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Identification of six new susceptibility loci for invasive epithelial ovarian cancer

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Genome-wide association studies (GWAS) have identified 12 epithelial ovarian cancer (EOC) susceptibility alleles. The pattern of association at these loci is consistent in *BRCA1* and *BRCA2* mutation carriers who are at high EOC risk. After imputation to the 1000 Genomes Project data, we assessed associations of 11 million genetic variants with EOC risk from 15,397 cases unselected for family history and 30,816 controls, 15,252 *BRCA1* mutation carriers and 8,211 *BRCA2* mutation carriers (3,096 with ovarian cancer), and combined the results in a meta-analysis. This new study design yielded increased statistical power, leading to the discovery of six new EOC susceptibility loci. Variants at 1p36 (nearest gene *WNT4*), 4q26 (*SYNPO2*), 9q34.2 (*ABO*) and 17q11.2 (*ATAD5*) were associated with

Author Contributions

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Websites

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The Cancer Genome Atlas Project -<http://cancergenome.nih.gov/>

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⁴⁸on behalf of the German Consortium of Hereditary Breast and Ovarian Cancer (GC-HBOC).

CIMBA QC guidelines-[http://ccge.medschl.cam.ac.uk/consortia/cimba/members/data%20management/CIMBA%20and%20BCAC](http://ccge.medschl.cam.ac.uk/consortia/cimba/members/data%20management/CIMBA%20and%20BCAC%20Quality%20Control%20November%202008%20v2.doc) [%20Quality%20Control%20November%202008%20v2.doc](http://ccge.medschl.cam.ac.uk/consortia/cimba/members/data%20management/CIMBA%20and%20BCAC%20Quality%20Control%20November%202008%20v2.doc)

EOC risk, and at 1p34.3 (*RSPO1*) and 6p22.1 (*GPX6*) specifically with the serous EOC subtype, at $p<5\times10^{-8}$. Incorporating these variants into risk assessment tools will improve clinical risk predictions for *BRCA1/2* mutation carriers.

The risk of developing invasive EOC is higher than the population average for relatives of women diagnosed with the disease^{1,2}, indicating the importance of genetic factors in disease susceptibility. Approximately 25% of the familial aggregation of EOC is explained by rare, high-penetrance alleles of *BRCA1* and *BRCA2*³ . Furthermore, population-based GWAS have identified common variants associated with invasive EOC at 11 loci^{4-9} but only six have also been evaluated in *BRCA1* and/or *BRCA2* mutation carriers. All loci displayed associations in mutation carriers that were consistent with the associations observed in the general population^{10–12}. In addition, the 4q32.3 locus is associated with EOC risk for *BRCA1* mutation carriers only¹³. However, the common genetic variants explain less than 3.1% of the excess familial risk of EOC so additional susceptibility loci are likely to exist.

Women diagnosed with EOC and unaffected women from the general population ascertained through the Ovarian Cancer Association Consortium (OCAC)14 and *BRCA1* and *BRCA2* mutation carriers from the Consortium of Investigators of Modifiers of *BRCA1/2* $(CIMBA)^{15}$ were genotyped as part of the Collaborative Oncological Gene-environment Study (COGS) using the iCOGS custom array. In addition, data were available for cases and controls from three EOC GWAS. We first evaluated whether the EOC susceptibility loci at 8q21.13, 10p12.31, 17q12, 5p15.33, and 17q21.31 recently identified by $OCAC^{7-9}$ also show evidence of association in *BRCA1* and *BRCA2* mutation carriers. Using data from $>$ 200,000 genotyped SNPs^{7,13,16}, we performed imputation of common variants from the 1000 Genomes Project data¹⁷ and evaluated the associations of these SNPs with invasive EOC risk in OCAC and in *BRCA1* and *BRCA2* mutation carriers from CIMBA. Given the strong evidence for a significant overlap in loci predisposing to EOC in the general population and those associated with risk in *BRCA1* and *BRCA2* mutation carriers, we carried out a meta-analysis of the EOC risk associations in order to identify novel EOC susceptibility loci.

Genotype data were available for imputation on 15,252 *BRCA1* mutation carriers and 8,211 *BRCA2* mutation carriers, of whom 2,462 and 631, respectively, were affected with $EOC^{13,16}$. From OCAC, genotyping data were available from 15,437 women with invasive EOC (including $9{,}627$ with serous EOC) and 30,845 controls from the general population⁷. Imputation was performed separately for *BRCA1* carriers, *BRCA2* carriers, OCAC-COGS samples and the three OCAC GWAS (Supplementary Tables 1-2; Supplementary Fig. 1; Supplementary Fig. 2). The meta-analysis was based on 11,403,952 SNPs (Supplementary Fig. 3).

