Analysis of **RAD51D** in Ovarian Cancer Patients and Families with a History of Ovarian or Breast Cancer

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**Abstract**

Mutations in **RAD51D** have been associated with an increased risk of hereditary ovarian cancer and although they have been observed in the context of breast and ovarian cancer families, the association with breast cancer is unclear. The aim of this current study was to validate the reported association of **RAD51D** with ovarian cancer and assess for an association with breast cancer. We screened for **RAD51D** mutations in **BRCA1/2** mutation-negative index cases from 1,060 familial breast and/or ovarian cancer families (including 741 affected by breast cancer only) and in 245 unselected ovarian cancer cases. Exons containing novel non-synonymous variants were screened in 466 controls. Two overtly deleterious **RAD51D** mutations were identified among the unselected ovarian cancer cases (0.82%) but none were detected among the 1,060 families. Our data provide additional evidence that **RAD51D** mutations are enriched among ovarian cancer patients, but are extremely rare among familial breast cancer patients.

**Introduction**

**RAD51** homolog D (S. cerevisiae) (**RAD51D/RAD51L3**; MIM#602954) is a component of the homologous recombination DNA repair pathway. The **RAD51D** protein forms a protein complex with **RAD51B**, **RAD51C** and **XRCC2** that binds to single stranded DNA (including single stranded gaps in double stranded DNA) and is required for the formation of **RAD51** foci in response to DNA damage [1,2]. Loveday et al [3] recently reported the identification of eight truncating mutations in **RAD51D** among 911 families with histories of breast and ovarian cancer, compared to one mutation among 1,060 population controls. They reported a significantly elevated risk of ovarian cancer (6.30, 95% CI 2.86–13.85) but did not detect a significantly elevated risk of breast cancer (1.32, 95% CI 0.59–2.96). They also reported that mutations are more prevalent in multiple case ovarian cancer families. **RAD51D** has subsequently been investigated in an additional series of 175 breast and ovarian cancer families, with an additional mutation being identified among the 51 families with at least two ovarian cancers (and among the 75 probands affected by ovarian cancer) [4]. Similarly, Pelttari et al [5] identified a splice site mutation (c.576+1G) in two breast cancer affected probands from 95 Finnish breast and/or ovarian cancer families. Pelttari et al then screened for the c.576+1G variant in an additional 2,200 breast and 553 ovarian cancer patients and overall identified 5/707 patients with a personal or family history of ovarian cancer compared to 2/2,105 breast cancer only patients/families.

Until recently, **BRCA1** and **BRCA2** were the only genes known to confer a considerable risk of ovarian cancer (in conjunction with breast cancer) with two recent studies reporting that 13.3–14.1% of unselected high grade ovarian cancers are accounted for by mutations in one of these two genes [6,7]. A further small proportion of unselected cases carry mutations in **RAD51C** [8,9]. Loveday et al [3] estimated that 0.6% of unselected ovarian cancer cases will carry **RAD51D** mutations. To validate the association of **RAD51D** mutations to ovarian cancer and assess if there is any risk for breast cancer risk, we screened all coding exons in germline DNA from an unselected cohort of 245 unselected ovarian cancer patients and **BRCA1/2**-unrelated index cases from 1,060 breast and/or ovarian cancer families. Exons containing novel, non-synonymous variants among these cases were screened in a panel of 466 cancer-naïve control samples.

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Materials and Methods

The unselected ovarian cancer cohort included 245 individuals with various histological subtypes of ovarian cancer (130 serous, 73 endometrioid, 35 mucinous, two clear cell, two granulosa cell tumours, two adenocarcinomas and one mixed mullerian tumour). These samples were obtained from patients presenting to hospitals in the south of England, UK [10]. Undocumented, verbal consent was obtained from patients as approved by the governing ethics committee at the time.

The familial cohort included 540 individuals with verified personal and family histories of breast and/or ovarian cancer who were previously assessed at the Peter MacCallum Cancer Centre Familial Cancer Centre (Australia), as well as index cases from 520 multiple case breast cancer families (with or without ovarian cancer) obtained from the Kathleen Cunningham Foundation Consortium for Research into Familial Breast Cancer (kConFab) [11]. kConFab families are recruited through Familial Cancer Centres throughout Australia and New Zealand. All families were recruited based on multiple affected, mutigenerational family and personal history of breast and/or ovarian cancer. The families fulfilled diagnostic criteria for BRCA testing, with no underlying BRCA1 or BRCA2 mutation having been identified. The ethnicity of the index cases was self-reported as Caucasian in the vast majority of cases. All individuals provided written, informed consent for genetic testing of the genetic causes of hereditary breast and ovarian cancer and subsequently tested negative for mutations in BRCA1 and BRCA2. This study was approved by the Peter MacCallum Cancer Centre Human Research Ethics Committee. In total, index cases from 1,060 families were examined in this study, including 16 with a family history of ovarian cancer only, and 303 with a family history of both breast and ovarian cancer. Of these index cases, 98 had a personal history of ovarian cancer. The remaining 741 families had a personal and family history of breast cancer only.

