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Integrated Analysis of Germline and Somatic Variants in Ovarian Cancer

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Abstract

We report the first large-scale exome-wide analysis of the combined germline-somatic landscape in ovarian cancer. Here we analyze germline and somatic alterations in 429 ovarian carcinoma cases and 557 controls. We identify 3,635 high confidence, rare truncation and 22,953 missense

COMPETING FINANCIAL INTERESTS

The authors declare no competing financial interests.

CONTRIBUTIONS

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variants with predicted functional impact. We find germline truncation variants and large deletions across Fanconi pathway genes in 20% of cases. Enrichment of rare truncations is shown in *BRCA1*, *BRCA2*, and *PALB2*. Additionally, we observe germline truncation variants in genes not previously associated with ovarian cancer susceptibility (*NF1*, *MAP3K4*, *CDKN2B*, and *MLL3*). Evidence for loss of heterozygosity was found in 100% and 76% of cases with germline *BRCA1* and *BRCA2* truncations respectively. Germline-somatic interaction analysis combined with extensive bioinformatics annotation identifies 237 candidate functional germline truncation and missense variants, including 2 pathogenic *BRCA1* and 1 *TP53* deleterious variants. Finally, integrated analyses of germline and somatic variants identify significantly altered pathways, including the Fanconi, MAPK, and MLL pathways.

INTRODUCTION

Ovarian cancer is diagnosed in ~22,000 women annually in the United States. The average five year survival is relatively poor at ~43%¹, which is primarily due to late-stage diagnosis. It is currently estimated that 20–25% of women have an inherited germline mutation that predisposes them to ovarian cancer.^{2,3} New strategies for the prevention and control of ovarian cancer will rely on a thorough understanding of the contributing genetic factors both at the germline and somatic levels.

High throughput sequencing technologies are rapidly expanding our understanding of ovarian cancer biology by providing comprehensive descriptions of genetic aberrations in tumors.⁴ The ability to rapidly sequence individual tumor and normal genomes allows for efficient discovery of candidate cancer-causing events and such work is already transforming risk assessment, diagnosis, and treatment. For example, targeted sequencing of 21 tumor suppressor genes in 360 cases of ovarian, peritoneal, fallopian tube, and synchronous ovarian/endometrial carcinomas recently revealed that 24% of cases harbored germline loss of function mutations in 1 of 12 genes: *BRCA1*, *BRCA2*, *BARD1*, *BRIP1*, *CHEK2*, *MRE11A*, *MSH6*, *NBN*, *PALB2*, *RAD50*, *RAD51C*, and *TP53.*³ In a different study, The Cancer Genome Atlas (TCGA) consortium analyzed somatic alterations in 316 serous ovarian carcinomas, identifying recurrent somatic *TP53* mutations in nearly all cases (96%) and finding recurrent somatic mutations in *NF1*, *BRCA1*, *BRCA2*, *RB1*, and *CDK12* in a minority of cases.⁴ Such work is deepening our understanding of genes involved in ovarian cancer.

Cancer genomics studies have most often focused on independent analyses of either somatic or germline mutations. However, studies that perform sequencing of matched tumor and normal samples have the advantage that data from the somatic and germline genomes can be ascertained and integrated to build a fuller picture of each genome's contribution to disease. In addition, the rapidly growing number of publicly available exome datasets from non-cancer populations now facilitates rare germline susceptibility variant discovery.

Here we describe the somatic and germline mutation spectrum in the tumor and normal exome data from 429 TCGA serous ovarian cancer patients. To identify likely pathogenic variants, we compare the frequency of germline mutations to those from a large control dataset of sequences of post-menopausal women from the Women's Health Initiative Exome

Sequencing Project (WHISP). We identify several novel candidate germline predisposition variants in known ovarian genes (e.g., *BRCA1*, *BRCA2*, *ATM*, *MSH3*, and *PALB2*) as well as several genes not previously associated with ovarian cancer (e.g., *ASXL1*, *RB1*, *NF1*, *CDKN2A*, and *EXO1*). We also characterize patterns of loss of heterozygosity in tumor suppressor genes, including *BRCA1*, *BRCA2*, *BRIP1*, *ATM*, *CHEK2*, and *PALB2*, and identify significantly mutated pathways, including Fanconi anemia, MAPK, and MLL. These results provide a foundation for future functional and clinical assessment of susceptibility variants in ovarian cancer.

RESULTS

Clinical Characteristics of samples

Of the 429 TCGA cases in this analysis, 90.2% were Caucasian (n=387), 4.9% were African American (n=21), 3.5% were Asian (n=15), and 0.5% (n=2) were American Indian/Alaska Native. Patients were diagnosed between 26–89 years (mean 59.4 ± 11.8 years), frequently at late stage (93% at stages 3–4), and 50.8% were deceased at the time of TCGA sample procurement (Table 1). Nineteen of twenty-three cases with unknown ethnicity information were assigned Caucasian (n=17) and African ancestry (n=2) using principal components analysis (Supplementary Fig. 1). We performed systematic germline variant and somatic mutation analyses for the sample set, as illustrated in Fig. 1.

Data for 614 samples from the National Heart Lung and Blood Institute (NHBLI) Women's Health Initiative Exome Sequencing Project (WHISP) was used for comparison of genetic variants to TCGA ovarian cancer cases. After extensive quality checks (Methods), 557 Caucasians with an average age of 63.3 years \pm 7.8 years (range 50–79 years) were selected as controls for downstream ovarian susceptibility variant analysis (Supplementary Data 1).

Somatic mutations and significantly mutated genes

We analyzed somatic mutations in 429 ovarian cancer cases. Of these, 142 were new TCGA cases and 287 cases were previously reported⁴; the remaining twenty-nine cases reported in that study⁴ did not meet our coverage requirement (20x coverage for at least 50% of target exons) and were excluded from this analysis. The average exome-wide coverage for the entire sample set was 68.1X with 99.5X and 96.5X average coverages for BRCA1 and BRAC2, respectively. We identified 11,479 somatic mutations in the 142 new TCGA cases. All of these mutations were manually reviewed, resulting in a total of 27,280 mutations in 429 cases (Fig. 1 and Supplementary Data 2 and 3). After removing genes with low or no RNA expression evidence from RNA-seq data, the significantly mutated genes (SMGs) identified by MuSiC⁵ include those previously reported: TP53, NF1, RB1, CDK12(CRKRS), and BRCA14, as well as the new SMG, KRAS (Supplementary Table 1). BRCA2 and RB1CC1 were near significance. We also identified 4 NRAS mutations, 3 NF2 mutations, and 3, 8, and 10 mutations in the known tumor suppressor genes: ATR, ATM, and APC, respectively. Somatic truncation mutations were also observed in histone modifier genes including: ARID1A, ARID1B, ARID2, SETD2, SETD4, SETD6, JARID1C, MLL, MLL2, and MLL3 as well as the DNA excision repair gene ERCC6 (Supplementary Data 3).

