Clinical Research Article

Progesterone Receptors Promote Quiescence and Ovarian Cancer Cell Phenotypes via DREAM in p53-Mutant Fallopian Tube Models

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Abbreviations: ATP, adenosine triphosphate; BrdU, bromodeoxyuridine; cAMP, cyclic adenosine monophosphate; ChIP, chromatin immunoprecipitation; DCC, dextran-coated charcoal; DMEM, Dulbecco’s Modified Eagle Medium; DYRK, dual-specificity tyrosine-regulated protein kinase; E2, estradiol; EV, empty vector; FBS, fetal bovine serum; FT, fallopian tube; FTE, fallopian tube epithelium; hFTE, human fallopian tube epithelium; HGSC, high-grade serous ovarian cancer; IHC, immunohistochemistry; IRB, institutional review board; NT, nontargeting; OC, ovarian cancer; PBS, phosphate-buffered saline; PR, progesterone receptor; qPCR, quantitative polymerase chain reaction; RT-qPCR, quantitative reverse transcription–polymerase chain reaction; RU486, mifepristone; SAβ-Gal, senescence-associated β-galactosidase; SDS-PAGE, sodium dodecyl sulfate–polyacrylamide gel electrophoresis; shRNA, short hairpin RNA; STIC, serous tubal intraepithelial carcinoma; STIL, serous tubal intraepithelial lesion.

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Abstract

Context: The ability of ovarian steroids to modify ovarian cancer (OC) risk remains controversial. Progesterone is considered to be protective; recent studies indicate no effect or enhanced OC risk. Knowledge of progesterone receptor (PR) signaling during altered physiology that typifies OC development is limited.

Objective: This study defines PR-driven oncogenic signaling mechanisms in p53-mutant human fallopian tube epithelia (hFTE), a precursor of the most aggressive OC subtype.
Methods: PR expression in clinical samples of serous tubal intraepithelial carcinoma (STIC) lesions and high-grade serous OC (HGSC) tumors was analyzed. Novel PR-A and PR-B isoform-expressing hFTE models were characterized for gene expression and cell cycle progression, emboli formation, and invasion. PR regulation of the DREAM quiescence complex and DYRK1 kinases was established.

Results: STICs and HGSC express abundant activated phospho-PR. Progestin promoted reversible hFTE cell cycle arrest, spheroid formation, and invasion. RNAseq/biochemical studies revealed potent ligand-independent/-dependent PR actions, progestin-induced regulation of the DREAM quiescence complex, and cell cycle target genes through enhanced complex formation and chromatin recruitment. Disruption of DREAM/DYRK1s by pharmacological inhibition, HPV E6/E7 expression, or DYRK1A/B depletion blocked progestin-induced cell arrest and attenuated PR-driven gene expression and associated OC phenotypes.

Conclusion: Activated PRs support quiescence and pro-survival/pro-dissemination cell behaviors that may contribute to early HGSC progression. Our data support an alternative perspective on the tenet that progesterone always confers protection against OC. STICs can reside undetected for decades prior to invasive disease; our studies reveal clinical opportunities to prevent the ultimate development of HGSC by targeting PRs, DREAM, and/or DYRKs.

Key Words: progesterone, fallopian tube epithelia, DREAM, ovarian cancer, STIC, DYRK1 kinases

Ovarian cancer (OC) only accounts for about 3% of all cancers diagnosed in women in the United States, yet it is the most lethal of all gynecologic cancers and is the fifth leading cause of cancer deaths (1). Epidemiological studies have implicated reproductive status and hormone exposure in the pathogenesis of this disease (2). Parity, breastfeeding, and the use of oral contraceptives or hormone replacement therapy have been examined as either risk-inducing or preventive factors (3, 4). Progesterone, in particular, is theorized to be the protective steroid in these cases, presumably due to elevated endogenous progestogen during pregnancy or exposure to the synthetic progestin components of oral contraceptives. Full-term pregnancies, for example, are associated with reduced risk, with the greatest reduction observed for the less aggressive, low-grade type I tumors (5). The current or recent use of contemporary combined hormonal contraceptives (eg, lower estrogen dose, newer progestin compounds) resulted in a reduced risk of any OC subtype, in particular, endometrioid, mucinous, and serous (4). Yet, other studies have found that the use of oral and nonoral progestin-only contraceptives (eg, norethisterone, levonorgestrel, desogestrel, medroxyprogesterone acetate) did not diminish risk (4), and that estrogen-only and estrogen-progestin combination hormone replacement therapy use by postmenopausal women may actually increase ovarian cancer risk (6, 7). Therefore, the ability of ovarian steroids such as progesterone or estrogens to modify OC risk still remains highly controversial and the potential mechanisms involved are poorly understood.

The most common and aggressive OC subtype, high-grade serous ovarian cancer (HGSC), expresses abundant estrogen receptors (ER; 76% (8);) and progesterone receptors (PR; 35%) and ex vivo explants of these tumors exhibit functional PR signaling (8-10). It is believed that HGSC, in contrast to other subtypes, originates predominantly within the distal fimbriae of the fallopian tube (FT). The FT secretory epithelia acquire DNA damage and TP53 mutations, forming early neoplasms known as serous tubal intraepithelial carcinomas (STICs) (11, 12). Recent evidence suggests that these lesions may be present for many years as “dormant” STICs (ie, serous tubal intraepithelial lesions—STIL) before progressing to an “active” STIC (12, 13). These “dormant” lesions possess TP53 mutations but often show little to no proliferative activity, similar to normal fallopian tube epithelia (FTE) (14). The evolution of these lesions to an “active” STIC is marked by enhanced proliferative capacity and the acquisition of additional genetic alterations that set the stage for seeding of metastases on the ovary and other organs and surfaces within the abdominal cavity. If such lesions also possess functional PR signaling, how this signaling differs from normal FTE and whether progestins modulate the survival and dissemination of these neoplasms is not known.

Signaling via nuclear PRs is mediated primarily by 2 distinct, co-expressed PR isoforms: the full-length PR-B (1-933
important to note that PR can sense input from steroid hor-
transcriptomes of phospho-PR target genes (18, 19). It is
Lys388 desumoylated PR species which drive distinct
signaling (15-17). For example, phosphorylation of critical
all contribute to the downstream cellular outcomes of PR
receptor modifications (ie, phosphorylation, sumoylation)
in the presence of potential ligands and the posttranslational
isoforms can occur in the absence (ie, ligand-independent)
and presence (ie, ligand-dependent) of hormone; therefore,
both states are critical to the functions of PR and many
other steroid receptors (21, 22).

The biological effects of PR signaling are intimately
connected to cell fate in normal and cancerous breast and
uterine epithelia. In these tissues, PR is known to be tightly
coupled to many cell cycle mediators, regulating cell cycle
progression (23). It is accepted that progesterone signaling
modulates fallopian tube epithelia (FTE) functions which
support gamete and zygote transport and survival (24).

Less well-studied are the progestins’ effects on FTE cell
fate. Immunohistochemical analyses and gene expression
profiling of 3-dimensional primary FTE cultures suggests
that these epithelia are quiescent, especially those in the
early/mid luteal phase of the menstrual cycle when PR
signaling is most prominent (25, 26). Yet, the molecular
mechanisms involved in such progestin-induced regulation
are poorly understood, most likely because development of
in vitro PR-expressing FTE models has been difficult—loss
of steroid receptor expression is common during the
immortalization process.

For all cells, including FTE, the entry into G0 arrest or
quiescence is thought to be associated with the active tran-
scriptional repression of cell cycle dependent genes by the
multimeric complex known as DREAM which consists of
the DREAM-specific proteins, Dimerization partner DP1/2,
Rb-like p130/p107, E2F4/5 plus the core complex MuvB
(LIN9, LIN37, LIN52, LIN54, RBBP4 proteins (27)). Such
quiescence or dormancy is a natural state of certain cell
types, especially stem cell progenitors, and has recently
been recognized as a common feature of many cancers.

Cycling between dormancy and proliferation may be es-

tential for the acquisition of new mutations, survival in a
suboptimal environment, metastasis, and chemoresistance
(28, 29). Interestingly, malignant OC tumor cells in multi-
cellular aggregates, as observed in ascites, appear to be in a
dormant state (30). Amplifications, copy number losses,
or gains in genes encoding subunits of the MuvB com-
plex are commonly present in HGSC tumors and can be
associated with poor prognosis (27, 31). These studies
highlight the importance of understanding how cells enter/
exit quiescence in HGSC progression. However, the mech-

anisms regulating DREAM and cell cycle progression have
not been evaluated in normal fallopian tube epithelia or in
p53-mutant models of early disease.

Our studies described here reveal that nuclear and focal
activated PR is robustly expressed in STIC lesions and inva-
sive HGSC. Using novel PR-expressing human fallopian
tube epithelial (hFTE) cell models, we show that proges-
tins are able to promote cellular phenotypes that could
support STIC lesion shedding and dissemination, including
cell-cell aggregation/spheroid formation and collagen in-
vasion. In the absence of progestins, PR-A isoforms act as
dominant inhibitors of DREAM, promoting permissive

cell cycle progression and migration, while both liganded
receptor isoforms associate with and activate DREAM as
well as dual-specificity tyrosine-regulated protein kinases
(DYRKs), driving cells into a quiescent state that may sup-
port early ovarian cancer progression.

Methods

General Reagents

R5020 (Perkin Elmer), progesterone (P4; Sigma-Aldrich),
estradiol (E2; Sigma-Aldrich), mifepristone (RU486,
Sigma-Aldrich), and onapristone (Context Therapeutics)
stocks were prepared in ethanol (EtOH). Harmine (Sigma-
Aldrich) stock was prepared in dimethyl sulfoxide.
Epidermal growth factor (EGF; Sigma-Aldrich) was pre-
pared in 10 mM acetic acid with 0.1% bovine serum al-
bumin (BSA).

