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#### **Priority Research Paper**

# Assessment of variation in immunosuppressive pathway genes reveals TGFBR2 to be associated with risk of clear cell ovarian cancer

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Keywords: ovarian cancer, immunosuppression, biomarkers, genetic variation, TGFBR2

Received: March 18, 2016 Accepted: May 29, 2016 Published: June 21, 2016

# ABSTRACT

Background: Regulatory T (Treg) cells, a subset of CD4+ T lymphocytes, are mediators of immunosuppression in cancer, and, thus, variants in genes encoding Treg cell immune molecules could be associated with ovarian cancer.

Methods: In a population of 15,596 epithelial ovarian cancer (EOC) cases and 23,236 controls, we measured genetic associations of 1,351 SNPs in Treg cell pathway genes with odds of ovarian cancer and tested pathway and gene-level associations, overall and by histotype, for the 25 genes, using the admixture likelihood (AML) method. The most significant single SNP associations were tested for correlation with expression levels in 44 ovarian cancer patients.

Results: The most significant global associations for all genes in the pathway were seen in endometrioid (p = 0.082) and clear cell (p = 0.083), with the most significant gene level association seen with *TGFBR2* (p = 0.001) and clear cell EOC. Gene associations with histotypes at p < 0.05 included: *IL12* (p = 0.005 and p = 0.008, serous and high-grade serous, respectively), *IL8RA* (p = 0.035, endometrioid and mucinous), *LGALS1* (p = 0.03, mucinous), *STAT5B* (p = 0.022, clear cell), *TGFBR1* (p = 0.021 endometrioid) and *TGFBR2* (p = 0.017 and p = 0.025, endometrioid and mucinous, respectively).

Conclusions: Common inherited gene variation in Treg cell pathways shows some evidence of germline genetic contribution to odds of EOC that varies by histologic subtype and may be associated with mRNA expression of immune-complex receptor in EOC patients.

## **INTRODUCTION**

Ovarian cancer is the leading cause of death due to gynecological cancers in the United States [1]. Although two-thirds of ovarian cancer patients initially respond to surgical debulking and chemotherapy [2], a majority eventually relapse [3, 4]. The five-year survival rate of ovarian cancer varies significantly across clinical stages, with almost 90% of stage I patients surviving, to just a little over 20% of advanced-stage patients surviving [5].

In recent years, host tumor immunosuppression has attracted research in ovarian cancer in hopes of identifying underlying biological mechanisms that determine the development and progression of ovarian cancer. Ovarian tumors have been found to induce migration of immunosuppressive cells into tumor tissue [6]. Thus, exploring molecular pathways underlying suppression of immune responses in ovarian cancer to identify novel targets for immunotherapy and/or to identify markers that can predict the risk of ovarian cancer may be a route to both treating this deadly disease and/or earlier identification.

An important pathway to consider in immune function is suppression of host immune response by regulatory T (Treg) cells, a subset of CD4+ T cells that maintain immune tolerance and inhibit the development of an antitumor immune response. In fact, higher prevalence of Treg cells has been found in various cancers [7-12], including ovarian cancer [13-16], compared to controls. Treg cells have been detected in ovarian tumors [15], as well as in malignant ascites [13] and peripheral blood [16] of ovarian cancer patients. Further, an association of ovarian cancer outcomes with genetic variation in Treg-related genes specific to induction, trafficking, or immunosuppressive function of Treg cells, also suggests a role for the Treg cell phenotype in ovarian cancer [17]. Given the importance of inherited factors in both ovarian cancer and Treg cells, we sought to characterize their role in ovarian cancer etiology. We conducted a comprehensive epidemiological study in which we investigated the significance of single nucleotide polymorphisms (SNPs) in the Treg cell pathway and mRNA expression profiles in epithelial ovarian cancer (EOC) etiology.

## RESULTS

The descriptive characteristics of the study population are presented in Table 1. The majority of EOC patients (n = 9,330) were of the serous histology. Compared to controls, cases were significantly older and more likely to report a family history of breast or ovarian cancer and a personal history of endometriosis. Conversely, pregnancy, tubal ligation, breastfeeding, and use of oral contraceptives (OCs) were more likely to be reported by controls.