Germline variant landscapes and significant germline events

We identified germline truncation variants (nonsense, nonstop, splice site, and frameshift indels) in these 429 matched tumor-normal cases using multiple algorithms.^{6–8} After removal of common variants, reference sequence errors, and recurrent artifacts, a total of 3,635 high confidence, rare (<1% population minor allele frequency) germline truncation variants were identified in 2,214 genes, 115 of which are in 40 known cancer genes (Fig. 1, Supplementary Fig. 2, Supplementary Data 4 and Methods).⁹ These 115 variants were validated using genomic DNA or a source of whole genome amplified DNA that differed from that used for discovery (Supplementary Data 5). We used several approaches to identify known and potentially pathogenic germline missense variants in the Caucasian subset (Table 1, n = 387). Specifically, a total of 22,953 missense variants in 3,637 genes were predicted to be functionally deleterious by Condel¹⁰ and also had population minor allele frequencies (MAFs) <1% in Caucasian data from the 1000 Genomes, and the current cohorts (TCGA ovarian cancer cases and WHISP exome controls) (Fig. 1, Supplementary Data 6, and Supplementary Fig. 3). After limiting our analyses to genes with an average expression RPKM >0.5 (Methods), we identified 17,348 missense variants in a total of 2,810 genes in this subset. We processed on 557 WHISP samples using the same software tools and filtering strategies and identified 7,889 rare (<1% minor allele frequency in the population and cohort) truncation variants and 30,335 rare missense variants defined as functional by Condel and in expressed genes (Supplementary Data 7 and 8).

Finally, although we performed a genome-wide germline copy number analysis using SNP array data, our manual review of the results indicated many false positives with very few passing our review criteria. Therefore, we focused our analysis of copy number alterations on *BRCA1*, *BRCA2*, and *TP53*, coupled with extensive manual review. Here, three high confidence germline deletion events in *BRCA1* were identified in three cases (TCGA-36-2539, TCGA-31-1959, and TCGA-23-1028) (Fig. 2). Two cases (TCGA-31-1959 and TCGA-23-1028) developed ovarian cancer at younger ages (50 and 43 years, respectively); information regarding age of diagnosis for TCGA-36-2539 was not available.

We used a right tailed CAST¹¹ burden test CAST_{greater} (personal communication, Qunyuan Zhang) to evaluate expressed genes (Methods) having significant enrichment of rare, potentially pathogenic missense variants in the TCGA Caucasian exomes versus the WHISP control group and, the test identified 24 genes that had significant enrichment (P < 0.0002, CAST_{greater}). As expected, *BRCA1* is one of the most significant genes on the list (P = 1.40 E-06, CAST_{greater}). A total of 9 unique *BRCA1* rare missense variants were detected in this ovarian cancer cohort; this list included two known pathogenic missense variants (R1699W and G1788V) and three singletons (V772A, L668F, and P1637L). It also included one known ovarian susceptibility gene (*FANCM*; P = 4.04–06, CAST_{greater}) as well as three cancer genes (*ARID1A*, *EGFR*, and *DNMT1*), not previously implicated in ovarian cancer (Supplementary Data 6 and 9). *ARID1A*, frequently mutated in endometrial cancer¹² and *EGFR*, a prominent oncogene involved in lung cancer¹³ and glioblastoma¹⁴, harbored 10 and 5 rare (1% MAF) unique missense variants in this ovarian sample set, respectively. Several other known cancer genes (e.g., *CREBBP*, *ASXL1*, *EZH2*, and *BRIP1*) were also

found to be in the top 100 and with $P_{CAST.greater} < 0.0015$. The significance of other top genes such as *EEF2K* requires additional investigation using larger sample sets.

We next focused on comparison of rare germline truncations in cancer genes between TCGA Ovarian cases and the WHISP control set. Three known ovarian cancer susceptibility genes were significant at the right tailed CAST test p 0.05 as a threshold (BRCA1 (P = 2E-08), BRCA2 (P = 8.89E-06), PALB2 (P = 0.042)) and two other known ovarian cancer susceptibility genes were among the highest ranked genes although they did not reach significance (CHEK2 (P = 0.11), and BRIP1 (P = 0.11)) (Supplementary Table 2). A total of 66 cases had truncations in one of these genes (Supplementary Data 4 and 5). It is worth noting that we have identified truncation mutations in USP6, ROPN1L, and RYR1, although their involvements in cancer are unclear. In addition, three truncation variants (T1222fs, Q645*, and L258fs) were detected in BLM that has recently been linked to familial breast cancer.¹⁵ Q645* and L258fs were previously reported in BLMbase (http://bioinf.uta.fi/ BLMbase/). The distribution of germline and somatic mutations in these genes is shown in Fig. 3. It is interesting to note that 11 cases had germline truncation variants in multiple cancer genes, including two cases with BRCA1 and BRCA2 variants (diagnosis ages 49 and 55 years), one case with BRCA2 and ERCC3 variants, one with PALB2 and ATM variants, and one with BLM and FANCD2 truncation variants. Finally, five cases had germline truncation variants in other genes on the cancer gene list, including: ERCC2 (n=1), TET2 (n=1), FANCD2 (n=2), and NF1 (n=1) while one case had a germline mutation in RAD51B which has recently been linked to breast cancer susceptibility¹⁶ and whose family members (RAD50, RAD51C, RAD51D) have previously been implicated in ovarian cancer susceptibility.¹⁷

When we combined missense and truncation variants in cancer genes for burden testing, known cancer susceptibility genes were among the most significant genes on the list (*BRIP1*^{3,18} and *BRCA1*). In addition, other established/suspected ovarian/breast cancer susceptibility genes were significant, including *BRCA2*², and *NF1*¹⁹; novel genes such as *ASXL1*, frequently mutated in myelodysplastic syndromes²⁰, myeloproliferative neoplasms²¹, and AML²²; *SETD2*, involved in clear cell renal cell carcinoma²³; and *MAP3K1*, a newly discovered breast cancer gene^{24,25} (Supplementary Data 10).