Of five EOC susceptibility loci that have not yet been evaluated in mutation carriers, two were associated with EOC risk for both *BRCA1* and *BRCA2* mutation carriers at p<0.05 (10p12.31 and 17q21.31) (Supplementary Table 3). Overall, seven of the twelve known EOC susceptibility loci provided evidence of association in *BRCA1* mutation carriers and six were associated in *BRCA2* mutation carriers. However, with the exception of 5p15.33 (*TERT*), all loci had hazard ratio (HR) estimates in *BRCA1* and *BRCA2* carriers that were in

the same direction as the odds ratio (OR) estimates for serous subtype EOC from OCAC (Fig. 1). Analysing the associations jointly in *BRCA1* and *BRCA2* carriers and serous EOC in OCAC provided stronger evidence of association, with smaller p-values for eight of the susceptibility variants compared to the analysis in OCAC alone.

Using the imputed genotypes, we observed no novel associations at $p<5\times10^{-8}$ in the analysis of associations in *BRCA1* or *BRCA2* mutation carriers separately. However, we identified seven previously unreported associations (p-values $<5\times10^{-8}$) in either OCAC alone, the meta-analysis of EOC associations in *BRCA1, BRCA2* carriers and OCAC, or in the metaanalysis in *BRCA1* and *BRCA2* carriers and serous EOC in OCAC (Supplementary Fig. 4; Supplementary Tables 4–5). SNPs in six of these loci remained genome-wide statistically significant after re-imputing genotypes with imputation parameters set to maximise accuracy (Table 1; Fig. 1). SNPs at 17q11.2 (near *ATAD5*) were found to be associated with invasive EOC in OCAC ($p<5\times10^{-8}$) (Table 1). For the lead SNP, chr17:29181220:I, the estimated HR estimate for *BRCA1* mutation carriers was significantly different from the estimate in OCAC (p=0.005); the association for *BRCA2* carriers was consistent with the OCAC OR estimate (*BRCA2-OCAC* meta-analysis p=2.6×10−9). SNPs at four loci were associated at $p<5\times10^{-8}$ with risk of all invasive EOC in the meta-analysis (Supplementary Fig. 5): 1p36, 1p34.3, 4q26, and 9q34.2. At 1p34.3, the most strongly associated SNP, rs58722170, displayed stronger associations in the meta-analysis of serous EOC for OCAC $(p=2.7\times10^{-12})$. In addition, SNPs at 6p22.1 were associated at genome-wide significance level in the meta-analysis of associations with serous EOC ($p=3.0\times10^{-8}$), but not in the meta-analysis of all invasive EOC associations ($p=6.8\times10^{-6}$).

The most significantly associated SNP at each of the six novel loci had high imputation accuracy $(r^2 \t0.83)$. At the 1p34.3, 1p36, and 6p22.1 loci, there was at least one genomewide significant genotyped SNP correlated with the lead SNP (pairwise r^2 0.73) (Supplementary Table 6; Supplementary Fig. 5; Supplementary Note). We genotyped the leading (imputed) SNPs of the three other loci in a subset of the samples using iPLEX (Supplementary Note). The correlations between the expected allele dosages from the imputation and the observed genotypes for the variants at $4q26$ and $9q34.2$, ($r^2=0.90$ and r^2 =0.84, respectively) were consistent with the estimated imputation accuracy (0.93 and 0.83 for CIMBA samples). The lead SNP at 17q11.2 failed iPLEX design. However, the risk allele is highly correlated with the *AA* haplotype of two genotyped variants on the iCOGS array (rs9910051 and rs3764419). This haplotype is strongly associated with ovarian cancer risk in the subset of samples genotyped using iCOGS (*BRCA2-OCAC* meta-analysis $p=8.6\times10^{-8}$ for haplotype, and $p=1.8\times10^{-8}$ for chr17:29181220:I) (Supplementary Table 7).

None of the regions contained additional SNPs that displayed EOC associations at p<10⁻⁴ in OCAC, *BRCA1* carriers or *BRCA2* carriers in multi-variable analyses adjusted for the lead SNP in each region, indicating that they each contain only one independent set of correlated highly associated variants (iCHAV). Relative to the 1000 Genomes Project data, we had genotyped or imputed data covering 91% of the genetic variation at 1p36, 84% at 1p34.3 and 83% at 4q26. The other three novel loci had coverage of less than 80% (Supplementary Note). There was evidence for heterogeneity at $p<0.05$ in the associations with histological

subtype in OCAC for the lead SNPs at 1p34.4 and 6p22.1, but not for at 1p36, 4q26, 9q34.2 and 17q11.2 (Table 2).