Human Tissues

Fresh frozen normal fallopian tube (FT) tissues were ac-
quired from patients undergoing salpingo-oophorectomy
or total abdominal hysterectomy with bilateral salpingo-

oophorectomy at the Fairview-University of Minnesota
Hospital. Biospecimen procurement protocols were ap-
proved as institutional review board (IRB)-exempt and ad-
ministered through the Clinical and Translational Sciences
Institute’s Biorepository & Laboratory Services BioNet
division. FT tissue was dissected from surrounding tissues,

snap-frozen on dry ice and stored at −80 °C until analysis.

Cases used for immunohistochemistry analyses were
selected from University of Pennsylvania Medicine patients
who were diagnosed with high-grade serous ovarian cancer
(HGSC) and showed evidence of serous tubal intraepithelial
carcinoma (STIC) lesions within the FT. Most of these cases
also exhibited invasive HGSC within FT stroma and/or
associated ovarian tissues; both STIC and HGSC sections were available for analysis. Procurement of biospecimens was covered by Penn Ovarian Cancer Research Center Biotrust Collection IRB protocol #702679. De-identified formalin-fixed paraffin-embedded blocks were obtained and sections cut and processed as outlined below.

Fresh normal FT tissue utilized for the establishment of the original UWFT.1681 cell line was collected from a University of Washington Seattle Medicine patient undergoing salpingo-oophorectomy. This patient provided informed consent under a protocol approved by the IRB of the University of Washington (#2872).

**Immunohistochemistry**

Four-micron thick sections of formalin-fixed tissue were used for immunoperoxidase analysis after baking at 60 °C for 1 hour, followed by deparaffinization and rehydration in successive xylene, ethanol, and water washes. The sections were blocked with 3% hydrogen peroxide in methanol and antigen retrieval was performed in a pressure incubator (Biocare Medical) at 123 °C in citrate buffer (DAKO Target Retrieval Solution). Slides were cooled and transferred to Tris-buffered saline. Primary antibodies and conditions are listed in Supplemental Table 1 (32). The secondary antibody was used per protocols and reagents in the DAKO Envision + System. The sections were developed using 3,3′-diaminobenzidine (DAB; Sigma Chemical Company) as substrate and counter-stained with Mayer's Hematoxylin. Bright-field images were acquired in a Huron Tissuescope LE equipped with a Nikon Plan Apo 20X Objective Lens NA 0.75 housed and managed by the University of Minnesota University Imaging Centers. Scans were subsequently analyzed, and images captured and calibrated in Huron Viewer software (Huron Digital Pathology). Identity of STICs was verified by presence of atypical histology and scoring for presumed p53 mutations, including p53 missense mutation characterized by strong staining, null p53 mutation with little to no staining, or wild-type p53 with blush staining (33). PR and p-PR staining was scored as nondetectable/low intensity (−/+ ) to high intensity (+++) relative to the analyzed tissues. Relative intensity of p-PR staining is illustrated in Supplemental Figure 1 (32). Case information, mutations, and staining intensity are summarized in Fig. 1C.

**Cell Lines and Culturing**

All cell lines were maintained at 37 °C under 5% CO2 in water-jacketed incubators. FT282 cells were cultured in Dulbecco's Modified Eagle Medium (DMEM)/F12 (Corning) supplemented with 10% charcoal stripped fetal bovine serum (DCC, Hyclone) and 1% penicillin-streptomycin (Gibco). Murine oviductal epithelium (MOE (34)) cells were gifted from Dr. Joanna Burdette (University of Illinois at Chicago) and were cultured in Minimum Essential Medium-Alpha (Corning) supplemented with 10% fetal bovine serum (FBS, Corning), 1% penicillin-streptomycin (Gibco), 1.1 μg/mL gentamicin (Abcam), 1.1% ITS (Corning), 2 ng/mL epidermal growth factor (Sigma-Aldrich), 20 ng/mL estradiol (Sigma-Aldrich) and 1% L-glutamine (Gibco). UWFT.1681 cells were cultured in MEGM (Lonza) supplemented with MEGM SingleQuots supplements (Lonza), 1% FBS, and 1% penicillin-streptomycin. All cell lines were routinely tested for mycoplasma using an e-Myco Plus Mycoplasma PCR Detection kit (Bulldog Bio Inc) and confirmed to be negative prior to experimentation.

**Stable Cell Line Generation**

Stable hFTE PR-expressing lines were generated by transducing FT282 cells with pLenti-CMV neo lentiviral vector containing PR-A or PR-B. Stable pools were selected and maintained in 0.2 mg/mL G418 (Corning) and then plated as single cells to generate empty vector (EV), PR-A, and PR-B clonal lines. Stable shDYRK1A (clones TRCN0000000253 and 524) expressing cell lines and shDYRK1B (clones TRCN0000002140 and 2141) expressing cell lines were generated by transducing FT282 PR-B+ and PR-A+ cells with pLKO.1 lentiviral vector containing target gene short hairpin RNA (shRNA) sequences for DYRK1A (AATACAAGAATCAAATGCTG and AAAGTCCAAGGTATTAGCAGC) or DYRK1B (TAGCAGCATTCCAGTCAAGG and ATATAGTACTTCATCTCCGTG). Stable pools were selected and maintained in 1 μg/mL puromycin and 0.2 mg/mL G418. Original UWFT.1681 cell line was established using dispersed FT epithelia from fresh primary FT tissues. Cells were immediately plated on 100 mm plates in MEGM media (Lonza) with 1% FBS. After 24-hour culturing with fibroblast adherence, supernatant was transferred serially to new plates for FTE cultures. Epithelial purity was confirmed by immunohistochemical evaluation of cytokeratins using the AE1/AE3 antibodies. Confluent FTE were infected with HPV E6/E7 as previously described (35).

**Immunoblotting**

Snap-frozen FT tissues were homogenized with a metal tissue pulverizer over dry ice. Protein was isolated with RIPA-lite lysis buffer (0.15 M NaCl, 6 mM Na2HPO4, 4 mM NaH2PO4, 2 mM EDTA, 0.1 M NaF, and 1% Triton-X 100 in H2O supplemented with 1x complete mini protease
Figure 1. Progesterone receptor expression in normal human fallopian tube, STIC lesions, and invasive HGSC. 

A. Western blot analysis of PR-A and PR-B isoform expression and phosphorylated PR at Ser294 (p-PR-A, p-PR-B) expression in normal FT tissues. T47D CO line treated with vehicle (veh) or R5020 (10nM) for 1 hour as a positive control. 
PAX8 as FT positive control and GAPDH as loading control.

B. Immunohistochemical (IHC) staining of normal FT tissues. Low (left) and high (right) magnification of hematoxylin-eosin (HE), PR and phosphoS294 PR (p-PR). Arrow indicates FT epithelia layer and star indicates FT stroma. 
Scale bars = 50 µm.

C. Patient case information and IHC staining scores. Fallopian tube tissues obtained from patients with HGSC diagnosis and presence of STIC lesions. Scoring of PR and p-PR proteins is indicated ranging from absence (−), light variable (−/+), low (+) to high (+++) intensity staining, based on relative intensity compared across the 8 cases. Presence of nuclear foci of p-PR staining within STICs is noted.

D. IHC of representative STIC (case 1) exhibiting p53 missense mutation. Red arrowheads indicate normal hFTE.
Scale bars = 50 µm.

E. STIC and invasive HGSC (case 5). Scale bars = 50 µm.

F. Nuclear foci of p-PR protein within STIC (case 1). Scale bars = 25 µm.
inhibitors [Roche], 1× PhosSTOP tablet [Roche], 25 mM β-glycerophosphate [BGP], 1 mM phenylmethanesulfonyl fluoride [PMSF], 20 µg/mL aprotinin [Fisher Bioreagents], 5 mM NaF, and 0.05 mM NaVO₄). Alternatively, adherent cells were washed with phosphate-buffered saline (PBS) and harvested with RIPA-lite lysis buffer. Lysates were cleared by centrifugation, quantified through Bradford assays using Bio-Rad reagent and equal protein concentrations were resolved on 8% sodium dodecyl sulfate–polyacrylamide gel electrophoresis (SDS-PAGE) gels. Proteins were then transferred to Immobilon-P polyvinylidene difluoride membranes (Millipore), probed with antibodies as listed in Supplemental Table 1 (32) and developed using SuperSignal West Pico Plus Chemiluminescent Substrate (Pierce).

Quantitative Reverse Transcription Polymerase Chain Reaction
Cells were plated in triplicate in phenol red–free DMEM/F12 supplemented with 5% DCC and treated the following day with vehicle or R5020 (10 nM). After the indicated time point, cells were washed with PBS and total RNA was harvested using TriPure Isolation Reagent (Roche) and isopropanol precipitation. According to the manufacturer’s instructions, 1 µg of RNA was then reverse transcribed to cDNA using qScript cDNA SuperMix (Quanta Biosciences). The quantitative polymerase chain reaction (qPCR) was performed using FastStart Essential DNA SYBR Green Master (Roche) on a LightCycler 96 Real-Time PCR Instrument (Roche). Human primer sequences are listed in Supplemental Table 2 (32). The qPCR cycling conditions were as follows: initial denaturation at 95 °C (10 minutes), denature at 95 °C (10 seconds), anneal at 60 °C (10 seconds) and extension at 72 °C (5 seconds) for 45 cycles. Target gene expression levels were normalized to the TATA-box binding protein gene (TBP).

Cell Proliferation Assays
Screening of the proliferative capacity of EV and PR-expressing clonal lines was initially determined using the CellTiter-Glo assay (Promega). Cells were seeded in triplicate in opaque-walled 96 well plates in phenol-free DMEM/F12 supplemented with 10% DCC. After the cells were settled, a baseline Day 0 adenosine triphosphate (ATP) concentration was determined using the CellTiter-Glo 2.0 Assay (Promega) according to the manufacturer’s instructions. Luminescence was measured using Synergy 2 (Biotek) and ATP concentrations were determined by generating an ATP standard curve with ATP disodium salt (Sigma). This process was repeated for the Day 2, 4, and 6 plates and the ATP concentrations were normalized by subtracting the Day 0 ATP concentration.