Germline variants that have been detected as somatically mutated in cancer might signal functional relevance of these variants. We compared our identified germline truncation and missense variants to those present in the COSMIC and OMIM databases to determine whether any were reported in other studies. Of the 3,635 exome-wide truncation variants, 84 and 10 germline variants matched precisely or within ±5 amino acids to reported variants in COSMIC and OMIM, respectively (Supplementary Data 11). Further analysis of 535 missense variants from cancer genes, using the same criteria applied for truncations, identified 35 and 14 missense events in COSMIC and OMIM, respectively (Supplementary Data 11). For example, the *ASXL1* germline variant G1397S that we identified in 6 of 387 ovarian cancer cases versus 2 of 557 WHISP non-cases and the *ASXL1* germline variant G643V identified in 1 of 387 cases vs. 0 of 557 WHISP non-cases have previously been found to be somatically mutated in hematologic malignancies.^{26,27} Although there was not an exact match of the germline variant P333L in *TET2* in COSMIC (observed in 1 of 387

cases vs. 0 of 557 WHISP non-cases), a somatic frameshift mutation, P333fs, was reported by Metzeler *et al.*²⁸ Another kinase domain germline variant, D837N, in *EGFR* was absent in WHISP controls but found in 5/387 ovarian cancer cases with a position matching a reported somatic mutation (D837G) in COSMIC.²⁹

Germline and somatic interactions in ovarian cancer

Since familial cancer predisposition genes are also often somatically mutated in non-familial cases³⁰, we examined previously characterized somatic SMGs (and BRCA2) that met our expression criteria for putative germline functional variants (truncation and predicted deleterious missense) in the germline data of ovarian cancer cases. As expected, a high frequency of germline truncation variants was observed in BRCA1 (n=32) and BRCA2 (n=25). We observed one germline truncation variant in NF1 (D290fs) in one case (age of diagnosis: 39 years). We similarly investigated somatically mutated protein tyrosine phosphatases and identified 8 germline truncation events in 4 genes (PTPN13, PTPRM, PTPRR, and PTPRH). Notably, 4 truncation events (two H942fs, one R199fs, and one T79fs) were found in PTPRH, a gene not previously linked to ovarian cancer (Fig. 3). Analysis of germline truncations in somatically mutated chromatin modifier genes also identified truncations in SETD4 (Y129fs), SETD6 (M264fs), MLL3 (e14-2), SMC5 (O810fs), and SMC6 (Y954*). This suggests a potential role for histone modifiers in ovarian susceptibility and motivates further study. Predicted functional germline missense variants having low frequencies were detected in several somatic SMGs, including BRCA1 (germline missense n=27), BRCA2 (n=13), NFI (n=8), RB1 (n=3), and TP53 (n=1) (Supplementary Table 3). The two patients having a germline V2148D variant in NF1, developed ovarian cancer at age 36 and 45 years.

We further investigated the interplay between germline variants (truncation and missense) and somatic mutations in ovarian cancer, discovering 18 patients with germline truncation variants and somatic mutations in the same gene (Supplementary Table 4). For instance, a patient with a germline frameshift mutation (M723fs) in *PALB2* also harbored a somatic nonsense mutation (Q378*) and another patient with a germline nonsense variant (Q153*) in *CDK5RAP1* acquired a somatic splice site mutation in that gene (e9-2). We also detected 8 patients with both germline missense and somatic mutations from the same cancer gene. This list includes 2 patients with *BRCA1* (Germline: R1347G and S1512I; somatic: E111* and G813fs), 1 *NF1* (germline: A2644G; somatic: I85fs), and 1 *TP53* (germline: G334R; somatic: P177R).

We investigated LOH in tumor samples for 535 missense variants in cancer genes and 2,214 genes having germline truncation variants (3,635) and found a total of 732 truncation variants (63 in cancer genes) that displayed LOH in the tumor samples (>20% increase of VAF over normal was used for defining LOH, considering the average 77% purity of the ovarian tumor cohort, false discovery rate = 22%, Supplementary Fig. 5 and Methods), suggesting their potential roles in ovarian cancer susceptibility (Figure 4a and 4b and Supplementary Data 12). Most notably, we observed at least a 20% increased VAF for 30/32 truncation mutations in *BRCA1* (all 32 having increased VAFs) and 13/25 in *BRCA2* (19 having increased VAFs) in the tumor samples when compared to the paired germline

samples (Figure 4c, and 4d). In *BRCA1*, 13 LOH events were associated with a loss of one copy in tumor (copy number segmentation mean 1.5), while 9 LOH events were associated with a single copy number loss for *BRCA2*. We also identified 14 *BRCA1* and 4 *BRCA2* copy number neutral LOH events in tumor samples (1.5 < copy number segmentation mean 2.5). A small number of cases carried germline truncation variants with clear evidence of somatic LOH (loss of the wild-type allele) in the tumor samples occurring in genes involved in cell-cycle checkpoint, Fanconi/DNA-repair pathways (e.g., *ATM*, *BRIP1*, *CHEK2*, *FANCA*, and *MSH3*), phosphatases (*PTPRH and PTPRM*), and a putative prostate cancer susceptibility gene, *ELAC2* (Figure 4e and Supplementary Data 12). This evidence suggests several additional genes may be associated with ovarian cancer susceptibility.

We examined LOH patterns indicating retained germline missense variants in BRCA1 Here we identified two known pathogenic missense variants, G1788V and R1699W³¹ (Supplementary Figure 4); R1699W has VAFs of 42% and 79% and G1788V has VAFs of 57% and 98% in the germline and tumor samples, respectively. For one variant of unknown significance (VUS), \$1521I, evidence indicating loss of the variant allele in the tumor was present in 3/3 cases, suggesting that S1521I is not pathogenic, in agreement with the BIC classification. Evidence of LOH was inconsistent for R1347G and R841W with 2/6 and 1/4 cases demonstrating LOH respectively. Three VUS (V772A, P1637L, L668F) identified in single cases showed LOH. The case with the V772A in BRCA1 was diagnosed with ovarian cancer at age of 49 years, however this case also carried a BRCA1 truncation variant. The case with the V1637L variant in BRCA1 also had a truncation in BRCA2 and V1637L has previously been predicted to be functionally neutral.³² For L688F that occurred in one ovarian cancer case and was not observed in the WHISP dataset no other truncation mutations were observed. None of the BRCA2 missense variants were classified as clinically important in the BIC BRCA2 database.^{31,33} Evidence of LOH for retaining some germline BRCA2 missense variants (S1172L, T2088I, K2434T, and A2951T) was observed (Figure 4d, Supplementary Figure 4, and Supplementary Data 13). The case harboring K2434T in BRCA2 was diagnosed at age of 37 years, however, further work is needed to confirm the functional relevance of such rare germline variants. We expanded our LOH analysis for all rare missense variants across cancer genes (Methods) and identified a total of 114 instances having a greater than 20% increase of VAF in the tumor compared to the germline (Figure 4d and Supplementary Data 13).