We carried out a competing risks association analysis in *BRCA1* and *BRCA2* mutation carriers in order to investigate whether these loci are also associated with breast cancer risk for mutation carriers (Supplementary Note). We used the most strongly associated genotyped SNPs for this purpose because the statistical method requires actual genotypes¹⁸. The EOC HR estimates were consistent with the estimates from the main analysis for all SNPs (Supplementary Table 8). None of the SNPs displayed associations with breast cancer risk at $p<0.05$.

At each of the six loci, we identified a set of SNPs with odds of less than 100 to 1 against being the causal variant; most are in non-coding DNA regions (Supplementary Table 9). None were predicted to have likely deleterious functional effects although some lie in or near chromatin biofeatures in fallopian tube and ovarian epithelial cells which may represent the functional regulatory targets of the risk SNPs (Table 3; Supplementary Table 10). We also evaluated the protein coding genes in each region for their role in EOC development, and as candidate susceptibility gene targets. Molecular profiling data from 496 HGSOCs performed by The Cancer Genome Atlas (TCGA) indicated frequent loss/deletion at four risk loci (1p36, 4q26, 9q34.2 and 17q11.2) (Supplementary Table 11). Consistent with this, *WNT4* and *ABO* were significantly down-regulated in ovarian tumours while *ATAD5* was up-regulated. Somatic coding sequence mutations in the six genes nearest the index SNPs were rare. We performed expression quantitative trait locus (eQTL) analysis in a series of 59 normal ovarian tissues (Supplementary Table 12) to evaluate the gene nearest the top ranked SNP at each locus. For the five genes expressed in normal cells, we found no statistically significant eQTL associations for any of the putative causal SNPs at each locus; neither did we find any significant tumour-eQTL associations for these genes based on data from TCGA (Supplementary Table 12). At the 1p36 locus, the most strongly associated variant, rs56318008, is located in the promoter region of *WNT4* which encodes a ligand in the WNT signal transduction pathway, critical for cell proliferation and differentiation. Using a luciferase reporter assay we found no effect of these putatively causal SNPs on *WNT4* transcription in iOSE4 normal ovarian cells (Fig. 2). Some of the putative causal SNPs at 1p36 are located in *CDC42* and *LINC00339*, and several are in putative regulatory domains in ovarian tissues (Supplementary Table 10; Fig. 2). *CDC42* is known to play a role in migration and signalling in ovarian and breast cancer^{19,20}. SNPs at 1p36 are also associated with increased risk of endometriosis and *WNT4, CDC42* and *LINC00339* have all been implicated in endometriosis²¹, a known risk factor for endometrioid and clear cell EOC^{22} .

The strongest associated variant at 1q34, rs58722170, is located in *RSPO1*, which encodes R-spondin 1, a protein involved in cell proliferation (Supplementary Fig. 6). RSPO1 is important in tumorigenesis and early ovarian development^{23,24}, and regulates *WNT4* expression in the ovaries²⁵. *SYNPO2* at 4q26 encodes myopodin which is involved in cell motility and growth²⁶ and has a reported tumour suppressor role^{27–30}. rs635634 is located upstream of the *ABO* gene (Supplementary Fig. 7). A moderately correlated variant (rs505922, r^2 =0.52) determines ABO blood group and is associated with increased risk of pancreatic cancer^{31,32}. Previous studies in OCAC also showed a modestly increased risk of

EOC for individuals with the A blood group³³. The moderate correlation between $rs635634$ and rs505922 and considerably weaker EOC association of rs505922 (p= 1.2×10^{-5}) suggests that the association with blood group is probably not driving the association with risk. The indel, 17:29181220:I, at 17q11.2 is located in *ATAD5* which acts as a tumour suppressor gene^{34–36} (Supplementary Fig. 8). ATAD5 modulates the interaction between RAD9A and BCL2 in order to induce DNA damage related apoptosis. Finally, rs116133110, at 6p22.1, lies in *GPX6* which has no known role in cancer.

The six novel loci reported in this study increase the number of genome-wide significant common variant loci so far identified for EOC to 18. Taken together, these explain approximately 3.9% of the excess familial relative risk of EOC in the general population, and account for approximately 5.2% of the EOC polygenic modifying variance in *BRCA1* mutation carriers and 9.3% in *BRCA2* mutation carriers. The similarity in the magnitude of associations between *BRCA1* and *BRCA2* carriers and population-based studies suggests a general model of susceptibility whereby *BRCA1* and *BRCA2* mutations and common alleles interact multiplicatively on the relative risk scale for EOC^{37} . This model predicts large differences in absolute EOC risk between individuals carrying many alleles and individuals carrying few risk alleles of EOC susceptibility loci for *BRCA1* and *BRCA2* mutation carriers^{13,16}. Incorporating EOC susceptibility variants into risk assessment tools will improve risk prediction and may be particularly useful for *BRCA1* and *BRCA2* mutation carriers.