#### Association of genetic variation by histotype

P-values for the gene burden test for each gene in the pathway and the Treg cell pathway (all SNPs analyzed together) by histotype (serous, high-grade serous, endometrioid, clear cell, invasive mucinous) are presented in Table 2. The most significant burden test (p =0.001) was seen with TGFBR2 and clear cell EOC. Other gene associations with histotypes at p < 0.05 included: *IL12B* (p = 0.005 and p = 0.008, serous and high-grade serous, respectively), IL8RA (p = 0.035, endometrioid and invasive mucinous), LGALSI (p = 0.03, invasive mucinous), STAT5B (p = 0.022, clear cell), TGFBR1 (p= 0.021, endometrioid) and TGFBR2 (p = 0.017 and p =0.025, endometrioid and invasive mucinous, respectively). The most significant global associations for all genes in the Treg cell pathway were seen in endometrioid (p =0.082) and clear cell (p = 0.083) EOC.

Single SNP associations for each gene are shown in Supplemental Table 1. The effective number of independent SNPs tested was 370; applying a bonferroni correction for testing 370 SNPs across 5 groups, yields  $p < 2.7 \times 10^{-5}$  as the significance threshold. No single SNPs remains significant after correction for multiple testing within histotype. The most single SNP association was seen with *TGFBR2* and clear cell; the T allele in rs3773636 was associated with a 21% increased risk of clear cell ovarian cancer (OR = 1.21, 95% CI = 1.10-1.33, p = 0.0001).

# eQTL in TGFBR2 associate with FCGR2B expression

TGFBR2 contained the SNP with the most significant association with risk of clear cell EOC and also contained several additional SNPs with suggestive associations with clear cell and mucinous EOC. Thus, SNPs in TGFBR2 were correlated with mRNA expression levels as measured by the 9,634 probes passing quality control (QC) and showing expression above the background in at least 25% of the samples [18]. Regression analyses showed the most significant association between rs1808602 and FCGR2B ( $P_{FDR} < .05$ ) with an adjusted  $r^2 = 0.51$  for a model including both SNP and histology; the variation attributable to the SNP alone was  $r^2 = 0.45$ . Each additional copy of the minor (G) allele (minor allele frequency (MAF) = 42.4%) was associated with an increase in mRNA expression level of 0.51 in FCGR2B (Figure 1). This SNP-gene association was the only association significant after correction for multiple testing.

### DISCUSSION

Treg cells have been shown to suppress tumor

# mRNA Expression by Genotype



**Figure 1:** Association of variant alleles in *TGFBR2* with circulating mRNA expression levels in *FCGR2B*. *FCGR2B* mRNA expression levels (y-axis) versus rs1808602 (x-axis). Each additional copy of the variant allele (G) in rs1808602 was associated with a significant increase in mRNA expression level after adjusting for age and histology.





Table 1: Descriptive characteristics	of 15,596	ovarian	cancer	cases	and	23,236	controls	from	the	Ovarian	Cancer
Association Consortium (OCAC)											