We further employed germline-somatic interaction analyses and extensive bioinformatics annotations to identify truncation and missense variants with high likelihood of having functional relevance. Specifically, we examined five aspects of each germline variant (3,635 truncations and 535 missense): pfam annotation, COSMIC/OMIM proximity match, LOH status, somatic SMG status, and somatic mutation in the same gene. When limiting our candidates to variants meeting at least two of the five criteria, the numbers of variants with putative function decreased to 302 truncation and 56 missense events, respectively. In addition, we limited our high confidence variants to genes expressed in ovarian cancer (RSEM (RNA-Seq by Expectation-Maximization) > 0.5) and those that had a lower frequency in cases than WHISP non-cases thereby obtaining 222 putative functional variants

(181 truncations and 41 missense) (Table 2 and Supplementary Data 14). After removing variants suspected to be non-pathogenic based on previous published findings (*ATM* F1463C³⁴, *BRCA1* L668F and P1637L³², *PALB2* H1170Y³⁵, *SMO*³⁶ and *TSC2*^{37,38}), the missense list includes variants from several genes including the two known pathogenic *BRCA1* variants (G1788V and R1699W), four *BRIP1* variants, three *ATM* variants, four NF1 variants, and one *TP53* variant previously identified in breast cancer³⁹ (Table 2). Notably some of the cases with variants identified through this analysis also had truncation variants in known ovarian cancer predisposition genes suggesting an alternative explanation or interacting risk alleles. Our integrated analysis of germline and somatic variants identifies a set of known ovarian cancer susceptibility variants and prioritizes a set of variants without previous association with ovarian cancer susceptibility.

Significant pathways in ovarian cancer

We performed pathway analysis using PathScan statistical test⁴⁰ including both germline truncation variants and somatic mutations and identified the KEGG Fanconi Anemia DNA repair pathway as significant (P = 4.2E-08) along with MAPK, Cell cycle, and TP53 signaling pathways (Fig. 5a and Supplementary Data 15). RB/RAS pathways were previously reported to be involved in ovarian cancer.⁴ Germline and somatic mutations in the Fanconi Anemia pathway affected a total of 40 genes in 37% (157/429) cases. Additional rare mutations detected but not shown occurred in *APITD1, EME1, ERCC1, HES1, MLH1, PMS2CL, POLK, POLI, RAD51, REV3L, RMI1, RPA1, RPA2, RPA4, TELO2, TOP3A, TOP3B, USP1*, and WDR48.

We used HotNet⁴¹ to identify subnetworks of a genome-scale protein-protein interaction network containing genes with significant numbers of somatic and germline variants. HotNet identified two such subnetworks (P < 0.01): one consisting of DNA repair and Fanconi Anemia genes (Fig. 5a and Supplementary Table 5) that is mutated in 33.1% (142/429) of samples. We combined Fanconi genes from PathScan and HotNet analyses and determined that 40.8% (175/429) of ovarian cancer patients in this study have germline/ somatic defects in the Fanconi pathway. As expected, we found that germline alterations in 47 Fanconi genes are significantly enriched in younger patients by a Wilcoxon Rank-Sum test (427 tumors with data, P-value=1.1878E-05, Fig. 5b).

A second subnetwork containing somatic mutations and germline variants in *EGFR*, *ERRB2*, *ERBB3*, and other genes is shown in (Fig. 5c and Supplementary Data 16). The frequency of somatic mutations in each of these genes is low (< 1.3%), as is the frequency of germline variants (< 0.3%). The significance of this subnetwork is thus derived from the combined analyses of somatic mutations, germline variants, and biological interactions among these proteins. Using more permissive parameters, HotNet identifies two additional subnetworks (See Methods), including a subnetwork containing *MLL*, *MLL3*, and *SETD1A* (Fig. 5d and Supplementary Data 16). Mutations in these histone methyltransferases have been previously reported in leukemias⁴², breast cancer²⁴, and renal carcinomas⁴³, but have not been widely reported in ovarian carcinoma.

DISCUSSION

We report here the first large-scale exome-wide analysis of the combined germline-somatic landscape of ovarian cancer. We used several analytic approaches to sift through millions of germline variants to discover both known and candidate cancer susceptibility genes and loss-of-function truncation and missense variants. As expected, we found enrichment of germline presumed loss-of-function truncation variants in the known ovarian cancer susceptibility genes, BRCA1, BRCA2, BRIP1, CHEK2, and PALB2. The average diagnosis age for patients with germline BRCA1/BRCA2 truncation variants was 53.4 years, significantly younger than either patients with somatic BRCA1/BRCA2 mutations (61.8 years, n=32, P = 0.0002, t-test) or the entire cohort (59.4 years, n=427, P=5.73E-06, t-test). Interestingly, patients harboring germline BRCA1/BRCA2 alterations have an average of 1.87 somatic mutations (n=60) in 127 SMGs from MuSiC analysis of 12 TCGA cancer types⁴⁴ (curated from doi:10.1038/nature12634) which is markedly lower than patients with somatic BRCA1/BRCA2 mutations (2.84 somatic mutations, n=32, P = 2.1E-05 t-test). Further, likely loss-of-function truncation variants were detected in several other genes/gene family members and syndromes (NF1) that have previously been associated with breast and/or ovarian cancer susceptibility including BLM¹⁵, FANCD2⁴⁵, NF1^{19,46}, RAD51B^{47,48}, FANCA⁴⁹, FANCB, FANCL, FANCM, ATRIP, and ATR⁵⁰. Notably, loss-of-function variants were dispersed across a set of genes, in particular, previously reported members of the Fanconi pathway⁵¹ and some novel members.

The identification of pathogenic missense variants in high-throughput sequencing data is challenging due to the large number of rare variants of unknown significance and inherent uncertainties associated with *in silico* based functional prediction. To identify a set of known and likely pathogenic missense variants, we used several complementary strategies including LOH, COSMIC/OMIM proximity match, PFAM domain, and case/control allele frequency analyses. We first applied the LOH analysis to germline truncation variants in *BRCA1* and *BRCA2* and a small set of other tumor suppressor genes, demonstrating a strong tendency to induce LOH of the wild-type allele in the tumor. For example, clear evidence for LOH of *BRCA1* wild-type alleles in the tumor was present in virtually all cases, similar to previous reports.^{3,52} Further, our analysis identified two pathogenic missense variants (G1788V, R1699W) as well as three with uncertain pathogenicity (L668F, V772A, P1637L) that demonstrated clear evidence of LOH. However, we note that the single cases with V772A and P1637L variants each had a *BRCA1* truncation variant suggesting an alternative explanation for these findings. LOH was also observed for several *BRCA2* missense variants.

Evidence for pathogenicity was also demonstrated for a number of variants in cancer genes including two pathogenic *BRCA1*, three *ATM*, and four *BRIP1* missense variants that met at least two of the five criteria for classifying candidate pathogenic missense variants. These results emphasize that integration of both somatic and protein domain information can facilitate identification of a set of known and potentially pathogenic missense variants among thousands of rare missense variants that informs functional assessment of variants of unknown significance.

Significance analysis of germline truncation and missense variants nominated a set of genes including *ASXL1*, *MAP3K1*, and *SETD2* as candidate novel ovarian susceptibility genes. COSMIC somatic mutation matches to *ASXL1* germline missense variant (G1397S) coupled with evidence for LOH support a potential role for this variant in ovarian cancer susceptibility. In addition, common variation in *MAP3K1*, another member of the MAP3K family, has been associated with breast cancer susceptibility⁵³, was recently identified as a target of frequent somatic breast cancer mutations^{24,25}, and was significant based on the burden test.