Immunofluorescence Assays
Cells were seeded on coverslips in phenol red–free DMEM/F12 supplemented with 5% DCC and treated the following day with vehicle or R5020 (10 nM). After 72 hours, cells were fixed with 4% paraformaldehyde and permeabilized with 0.1% Triton-X-100. All antibodies and blocking conditions are listed in Supplemental Table 1 (32). Coverslips were then mounted with ProLong Gold Antifade with DAPI (Molecular Probes) and analyzed using a Leica DM40000 B microscope. For quantitation of Ki67 staining, the percentages of Ki67+ cells were analyzed and calculated for ~10 fields per coverslip using a Leica DM40000 B microscope.

Senescence-Associated β-Galactosidase Activity Assays
Cells were seeded on coverslips in phenol red–free DMEM/F12 supplemented with 5% DCC and treated the following day with vehicle or R5020 (10 nM). After 72 hours, cells were washed with PBS, fixed with 4% paraformaldehyde, and stained for senescence-associated β-galactosidase (SAβ-Gal) activity using the Senescence β-Galactosidase Staining Kit (Cell Signaling Technology) according to the manufacturer’s instructions. Cells were analyzed using a Leica DM40000 B microscope. The percentages of β-galactosidase+ cells were analyzed and averaged for ~10 fields per coverslip using a Leica DM40000 B microscope.

5-Bromo-2′-Deoxy-Uridine Immunofluorescence Assays
Cells were seeded on coverslips in phenol red–free DMEM/F12 supplemented with 5% DCC and treated the following day with vehicle or R5020 (10 nM). After 72 hours, cells were incubated with bromodeoxyuridine (BrdU), fixed with ethanol at −20 °C and stained for BrdU using the 5-Bromo-2′-deoxy-uridine Labeling and Detection Kit I (Sigma-Aldrich) according to the manufacturer’s instructions. Percentages of BrdU+ cells were analyzed and averaged for ~10 fields per coverslip using a Leica DM40000 B microscope.

Migration Assays
Cells were harvested with 0.25% trypsin-EDTA (Invitrogen), washed and resuspended in phenol red–free DMEM/F12, and seeded in triplicate in 8-µm pore 24-well Transwell inserts (Corning). Medium (10% FBS or serum-free control medium) was placed in the bottom chamber of the 24-well plates and the cells were incubated for 18 hours. Membranes were then fixed with 4% paraformaldehyde,
cut out of the chamber and mounted with ProLong Gold Antifade with DAPI (Molecular Probes) and analyzed using a Leica DM40000 B microscope. The numbers of migrated cells were counted for 6 representative fields per chamber and averaged between conditions.

3D Spheroid Culture Assays

Cells were harvested with 0.25% trypsin-EDTA, washed, and resuspended in phenol red–free DMEM/F12 supplemented with 10% DCC. The cells were then sieved through 40-µm pore cell strainers (Falcon) and 8 single-cell suspensions (~1000 cells/well) were seeded per condition in 96-well U-bottom plates coated with 6 mg/mL poly(2-hydroxyethyl methacrylate) (Sigma). After 72 hours, single spheroids were imaged and measured on a Nikon Eclipse Ts3 microscope. Spheroid diameter was quantified for approximately 8 wells/data point.

3D Spheroid Invasion Assays

Collagen Type I (Corning) was neutralized to a pH of 7.4 and diluted to 2 mg/mL with phenol red–free DMEM/F12 supplemented with 10% DCC. Then 50 µL of diluted collagen was transferred into the wells of 96-well plates and incubated for 30 minutes at 37 °C to solidify the collagen. Using cold-cut pipette tips, single spheroids were individually harvested, resuspended in 100 µL collagen, and transferred on top of the solidified collagen layers. The spheroid-collagen layer was incubated for 1 hour at 37 °C to solidify before spheroid culture medium containing experimental treatments was plated on top. Spheroids were initially imaged using a Nikon Eclipse Ts3 microscope at Day 0 and then allowed to invade for 72 hours. Each spheroid was again imaged at the end of this 72-hour incubation. Areas (µm²) for each spheroid at Day 0 and Day 3 were calculated from these digital images using ImageJ. The area of collagen invaded was calculated by subtracting the initial area of that spheroid at Day 0 from the final Day 3 area of that same spheroid to determine the relative invasion area for that single spheroid/well. Invasion area was quantified for approximately 6 to 8 wells per average data point.

Gene Expression Profiling

For RNAseq studies, clonal lines were pooled (EV #3,11; PR-A #7, 9; PR-B # 20, 24) and plated in triplicate. RNA was isolated using RNAeasy kit (Qiagen) according to the manufacturer’s instructions. RNA quantity, quality, and size were determined by RiboGreen and Agilent BioAnalyzer. Strand-specific RNAseq libraries were created (Illumina TruSeq), quality/quantity verified (BioAnalyzer) followed by ~20 million 50 paired-end sequencing reads performed per library (HiSeq 2500 high-output mode) at the University of Minnesota Genomics Center. Bioinformatics analyses of RNAseq results was provided by Artificial Intelligene (Intelligene Technologies, Kenosha, WI; www.artificialintelligene.com). Briefly, each sample was aligned, BAM files sorted/indexed, and transcript abundance files created. Reads were aligned to the hg19 human genome using STAR (36). Cufflinks software (version 2.2.1) was used to generate transcript assemblies (37). Identification of differentially expressed genes using the reference and experimental groups of interest was performed using DESeq2 with a cutoff of ≥ 2-fold (38). Cluster analyses on sample groups was calculated where union of all the genes and their expression fragments per kilobase of exon per million reads (FPKM) values within that group were generated to build a read count matrix for the groups of interest. Unsupervised and other machine learning techniques were applied to this composite read count matrix of interest. R packages, including ggplot2, heatmap.2, and Pheatmap, were used to build various heatmaps with sample-feature heatmaps representing the signal intensity of a feature for any given sample. The volcano plots were created using data from differential expression values downloaded from the Artificial Intelligene interface and then R software (v 3.5.2) was implemented for the creation of these plots. Functional pathway enrichment in the gene list of interest was performed using gene set enrichment analyses (39) on Gene Ontology, Broad Institute, and Reactome input datasets from Artificial Intelligene (40). RNAseq data is available through the Gene Expression Omnibus (GEO) database (41).

Immunoprecipitations

Cells were seeded in phenol red–free DMEM/F12 supplemented with 5% DCC and treated the following day with vehicle or R5020 (10 nM). After the indicated time point, cells were washed with PBS and harvested with PBS supplemented with Protease-inhibitor Cocktail Set 1 (Calbiochem, 1:100) and Protease-inhibitor Cocktail Set 2 (Calbiochem, 1:500). Cells were pelleted through centrifugation and snap-frozen on dry ice. For immunoprecipitations, cell pellets were lysed in EBC buffer (Boston Bioproducts) supplemented with Protease-inhibitor Cocktail Set 1 (1:100), Protease-inhibitor Cocktail Set 2 (1:500) and 2-mercaptoethanol (1:10000). To immunoprecipitate DREAM and B-MYB/ MMB complexes, 2 mg of lysate were incubated with 1 µg LIN37 antibody (Bethyl) and protein A sepharose
CL-4B beads (GE Healthcare) overnight at 4 °C. The samples were then washed with EBC, eluted with SDS-PAGE gel loading buffer and resolved on SDS-PAGE gels using standard protocols. After transfer to nitrocellulose membrane (Bio-Rad), the proteins of interest were probed with antibodies listed in Supplemental Table 1 (32) and protein band densities were calculated using ImageJ software as previously described (31).

Flow Cytometry
Cells were seeded in phenol red–free DMEM/F12 supplemented with 5% DCC and treated the following day with vehicle or R5020 (10 nM). After 24 hours, cells were collected, fixed, and stained as described (42). Cells were stained with Hoechst 33342 DNA-specific dye (Invitrogen). Acquisition and separation of the phases of the cell cycle were performed on a BD LSRFortessa Flow Cytometer System (BD Biosciences). Data analysis was performed using FlowJo v10 software (BD Biosciences).

Chromatin Immunoprecipitation
Cells were seeded in phenol-free DMEM/F12 supplemented with 5% DCC and treated the following day with vehicle or R5020 (10 nM) for 3 hours. Cells were then fixed, harvested, and lysed according to optimized manufacturer’s instructions using the ChiP-IT Express Magnetic Chromatin Immunoprecipitation Kit (Active Motif). Samples were homogenized using a Bioruptor sonicator (Diagenode, Inc.). Chromatin immunoprecipitation (ChiP) reactions were incubated overnight on an end-to-end rotator using antibodies listed in Supplemental Table 1 (32). Samples were washed, eluted, reverse cross-linked, and treated with Proteinase K according to manufacturer’s instructions (Active Motif). DNA was analyzed by reverse transcriptase-qPCR (RT-qPCR) using primers and chromosomal coordinates listed in Supplemental Table 3 (32).

Statistical Analyses
For all datasets, a Shapiro-Wilk normality test for normal distribution was performed followed by, as appropriate, a 1-way or 2-way analysis of variance (ANOVA) in combination with the Tukey multiple comparisons (Prism 8; GraphPad software). Statistical significance was set at $P < 0.05$ and graphs of continuous variable data show the mean ± standard deviation ($n =$ sample size). All data presented is representative of at least 3 independent replicate experiments.

Results

Progesterone Receptor Expression in Normal Fallopian Tube, STIC Lesions, and Invasive HGSC
Little is known about PR or the PR phosphospecies expressed in normal fallopian tube (FT) epithelia or in neoplasms of these cells. Therefore, we examined the expression of both PR and phospho-Ser294 PR in normal FT and STIC lesions. Immunodetection of phospho-S294 PR was performed using a custom polyclonal antibody developed in our laboratory and previously validated for cellular protein assays and immunohistochemistry (IHC) of breast cancer tissue microarrays (19). Detection of “total” PR (ie, the combination of both unphosphorylated and multiple phosphospecies of both isoforms) was conducted with a commercial monoclonal antibody routinely used by our lab (SCBT sc-166169; Supplemental Table 1 (32)) as well as a clinical monoclonal antibody (Dako 3569 cl PgR636; Supplemental Table 1) (32). Western blot analyses of normal human FT tissues confirmed robust expression of both total PR-A and PR-B isoforms in all samples, as well as strong (3/5 cases) to weak (2/5 cases) expression of phosphorylated PR protein at Ser294 (p-S294, p-PR; Fig. 1A). This expression is localized primarily to the epithelial layer lining the FT, with nuclear staining of both PR isoforms and the phosphorylated S294 proteins of both isoforms (Fig. 1B).