Cancer-naive control DNA samples were obtained from kConFab (231 age- and ethnicity-matched best friend controls) and from the Princess Anne Hospital, UK (235 Caucasian female volunteers, as described previously) [12]. kConFab control individuals provided written, informed consent. Controls from the Princess Anne Hospital provided undocumented, verbal consent as approved by the governing ethics committee at the time.

DNA for mutation screening underwent whole genome amplification (WGA) using the Repli-G amplification system (Qiagen). Ten primer pairs were designed to amplify the ten coding exons of RAD51D with amplicons ranging in size from 215–277 bp for high resolution melt (HRM) analysis (Table 1). HRM analysis and DNA resequencing were performed as described previously [13]. Variant positions were determined with reference to GenBank reference sequence NM_002878.3. Nucle-
### Table 2. Summary of germline RAD51D variants.

| Site | Nucleotide changea | Protein changeb | dbSNP (v135) | Previously described in RAD51D mutation studies? | 1000 G MAF (%)c | EVS MAF (%)d | BC families (n = 741) | BC/OC families (n = 303) | OC families (n = 303) | Controls (n = 1066) | Grantham score | Mutation Taster | MutPred | POLYPHEN-2 | SNP & GO |
|------|--------------------|----------------|--------------|-----------------------------------------------|-----------------|--------------|-------------------|----------------|-------------------|----------------|----------------|----------------|----------------|---------------|-----------|-----------|
| Truncating | | | | | | | | | | | | | | | | | |
| Ex4 | c.270_271dup | p.(Lys91Ilefs*13) | no | Yes [3] | - | - | - | - | - | 1 | - | - | - | - | - |
| Ex6 | c.556C>T | p.(Arg186*) | no | Yes [3,4] | - | - | - | - | - | 1 | - | - | - | - | - |
| Ex9 | c.803G>A | p.(Trp268*) | no | Yes [3] | - | - | - | - | - | 1 | - | - | - | - | - |
| Coding non- synonymous | | | | | | | | | | | | | | | | | |
| Ex1 | c.47T>C | p.(Met16Thr) | no | - | - | - | 1 | - | - | - | 81 | Polymorphism | 0.587 | Neutral | Benign | Neutral |
| Ex4 | c.286G>T | p.(Gly96Cys) | no | - | - | - | 1 | - | - | - | 159 | Disease causing | 0.938 | Pathological | Probably damaging | Disease |
| Ex5 | c.394G>A | p.(Ala132Ile) | rs150498754 | - | 0.05 | 0.01 | 1 | - | - | N/A | 29 | Disease causing | 0.487 | Neutral | Probably damaging | Neutral |
| Ex9 | c.793G>A | p.(Gly265Arg) | rs140285068 | Yes (control) [3,4] | - | 0.02 | 1 | - | - | - | 125 | Disease causing | 0.924 | Neutral | Probably damaging | Disease |
| Ex9 | c.796G>T | p.(Arg266Cys) | no | - | - | - | 1 | - | - | - | 180 | Disease causing | 0.621 | Pathological | Probably damaging | Neutral |
| Coding synonymous | | | | | | | | | | | | | | | | | |
| Ex2 | c.117A>T | p.(=) | no | - | - | - | 1 | - | - | - | N/A | - | - | - | - | - |
| Ex9 | c.864C>T | p.(=) | rs138557828 | - | - | 0.01 | - | - | - | 1 | - | - | - | - | - |
| Ex9 | c.879G>A | p.(=) | no | - | - | 0.01 | 1 | - | - | - | - | - | - | - | - |
| Ex9 | c.900A>G | p.(=) | no | Yes [3] | - | - | - | - | - | 1 | - | - | - | - | - |
| Non-coding | 5’UTR | c.-39C>T | p.(=) | no | - | - | - | 1 | - | - | - | - | - | - | - | - |
| IVS1 | c.82+60C>T | p.(=) | no | - | - | - | - | 1 | - | - | - | - | - | - | - | - |
| IVS1 | c.83-20T>C | p.(=) | rs182793287 | - | 0.05 | - | - | - | 1 | N/A | - | - | - | - | - |
| IVS3 | c.264-6C>T | p.(=) | no | - | - | - | 1 | - | - | - | - | - | - | - | - | - |

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*a* Exon (Ex), intervening sequence (IVS).