Pathway and network analyses of the integrated collection of germline and somatic variants revealed pathways with significant enrichment of variants including the Fanconi anemia/DNA repair pathway, MAPK pathway, and histone methyltransferases. In most cases, the individual genes in these pathways are altered rarely by either germline or somatic variants, and it is only through the combined analysis of both types of variants across many genes that the alteration of these pathways becomes apparent. This further emphasizes the extensive genetic heterogeneity in serous ovarian carcinoma, as suggested by the relatively small number of genes found to be recurrently mutated by somatic mutations in TCGA study.⁴

We are mindful of limitations of TCGA and WHISP data for germline analyses and the analysis of rare variants in general including lack of family history information in TCGA cases that would further inform these results, exclusion of women with a prior malignancy that required systemic therapy from the TCGA case set that might lead to an underestimation of the frequency of germline susceptibility alleles in the population, lack of personal cancer history information in WHISP controls, differences in sequencing platforms used to generate the TCGA and WHISP exome sequence data, and detection of rare germline variants that are extremely rare/private and have no pathogenic significance. With respect to differences in sequencing platforms between the case and control datasets, more variants were called in the WHISP data than the TCGA data, which would reduce our ability to detect significantly higher frequencies of rare deleterious germline variants in TCGA cases compared to WHISP controls. In addition, it is worth noting that the WHISP controls were older on average than TCGA cases and were assembled for the purpose of examining genetic susceptibility to non-cancer outcomes. Therefore pathogenic germline variants would most likely be under-represented in this cohort, which would increase our ability to identify pathogenic variants in TCGA OV cancer cases.

In conclusion, this is the first large scale and comprehensive analysis of both germline and somatic exome variants in ovarian cancer. Our exome-wide analysis strongly supports and extends results from previous studies employing candidate gene approaches for discovery of ovarian cancer genes, and is in line with previous reports by identifying Fanconi anemia pathway genes as the most frequent targets of germline and somatic mutations. Our integrated analyses of somatic and germline data indicate additional genes and variants of potential importance in ovarian cancer susceptibility for further investigation. In addition, we emphasize that candidate variants and genes nominated by our study will require extensive experimental functional validation as well as replication in additional ovarian

cancer datasets. Functionally validated variants will have important implications for the development of screening strategies to evaluate ovarian cancer predisposition.

METHODS

Study population

We obtained approval from the database of Genotypes and Phenotypes (dbGaP) to access the exome sequence and clinical data from TCGA ovarian cancer cases for this study (document #3281 Discover germline cancer predisposition variants). We selected a total of 460 ovarian cancer cases (316 cases previously reported⁴ and 144 new ovarian cases) with their germline and tumor DNA sequenced by exome capture followed by next generation sequencing on Illumina or SOLiD platforms. Of the 460 cases, 429 met our inclusion criteria of 50% coverage of targeted exome having at least 20X coverage in both germline and tumor samples. 74% of targets reached 20X coverage for 80% of breadth. Population estimates of allele frequencies were obtained from a control group of 3,505 European individuals from the National Heart, Lung, and Blood Institute (NHLBI) exome data set (https://esp.gs.washington.edu/drupal/), and from 379 European, 246 African, 286 ASN, and 181 AMR descent individuals from the 1000 genomes project.⁵⁶ The global minor allele frequencies were obtained from dbSNP release 137, based on the 1000 Genomes phase 1 genotypes for 1094 individuals, released in May 2011.

Ancestry classification using PLINK

TCGA ovarian cancer cases were classified with respect to ancestry using their SNP array data⁴ and the multi-dimensional scaling (MDS) analysis program in PLINK. (http:// pngu.mgh.harvard.edu/~purcell.plink/, version 1.07). Five clusters were used for MDS analysis. Twenty-three TCGA cases had unknown ethnicity information; we were able to assign ethnicity for 19 of these as Caucasian (n=17) and African American (n=2), respectively, using principal components analysis (Supplementary Fig. 1).

Control cohort

WHISP (Women's Health Initiative Exome Sequencing Project, part of the Women's Health Initiative (WHI)) data for 614 samples were downloaded from dbGaP (dbGaP Study Accession: phs000281.v4.p2), verified for file integrity, and then imported as BAM files into our data warehouse. The WHISP data was collected as part of the NHBLI Exome Sequencing Project that has the objective of detecting genetic variants related to heart, lung, and blood diseases as described at http://www.ncbi.nlm.nih.gov/projects/gap/cgi-bin/ study.cgi?study_id=phs000281.v4.p2. Women included in WHISP were a subset of women who were part of the WHI ⁵⁷. To minimize batch differences between the ovarian dataset and these controls, we processed imported samples through the same pipeline, including alignment to the GRCh37-lite reference sequence with BWA⁵⁸ v0.5.9 with parameters –t 4 – q 5 and marking of duplicates by Picard v1.46. SNVs and indels were called using VarScan v2.2.9 (with parameters --min-coverage 3 --min-var-freq 0.20 --p-value 0.10 --strand-filter 1 --map-quality 10) with the false-positive filter⁵⁹ and GATK⁶⁰ v5336 (with parameters -T IndelGenotyperV2 --window_size 300). Variant calls were restricted to the ~34 Mbp CDS target region.⁴

To remove outliers in data quality, we required that WHISP samples have read mapping rates <80%, duplication rates <40%, and at least 10,000 SNVs called in the target region. The 557 WHISP samples that met these criteria had, on average, mapping rates of ~95%, duplication rates of ~9%, and ~18,000 SNVs called in the target region. 81% of targets have reached 20X coverage for 80% of breadth. These were used as controls in the downstream analysis.