We first reported that ~35% of metastatic HGSC tumors express abundant progesterone receptors (8). Herein, FT and associated HGSC tumor tissues from each HGSC patients with confirmed STIC lesions and invasive HGSC were stained for p53, total PR, and p-PR expression (Fig. 1C-1F; Supplemental Fig. 1 (32)). Both PR and p-PR staining was observed in STICs (Fig. 1C-1D) and this expression was retained in invasive HGSC tissue when present, whether localized within the FT stroma or ovarian tissue (Fig. 1E). Interestingly, p-PR staining was often localized within punctate nuclear foci (Fig. 1F); a phenotype indicative of active transcriptional complexes (43, 44). Such nuclear foci were more commonly observed in STIC lesions but were occasionally seen in adjacent normal FTE. As expected, the presence or intensity of the total vs phosphorylated PR protein was sometimes discordant; p-PR protein was consistently present when total PR was sometimes low or nondetectable in the same tissue. Additional staining with a diagnostic PR monoclonal antibody revealed the same staining patterns (Dako 3569 cl PgR636; data not shown). We have reported differential detection of total PR and phospho-PR proteins in breast cancer tissues using similar antibodies and antibody protocols (19, 32). This may be due to the inherent differences in the recognition

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of multiple epitopes by polyclonal antisera versus single epitope recognition by a monoclonal antibody, as well as the potential for epitope masking due to phosphorylation or other posttranslational modifications of PRs present in multiprotein nuclear complexes. In addition, we and others have reported differing turnover rates of PR isoforms relative to their PR phosphospecies (45, 46).

Overall, these IHC analyses demonstrate that activated progesterone receptor species (ie, phospho-S294 PR) can be reliably detected and are well-expressed in STICs as well as in HGSC tumors localized to the fallopian tube and the ovary.

Generation of PR-Expressing Human Fallopian Tube Epithelial Models

The lack of robust cell models of human FT epithelia that express ovarian steroid receptors, such as PRs, has hampered mechanistic studies of steroid receptor signaling. Immortalization procedures and 2-dimensional culturing invariably leads to loss of endogenous receptor expression. To facilitate our studies, the FT secretory cell line which stably expresses the p53 missense mutation R175H (47), was genetically engineered to express either the PR-A or PR-B isoform. Multiple clones of empty vector controls (EV; #3, #11), PR-A+ (#7, 8, 9) and PR-B+ (#15, 20, 24) cells were isolated, which showed similar protein expression to T47D CO breast carcinoma cells expressing endogenous PRs (Fig. 2A). Only the PR+ clones exhibited rapid activation of PR isoforms with phosphorylation of the Ser294 site following treatment with R5020, a stable progestin that binds specifically to PR (Figs. 2B-2C). As expected, this potent PR agonist shows activation of phosphorylation at concentrations between 1 nM and 10 µM whereas the natural ligand, progesterone, shows activation at a slightly narrower concentration range of 10 nM to 10 µM (Supplemental Fig. 2A (32)). Additional sites important for kinase modulation of PR signaling such as Ser345 were also rapidly phosphorylated in response to R5020 and progesterone (Fig. 2C; Supplemental Fig. 2B (32)), suggesting that PR isoforms expressed in hFTE cells were properly folded and functional with regard to rapid induction of signaling pathways and subsequent phosphorylation events.

To verify downstream ligand-induced transcriptional repression/activation in these lines, the regulation of select PR target genes previously identified by our laboratory in ES-2 OC cell models was examined (8). Quantitative RT-PCR analyses showed concentration-dependent progestin- or progesterone-induced downregulation (ie, transcriptional repression) of interleukin-1β mRNA in PR-A+ and PR-B+ hFTE cells at 6 hours and upregulation (ie, transcriptional activation) of NEDD9 mRNA at 48 hours, with no change observed in EV cells at either time point (Supplemental Fig. 2C (32)). Analyses of multiple PR+ clones showed similar regulation of these genes and revealed PR-A-dominant downregulation of interleukin-8 mRNA at 6 hours (Supplemental Fig. 2D (32)) and PR-B-specific upregulation of FOXO1 mRNA at 48 hours of progesterin or progesterone treatment (Supplemental Fig. 2E (32)).

PR and Progestins Promote Cell Behaviors That Support Cancer Phenotypes

Mutations acquired within fallopian tube epithelia can lead to cell survival, proliferation, cell-aggregation/disaggregation, and other cell behaviors that can support disease progression (11, 12, 48). The effect of PR expression and progestin treatment on phenotypes that could promote such processes was examined. The initial characterization of proliferation was analyzed using an ATP-based assay that allowed for high-throughput measurement of rate of change over time in numerous clones. The expression of PR-A alone led to enhanced proliferation as compared to EV and PR-B+ hFTE (Fig. 2D). In addition, these cells also exhibited greater migratory capacity in transwell migration assays in the presence of 10% FBS (ie, as the chemo-attractant; Fig. 2E). Signaling in the absence of hormone (ie, ligand-independent; unliganded) is a common feature of steroid receptors, including PR, and is as important as their ligand-dependent (liganded) actions (21, 22).

The ability of progestins to mediate FTE cell fate through the regulation of cell cycle progression was studied in these hFTE lines. Cellular analyses of indices of proliferation (Ki67), senescence (β-galactosidase; β-gal), and DNA synthesis (BrdU incorporation) were employed in parallel samples using immunofluorescence assays. Treatment with R5020 for 48 hours resulted in a decrease in proliferative cells in the PR+ hFTE, visualized by a decline in Ki67+ cells along with an increase in senescence-associated β-galactosidase (SAβ-gal) cells (Fig. 2F; immunofluorescence images-Supplemental Fig. 3A (32)). A coincident decrease in DNA synthesis also occurred, observed as a decline in BrdU+ cells (Fig. 2F; immunofluorescence images-Supplemental Fig. 3B (32)). Additionally, flow cytometry analyses showed that, within 24 hours, these cells were pushed into G0/G1 with concomitant decrease in the percentage of cells in S phase (Fig. 2G). This inhibition, however, did not appear to be a state of terminal arrest (senescence) since serum stimulation for 24 to 48 hours following progestin treatment resulted in a recovery of proliferative Ki67+ cells (Fig. 2H). Exposure to serum for several days following this cell cycle arrest led to a steady increase...
**Figure 2.** Novel PR-expressing human fallopian tube epithelia (hFTE) show functional PR signaling and progestin-mediated cell cycle arrest. 

A, PR expression in empty vector (EV), PR-A+, and PR-B+ hFTE clonal lines; T47D CO cells as a positive control.

B, Phosphorylation of PR at Ser294 in hFTE PR-A+ (#7) and PR-B+ (#24) clones following vehicle (−) or varying concentrations of R5020 (1 hour).

C, Phosphorylation of PR at Ser294 and Ser345, in hFTE following vehicle (−) or R5020 (+; 10nM; 1 hour).

D, Proliferation of hFTE EV #1, PR-A #7, #8, and PR-B #15, #24 over 0, 2, 4, and 6 days.

E, Transwell migration of hFTE EV #1, PR-A #7, and PR-B #24 in response to 0% (control) or 10% FBS for 18 hours.

F, Percentage of hFTE cells positive for Ki67, SAβ-gal, and BrdU (cells/field) following treatment with vehicle (veh) or R5020 (10nM; 72 hours).

G, Cell cycle analyses (FACS) of hFTE treated with vehicle (veh) or R5020 (10nM; 24 hours).

H, Percentage of positive Ki67 hFTE cells after 0 to 2 days serum stimulation. Treatment of cells for 3 days with vehicle or R5020, followed by a change to media containing FBS (no R5020) for 0-2 days (schematic). Graphs represent the mean ± SD, ***P < 0.001 (n = 3).
in cell numbers (Supplemental Fig. 3C (32)). These studies demonstrate that progestins are able to promote G0 cell cycle arrest in PR+ hFTE that is fully reversible.

The ability of progestins to support the formation of tumor emboli (ie, spheroids), a process important for dissemination, was also examined. R5020 treatment robustly promoted cell-cell aggregation with larger, compact spheroids observed for PR-A+ and PR-B+ hFTE (Fig. 3A). In contrast, no PR expression (EV hFTE) or the absence of progestins (vehicle) resulted in looser, smaller structures (Fig. 3A). To simulate the disagggregation/reaggregation thought to occur during dissemination of early FTE lesions (49, 50), these structures were dissociated and reseeded without treatments. Spheroid formation under these conditions only occurred in the PR+ hFTE cells treated with R5020 during the initial aggregation (Fig. 3B). No discernable structures were evident for EV hFTE or for PR-A+ or PR-B+ hFTE cultured with vehicle only. Importantly, aggregation experiments were also conducted with an additional cell model. Immortalized murine oviductal epithelial (MOE) cells, expressing low endogenous levels of both PR-A and PR-B (inset, Fig. 3C), also exhibited enhanced spheroid formation with R5020 treatment (Fig. 3C).

