation may contain reactive oxidants that act on the tubal epithelium. The cumulus–oocyte complexes in the oviduct may release cytokines that promote further invasion of the macrophages into the tubal epithelium from local blood vessels (Zolli et al. 1991). Through release of cytokines and increase in cyclooxygenase (COX) enzymes associated with leukocyte infiltration into the ovary, ovulation increases levels of reactive oxidants, leading to formation of DNA adducts, base modifications, and strand breaks (Croteau & Bohr 1997, Murdoch 2008, Jabbour et al. 2009). This is the first study to show that ovulation can influence DNA damage in the TEC, perhaps due to the close proximity to the fallopian tube epithelium of the ovary and inflammatory factors released during ovulation. Further studies are needed to conclusively determine whether inflammation following ovulation causes DNA damage in the fallopian tube.

TECs are susceptible to DNA damage following ovulation, but the fate of the damaged cells is unknown. Although little or no apoptosis was observed in TEC following ovulation (data not shown), it is likely that some or all of the DNA damage is repaired. This may be reflected in a lack of increase in the percentage of TEC exhibiting H2A.X foci at 12 h post-hCG compared with 16 h post-hCG, despite an increase in macrophage association with the TEC. It is possible that by 16 h after hCG treatment, TECs have begun to repair the double-strand DNA breaks observed following ovulation. Cells with unrepaired double-strand DNA breaks or DNA replication errors as a result of oxidative damage at critical sites, such as in tumor suppressor genes, may serve as progenitor cells for neoplastic lesions (Schildkraut et al. 2010). It is possible that the OSE, which undergoes cyclic remodeling during monthly ovulations, is better equipped to process cells damaged in response to ovulation compared to TEC. Investigation into the pathways necessary to repair DNA strand breaks may provide valuable information regarding the mechanisms of p53 protein regulation as part of the tubal ‘precursor signature’ and how mutations in BRCA increase the risk of high-grade serous cancer.

In summary, ovulation impacts the fallopian tube epithelium as well as the OSE, lending support to the correlation between a reduction in the number of lifetime ovulations and a reduced risk of ovarian cancer. Ovulation and its associated hormones did not increase proliferation or directly induce DNA lesions using novel organ culture systems and a newly generated tubal cell line. However, in vivo ovulation induced macrophage infiltration into the fallopian tube epithelium and enhanced DNA damage in the TEC similar to that observed with reactive oxidants in vitro. This study examines the three major hypotheses connecting a long-standing correlation between ovulation and the risk of ovarian cancer in light of emerging evidence suggesting that the fallopian tube epithelium may be a site of origin for serous carcinomas.
Supplementary data
This is linked to the online version of the paper at http://dx.doi.org/10.1530/ERC-11-0107.

Declaration of interest
The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.

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Author contribution statement
S M King carried out immunohistochemistry and immunofluorescence analyses, participated in chemically induced ovulation studies, carried out organ culture studies, performed western blot analyses and RT-PCR, and drafted the manuscript. T S Hilliard carried out MTS analyses and immunofluorescence analyses. L Y Wu participated in immunohistochemistry analyses. R C Jaffe performed luciferase analysis of TEC40 cells and critical reading of the manuscript. A T Fazleabas provided baboon tissue and critical reading and revision of the manuscript. J E Burdette conceived of the study design, participated in its design and coordination, and involved in manuscript preparation and revision. All authors read and approved the final manuscript.

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