Germline variant calling and filtering

Sequence data from paired tumor and germline samples were aligned independently to NCBI Build 36 of the human reference using BWA 0.5.9 and de-duplicated using Picard 1.29. Germline SNPs and indels were identified in paired BAMs using VarScan2 with the following parameters: min-coverage = 30, min-var-freq = 0.08, normal-purity = 1, p-value = 0.10, somatic-p-value = 0.001, and validation = 1. Additional germline SNPs were identified using Samtools (Version 0.1.7a (revision #599) and additional germline indels were identified using GATK (Version 1.0 (revision 5336). All predicted variants were filtered to remove false positives related to potential homopolymer artifacts (variants found in homopolymers having sequence length 5 were removed), strand specific sequence artifacts, ambiguously mapped data (average mapping quality difference between the reference supporting reads and variant supporting reads 30), and low quality data at the beginning and end of reads (variants supported exclusively by bases observed in first or last 10% of the reads). Variants having an allele frequency < 8% were removed. Initial variant transcript annotation was based on a combined database, including NCBI Refseq (May 2009) and Ensembl (version 54). All variants were additionally annotated using (Version 2.2) of Ensembl variant effect predictor (VEP).⁶¹ Variants that occurred outside of tier 1 (coding exons, canonical splice sites, and RNA genes) and variants that did not change the amino acid sequence were not included in the downstream analysis. Putative variants with translational effect were filtered in the multistep process shown in Supplementary Figure 2 and described below. Variants were filtered if they either could not be mapped uniquely from NCBI build 36 to GRCh37, were protein altering in a rare transcript that was exclusive to either the NCBI or Ensembl database, or if they were nonsynonymous only in transcripts that lacked a valid open reading frame due to internal frame shifts, missing start codons, and/or missing stop codons. In addition, all variants were discarded from genes suspected to have pseudogenes or other paralogs missing from the human reference sequence, such as PDE4DIP, CDC27, MUC4, DUX4, and XPC. We additionally filtered variants that occurred exclusively in non-coding RNA genes, those that affected only predicted, hypothetical, or olfactory genes, those that had a frequency >1% in the Caucasian population in the NHLBI GO exomes sequence data, those exclusively within a transcript annotated as a pseudogene or processed pseudogene based on Ensembl release (64) annotation downloaded via Biomart, and lastly those that were reported as a validated somatic mutation in the same sample. Sequence data supporting all remaining germline truncational variants were visually examined with the Integrative Genomics Viewer⁶² and any data that appeared to be supported by potential sequencing, amplification, or alignment artifacts were discarded. Additional validated germline variants reported in BRCA1, BRCA2 were recovered, followed by removal (filtering) of any remaining nonsynonymous germline variants that were recurrent at the same position in more than 2% of the cohort (more than 8 samples at

the same position). Finally, for the analysis of significantly mutated genes, genes not typically expressed in ovarian adenocarcinoma tumor samples were filtered if they had an average RPKM 0.5. For the RNA-seq based gene expression analysis, we used the Pancan12 per-sample log2-RSEM matrix from doi:10.7303/syn1734155.1. A gene qualified as expressed if it had at least 3 reads in at least 70% of samples. For every gene, the average per-sample RSEM value was calculated across samples from the same tumor-type. The genes that had an average RSEM < 0.5% were considered to be low expressed genes. Of the 20,239 genes that had an expression value in ovarian cancer, 4,957 were low expressed genes.

Cancer gene list

The cancer gene list (Supplementary Data 17) was comprised of a total of 672 unique genes of interest that included 436 genes from the Sanger Cancer Gene⁹ list (http:// www.sanger.ac.uk/genetics/CGP/Census/ as downloaded on December 1, 2010), 41 uterine and endometrial cancer genes that we previously identified as having recurrent somatic mutations¹², and 50 genes that have been identified in genome wide association studies as containing common cancer susceptibility variants to ovarian or breast cancer (HugeNet, http://www.cdc.gov/genomics/about/index.htm). Of note, the 436 genes on the Sanger cancer gene list contained gene clusters (IGH@, IGK@, IGL@). Individual genes from these clusters were extracted. Any genes on the list that represented common fusion products of translocation or any gene that could not be identified based on Ensembl release 58 and the corresponding release of NCBI Refseq from the same time point were excluded. This process resulted in a total of 616 putative cancer related genes.

Validation of truncation variants in cancer genes

We designed validation PCR primers pairs using Primer3 and tailed the sequences with universal forward and reverse primer sites. Primer pairs for PCR were selected to favor products with an optimal size of 200 to 300 bp. (Supplementary Data 19 and 20) Larger or smaller products were allowed to avoid problematic sequences. Alternate sources of WGA-amplified or original source genomic DNA samples from tumor and normal pairs were amplified with PCR using a single primer pair and each individual PCR product was sequenced with BigDye Terminators using universal primers. Products were purified and then loaded on an ABI 3730. Resulting reads were base called using Phred and aligned to genomic sequence representative of the PCR products using Crossmatch. PolyScan⁶³ and PolyPhred⁶⁴ were used to identify SNPs and Indels. Predicted putative rare germline variants were visually reviewed using Consed to determine the exact position and sequene of indel events and eliminate false positives due to data quality, LOH in the tumor sample, artifacts resulting from sequence context, paralog amplification, or WGA or Illumina library generation or sequencing artifacts.

Missense germline variant analysis

Missense germline variants were filtered using the same methods (Supplementary Figure 3) previously described for germline truncations. To minimize the number of variants tied with ancestral origins, only missense germline variants from individuals classified as Caucasian by Plink were used for downstream significance testing. Missense germline variants were

further filtered to retain only those identified as deleterious by the Ensembl implementation of Condel, a software program that employs a weighted approach to calculate the functional impact of missense variants from scores calculated by SIFT⁶⁵ and PolyPhen-2.⁶⁶ We then removed missense germline variants that occurred at >1% frequency in the ovarian cancer cases and followed that by removing germline predicted missense variants that were better classified as somatic variants. Variants with population minor allele frequencies 1% in NHLBI ESP GO Exomes or 1000 Genomes were also filtered. Remaining sites were annotated using the Ensembl VEP instance of Condel and remaining predicted deleterious variants were retained for burden analysis. Sites were further filtered to only retain expressed variants in cancer genes (as described above). In addition, we have performed internal unbiased validation of all rare variants identified in 11 cases using available whole genome sequencing data that were independently generated. It is worth noting that whole genome sequencing data for 2 cases were generated using the SOLiD platform, furnishing orthogonal validation of the variants discovered using Illumina sequencing data. (Supplementary Data 18)

We applied a modified version of the cohort allelic sums test (CAST)¹¹ to the final list of germline missense variants in the ovarian cancer dataset to determine the statistical significance of deleterious variants in genes that were over-represented in ovarian cases vs. control exomes from the WHISP. A one-tailed CAST test was used to identify only the genes with higher burden frequency in cases than in controls.

Germline copy number alterations analysis

Segmented copy number deletion events were extracted from GISTIC (10.1073/pnas. 0710052104) analysis of Affymetrix 6.0 SNP array data for a total of 426 exome sequenced tumor-normal sample pairs with available array genotype data. Matched tumor and normal samples were processed in parallel to identify putative germline copy number variations (CNV) with overlapping deletion segments defined by 8 consecutive probes in both tumor and normal. Potentially truncating CNV deletion events in the 672 cancer-related genes list (Details at http://goo.gl/zLk8i) were extracted from the total list. Graphical plots were visually examined to identify and filter suspected artifacts and somatic copy-number events. All CNV deletion events were annotated to identify those overlapping coding exons and those that were intronic, intergenic, or affected UTR exons were removed. Matched tumor-normal exome capture BAMs were examined to identify any heterozygous SNPs refuting germline copy number deletions or, alternatively to identify coverage anomalies supporting the presence of germline deletion events. Finally, individual probe intensities were plotted and reviewed to remove additional artifacts.