Notably, treatment of cells with the PR antagonists, onapristone (Ona) or RU486, did not block PR-driven spheroid formation and, similar to R5020 (ie, a PR agonist), both compounds supported cell aggregation (Supplemental Fig. 4A (32)). Analysis of S294 phosphorylation revealed that onapristone and RU486 were unable to block R5020-induced PR phosphorylation and acted instead as partial agonists, inducing phosphorylation at the S294 residue (Supplemental Fig. 4B (32)). These antagonists are effective in blocking R5020 effects in breast cancer lines (19), suggesting unique actions of these compounds that may be specific to the hFTE cell type and/or the state of these cells. Interestingly, it is well known that PR antagonists like RU486 acquire partial agonist activity in a background of elevated cyclic adenosine monophosphate (cAMP) activity (51) and that selective estrogen receptor modulators such as tamoxifen also tend to have agonistic actions in reproductive tissue like the uterus, contrary to its antagonistic actions in the breast (52). The basis for this reversal of antagonist to agonist action is unknown, but likely involves activation of rapid signaling pathways (cAMP, MAPKs) known to modify nuclear steroid receptors, such as hormone-binding membrane estrogen receptors (mERs, GPER) and membrane PRs (mPRs or PAQRs, PGRMC1) also highly expressed in the reproductive tract (53, 54).

Epithelial cells within FT lesions can be shed, aggregate into emboli, and potentially go on to invade the layer lining the peritoneal cavity (55, 56). Interaction with the abundant collagen type I layer beneath the mesothelium is important for OC cell attachment, motility, and invasion (50, 57). To determine the effect of PR signaling on a similar cell behavior, we utilized a simplified model of collagen invasion where our hFTE lines were cultured in the absence or presence of R5020 to form spheroids, encased in a collagen type I matrix, then allowed to invade for 72 hours. Remarkably, we observed heightened invasive behavior that was fully dependent on PR expression and the presence of R5020 during spheroid formation (Fig. 3D). Similar to what was observed with spheroid formation, the activation of PR by progestin was necessary to support this invasive behavior.

### Gene Expression Profiling of PR+ hFTE Highlights Unique Regulation

To define the potential pathways transcriptionally regulated by PR and progestins that could drive these cancer-like phenotypes, RNAseq analyses was performed on EV and PR+ hFTE treated with vehicle or R5020 for 6 and 48 hours. Representative Venn diagrams (Fig. 4A) and heat maps were created to analyze the resulting transcriptomes via either unsupervised (Supplemental Fig. 5A (32)) or supervised clustering (ie, by treatment; Supplemental Fig 5B (32)). In the absence of progesterin, cells expressing either PR-A or PR-B regulate distinct gene clusters relative to the same parental cells expressing EV. Surprisingly, however, simply the expression of PR-A in the absence of R5020 dramatically altered the transcriptome, activating over 400 genes and repressing over 500 genes at both time points relative to both untreated EV or PR-B+ cells (Fig. 4A; PR-A = blue, PR-B = yellow). The addition of ligand to PR-A+ cells induced only subtle further changes in gene expression; compare all PR-A gene sets to all EV or PR-B gene sets (Supplemental Fig. 5A (32)) and compare gene sets in lanes 1 and 2–3 or lanes 6 and 7–8 (Supplemental Fig. 5B (32)).

Ligand-independent actions of PR-A as measured by changes in global gene expression have also been reported in breast (18) and ovarian cancer (9) models but unliganded PR-A was predominantly repressive in these cancer contexts. In sharp contrast to PR-A expression, the expression of PR-B in the absence of ligand was most similar to EV (Supplemental Fig. 5A (32); far right cluster). Strikingly, the addition of R5020 to PR-B+ cells, especially at 48 hours, dramatically altered gene expression (Fig. 4A; 808 upregulated genes, 572 downregulated genes) relative to vehicle controls. This is also evident in Supplemental Fig. 5B (32); compare supervised clustering of lanes 4 and 5 or lanes 9 and 10. These data suggest that similar to both breast and ovarian cancer models (9, 58), PR-B is the dominant hormone-regulated (ie, hormone-sensitive) isoform,
Figure 3. PR and progestins promote cell behaviors that support cancer phenotypes. A-B, Representative bright-field images and average diameter of primary aggregated (A) and secondary dispersed and reaggregated (B) spheroid cultures of hFTE treated with vehicle (veh) or R5020 (10nM; 72 hours). No treatments were included in secondary cultures. C, Bright-field images and average area of murine oviductal epithelium (MOE) spheroid cultures treated with vehicle (veh) or R5020 (10nM; 72 hours) graphed. Inset, PR expression in MOE cells with T47D CO cells as positive control. ERK1/2 shown as loading control. D, Collagen invasion assay of hFTE spheroid cultures generated under vehicle (veh) or R5020 (10nM; 72 hours) treatment, then embedded into collagen for 72 hours without treatments. Bright-field images after 0 and 3 days of invasion. Inverse Day 3 images below. Normalized invasion area was calculated as described in methods and represented in graph. Graphs represent mean ± SD, ***P < 0.001 (n = 3).
Figure 4. Gene expression profiling of hFTE highlights unique regulation. A, Venn diagrams representing common, isoform-specific and intersecting gene sets in the absence of progestins (LEFT: vehicle; unliganded) or presence of progestins (RIGHT: R5020; liganded) at 6 and 48 hours. BLUE circles depict PR-A gene sets and YELLOW circles depict PR-B. Upregulated sets are on TOP; downregulated on BOTTOM. Cutoff for DEGs was ≥2 fold. B, Osteoprotegerin (TNFRSF11B), LGR5, FOXO4 and secretory leucocyte peptidase inhibitor (SLPI) mRNA expression in hFTE pools, vehicle (veh) or R5020 (10nM) for 6 or 48 hours treatments. Graphs represent mean ± SD, *** P < 0.001 (n = 3). C, Enriched pathways analyses of 6-hour and 48-hour vehicle-treated (unliganded) PR-A+ hFTE (PR-Aveh vs EVveh; table) and DREAM and Cell Cycle volcano plots for PR-A+ hFTE (unliganded; veh, 6
while PR-A mediates significant ligand-independent actions. The major contrast between hFTE and breast or ovarian cancer models lies in the direction and magnitude of gene regulation, with unliganded PR-A strongly activating a significant number (>400 at both time points) of genes and liganded PR-B strongly repressing a large subset (~572 at 48 hours) of genes in PR+ hFTE models (Fig. 4A and Supplemental Fig. 5 (32)); PR-B primarily activates genes in the presence of ligand while PR-A is dominantly repressive regardless of ligand and ovarian cancer models (9).

The progestin and PR-dependent regulation of known PR target genes was independently validated by RT-qPCR (Fig. 4B). Importantly, both common and isoform-specific genes are progestin-regulated, as illustrated in Fig. 4B and Supplemental Fig. 6A (32). Downregulation of the soluble decoy receptor of RANK, osteoprotegerin (TNFRSF11B gene; OPG), was observed for both isoforms, in contrast to PR-B-specific downregulation of the canonical Wnt receptor, LGFR5, and greater upregulation of the forkhead transcription factor, FOXO4, as compared to PR-A. PR-A isoform-specific gene regulation was also observed, including such genes as the secretory leucocyte peptidase inhibitor, SLPI, which is known to be highly expressed in normal cervix and fallopian tube (59). These observations suggest that exogenously expressed PRs in hFTE cells appropriately model endogenous PR actions at known PR target genes (60-62).

Regulation of both common and isoform-specific PR target genes in hFTE cells suggests important shared functions as well as significant functional differences between PR isoforms. Pathway enrichment analysis was performed to determine what specific biological processes were significantly regulated by either PR-A or PR-B. Interestingly, cell cycle progression pathways were greatly enriched in PR+ hFTE cells (Fig. 4C, D; Supplemental Fig. 6B (32)). A top pathway observed in unliganded PR-A+ hFTE, as well as liganded PR-B+ hFTE, was the DREAM pathway, encompassing the target genes regulated by the DREAM complex which modulates G0 cell cycle arrest. Notably, PR-A and PR-B isoforms exhibited opposing effects: in the absence of progestin, DREAM target genes, including cell cycle pathways, were activated in PR-A+ hFTE cells, consistent with permissive cell cycle progression (see Fig. 2D). However, these pathways were potently deactivated in progestin-treated PR-B+ hFTE (and to a lesser extent in progestin-treated PR-A+ hFTE), indicative of cell cycle blockade, perhaps via DREAM activation. Activated DREAM complexes collectively target and repress ~900+ genes which include critical G1/S and G2/M cell cycle genes, p53 target genes, as well as many other cell cycle dependent genes (63). Volcano plots of Fischer DREAM and Cell Cycle gene sets illustrate the enrichment of such target genes, along with their isoform-specific (ie, opposing directions) and shared hormone-dependent regulation (Fig. 4C and 4D). In addition, a heatmap of differentially expressed genes critical for DREAM function reveals that PR expression (PR-A) and progestin treatment (PR-B) can regulate genes encoding the protein components of the DREAM complex itself (RBBP4, E2F4/5, LIN9/37/52/54, RBL1/2), genes encoding the kinases that modulate the formation of this complex (DYRK1A/B), along with genes for the components of those complexes that promote the re-entry into the cell cycle (FOXM1, MYBL2; Fig. 4F; schematic of cell cycle complexes Fig. 4E). Of note, is the strong upregulation of DREAM target genes such as FOXM1 and MYBL2 and other proteins of the MuvB complex in unliganded PR-A+ cells—such regulation would be expected to enhance the progression through G1/S/G2 phases of the cell cycle (left heatmap; PR-A vehicle vs EV control). In contrast, many of these same genes are potently repressed in liganded PR-B hFTE with R5020 treatment, presumably promoting cell cycle arrest (right heatmap; PR-B R50 vs PR-B vehicle).