METHODS

Study populations

We obtained data on *BRCA1* and *BRCA2* mutation carriers through CIMBA. Eligibility in CIMBA is restricted to females 18 years or older with pathogenic mutations in *BRCA1* or *BRCA2*. The majority of the participants were sampled through cancer genetics clinics¹⁵, including some related participants. Fifty-four studies from 27 countries contributed data. After quality control, data were available on 15,252 *BRCA1* mutation carriers and 8,211 *BRCA2* mutation carriers, of whom 2,462 and 631, respectively, were affected with EOC (Supplementary Table 1).

Data were available for the stage 1 of three population-based EOC GWAS. These included 2,165 cases and 2,564 controls from a GWAS from North America ("US GWAS")39, 1,762 cases and 6,118 controls from a UK-based GWAS ("UK GWAS")⁶, and 441 cases and 441 controls from the Mayo GWAS. Furthermore, 11,069 cases and 21,722 controls were genotyped using the iCOGS array ("OCAC-iCOGS" stage data). Overall, 43 studies from 11 countries provided data on 15,347 women diagnosed with invasive epithelial EOC, 9,627 of whom were diagnosed with serous EOC, and 30,845 controls from the general population.

All subjects included in this analysis were of European descent and provided written informed consent as well as data and blood samples under ethically approved protocols. Further details of the OCAC and CIMBA study populations as well as the genotyping, quality control and statistical analyses have been described elsewhere^{7,13,16}.

Genotype data

Genotyping and imputation details for each study are shown in Supplementary Table 1.

Confirmatory genotyping of imputed SNPs

To evaluate the accuracy of the imputation of the SNPs we found to be associated with EOC risk, we genotyped rs17329882 (4q26) and rs635634 (9q34.2) in a subset of 3,541 subjects from CIMBA using Sequenon's iPLEX technology. The lead SNP at 17q11.2, chr17:29181220:I failed iPLEX design. We performed quality control of the iPLEX data according to the CIMBA guidelines. After quality control, we used the imputation results to generate the expected allele dosage for each genotyped sample and computed the Pearson product-moment correlation coefficient between the expected allele dosage and the observed genotype. The squared correlation coefficient was compared to the imputation accuracy as estimated from the imputation.

Quality control of GWAS and iCOGS genotyping data

We carried out quality control separately for *BRCA1* carriers, *BRCA2* carriers, the three OCAC GWAS, and OCAC-iCOGS samples, but quality criteria were mostly consistent across studies. We excluded samples if they were not of European ancestry, if they had a genotyping call rate < 95%, low or high heterozygosity, if they were not female or had ambiguous sex, or were duplicates (cryptic or intended). In OCAC studies, one individual was excluded from each pair of samples found to be first-degree relatives and duplicate samples between the iCOGS stage and any of the GWAS were excluded from the iCOGS data. SNPs were excluded if they were monomorphic, had call rate<95%, showed evidence of deviation from Hardy-Weinberg equilibrium or had low concordance between duplicate pairs. For the Mayo GWAS and the UK GWAS, we also excluded rare SNPs (MAF<1% or allele count <5, respectively). We visually inspected genotype cluster plots for all SNPs with P<10⁻⁵ from each of the newly identified loci. We used the R GenABEL library version 1.6.7 for quality control⁴⁰.

Genotype data were available for analysis from iCOGS for 199,526 SNPs in OCAC-iCOGS, 200,720 SNPs in *BRCA1* mutation carriers, and 200,908 SNPs in *BRCA2* mutation carriers. After QC, for the GWAS, data were available on 492,956 SNPs for the US GWAS, 543,529 SNPs for the UK GWAS and 1,587,051 SNPs for the Mayo GWAS (Supplementary Table 2).

Imputation

We performed imputation separately for *BRCA1* carriers, *BRCA2* carriers, OCAC-iCOGS samples and each of the OCAC GWAS. We imputed variants from the 1000 Genomes Project data using the v3 April 2012 release¹⁷ as the reference panel. For OCAC-iCOGS, the UK GWAS and the Mayo GWAS, imputation was based on the 1000 Genomes Project data with singleton sites removed. To improve computation efficiency we initially used a twostep procedure, which involved pre-phasing in the first step and imputation of the phased data in the second. We carried out pre-phasing using the SHAPEIT software⁴¹. We used the IMPUTE version 2 software for the subsequent imputation⁴² for all studies with the

exception of the US GWAS for which the MACH algorithm implemented in the minimac software version 2012.8.15, mach version 1.0.18 was used. To perform the imputation we divided the data into segments of approximately 5Mb each. We excluded SNPs from the association analysis if their imputation accuracy was $r^2 < 0.3$ or their minor allele frequency (MAF) was ≤ 0.005 in *BRCA1* or *BRCA2* carriers or if their accuracy was $r^2 \leq 0.25$ in OCACiCOGS, the UK GWAS, UK GWAS or Mayo GWAS.