Variable	Case <i>N</i> = 15596	<b>Control</b> <i>N</i> <b>= 23236</b>	P value
Age <sup>1</sup>	57.36 (11.70)	55.61 (11.90)	< 0.0001
Ethnicity <sup>2</sup>			
Non-Hispanic	13847 (99.6)	21539 (99.7)	0.03
Hispanic	56 (.4)	59 (.3)	
Missing	1693	1638	
Family history of ovarian cancer <sup>2</sup>			
No	5891 (91.6)	7643 (95.7)	< 0.0001
Yes	543 (8.4)	343 (4.3)	
Missing	9162	15250	
Height <sup>1</sup>	1.64 (0.07)	1.63 (0.06)	< 0.0001
Missing	4571	6596	
Weight <sup>1</sup>	57.2 (9.80)	56.4 (8.71)	1
Missing	6600	9277	< 0.0001
Body Mass Index (BMI) <sup>1</sup>	21.29 (3.43)	21.18 (3.06)	0.01
Missing	6642	9311	Ì
Age at menarche <sup>1</sup>	12.8 (1.60)	12.9 (1.68)	0.02
Missing	4914	7195	1
Total number of pregnancies <sup>1</sup>	2.37(1.80)	2.63(1.74)	< 0.0001
Missing	4879	7066	
Breast feeding <sup>2</sup>			1
No	3070 (40.5)	4078 (30.2)	< 0.0001
Yes	4502 (59.5)	9426 (69.8)	
Missing	8024	9732	
Menopausal status <sup>2</sup>			
Pre/perimenopausal	3585 (32.4)	4519 (28)	< 0.0001
Post-menopausal	7491 (67.6)	11640 (72.0)	0.0001
Missing	4520	7077	
HRT <sup>2</sup>			
No	2675 (44 3)	3237 (44 6)	0.73
Yes	3366 (55 7)	4025 (55 4)	
Missing	9555	15974	
$OC use^2$			
Never	4465 (41.9)	6054 (37 4)	< 0.0001
Ever	6191 (58 1)	10152 (62.6)	0.0001
Missing	4940	7030	
OC use in months <sup>1</sup>	38 21 (59 83)	49 40( 69 27)	< 0.0001
Missing	5164	7209	0.0001
Tubal ligation <sup>2</sup>			
No	8420 (84.4)	8278 (76.7)	< 0.0001
Yes	1562 (15 7)	2514 (23 3)	
Missing	5614	12444	İ
Endometriosis <sup>2</sup>			
No	7435 (90.8)	10030 (93.2)	< 0.0001
Yes	755 (9.2)	731 (6.8)	
Missing	7406	12475	1
Hysterectomy <sup>2</sup>	,		
No	7352 (68 3)	13103 (81 2)	< 0.0001
Ves	3413 (31 7)	3025 (18.8)	-0.0001
Missing	4831	7108	1
111000115	1031	/100	1

Clinical characteristics Histology <sup>2</sup>		
Serous	9330 (59.8)	
Mucinous	1592 (10.2)	
Endometrioid	2099 (13.5)	
Clear cell	1033 (6.6)	
Mixed Cell	505 (3.2)	
Other	1037 (6.7)	
Behavior <sup>2</sup>		
LMP	1724 (11.1)	
Invasive	13872 (88.9)	
FIGO stage <sup>2</sup>		
1	3488 (31.7)	
2	1147 (10.4)	
3	5412 (49.2)	
4	954 (8.7)	
Grade <sup>2</sup>		
Well differentiated	1240 (12.5)	
Moderately differentiated	2427 (24.4)	
Poorly differentiated	5591 (56.2)	
Undifferentiated	699 (7.0)	
Missing	5639	

<sup>1</sup>Mean (standard deviation),  $^{2}N(%)$ , CI = Confidence interval, BMI = body mass index, HRT = hormone replacement therapy, OC = oral contraceptive, LMP = low malignant potential

antigen specific immunity in ovarian cancer, in vitro and in vivo [13]. However, the role of Treg cells in the etiology of ovarian cancer is not well established. We attempted to evaluate robust genetic biomarkers associated with Treg cells in relation to EOC in a large sample pooled from the Ovarian Cancer Association Consortium (OCAC). We hypothesized that SNPs in genes that regulate the function of Treg cells could potentially be associated with variation in immune response to ovarian tumors. Hence, in this study we evaluated SNPs in 25 genes thought to govern the function of Treg cells to determine their association to EOC. We found a modest association between TGFBR2 and invasive clear cell EOC. SNPs in this gene have been found to be associated with other pathological conditions, including gastric and colorectal cancer [19, 20]. The TGF-β family of cytokines plays an important role in proliferation, differentiation, and apoptosis of many cell types [21]. However, some tumors, such as ovarian tumors, evade the anti-proliferative effects of TGF-β by acquiring mutations in TGF- $\beta$  signaling pathway [22]. Furthermore, the TGF- $\beta$  signaling pathway plays a paradoxical role in tumorigenesis, initially suppressing and later promoting tumor growth and metastasis [23].