In addition to cell cycle progression pathways, other pathways that may be relevant to our observed cellular phenotypes were enriched in our PR+ hFTE models including cell-cell/cell-matrix adhesion, cell morphogenesis, Wnt signaling, oxidation-reduction activity, and stem cell differentiation (Supplemental Fig. 6B (32)). For example, PR-A+ enriched pathways included DNA replication in the absence of progestin and stem cell differentiation in the presence of progestin. This is consistent with our finding that PR-A+ hFTE cells have an increased rate of growth relative to EV cells (Fig. 2). PR-B+ enriched pathways included numerous pathways associated with cell shape, regulation of

Figure 4: continued
Figure 5. Progestins promote quiescence in hFTE cells through modulation of the DREAM complex. A, Immunoprecipitated proteins (IP) of LIN37 in hFTE following 24 hours and B, 72 hours of treatment with vehicle (-) or R5020 (+; 10nM). Left blot = input cell lysates; right blot = IP from these lysates plus IgG control. Lanes L to R: hFTE EV vehicle (-) or R5020 (+); hFTE PR-A (-)/(+); hFTE PR-B (-)/(+). GAPDH shown as loading control. Expression of LIN9 and LIN37 as core MuvB complex proteins, p130 as DREAM-specific protein, and BMYB as G1/S complex protein was determined. DREAM complexes (G0 state) were quantified using the band density ratio of p130/LIN37. B-MYB/MMB complexes (G1/S progression) were quantified using...
cell projections and adhesions in the absence of ligand and amino acid metabolism and cytokine signaling in the presence of progestin. Again, these findings are consistent with our finding that PR-B+ hFTE cells are more invasive relative to PR-A+ cells (Fig. 3D). Interestingly, consistent with strong enrichment of cell cycle pathways, both isoforms also regulate p53- and BRCA2-associated pathways in the absence (PR-A) or presence (PR-B) of progestin.

**Progestins Modulate DREAM Complex Formation and Recruitment**

RNAseq analyses indicated that the target genes of the DREAM complex were robustly enriched in PR+ hFTE models. During arrest in G0, the expression of these cell cycle-dependent genes is actively repressed by the DREAM complex, which consists of the DREAM-specific proteins, Dimerization partner DP1/2, Rb-like p130/p107, E2F4/5 and the MuvB core complex (LIN9, LIN37, LIN52, LIN54, RBBP4 proteins (27)). During the transition from G0 into late G1/early S, this transcriptional repression is lifted by dissociation of DP1/2, p130 and E2F4/5 from the MuvB core. This is followed by association of the MuvB core with B-MYB, together known as the B-MYB/MBB complex, in G1/S phase. In order to determine if progestins induce cell cycle arrest/quiescence (see Fig. 2) via regulation of the formation of the DREAM and B-MYB/MBB complexes, EV and PR+ hFTE cells were treated with R5020 and complexes were immunoprecipitated using a LIN37 antibody. The relative formation of DREAM (p130/LIN37 ratio) and B-MYB/MBB complexes (B-MYB/LIN37 ratio) was quantified by blotting for p130, B-MYB, LIN9, and LIN37. Progestins (24-hour) enhanced the formation of DREAM complexes and decreased B-MYB/MBB complexes in PR-A+ hFTE, while progestin-treated PR-B+ cells followed a similar trend (Fig. 5A). At 72 hours, progestin treatment resulted in further enhanced DREAM complex formation, with a total loss of B-MYB/MBB complexes (Fig. 5B). In addition, at this time point, the amount of B-MYB protein was dramatically reduced in the input pellets, along with LIN9, supporting the RNAseq results, which revealed that activation of PR signaling modulates expression of DREAM and B-MYB/MBB complex subunits (see Fig. 4E and 4F). In contrast, at 24 hours, the amount of input B-MYB protein was unchanged in vehicle and R5020 treated cells yet B-MYB/MBB complex formation was reduced with progestins, supporting the actions of activated PR isoforms to influence complex formation.

When these complexes were analyzed in PR+ hFTE that were treated with serum for 24 hours to initiate the reversal of the progestin-induced G0 arrest (see schematic in Fig. 2H), the slower growing isofrom PR-B+ hFTE shows a further increase in DREAM complexes (Supplemental Fig. 7A (32)). A basal level of DREAM complexes was observed in the unliganded PR-A+ and PR-B+ hFTE lysates. This could reflect the ability of PR, in the absence of progestins, to modulate DREAM complex interactions with its own subunits and/or other proteins, potentially inhibiting DREAM-mediated repression.

RT-qPCR analyses of R5020-treated PR-A+ and PR-B+ hFTE cells confirmed that MYBL2 mRNA, the gene encoding the B-MYB protein, and LIN9 mRNA were downregulated as well as FOXM1 mRNA, which encodes the transcription factor that associates with MuvB during the transition to G2/M phase (Supplemental Fig. 7B (32)). In addition, DYRK1A mRNA, which encodes a kinase that regulates DREAM complex assembly, was upregulated in PR-B+ hFTE. ChIP analyses confirmed that, following progestin treatment for 3 hours, PRs are recruited to progesterone receptor binding sites (PRE) in the promoters of LIN9, MYBL2, and DYRK1A (Fig. 5C). No change in relative mRNA levels or PR recruitment was observed for these genes in the EV hFTE cells (Fig. 5C).

Interestingly, additional ChIP analyses revealed that PR is co-recruited along with DREAM complex proteins to target genes. A PRE site in both the DREAM target genes, MYBL2 and BIRC5, overlaps with an E2F binding site and, as expected, PR and E2F4 are recruited to the same region following a 3-hour R5020 treatment, with no change observed in p130 (Fig. 5D schematic—site #1; data—Supplemental Fig. 7C (32)). A secondary site within the promoter of these genes contains consensus E2F binding elements and is known to recruit only E2F4, with p130 acting as a cofactor. This unexpected recruitment could also regulate p53- and BRCA2-associated pathways in the presence or absence of progestins.

**Figure 5: continued**

the band density ratio of BMYB/LIN37. See Fig. 4E for schematic of complexes. C, Chromatin immunoprecipitation (ChIP) assays of PR recruitment to progesterone response elements (PRE) present in LIN9, MYBL2 and DYRK1A genes after vehicle (veh) or R5020 (10nM; 3-hour) treatment. (D-E) ChIP assays of PR, p130, and E2F4 recruitment to shared and unique sites within DREAM target genes. D, Schematic of the MYBL2 and BIRC5 genes with site #1 containing overlapping response elements (RE) for PR and E2F4 and site #2 containing RE for E2F4 only with p130 acting as a known cofactor. Approximate distance between sites and the promoter of each gene is indicated. E, ChIP assay of site #2 following treatment with vehicle (veh) or R5020 (both D & E: 10nM; 3 hours). F, Progestins promote quiescence in PR+ hFTE by increasing DREAM complex formation and increasing recruitment to DREAM complex target genes, resulting in a repression of cell cycle gene transcription. Graphs represent mean ± SD, *P < 0.05, **P < 0.01, ***P < 0.001 (n = 3).
asterisk). Yet, our data demonstrated that all 3 proteins show enhanced recruitment to this site with R5020 treatment (Fig. 5E). No significant change in recruitment was observed in the EV hFTE cells. In addition, analyses of 72-hour DREAM immunoprecipitations indicates that PR protein is present and progestin treatment enhanced PR association, in particular, for the PR-B isoform (Supplemental Fig. 7D (32)).

Taken together, these results suggest that progestins promote G0 arrest through modulation of and interaction with the DREAM complex assembly (Fig. 5F). Activation of PR by the progestin, R5020, leads to increased DREAM complex formation and increased recruitment of the repressive DREAM complex, including PR, to the regulatory elements of the DREAM target genes. PR recruitment to these genes also results in the regulation of the expression of critical DREAM and B-MYB/MMB complex proteins, which, in concert with enhanced DREAM complex formation, leads to potent repression of cell cycle gene sets and the observed block in cell cycle progression.

Manipulation of DREAM Function and DYRKs Attenuates the Effects of PR Signaling in hFTE Cells

To validate the above findings in an independent FTE model and further explore the connection between the progestins’ effects on FTE and DREAM complex modulation, we created an additional model of PR-expressing FTE using the human fallopian tube epithelial cell line UWFT.1681, originally immortalized with the HPV E6/E7 proteins (Fig. 6A). The HPV E7 oncprotein is known to interfere with DREAM function by promoting the degradation of the p130 protein (64, 65). Western analysis of DREAM and B-MYB/MMB complex proteins in either whole cell lysates (input) or Lin37 immunoprecipitates verified that these lines express very low p130 protein as compared to the hFTE model (Supplemental Fig. 8A (32); red arrowheads, left hFTE vs right UWFT). Given the loss of p130 in this model, we anticipated that progestins’ actions should be attenuated or abolished in these cells, if dependent on DREAM function. Treatment of EV, PR-A+, and PR-B+ UWFT.1681 cells with R5020 (10 nM, 24 hours) failed to halt cell cycle progression in these E6+/E7+ models. In addition, progestins enhanced collagen invasion for both PR-A+ and PR-B+ UWFT.1681 cells (Fig. 6E). These data indicate that PR exerts similar effects on cell aggregation/spheroid formation and collagen invasion in multiple independently derived hFTE cell models. Remarkably, loss of p130 (ie, via HPV infection/expression of E6/E7) abrogates PR-driven cell cycle arrest but allows the expression of other progesterone/PR-driven ovarian cancer cell phenotypes.

Alterations in the expression of genes encoding members of the Myb-MuvB complex are associated with prognostic markers of aggressiveness in some cancers (27). In addition, such dysregulation of the DREAM complex may be tied to disease recurrence due to its role in maintaining quiescence. As such, the value of targeting transcriptional complexes that regulate cell cycle exit, such as the DREAM complex, is being actively explored (66, 67). Dual-specificity tyrosine phosphorylation-regulated kinases (DYRKs) are enzymes which are activated by autophosphorylation and go on to phosphorylate serine and threonine residues of target proteins (68). Like p130, the class I DYRKs, DYRK1A, and DYRK1B, are negative regulators of the cell cycle. DYRK1A, in particular, is known to phosphorylate LIN52 (a MuvB subunit protein) allowing for the association of p130 and MuvB and subsequent DREAM complex assembly (69). The actions of DYRK1B also support the quiescent state by phosphorylating LIN52 (69, 70). As a pharmacologic means to block DREAM and determine if DYRK1 kinases are required mediators of progestin-induced quiescence, we tested the potential chemotherapeutic drug, harmine, in the PR+ hFTE lines. Harmine is a naturally occurring β-carboline alkaloid and inhibitor of class I DYRK kinase activity (71) that interferes with DREAM assembly and entrance into G0 (72). As expected, harmine treatment inhibited the accumulation in G0/G1 observed in both PR-A+ and PR-B+ hFTE following progestin treatment (Fig. 7A). This treatment also resulted in a greater percentage of cells in G2/M. The dependence of PR transcriptional signaling on DYRK1 activity is also supported by harmine’s ability to lift the R5020-dependent transcriptional repression of
Figure 6. Progestin-induced effects are altered in a cell model deficient in DREAM complex formation. 