We performed more accurate imputation for the regions around the novel EOC loci from the joint analysis of the data from *BRCA1* and *BRCA2* carriers and the general population (any SNP with P<5×10⁻⁸). The boundaries of these regions were set +/− 500kb from any significantly associated SNP in the region. As in the first run, the 1000 Genomes Project data v3 were used as the reference panel and the software IMPUTE2 was applied. However, for the second round of imputation, we imputed genotypes without pre-phasing in order to improve accuracy. To further increase the imputation accuracy we changed some of the default parameters in the imputation procedure. These included an increase of the MCMC iterations to 90 (out of which the first 15 were used as burn-in), an increase of the buffer region to 500kb and an increase of the number of haplotypes used as templates when phasing observed genotypes to 100. These changes were applied consistently for all data sets.

Statistical analyses

Association analyses in the unselected ovarian cancer cases and controls from OCAC—We evaluated the association between genotype and disease using logistic regression by estimating the associations with each additional copy of the minor allele (logadditive models). The analysis was adjusted for study and for population substructure by including the eigenvectors of the first five ancestry specific principal components as covariates in the model. We used the same approach to evaluate the SNP associations with serous ovarian cancer after excluding all cases with any other or with unknown tumour subtype. For imputed SNPs we used expected dosages in the logistic regression model to estimate SNP effect sizes and p-values. We carried out analyses separately for OCACiCOGS and the three GWAS and pooled thereafter using a fixed effects meta-analysis. We carried out the analysis of re-imputed genotypes of putative novel susceptibility loci jointly for the OCAC-iCOGS and GWAS samples. All results are based on the combined data from iCOGS and the three GWAS. We used custom written software for the analysis.

Associations in BRCA1 and BRCA2 mutation carriers from CIMBA—We carried out the ovarian cancer association analyses separately for *BRCA1* and *BRCA2* mutation carriers. The primary analysis was carried out within a survival analysis framework with time to ovarian cancer diagnosis as the endpoint. Mutation carriers were followed until the age of ovarian cancer diagnosis, or risk-reducing salpingo-oophorectomy (RRSO) or age at last observation. Breast cancer diagnosis was not considered as a censoring event. In order to account for the non-random sampling of *BRCA1* and *BRCA2* mutation carriers with respect to their disease status we conducted the analyses by modelling the retrospective likelihood of the observed genotypes conditional on the disease phenotype¹⁸. We assessed the associations between genotype and risk of ovarian cancer using the 1 degree of freedom

score test statistic based on the retrospective likelihood^{18,43}. To account for the nonindependence among related individuals in the sample, we used an adjusted version of the score test statistic, which uses a kinship adjusted variance of the score⁴⁴. We evaluated associations between imputed genotypes and ovarian cancer risk using a version of the score test as described above but with the posterior genotype probabilities replacing the genotypes. All analyses were stratified by the country of origin of the samples.

We carried out the retrospective likelihood analyses in CIMBA using custom written functions in Fortran and Python. The score test statistic was implemented in R version $3.0.1^{45}$.

We evaluated whether there is evidence for multiple independent association signals in the region around each newly identified locus by evaluating the associations of genetic variants in the region while adjusting for the SNP with the smallest meta-analysis p-value in the respective region. This was done separately for *BRCA1* carriers, *BRCA2* carriers and OCAC.

For one of the novel associations, it was not possible to confirm the imputation accuracy of the lead SNP chr17:29181220:I at 17q11.2 through genotyping. Therefore, we inferred twoallele haplotypes for rs9910051 and rs3764419, highly correlated with the lead SNP $(r^2=0.95)$, using an in-house program. These variants were genotyped on the iCOGS array and therefore this analysis was restricted to 14,733 ovarian cancer cases and 9,165 controls from OCAC-COGS, and 8,185 *BRCA2* mutation carriers that had available genotypes for both variants based on iCOGS. The association between the AA haplotype and risk was tested using logistic regression in OCAC and using Cox regression in *BRCA2* mutation carriers.