The significant association of rs1808602 in TGFBR2 with lymphoblastoid cell line (LCL) mRNA expression of FCGR2B (Fc $\gamma$ RIIB) adds evidence for an immune component in ovarian carcinogenesis. FCGR2B binds to the Fc component of the antigen-IgG immune complex, suppressing immune response

through several mechanisms, including inhibition of antigen presentation to T lymphocytes as well as reduced phagocytosis by neutrophils [24]. The only inhibitory receptor among members of the FcGR family in humans, FCGR2B, expressed on B lymphocytes [25] and follicular dendritic cells, is thought to be critical for maintenance of humoral immune response [26, 27]. The modest correlation between the TGFBR2 polymorphism and mRNA expression of FCGR2B observed suggests that TGF-β cytokine signaling pathway may, directly or indirectly through Treg cells, regulate the expression of FcGR, thereby potentially altering the balance between pro-inflammatory and anti-inflammatory immune response. Furthermore, the downstream inhibitory effect of FCGR2B expression is not limited to immune cells. Experimental models have demonstrated the potential of FCGR2B to promote tumorigenesis when expressed on non-lymphoid tumor cells [28, 29]. FCGR2B expression is thought to be a mechanism of immune escape by tumor cells [30]. Thus, our findings indicate that polymorphisms in TGFBR2 may potentially affect inter-individual variation in anti-tumor immune response through FcG receptor modulation. Additional evidence for Treg-cellrelated eQTL SNPs has been seen with survival in ovarian cancer [31, 32]. Specifically, genetic variation in CD80 was associated with poorer survival of endometrioid cases and with increased tumor CD80 expression. The above findings suggest that inherited factors contributing to ovarian cancer etiology and outcome may, in part, drive

Gene	Serous ( <i>n</i> = 9,330)	High-grade serous $(n = 5,792)$	Endometrioid $(n = 2,060)$	Clear cell $(n = 1,021)$	Invasive Mucinous (n = 933)
CTLA4	0.612	0.984	0.337	0.471	0.178
FCRL3	0.426	0.388	0.464	0.546	0.110
FOXP3	0.362	0.254	0.630	0.525	0.287
GZMB	0.484	0.203	0.220	0.931	0.847
HDAC9	0.679	0.864	0.212	0.398	0.990
IL12B	0.005	0.008	0.127	0.915	0.088
IL17RA	0.269	0.243	0.974	0.831	0.652
IL23A	0.137	0.111	0.990	0.431	0.561
IL23R	0.423	0.903	0.470	0.101	0.221
IL2RA	0.948	0.960	0.153	0.281	0.148
IL7	0.915	0.933	0.339	0.822	0.670
IL7R	0.558	0.562	0.296	0.459	0.670
IL8RA	0.118	0.084	0.035	0.344	0.035
LGALS1	0.222	0.054	0.841	0.520	0.030
LGALS9	0.958	0.949	0.649	0.885	0.081
PRKCQ	0.511	0.862	0.879	0.528	0.729
STAT5A	0.283	0.463	0.556	0.117	0.442
STAT5B	0.721	0.873	0.412	0.022	0.297
TGFB1	0.864	0.908	0.864	0.966	0.168
TGFB2	0.739	0.418	0.481	0.087	0.672
TGFB3	0.335	0.250	0.139	0.354	0.438
TGFBR1	0.378	0.398	0.021	0.504	0.493
TGFBR2	0.644	0.242	0.017	0.001	0.025
TGFBR3	0.068	0.256	0.446	0.295	0.366
TNFSF14	0.742	0.521	0.964	0.981	0.848
Treg cell gene pathway	0.444	0.719	0.082	0.083	0.632

 Table 2: Admixture maximum likelihood gene burden *p*-values for each gene in the Treg cell pathway and overall considering all genes

the expression of important immune-related genes.

Further evaluation of the structure of *TGFBR2* showed that the rs3773636 SNP is in strong linkage disequilibrium ( $r^2 = 1$ ) with a SNP (rs995435) that is thought to likely affect binding of proteins such as *HNF4A*, *EP300*, and *GATA2*, all associated with the balance of cell differentiation [33] (Figure 2). This SNP resides in *SMAD4* and *ELF5* (an ETS-related transcription factor) motifs in a relatively important position. In addition, we find that rs1463535 in *TGFBR2*, ~2 Mb from rs3773636 and independent of rs3773636, is associated (p < 8e-05) with expression of *TGFBR2* in lymphoblastoid cell lines (p < 8e-05) [34].