A, PR expression in empty vector (EV), PR-A+, and PR-B+ UWFT.1681 pools. GAPDH shown as loading control.

B, Cell cycle analyses (FACS) of UWFT.1681 cells treated with vehicle (veh) or R5020 (10nM; 24 hours).

C, Transcriptional regulation of MYBL2, LIN9, and FOXM1 mRNA in PR+ UWFT.1681 with vehicle (veh) or R5020 (10nM) for 72 hours.

D, Average diameter of secondary reaggregated spheroid cultures of UWFT.1681 cells treated as in C.

E, Collagen invasion assay of UWFT.1681 spheroid cultures generated under vehicle (veh) or R5020 (10nM; 72 hours) treatment, then embedded into collagen for 72 hours without treatments. Representative bright-field images after 0 and 3 days of invasion and normalized invasion area shown. Graphs represent the mean ± SD, *P < 0.05, **P < 0.01, ***P < 0.001, ****P < 0.0001 (n = 3).
Figure 7. DYRK1 inhibition interferes with PR-driven cell phenotypes in hFTE models. A, Cell cycle analyses (FACS) of hFTE, vehicle (veh), R5020 (10nM) or R5020+harmine (R/H) for 24 hours. B, Gene regulation in absence/presence of harmine: Left: regulation of MYBL2 mRNA vehicle (veh), harmine (10µM), R5020 (10nM) or R5020+harmine (R/H) for 72 hours. Right 3 graphs: ChIP assays of PR, p130 and E2F4 recruitment to the E2F4 binding site within the MYBL2 promoter (site #2), following 3-hour treatment with vehicle, harmine (30 minutes pretreatment; 10µM), R5020 (10nM)
the MYBL2 gene by interfering with the recruitment of DREAM complex members (E2F4, p130) and PR-B to these genes (Fig. 7B).

Additionally, harmine treatment included during primary spheroid formation led to a slight decrease in spheroid diameter (Supplemental Fig. 9A (32)) whereas reaggregated spheroid cultures resulted in no viable spheroids observed in harmine alone for both isoforms or for combined harmine and R5020 treatment in the PR+ hFTE (Supplemental Fig. 9B (32)). A significant reduction in spheroid diameter was also observed in harmine-treated PR-B+ hFTE. Interestingly, spheroid formation in the presence of R5020 followed by encasement in collagen I matrix for 72 hours, in the absence (vehicle) or presence of harmine, resulted in an almost complete blockage of collagen invasion, especially for highly invasive PR-B+ cells (Fig. 7C).

To implicate a specific DYRK1 isoform(s) in PR regulation of DREAM, knockdown of either DYRK1A or DYRK1B was performed in PR+ hFTE and subsequent analyses of PR-induced gene regulation and cancer-associated cell phenotypes were repeated. The mRNA and protein expression of each kinase (DYRK1A or DYRK1B) was depleted singly in each cell line (Fig. 7D; Supplemental Fig. 10A (32)). As with harmine, DYRK knockdown altered transcriptional regulation of DREAM-relevant genes. A strong attenuation of the basal (vehicle) expression of MYBL2, LIN9, and FOXM1 was observed in both PR-A+ and PR-B+ hFTE in the DYRK1A-depleted state (Fig. 7E). In the presence of progestin (R5020 treated), we previously observed that both PR isoforms strongly repress the expression of these genes (see Supplemental Fig. 7B (32)) as observed in the nontargeting control (NT). This may indicate that progestin regulation of these genes is lost or nonresponsive due to the lower basal levels of these genes in the DYRK1A-depleted state. DYRK1B-depleted PR-A+ hFTE treated with R5020 showed an attenuation of this repression, exhibiting greater relative expression levels as compared with control NT cells treated with R5020 (Fig. 7E). Similarly treated, DYRK1B-depleted PR-B+ hFTE showed no significant change. More importantly, depletion of DYRK1A or DYRK1B also attenuated the PR-induced enhancement of cell aggregation and reaggregation in both PR-A+ and PR-B+ hFTE (Supplemental Fig. 10B and 10C (32)) with reaggregation showing the most robust effect. Interestingly, for some of the biological readouts, the effect of silencing each kinase was highly PR isoform-specific. In collagen invasion assays, for example, depletion of DYRK1A resulted in an almost total block of PR-A+ cell invasion whereas DYRK1B shRNA had no effect (Fig. 7F). In contrast, DYRK1B depletion reduced PR+B+ cell invasion; DYRK1A shRNA had no effect (Fig. 7F). These data support a novel role for DYRK1 isoforms as mediators of both PR-A (DYRK1A) and PR-B (DYRK1B) transcriptional responses and cancer cell-associated behaviors.

Taken together, the data presented here suggest a model wherein PR signaling promotes changes in hFTE cell fate that enable early stages of HGSC progression (Fig. 7G). In the absence of progestin, unliganded PR-A inhibits DREAM/DYRK1 actions, supporting permissive proliferation. In the presence of progestins, PRs mediate the transcriptional regulation of DREAM complex proteins and promote formation of active (ie, repressive) DREAM complexes that require p130 and DYRK1. Thus, liganded PRs support the DREAM/DYRK1-mediated repression of cell cycle-dependent genes and the subsequent arrest in G0. This quiescent state, along with other direct actions of PRs (ie, in part mediated via PR isoform specific use of DYRKs) may promote cell survival by supporting cell aggregation and spheroid formation following shedding from the fallopian tube. Once PR+ tumor emboli are circulating within the abdominal cavity, a hormone-rich microenvironment may promote their subsequent invasion of the mesothelial layer lining the peritoneal cavity. Further imbalance of DREAM or PR isoforms (ie, loss of p130 or PR) may permit cell cycle re-entry at distant sites.

**Discussion**

The actions of ovarian steroid hormones such as progesterone are known to be highly contextual and complex; they are dependent on the tissue, the specific cell type, as well as the local hormonal milieu. Within breast tissue, progesterone can be tumor-promoting or protective; its effects greatly influenced by the hormone concentration, the duration of exposure, presence of other steroids and signaling molecules, and the age of the woman (15, 73). In reproductive tract tissues such as the uterus, progesterone has traditionally been viewed as a protective factor due to its...
antagonistic actions on estrogen-induced endometrial proliferation (15). The work presented here shows another facet of progesterone’s actions in the reproductive tract—this hormone, in the context of mutations within early fallopian tube lesions, may drive reversible cell cycle arrest and associated cell behaviors that could contribute to OC progression.

The functions of PR are intimately connected to cell fate and stemness. PR senses inputs from sex hormone and growth factor–initiated signals, integrating these with activation of multiple kinase pathways that ultimately modulate cell cycle progression (16, 74, 75). From studies conducted in breast, uterine, and ovarian carcinoma models, PR is known to be tightly coupled to cell cycle mediators via cell cycle dependent PR phosphorylation (76), direct PR-cyclin (D1, A, and E1), or PR-cyclin-dependent kinase (CDK2) interactions along with transcriptional regulation of critical cell cycle genes (76-78). The connection between PR actions and DREAM complex function revealed by our studies is a novel mechanism by which progesterone can manipulate cell fate—unique in the context of fallopian tube epithelia and previously unreported as a downstream effect of PR signaling. Studies in breast carcinoma lines have shown that the estrogen antagonist, ICI 182780, can induce an increase in p130 (RBL2) protein and p130/E2F4 complex accumulation (79). Progesterone was able to reduce phosphorylation of p107 (RBL1), inhibiting estrogen-induced proliferation in uterine epithelia (80) and synthetic progestins alone showed similar effect on p107 phosphorylation in T47D breast cancer cells (81). None of these studies connected the actions of either estrogen or progesterone or their receptors to direct modulation of DREAM complexes, at both the level of protein complex formation and transcriptional regulation of DREAM complex components as we have shown.

Multiple experimental approaches were utilized in our studies to interfere with DREAM complex, including E6/E7 immortalized cell line (ie, in which p130 is naturally depleted via degradation), chemotherapeutic blockade (harmine) of DYRK1 kinase function, and depletion (shRNA knockdown) of DYRK1A/1B kinase expression. These approaches consistently prevented the progestin-induced cell cycle arrest and totally abolished or attenuated the transcriptional regulation of genes encoding DREAM complex proteins and/or cell cycle dependent proteins known as DREAM targets. This suggests that progestins induce G0 cell cycle arrest, in part, through the modulation of DREAM function. The cell behaviors of cell-cell aggregation and collagen invasion exhibited more variable attenuation/abolishment upon DREAM interference, implying that the G0 state mediated by DREAM/DYRK1 kinases contributes to, but is not the sole regulator of, these PR/progestin-mediated phenotypes (ie, inhibition of proliferation was separable from other cancer-associated phenotypes in the E6/E7 immortalized cells). Therefore, direct effects of PR expression and progestins also drive these behaviors (cell aggregation/invasion) even when cell growth inhibition is disabled by loss of p130.