Meta-analysis—We conducted a meta-analysis of the EOC associations in *BRCA1, BRCA2* carriers and the general population for genotyped and imputed SNPs using an inverse variance approach assuming fixed effects. We combined the logarithm of the perallele hazard ratio estimate for the association with EOC risk in *BRCA1* and *BRCA2* mutation carriers and the logarithm of the per-allele odds ratio estimate for the association with disease status in OCAC. For the associations in *BRCA1* and *BRCA2* carriers, we used the kinship adjusted variance estimator 44 which allows for inclusion of related individuals in the analysis. We only used SNPs with results in OCAC and in at least one of the *BRCA1* or the *BRCA2* analyses. We carried out two separate meta-analyses, one for the associations with EOC in *BRCA1* carriers, *BRCA2* carriers and EOC in OCAC, irrespective of tumour histological subtype, and a second using only the associations with serous EOC in OCAC. The number of *BRCA1* and *BRCA2* samples with tumour histology information was too small to allow for subgroup analyses. However, previous studies have demonstrated that the majority of EOCs in *BRCA1* and *BRCA2* mutation carriers are high-grade serous^{49–53}. Metaanalyses were carried out using the software "metal", $2011-03-25$ release⁵⁴.

Candidate causal SNPs in each susceptibility region—In order to identify a set of potentially causal variants we excluded SNPs with a likelihood of being causal of less than 1:100, by comparing the likelihood of each SNP from the association analysis with the one

of the most strongly associated SNP^{46} . The remaining variants were then analysed using pupasuite 3.1 to identify potentially functional variants (Supplementary Table 9).

Functional analysis

Expression quantitative trait locus (eQTL) analysis in normal OSE and FTSE

cells—Early-passage primary normal ovarian surface epithelial cells (OSECs) and fallopian tube epithelial cells were harvested from disease-free ovaries and fallopian tubes. Normal ovarian epithelial cells were collected by brushing the surface of the ovary with a sterile cytobrush, and were cultured in NOSE-CM55. Fallopian tube epithelial cells were harvested by Pronase digestion as previously described⁵⁶, plated onto collagen-coated plastics (Sigma) and cultured in DMEM/F12 (Sigma-Aldrich) supplemented with 2% Ultroser G (BioSepra) and $1\times$ penicillin/streptomycin (Lonza). By the time of RNA harvesting, fallopian tube cultures tested consisted of PAX8 positive fallopian tube secretory epithelial cells (FTSECs), consistent with previous observations that ciliated epithelial cells from the fallopian tube do not proliferate *in vitro*.

For gene expression analysis, RNA was harvested from 59 early passage samples: 54 OSECs and 5 FTSECs from cell cultures harvested at ~80% confluency using the QIAgen miRNAeasy kit with on-column DNase 1 digestion. 500ng RNA was reverse transcribed using the Superscript III kit (Life Technologies). We preamplified 10ng cDNA using the TaqMan® Preamp Mastermix; the resulting product was diluted 1:60 and used to quantify gene expression using the following TaqMan® gene expression probes: WNT4, Hs01573504_m1; RSPO1, Hs00543475_m1; SYNPO2, Hs00326493_m1; ATAD5, Hs00227495 m1 and GPX6, Hs00699698 m1. Four control genes were also included: ACTB, Hs00357333_g1; GAPDH, Hs02758991_g1; HMBS, Hs00609293_g1 and HPRT1 Hs02800695_m1 (all Life Technologies). Assays were run on an ABI 7900HT Fast Real-Time PCR system (Life Technologies).

Data Analysis: Expression levels for each gene were normalized to the average of all four control genes. Relative expression levels were calculated using the δδCt method. Genotyping was performed on the iCOGs chips, as described above. Where genotyping data were not available for the most risk-associated SNP, the next most significant SNP was used: rs3820282 at 1p36, rs12023270 at 1p34.3, rs752097 at 4q26, rs445870 at 6p22.1, rs505922 at 9q34.2 and rs3764419 at 17q11.2. Correlations between genotype and gene expression were calculated in 'R'. Genotype specific gene expression in the normal tissue cell lines (eQTL analysis) was compared using the Jonckheere-Terpstra test. IData were normalized to the four control genes and we tested for eQTL associations, grouping OSECs and FTSECs together. Secondly, OSECs were analysed alone. eQTL analyses were performed using 3 genotype groups, or two groups (with the rare homozygote samples grouped together with the heterozygote samples).

eQTL analysis in primary ovarian tumours—eQTL analysis in primary tumours was based on the publicly available data available from The Cancer Genome Atlas (TCGA) project, which includes 489 primary high grade serous ovarian cancers. The methods have been described elsewhere⁵⁷. Briefly, we determined the ancestry for each case based on the

germ line genotype data using EIGENSTRAT software with 415 HapMap genotype profiles as a control set. Only populations of Northern and Western European ancestries were included. We first performed a *cis*-eQTL analyses using a method we described previously, in which the association between 906,600 germline genotypes and the expression levels of mRNA or miRNA (located within 500Kb on either side of the variant) were evaluated using linear regression model with the effects of somatic copy number and CpG methylation being deducted (For miRNA expression, the effect of CpG methylation is not adjusted for since the data are not available). To adjust for multiple tests, we adjusted the test P values using Benjamini-Hochberg method. A significant association was defined by a false discovery rate (FDR) of less than 0.1.