*b* Variant positions are reported in reference to NCBI RefSeq NM_002878.3 (mRNA) and NP_002869 (protein).

*c* In addition to the variants listed, common variants rs4796033, rs28363284 (non-synonymous) and rs9901455 (synonymous) were detected at high frequency in both cases and controls.

*d* All variants were queried against 1000 Genomes (1000 G) data using the 1000 Genomes Browser which integrates SNP and indel calls from 1,092 individuals (data release 20110521 v3). The minor allele frequency (MAF) is provided here.

*e* All variants were queried against Exome Variant Server (EVS), NHLBI Exome Sequencing Project (ESP). EVS contains SNP information from 5,379 individuals (data release ESP5400). The minor allele frequency (MAF) is provided here.

*f* Breast cancer only family (BC), ovarian cancer only family (OC), breast and ovarian cancer family (BC/OC).

*g* Only breast cancer family (BC). In addition to the variants listed, common variants rs791040, rs28463284 (non-synonymous) and rs9901455 (synonymous) were detected at high frequency in both cases and controls.

*h* All variants were queried against 1000 Genomes (1000 G) data using the 1000 Genomes Browser which integrates SNP and indel calls from 1,092 individuals (data release 20110521 v3). The minor allele frequency (MAF) is provided here.

**Note:** Variants annotated with N/A were not assessed in control samples.

[1] doi:10.1371/journal.pone.0054772.t002
otide numbering reflects cDNA numbering with +1 corresponding to the A of the ATG translation initiation codon in the reference sequence, according to HGVS guidelines (www.hgvs.org/mutnomen). All novel variants were verified by Sanger resequencing of non-WGA DNA. Tumour cells were needle dissected from 10 mm sections to obtain tumour DNA, which was subsequently whole genome amplified.

The following in silico prediction tools were used to assess the likely functional effect of the missense variants identified in this study: PolyPhen-2 [14], SNPs&Go [15], MustPred [16], PMut [17] and MutationTaster [18]. Human Splicing Finder (HSF) was used to assess the effect of all non-truncating variants on splice sites [19].

Results and Discussion

Two previously reported truncating mutations, p.(Arg186*) and p.(Trp268*) were identified among a series of 245 unselected ovarian cancer patients (0.82%). The p.(Arg186*) variant was detected in a patient diagnosed with a grade 2 papillary serous cystadenocarcinoma at 66 years of age. DNA sequence analysis of tumour tissue obtained from this tumour showed reduction of the wildtype allele consistent with loss of heterozygosity (LOH) (Figure 1). The p.(Trp268*) variant was detected in a patient diagnosed with an endometrioid carcinoma (no grade information) at 70 years of age. No tumour tissue was available for LOH analysis. No family history information is available from either case. The histology of the two ovarian cases with truncating RAD51D mutations (i.e. high grade serous and endometrioid) is consistent with the majority of mutations reported in other RAD51D studies [3,4,5,20], and with other ovarian cancers associated with mutations in double strand break DNA repair genes (e.g. BRCA1 and BRCA2), but the number of mutations in RAD51D identified to date is too few to determine the significance of this observation. A third truncating mutation, p.(Lys91Ilefs*13), was identified in one of 466 control samples (0.21%). All three of these mutations have previously been reported [3,4], and may represent founder mutations. However, there is no overlap with variants reported in more recent studies by Pelttari et al. or Wickramanyake et al. [3,20]. To date, no truncating mutations have been detected among 1,092 individuals in the 1000 genomes cohort (data release 20100521 v3) [21] or 5,379 individuals in the Exome Variant Server (release ESP5-100; NHLBI Exome Sequencing Project (ESP), Seattle, WA (http://evs.gs.washington.edu/EVS/)) [June 2012].

The absence of truncating mutations in 741 breast cancer only families (or 962 breast cancer-affected probands) provides further evidence that RAD51D mutations do not contribute significantly to breast cancer risk.

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Author Contributions

Conceived and designed the experiments: ET IC. Performed the experiments: SR. Analyzed the data: SR SS ET. Contributed reagents/materials/analysis tools: PJ GM AT DE kConFab. Wrote the paper: ET IC.

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