Progestins are known to regulate cell migration and invasion in advanced cancer models. In multiple breast cancer lines, PR signaling enhances migration and invasion through stabilization of the RhoA complex, modulation of focal adhesions, and transcriptional regulation of key genes (82-84). Progesterone, allopregnanolone, and mifepristone (RU486) have been shown to increase migration in ovarian carcinoma lines (85, 86). Pathway analyses of our PR+ hFTE showed that PR expression modulated genes associated with cell morphogenesis, cell-cell and cell-matrix adhesion, as well as Rho signaling (Supplemental Fig. 6B (32)). Studies have observed that p53 mutations can support enhanced cell adhesion and mesothelial invasion in immortalized “normal” hFTE cell lines (48). Though interplay between PR activation and wild-type p53 expression and their transcriptional effects has been reported (87, 88), an understanding of the mechanisms of this interaction and subsequent modulation of metastatic phenotypes is limited. It is interesting to note that recent work has revealed several p53 mutant species, including R175H, can disrupt the progesterone-activated PR-A/p53 complexes that regulate p27 expression, but no downstream phenotypic effects were explored (87). It remains to be proven if mutant p53 species in FTE can synergize with progestins to support these cell behaviors. Further research will be needed to determine how additional commonly observed genetic alterations in STICs, such as Cyclin E1 (CCNE1) amplification (47, 89), BRCA1/2 mutations (12), or viral (HPV) infection (90, 91) could further modify PR signaling. Notably, both cyclins/CDKs (76, 92) and BRCA1/2 (93) are known PR-binding proteins.

The ratio of PR isoforms in a cell, the presence of potential ligands, and posttranslational receptor modifications (ie, phosphorylation) all contribute to the downstream cellular outcomes of PR signaling (15-17). In our hFTE models, PR-A+ cells without progestins (ie, ligand-independent) were more proliferative and migratory, whereas PR-B+ cells with progestins (ie, ligand-dependent) were more invasive. Such distinct effects have been observed in other advanced cancer models and could modulate PR signaling in normal FTE as well as during the progression to early neoplasms. For example, PR-A actions drive stemness in breast carcinoma models while PR-B primarily promotes proliferation; opposite isoform effects compared with our hFTE models (94, 95). In addition, recent mouse models of...
constitutive PR isoform overexpression have revealed that PR-B was the stronger driver of proliferation in the development and progression of ovarian neoplasms originating from ovarian luteal cells (78). In normal human and mouse fallopian tube, these isoforms appear to be equally expressed, but the question of whether changes (ie, imbalance) in isoform expression occur in early STIC lesions is currently unknown. IHC studies of PR isoform staining across advanced OC tumor subtypes would suggest a loss of PR-A protein expression during disease progression, opposite to the PR-A dominance (ie, loss of PR-B protein) observed in breast cancer (96-98). While PR isoform imbalance is a hallmark of hormone-driven cancers (15), it is possible that loss of PR isomorph IHC epitopes, as commonly measured using clinical monoclonal antibodies, represents the presence of highly modified/activated receptors (18, 46). In addition, clinical-grade PR monoclonal antibodies may exhibit unequal detection of PR isoforms (99, 100). Therefore, the variability of PR protein expression previously reported in STICs (101) and invasive HGSC (102, 103) using such antibodies could be an artifact of the limitations of monoclonal antibodies and is most likely missing an important component of PR signaling, wherein p-PR is a biomarker of activated PR. The opposing actions of unliganded PR-A relative to liganded PR-B are likely highly context-dependent. Our data support a model whereby in FTE exposed to low/no progesterone (ie, postmenopausal or hormone-ablated contexts), PR-A may dominantly repress DREAM (PR-A+ hFTE cells proliferate freely), while in the presence of abundant progesterone, both PRs, but especially PR-B, may dominantly activate DREAM (PR+ hFTE cells exit the cell cycle). This relationship between PR isoforms perhaps ensures a decisive or “switch-like” and robust response to hormonal cycles. Notably, while STICs give rise to invasive serous OC, they appear to preexist for years to decades as relatively dormant (STIL) lesions. In light of our IHC analyses, activated (ie, phosphorylated) PRs may drive the enhancement of DREAM and thus maintain cellular quiescence in early lesions (ie, when progesterone is present). The decline of ovarian progesterone during peri/postmenopausal transitions, in addition to the accumulation of genetic alterations (discussed above), may “release” PR-mediated cell cycle blocks and instead enable proliferation, cell-cell aggregation, and invasion in PR+ cells in STILs thereby facilitating transition to STICs. Therefore, future IHC studies of phospho-PR species and quiescence markers in STILs/STICs and associated invasive lesions from a larger, more diverse cohort of both pre- and postmenopausal patients could help differentiate between the PR isoforms present and their signaling potential, clarifying the role of changing PR signaling as well as altered expression of DREAM components (ie, a release of PR-dependent repression) during HGSC progression.

The distinct isoform-specific effects of DYRK1A or DYRK1B kinase depletion in the unliganded and liganded PR-A vs PR-B hFTE were unexpected. For example, we observed that collagen invasion was abolished in PR-A+ hFTE only when DYRK1A was depleted whereas there was no effect of DYRK1B depletion. Interestingly, both PR-A+ and PR-B+ hFTE exhibited unique attenuation of basal gene expression in an unliganded state (vehicle) with DYRK1A depletion, potentially revealing a more prominent role of this kinase in unliganded PRs regulation of cell cycle dependent transcription. In contrast, DYRK1B depletion exposed a potential role of DYRK1B in liganded PR regulation, especially for PR-A, since loss of this kinase partially reversed the strong R5020-mediated repression normally seen in unaltered hFTE cells. No published studies have examined these kinases in the context of FTE and/or PR signaling. Both kinases can promote G0 arrest through DREAM and other mechanisms, and inhibition or depletion of either can induce cell cycle re-entry in normal and cancer cells (66, 69). But recent studies are expanding the function of these kinases, in particular, their role in DNA damage repair. DYRK1A is proposed to be involved in DNA double-stranded break repair by modulating 53BP1 recruitment, thereby supporting a shift to error-prone nonhomologous end joining repair pathways (49). This is intriguing, since preliminary studies in our laboratory suggest that the progestins enhance DNA damage (unpublished results). Therefore, it will be interesting to further explore the regulation and activity of DYRK1 kinases and the potential interplay between PR signaling and kinase actions in early lesions.

The evolution of HGSC makes a PR-driven quiescent state, and its potential to promote pro-survival/pro-dissemination phenotypes, highly relevant to our understanding of disease etiology. The presence of total and nuclear “focal” and phosphorylated PRs observed in STICs and invasive tumors revealed in our studies suggests that active PR signaling in these lesions could be promoting cell dormancy. Such promotion of dormancy by these activated PR could be perceived as protective—for example, minimizing the effects of cellular stressors that bathe the distal fimbriae of the fallopian tube (104). However, research in cancer, aging, and stem cell dynamics has revealed that quiescence comes at a cost. Slow-cycling/quiescent cells must rely on nonhomologous end joining, an error-prone mechanism for DNA damage repair that can lead to misreplicated double-stranded breaks, mutagenesis, and genomic instability (105). In addition, dormant cancer cells often display more aggressive migratory and invasive behaviors.
and exhibit chemoresistance to therapeutics that target proliferating cells (29). Blocking DREAM formation and quiescence in advanced OC models, through depletion of DYRK1A, DYRK1B, or p130/RB2, leads to lack of spheroid formation, reduced viability, increased apoptosis, and sensitization to platin therapeutics (30, 72).

A quiescent cell can be stimulated to re-enter the cell cycle and, depending on the “health” of that cell (eg, DNA damage, levels of reactive oxygen species), go on to proliferate, enter senescence (which potentially can be reversible) or undergo regulated cell death (29). Previous work in our laboratory has revealed progesterin-mediated, PR-A- and PR-B-dependent cellular senescence in advanced OC cell models (9). Other researchers have suggested that progesterone can cause apoptosis (108) or necroptosis (109); these studies often are complicated by high hormone treatment levels and lack of nuclear receptor expression. Taken together, it is probable that during OC progression, the effects of PR and progestins will be nuanced, depending on signaling inputs resulting from genetic alterations and microenvironment changes. Such shifts in PR actions have been observed in breast cancer, where progesterone will drive migration in early lesions, prior to a detectable tumor, yet promote proliferation during metastasis (110). In addition, the ligand-independent effects that our studies have revealed suggests that although certain physiological states exhibit low or no progesterone (ie, postmenopausal), this should not negate the capacity of these receptors to drive cellular outcomes in early and advanced stages of HGSC. The studies presented here provide evidence for a novel mechanism of PR signaling in healthy fallopian tube epithelia as well as those in a compromised state that could lead to disease initiation and progression. For example, in HPV-infected FTE, the cell cycle would be predicted to be “released” from progesterone blockade, while other PR-driven cancer phenotypes (cell aggregation/invasion) are robustly promoted.

Our findings represent an alternative perspective on the idea that progesterone always confers protection against ovarian cancers. In our studies, PRs support a quiescent FTE cell fate through modulation of DREAM/DYRK1 function and such a state could be protective. But, over an extended period, quiescence can also be associated with many of the hallmarks of cancer such as the acquisition of new mutations, the survival in a suboptimal environment, metastasis, and chemoresistance (28, 29). In addition, the potential for coincident dysregulation of DREAM components in FTE, an event often observed in cancers (27, 31), along with the emerging role of DYRK1 kinases in DNA damage repair (111), the cellular outcomes of PR signaling will be nuanced and highly contextual, like other sex steroids. This idea that PR signaling could promote cellular states that enhance ovarian cancer risk is supported by recently published research showing progesterone-driven ovarian tumor progression in transgenic mouse models (78). Taken together, we might consider an expanded view of another role of progesterone/PR as contributing factors in the development of STICs that progress to HGSC. Considering that occult STICs can reside undetected within FT for decades prior to established invasive disease, there are now significant clinical opportunities to prevent the ultimate development of HGSC by targeting PRs, DREAM, and/or DYRKs (ie, as with harmine). Since existing antiprogestins (onapristone, RU486) behave as PR agonists in hFTE (Supplemental Fig. 4 (32)), we suggest that trials of clinical interventions directly targeting PRs as a means to eliminate STICs in high-risk women proceed with caution until more effective (ie, pure antagonists or PR degraders) are developed.

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Data Availability: Some or all data generated or analyzed during this study are included in this published article or in the data repositories listed in the References. RNAseq data collected in these studies has been deposited in NCBI Gene Expression Omnibus (GEO) database (41). Supplemental figures are available at Figshare digital repository (32).
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