Having established a genome-wide *cis*-eQTL associaitions in this series of tumours, we then evaluated *cis*-eQTL associations for the top risk associations between each of the six new loci and the gene in closest proximity to the risk SNP. For each risk locus, we retrieved the genotype of all SNPs in ovarian cancer cases based on the Affymetrix 6.0 array. Using these genotypes and the impute2 March 2012 1000 Genomes Phase I integrated variant cosmopolitan reference panel of 1,092 individuals (Haplotypes were phased via SHAPEIT), we imputed the genotypes of SNPs in the 1000 Genomes Project in the target regions for TCGA samples⁵⁸. For each risk locus where data for the most risk-associated variant were not available, we retrieved the imputed variants tightly correlated with the most riskassociated variant. We then tested for association between imputed SNPs and gene expression using the linear regression algorithm described above, where each imputed SNP was coded as an expected allele count. Again, significant associations are defined by a false discovery rate (FDR) of less than 0.1.

Regulatory profiling of normal ovarian cancer precursor tissues—We performed genome-wide formaldehyde assisted regulatory element (FAIRE) and ChIP seq with histone 3 lysine 27 acetylation (H3K27ac) and histone 3 lysine 4 monomethylation (H3K4me) for two normal OSECs, two normal FTSECs and two HGSOC cell lines (UWB1.289 and CAOV3) [Shen et al. in preparation]. These datasets annotate epigenetic signatures of open chromatin, and collectively indicate transcriptional enhancer regions. We analysed the FAIRE-seq and ChIP-seq datasets and publically available genomic data on promoter and UTR domains, intron/exon boundaries, and positions of non-coding RNA transcripts to identify SNPs from the 100:1 likely causal set that align with biofeatures that may provide evidence of SNP functionality.

Candidate Gene Analysis Using Genome Wide Profiling of Primary Ovarian Cancers

Data Sets: The Cancer Genome Atlas (TCGA) Project and COSMIC Datasets

TCGA has performed extensive genomic analysis of tumours from a large number of tissue types including almost 500 high-grade serous ovarian tumours. These data include somatic mutations, DNA copy number, mRNA and miRNA expression and DNA methylation. COSMIC is the catalogue of somatic mutations in cancer that collates information on mutations in tumours from the published literature⁵⁹. They have also identified The Cancer

Gene Census, which is a list of genes known to be involved in cancer. Data are available on a large number of tissue types, including 2,809 epithelial ovarian tumours.

Somatic coding sequence mutations: We analysed all genes for coding somatic sequence mutations generated from either whole exome or whole genome sequencing. In TCGA, whole exome sequencing data were available for 316 high-grade serous EOC cases. In addition, we determined whether mutations had been reported in COSMIC⁵⁹ and whether the gene was a known cancer gene in the Sanger Cancer Gene Census.

mRNA expression in tumour and normal tissue: Normalized and gene expression values (Level 3) gene expression profiling data were obtain from the TCGA data portal for three different platforms (Agilent, Affymetrix HuEx and Affymetrix U133A). We analysed only the 489 primary serous ovarian tumour samples included in the final clustering analysis⁵⁸ and eight normal fallopian tube samples. The boxplot function in R was used to compare ovarian tumour samples to the fallopian tube for 91 coding genes with expression data on any platform within a 1MB region around the most significant SNP at the six loci. A difference in relative expression between EOC and normal tissue was carried out using the Wilcoxon rank-sum test.

DNA copy number analysis: Serous EOC samples for 481 tumours with log2 copy number data were analysed using the cBio portal for analysis of TCGA data^{60,61}. For each gene in a region the classes of copy number; homozygous deletion, heterozygous loss, diploid, gain, and amplification were queried individually using the advanced onco query language (OQL) option. The frequency of gain and amplification were combined as "gain", and homozygous deletion and heterozygous loss were combined as "loss".

Analysis of copy number vs mRNA expression: Serous EOC samples for 316 complete tumours (those with CNA, mRNA and sequencing data) were analysed. Graphs were generated using the cBio portal for analysis of TCGA data and the setting were mRNA expression data Z-score (all genes) with the Z-score threshold of 2 (default setting) and putative copy number alterations (GISTIC). The Z-score is the number of standard deviations away from the mean of expression in the reference population. GISTIC is an algorithm that attempts to identify significantly altered regions of amplification or deletion across sets of patients.