Loss of heterozygosity analysis

Loss of heterozygosity (LOH) Analysis was performed by calculating the variant allele frequency of both SNV and short indels using our internally developed tool bam-readcount (https://github.com/genome/bam-readcount) for SNVs and Samtools mpileup⁶/VarScan⁷ for indels. Significance testing was done on the basis of generating an approximate empirical distribution of the actual population null distribution using a resampling method (bootstrapping with replacement). We corrected each case for tumor purity using

$$VAF_{tumor,C} = \frac{VAF_{tumor,U} - (1 - P_{tumor}) \cdot VAF_{normal}}{P_{tumor}} \quad (1)$$

where VAF_{tumor,C} and VAF_{tumor,U} are the corrected and uncorrected tumor variant allele fractions, respectively, P_{tumor} is tumor purity, and VAF_{normal} is variant allele fraction in the normal. This equation is an algebraic consequence of assuming that foreign variant and reference reads in the tumor are proportional to their corresponding numbers in the normal sample. The distribution converged within 10^8 trials (Supplementary Figure S4) and this, in turn, agreed well with another distribution model obtained by full enumeration of all possible VAF differences within the data set. A threshold of 20, i.e. Ptumor x (VAF_{tumor} – VAF_{normal}) 20%, was taken as significant and this threshold incurs a false-positive error rate of roughly $\alpha = 22\%$. The actual error rate may be slightly less because VAF differences above 50 are, strictly speaking, spurious and likely due to contamination in the normal.

Pathway analysis using HotNet

We applied HotNet⁶⁷ to identify subnetworks in a genome-scale protein-protein interaction network, each containing genes with significant numbers of somatic and germline aberrations. HotNet identifies a list of subnetworks, each containing at least *s* genes, and employs a two-stage statistical test to assess the significance of the list of subnetworks. We used HotNet version 1.1 and an interaction network from iRefIndex 9⁶⁸ containing 212,746 interactions among 14,384 proteins, using parameter t = 0.05 to derive the influence graph. With parameter $\delta = 0.02$, we find 2 subnetworks (Supplementary Table S5), each containing at least 6 genes (P = 0.0005). With parameter $\delta = 0.02$, we find 4 subnetworks (Supplementary Data 16), each containing at least 4 genes (P = 0.1555).

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Figure 1. Overview of the integrated analysis of germline and somatic variants in 429 TCGA serous ovarian cases

A total of 27,280 somatic mutations were identified, including 6 SMGs (blue shaded area). germline variants included a total of 3 BRCA1 large-scale deletions, Following filtering of variants with >1% MAF in the population, TCGA ovarian cancer cases, and WHISP controls, a total of 3,635 truncation variants, and 22,953 missense variants (17,348 in expressed genes) remained for TCGA cases. For WHISP controls a total of 10,443 truncation and 30,335 missense variants (in expressed genes) remained. After applying the burden test using WHISP exome sequence data, a total of 6 and 24 genes were significantly enriched for truncation analysis (purple shaded area) that retained variants in expressed genes in ovarian cancer that met 2 out of 5 criteria identified a total of 237 candidate germline susceptibility variants. The pathway analysis identified three significant pathways involved in ovarian cancer pathogenesis, Fanconi, MAPK, and MLL.





Shown are three germline copy number deletion variants affecting *BRCA1* in three ovarian tumor pairs. Normal samples appear above the corresponding tumor samples. Red lines indicate normalized copy number segments based on a minimum of eight probes and blue dots indicate individual probe intensities from Affymetrix 6.0 SNP arrays within the region.

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Figure 3. Lolliplots showing the distribution of germline truncation variants and somatic mutations

Somatic mutations in *BRCA1*, *BRCA2*, *PALB2*, *CHEK2*, *BRIP1*, *BLM*, *MAP3K15*, *and PTPRH* are shown in blue and germline truncation variants are in orange. Two known pathogenic *BRCA1* germline missense variants are also shown (G1788V and R1699W).



Figure 4. Loss of heterozygosity analysis in tumor samples

(a) Scatter plot displaying variant allele frequencies for all germline truncation variants in normal and tumor samples. Truncation variants in *BRCA1* and *BRCA2* are highlighted in red and blue, respectively. (b) Scatter plot displaying variant allele frequencies for germline missense variants from cancer genes in normal and tumor samples. Germline missense variants in *BRCA1* and *BRCA2* are highlighted in red and blue, respectively. (c) VAFs for the 32 samples showing LOH truncation in *BRCA1*, (d) VAFs for 25 samples showing LOH in *BRCA2*, (e) VAFs in *ATM*, *BLM*, *BRIP1*, *CHEK2*, *ERCC2*, *FANCA*, and *PALB2*. Overall, 100% (32/32) and 76% (19/25) of respective germline *BRCA1* and *BRCA2* truncation

variants showed increased VAFs in the tumor. All germline truncation variants in *BRIP1* and *CHEK2* also showed increased VAFs in corresponding tumors.





Figure 5. Significant pathways and subnetworks in ovarian cancer

(a) Oncoprint of genes with germline truncation variants and somatic mutations found in the Fanconi subnetwork identified as significant by HotNet. Genes in the iRefIndex database 58 are underlined. (b) The age distribution for patients with or without germline alterations in Fanconi genes (genes include: Panel a). The Horizontal red line indicates the median age of the group and the blue whiskers represent the age of the individual sample. (c) Oncoprint of genes with germline truncation variants and somatic mutations found in the MAPK subnetwork identified as significant by HotNet. Additional genes in the MAPK pathway with somatic mutations and/or germline truncation variants are included. (d) Oncoprint of

genes with germline truncation variants and somatic mutations found in a subnetwork including *MLL*, *MLL3*, and *SETD1A* identified as significant by HotNet. Additional chromatin modifiers with somatic mutations and/or germline truncation variants are included.

Table 1

Clinical Characteristics of TCGA cases.