Although we find relatively weak associations between SNPs in the Treg cell pathway and EOC etiology, we do see modest evidence that *TGFBR2* contains an eQTL that is perhaps modulating expression of inhibitory immune-complex receptor genes. Thus, the Treg cell genetic hypothesis perhaps merits further investigation in a larger, more diverse population.

# **MATERIALS AND METHODS**

# **SNP** selection

An extensive literature review of studies examining the role of regulatory T cells in immune response was conducted in 2010, and genes relevant to the function of Treg cells were identified. Tag SNPs in 25 genes (MAF  $\geq$ 0.05),were selected using the SNP database on Genome Variation Server [35]. SNP selection parameters included an r<sup>2</sup> > = 0.8 and the Centre d'Etude du Polymorphisme Humain (CEPH) reference population. The genomic region was expanded upstream and downstream (5 Kb) of each gene using linkage disequilibrium block structure to capture tag SNPs in regulatory regions. Tag SNPs were then assessed for design scores using Illumina's Assay Design Tool for Infinium, and SNPs with a design score < 0.4 were excluded. SNPs were also excluded if the call rate was < 95%, if the test for deviation from Hardy Weinberg equilibrium proportions in controls was  $p < 10^{-4}$ , or if greater than 2% discordance in duplicate pairs was observed. Of the 1,358 SNPs from the Treg cell pathway that were included for genotyping, a total of 1,351 passed QC and were included in the analysis presented in this paper (Supplemental Table 2).

### Study population, genotyping, and quality control

Germline DNA (250 ng genomic or 750 ng wholegenome amplified) from a total of 15,596 ovarian cancer cases and 23,236 controls from 40 studies in the OCAC (Supplemental Table 3) was genotyped on a custom Illumina iSelect BeadArray. OCAC is an international, multidisciplinary consortium, comprising populationbased, hospital-based and nested case-control, and caseonly studies of ovarian cancer, conducted in the United States, Europe, Asia, and Australia. Genotype calling and quality control procedures were described previously [36, 37]. Samples with a genotype call rate of < 95% were excluded. Hap Map samples from European (CEU, N =60), African (YRI, N = 53), and Asian (JPT+CHB, N =88) populations were used to estimate intercontinental ancestry for each individual using the Local Ancestry in Admixed Population (LAMP) program [38], and variation in population substructure was estimated using principal components (PCs). Only individuals with a LAMP score greater than 90% European ancestry were included in the present analyses.

### Statistical analyses

Logistic regression analyses in PLINK were used to test for evidence of additive associations of SNPs by histotype and restricted to invasive tumor behavior [39]. Evaluation of the scree plot of eigenvectors, derived using Eigenstrat, revealed that five PCs explained most of the variation in population substructure; the logistic regression models were adjusted accordingly for PCs, along with age. PC analysis was done using an in-house program written in C++ using the Intel MKL libraries for eigenvectors (available at http://ccge.medschl.cam. ac.uk/software/) [40]. We used the approach of Li et al. to calculate the effective number of independent SNPs tested, and this value was then used in a Bonferroni correction to determine single SNP significance [41, 42]. Regional association plots for SNPs with significant associations were constructed using LocusZoom software [43].

Both gene-level tests of association and global Treg cell pathway analyses by ovarian cancer histotypes were conducted using the admixture likelihood (AML) method [40, 44]. The AML method assumes a proportion of variants in each gene or pathway ( $\alpha$ ) is associated with outcome. The effect size of each SNP is assumed to be on a non-central  $\chi^2$  distribution with non-centrality parameter n, which approximately captures that SNP's contribution to the total genetic variance of the outcome. To accommodate the correlation between SNPs in each gene, AML uses a pseudo-maximum likelihood method to estimate the  $\alpha$  and  $\eta$ . For each gene-level and pathwaylevel test, we performed 1,000 simulations, assuming that the maximum proportion of associated SNPs in each gene or pathway was 0.20. We report p-values for the AML trend test.