Luciferase Reporter Assay—The putative causal SNPs at the 1p36 locus lie in the *WNT4* promoter and so we tested their effect on transcription in a luciferase reporter assay (Fig. 2D). Wild-type and risk haplotype (comprising five correlated variants) sequences corresponding to the region bound by hg19 co-ordinates chr1:22469416-22470869 were generated by Custom Gene Synthesis (GenScript Corporation), and then sub-cloned into pGL3-basic (Promega). Equimolar amounts of luciferase constructs (800 ng) and pRL-TK Renilla (50 ng) were co-transfected into -8×10^4 iOSE4⁶² normal ovarian cells in triplicate wells of 24 well plates using LipoFectamine 2000 (Life Technologies). Independent transfections were repeated three times. The Dual-Glo Luciferase Assay kit (Promega) was used to assay luciferase activity 24 hours post transfection using a BioTek Synergy H4 plate

reader. The iOSE-4 cell line (derived by K. Lawrenson) was maintained under standard conditions and routinely tested for *Mycoplasma* and short tandem repeat profiled.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Figure 1.

Hazard ratios for the association with EOC of 12 previously reported epithelial ovarian cancer susceptibility variants and the six novel susceptibility variants for OCAC, *BRCA1* mutation carriers and *BRCA2* mutation carriers

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Figure 2.

The 1p36 epithelial ovarian cancer susceptibility locus

A) The Manhattan plot depicts the strength of association between all imputed and genotyped SNPs across the region bound by hg19 co-ordinates chr1:21922893-22991643. The dotted line represents the genome-wide significance level 5×10^{-8} . Additional tracks show genes and enhancers in ovary as described in Hnisz et al^{38} . Positions of SNPs for which imputation $r^2 < 0.3$ and/or minor allele frequency < 0.005 are shown in the bottom track as 'untyped' SNPs.

B) The shaded iCHAV from (A) is shown depicting the genes and the location of the *WNT4* promoter construct as a red box. Red ticks show the positions of the putative causal variants following likelihood ratio testing. Signals from FAIRE-seq data derived from ovarian cells are represented by black marks, and the locations of predicted *CDC42* enhancers³⁸ as blue boxes. The positions of genotyped SNPs, and those that were neither genotyped nor well imputed ('untyped'), are shown.

C) Normalised luciferase reporter activity following triplicate transfections of wildtype and risk haplotype *WNT4* promoter constructs in iOSE4 cells. Error bars represent standard error from three independent experiments.

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Table 1

each locus Association test results for loci associated at p<5×10⁻⁸ in the second imputation stage. Results reported for ovarian cancer in *BRCA1* and *BRCA2* mutation carriers, ovarian cancer as well as serous subtype in Association test results for loci associated at p<5×10−8 in the second imputation stage. Results reported for ovarian cancer in *BRCA1* and *BRCA2* mutation carriers, ovarian cancer as well as serous subtype in OCAC, the meta-analysis for ovarian cancer, and for the meta-analysis for all tumour histologies in *BRCA1* and *BRCA2* and serous ovarian cancer in OCAC. SNP with smallest p-value reported for each locus $H \circ A$ for $\frac{1}{2}$ ~ 11 \therefore in OCAC SND with \circ Ą $\frac{1}{2}$ bietologies in RPCA1 and RPCA2 and $\frac{1}{2}$ ~ 1 and ~ 10 fm. \cdot \overline{d} for the $\ddot{}$ $\frac{1}{2}$ $\ddot{}$ NOAC_{the}

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*4*rs116133110 listed as rs6456822 on dbSNP

 4 rs116133110 listed as rs6456822 on dbSNP

*5*chr17:29181220:I listed as rs199661266 on dbSNP

 5 chr
17:29181220:1
listed as rs199661266 on dbSNP

*6*Reference and effect allele

 $6\,$ Reference and effect allele

*7*Effect allele frequency

 $7\!\!$ Effect allele frequency

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p-value for the heterogeneity in associations with different tumour subtypes p-value for the heterogeneity in associations with different tumour subtypes

Summary of data on SNPs, closest gene and all genes in 1 MB region for each locus Summary of data on SNPs, closest gene and all genes in 1 MB region for each locus

Biofeatures defined as open chromatin, H3K4me3 or H3K27ac marks detected in normal ovarian and/or fallopian cells Biofeatures defined as open chromatin, H3K4me3 or H3K27ac marks detected in normal ovarian and/or fallopian cells

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*** There are 16 genes in this region; *ABO, SURF6, MED22, RPL7A, SNORD24, SNORD36B, SNORD36A, SNORD36C, SURF1, SURF2, SURF4, C9orf96, REXO4, ADAMTS13, CACFD1, SLC2A6,* however all SNPs are within or upstream of *ABO* or upstream of *SLC2A6*.

**** Trend p=0.067