	Category	No. (%)
Ethnicity ^a	Caucasians	387 (90.2)
	African American	21 (4.9)
	Asian	15 (3.5)
	American Indian	2 (0.5)
	Unknown	4 (0.9)
Survival	Living	207 (48.3)
	Deceased	218 (50.8)
	Unknown	4 (0.9)
Age	45	57 (13.3)
	46–69	267 (62.2)
	70	103 (24.0)
	Unknown	2 (0.5)
Stage	IA–IC	5 (1.2)
	IIA–IIC	20 (4.7)
	IIIA–IIIC	338 (78.8)
	IV	62 (14.5)
	Unknown	4 (0.9)

 $^a\mathrm{Number}$ assigned to each category after PCA analysis (Supplementary Fig. S1)

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	LOF ^{fg} Fanc		BRCAI	BLM/FANCI			BRCAI								BRIPI					BRCA2		BRIPI	BRCAI			
	Contol Freq ^f	0	0	0	0	0	0	0	0	0	0	0	1/557 (0.002)	0	0	1/557 (0.001)	0	0	0	2/557 (0.003)	2/557 (0.003)	0	0	0	1/557 (0.002)	1/557 (0.002)
	Case Freq^f	1/387 (0.003)	1/387 (0.003)	1/387 (0.003)	1/387 (0.003)	1/387 (0.003)	1/387 (0.003)	1/387 (0.003)	1/387 (0.003)	1/387 (0.003)	1/387 (0.003)	1/387 (0.003)	1/387 (0.003)	1/387 (0.003)	1/387 (0.003)	1/387 (0.003)	3/387 (0.008)	3/387 (0.008)	3/387 (0.008)	2/387 (0.005)	2/387 (0.005)	1/387 (0.003)	1/387 (0.003)	1/387 (0.003)	1/387 (0.003)	1/387 (0.003)
	RNA Reads	NA	1	NA	10	46	ΥN	NA	3	ΥN	ΥN	1	12	5	2	379	NA	NA	31	103	73	247	27	1	ΥN	665
	RNA VAF ^e	NA	100	NA	70	95.65	NA	NA	100	NA	NA	0	100	20	100	98.15	NA	NA	48.39	58.25	97.26	90.28	88.89	0	NA	89.68
	Exome Reads	105	57	129	81	217	292	14	56	125	52	377	436	411	205	72	101	95	63	54	22	235	75	94	95	78
	Exome VAF ^e	91.43	75.44	92.25	79.01	98.16	91.44	7.14	94.64	82.4	13.46	76.66	97.71	13.87	88.29	93.06	75.25	57.89	53.97	44.44	68.18	70.64	69.33	41.49	43.16	93.59
	HGMD pheno ^d	NR	NR	NR	Breast and Colorectal Cancer susceptibility	Ovarian Cancer	Breast Cancer	- 1	- 1	1	1	1	Breast Cancer	Fanconi anemia	1	- 1	Trichothio-dystrophy	Trichothio-dystrophy	Trichothio-dystrophy	1	1	I	1	1	-	Enstein syndrome
	HGMD ^c	NR	NR	NR	DM	DM	DM	NR	NR	NR	NR	NR	DM	DM	NR	NR	DM	DM	DM	NR	NR	NR	NR	NR	NR	DM?
	BIC^{b}	NA	NA	NA	Clinically Important	Clinically Important	Unknown	Unknown	NR	Unknown	NR	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
	r0VD ^a	NR	NR	NR	1-/2, 10 2/2, 8+/?	1-/2, 10 2/2, 8+/?	4 -/?, 2 ?/?, 1 +/?	NR	NR	NR	NR	NR	NR	1 + ?	NR	NR	NR	NR	NR	NR	NR	NR		NR	NR	NR
	Annotation	p.R2459G	p.L480F	p.P1112A	p.R1699W	p.G1788V	p.V772A	p.A1996T	p.T2088I	p.K2434T	p.F1241L	p.N370S	p.P47A	p.A349P	p.K703I	p.R1498H	p.R616P	p.R616P	p.R616P	p.A635V	p.A635V	p.G76V	p.T62M	p.R448H	p.R581W	p.R1400W
	Gene	ATM	ATM	ATM	BRCAI	BRCAI	BRCAI	BRCA2	BRCA2	BRCA2	BRCA2	BRIPI	BRIPI	BRIPI	BRIPI	CLTC	ERCC2	ERCC2	ERCC2	ERCC2	ERCC2	FRG1	HIPI	ITK	ITK	0 H M H H H H H H H H H H H H H H H H H
		-	-	-	-	-	-	-	-	-	-	-	-	-	_	-	_	-	-	_	_		_	_	_	_

Thirty-five known and candidate functional missense variants

These variants were identified using a combination of integrated germline and somatic analysis and bioinformatics annotation.

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Table 2

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LOF^{Jg} Fanconi

qf Contol Freqf	03) 1/557 (0.002)	03) 0	05) 0	05) 0	03) 0	03) 0	03) 0	05) 1/557 (0.002)	05) 1/557 (0.002)	03) 0	03) 1/557 (0.002)	03) 0	03) 0	03) 0
Case Fre	1/387 (0.00	1/387 (0.00	2/387 (0.0	2/387 (0.0	1/387 (0.00	1/387 (0.00	1/387 (0.00	2/387 (0.00	2/387 (0.00	1/387 (0.00	1/387 (0.00	1/387 (0.00	1/387 (0.00	1/387 (0.0
RNA Reads	NA	55	61	76	83	11	17	414	241	NA	157	2	5	NA
RNA VAF ^e	NA	92.73	0	0	10.84	81.82	100	49.28	92.95	NA	58.86	100	100	NA
Exome Reads	115	14	12	14	145	146	542	87	82	<i>LL</i>	388	29	35	81
Exome VAF ^e	86.96	85.71	41.67	35.71	8.28	89.04	95.2	36.78	86.59	44.16	39.95	82.76	85.71	83.95
HGMD pheno ^d		-	-	-	-	-	-	-	-	-	-	-	-	-
HGMD ^c	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR	NR
BIC^p	NA	ΝA	ΝA	ΝA	NA	NA	NA	NA	NA	NA	ΥN	ΝA	ΝA	ΥN
гоур ^а	NR		NR	NR	NR	NR	NR	NR	NR	NR	5 3/5	NR	NR	NR (IARC)
Annotation	p.D507N	p.R677H	p.V2148D	p.V2148D	p.A2644G	p.P1421L	p.R765H	p.H2032N	p.H2032N	p.I831V	p.R656W	p.P978L	p.V824L	p.G334R
Gene	6НХМ	NCKIPSD	NFI	NFI	NFI	NFI	NFI	<i>NOTCH2</i>	<i>NOTCH2</i>	RBI	RBI	RNF213	SLC4A7	TP53

BLM

0

1/387 (0.003)

37

81.08

51

76.47

E285X DM for Wiskott- Aldrich

¥ź

ΝA

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p.E285Q

WAS

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NR=Not reported, NA=Not available,

^dLeiden Open Variation Database (LOVD)⁵⁴ key: Numbers indicate number of LOVD reports. Path: Variant pathogenicity, in the format Reported/Concluded; +' indicating the variant is pathogenic, +?' probably pathogenic, '-' no known pathogenicity, '-?' probably no pathogenicity, '?' effect unknown,

 b Breast Cancer Information Core (BIC)³¹ report (*BRCA1* and *BRCA2* only),

^cHuman Gene Mutation Base (HGMD)⁵⁵ status reported pathogenicity (DM=disease causing mutation),

 d Human Gene Mutation Base (HGMD)⁵⁵ phenotype

^eVAF=Variant Allele Frequency

 $f_{\rm Freq} = {\rm Frequency}$

 ${}^{g}_{\rm Loss}$ of function truncation mutations in Fanconi Pathway