# Expression quantitative trait loci (eQTL) analysis in ovarian cancer patients

We measured *trans* and *cis* genotype associations with mRNA expression levels in LCL collected pretreatment from unrelated EOC cases enrolled in the Gilda Radner Ovarian Family Cancer Registry (GRR) at Roswell Park Cancer Institute (RPCI), a part of the larger OCAC study described above. Microarraybased gene expression was assayed using the Illumina HumanHT-12v3 Gene Expression Beadchip, with almost 50,000 probes derived from the National Center for Biotechnology Information Reference Sequence (NCBI) RefSeq (Build 36.2, Rel 22) and the UniGene (Build 199) databases [45]. Beadscan was used to scan and extract the raw intensity and the data corrected by local background subtraction in GenomeStudio module. A quantile normalization algorithm in the lumi package in the R-based Bioconductor Package was used to normalize the log, transformed intensity data. For data QC, we excluded the probes with detection P value > 0.05 (the P values were generated in BeadStudio software) in at least 25% of the samples, yielding 9,634 genes (18). Both LCL mRNA levels and genotype data were available on 44 patients with EOC from the GRR. Genes containing the SNPs most significantly associated with risk of EOC were selected for SNP-mRNA expression level analyses using linear regression adjusted for patient age and histotype. All analyses were corrected for multiple testing [46].

### ACKNOWLEDGMENTS

The Australian Ovarian Cancer Study Management Group (D. Bowtell, G. Chenevix-Trench, A. deFazio, D. Gertig, A. Green, P. Webb) and ACS Investigators (A. Green, P. Parsons, N. Hayward, P. Webb, D. Whiteman) thank all the clinical and scientific collaborators (see http:// www.aocstudy.org/) and the women for their contribution. The Belgian Ovarium Cancer Study wished to thank Gilian Peuteman, Thomas Van Brussel and Dominiek Smeets for technical assistance. The German Ovarian Cancer Study (GER) thanks Ursula Eilber and Tanja Koehler for competent technical assistance. The Helsinki Ovarian Cancer Study study was supported by the Helsinki University Central Hospital Research Fund. The Mavo Clinic Ovarian Cancer Case-Control Study, for iCOGS thanks C. Hilker, S. Windebank, and J. Vollenweider for iSelect genotyping. The Nurses Health Study and Nurses Health Study II thank the following state cancer registries for their help: AL, AZ, AR, CA, CO, CT, DE, FL, GA, ID, IL, IN, IA, KY, LA, ME, MD, MA, MI, NE, NH, NJ, NY, NC, ND, OH, OK, OR, PA, RI, SC, TN, TX, VA, WA, and WY. The Study of Epidemiology and Risk Factors in Cancer Heredity (SEARCH) thanks the SEARCH team, Craig Luccarini, Caroline Baynes, Don Conroy. Thanks to all members of Scottish Gynaecological Clinical Trails group and SCOTROC1 investigators. United Kingdom Ovarian cancer Population Study thanks I. Jacobs, M.Widschwendter, E. Wozniak, A. Ryan, J. Ford and N. Balogun for their contribution to the study. The UK Familial Ovarian Cancer Registry thanks Carole Pye.

# **CONFLICTS OF INTEREST**

D. Cramer reports a financial relationship with Beasley Allen Crow. E. Goode reports a relationship with Johnson & Johnson. M.T. Goodman is a consultant/ advisory board member for Johnson & Johnson. No additional conflicts of interest were reported.

# FINANCIAL SUPPORT

This study was supported by funding from several sources including the Ovarian Cancer Research Fund thanks to donations by the family and friends of Kathryn Sladek Smith (PPD/RPCI.07); the Genetic Associations and Mechanisms in Oncology (GAME-ON): a NCI Cancer Post-GWAS Initiative (U19-CA148112 and U19-CA148537); the European Community's Seventh Framework Programme under grant agreement n° 223175 (HEALTH-F2-2009-223175); the Canadian Institutes for Health Research (CIHR) MOP-86727 and the CIHR Team in Familial Risks of Breast Cancer; the American Cancer Society (CRTG-00-196-01-CCE); the California Cancer Research Program (00-01389V-20170, N01-CN25403, 2II0200); Cancer Council Victoria; Cancer Council Queensland; Cancer Council New South Wales; Cancer Council South Australia; Cancer Council Tasmania; Cancer Foundation of Western Australia; the Cancer Institute of New Jersey; Cancer Research UK (C490/ A6187, C490/A10119, C490/A10124, C536/A13086, C536/A6689, C1287/A10118, C1287/A 10710, C12292/ A11174, C5047/A8384, C5047/A15007, C5047/A10692); the Celma Mastry Ovarian Cancer Foundation; the Danish Cancer Society (94-222-52); the ELAN Program of the University of Erlangen-Nuremberg; the Eve Appeal; the Helsinki University Central Hospital Research Fund; Helse Vest; Imperial Experimental Cancer Research Centre (C1312/A15589); the Norwegian Cancer Society; the Norwegian Research Council; the Ovarian Cancer Research Fund; Nationaal Kankerplan of Belgium; the L. & S. Milken Foundation; the Polish Ministry of Science and Higher Education; the US National Institutes of Health/National Center for Research Resources/General Clinical Research Center grant MO1-RR000056; the US National Cancer Institute (RPCI-UPCI Ovarian Cancer SPORE P50CA159981-01A1, K07-CA095666, K07- K07-CA80668, CA143047, K22-CA138563, N01-CN55424, N01-PC067010, N01-PC035137. P01-CA017054. P01-CA087696, P20-GM103418, P30-CA072720, P30-CA15083. P30-CA168524, P30-CA008748. P50-CA105009, P50-CA136393, R01-CA014089, R01-CA016056, R01-CA017054, R01-CA049449, R01-CA050385, R01-CA054419, R01-CA058598, R01-CA058860, R01-CA061107, R01-CA061132. R01-CA063682, R01-CA064277, R01-CA067262, R01-CA071766, R01-CA074850, R01-CA076016, R01-CA080742, R01-CA080978, R01-CA128978, R01-CA083918, R01-CA087538, R01-CA092044, R01-095023. R01-CA106414, R01-CA122443. R01-CA61107, R01-CA112523, R01-CA114343, R01-CA126841, R01-CA136924, R01-CA149429. R03-CA113148, R03-CA115195, R21-GM86689, R37-CA070867, R37-CA70867, U01-CA069417, U01-CA071966, CA58860, CA92044, PSA042205, UM1-CA186107, P01-CA87969, R01-CA49449, UM1-CA176726, R01-CA67262 and Intramural research funds); National Institute of Environmental Health Sciences (T32ES013678); the US Department of Defense Ovarian Cancer Research Program (W81XWH-07-0449); the US Army Medical Research and Material Command (DAMD17-98-1-8659, DAMD17-01-1-0729, DAMD17-02-1-0666, DAMD17-02-1-0669, W81XWH-10-1-0280, W81XWH-10-1-0341); the National Health and Medical Research Council of Australia (199600, 400413, and 400281); the German Federal Ministry of Education and Research of Germany Programme of Clinical Biomedical Research (01 GB 9401); the state of Baden-Württemberg through Medical Faculty of the University of Ulm (P.685); the Minnesota Ovarian Cancer Alliance; the Mayo Foundation; the Fred C. and Katherine B. Andersen Foundation; the Lon V. Smith Foundation (LVS-39420); the Polish Committee for Scientific Research (4P05C 028 14 and 2P05A 068 27); the Oak Foundation; the OHSU Foundation; the Mermaid I project; the Rudolf-Bartling Foundation; the UK National Institute for Health Research Biomedical Research Centres at the University of Cambridge; Imperial College London; University College Hospital "Womens Health Theme" and the Royal Marsden Hospital; WorkSafeBC; Komen Foundation for the Cure; and the Breast Cancer Research Foundation; the Lon V Smith Foundation grant LVS-39420.

G. Chenevix-Trench and P.M. Webb are supported by the Australian National Health and Medical Research Council; B. Karlan holds an American Cancer Society Early Detection Professorship (SIOP-06-258-01-COUN); and A. Berchuck holds the Barbara Thomason Ovarian Cancer Research Professorship from the American Cancer Society (SIOP-06-090-06). Cytometry services were provided by the Flow and Image Cytometry Core facility at the Roswell Park Cancer Institute which is supported in part by the NCI Cancer Center Support Grant 5P30 CA016056.

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