Targeted RNA-seq successfully identifies normal and pathogenic splicing events in breast/ovarian cancer susceptibility and Lynch syndrome genes

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A subset of genetic variants found through screening of patients with hereditary breast and ovarian cancer syndrome (HBOC) and Lynch syndrome impact RNA splicing. Through target enrichment of the transcriptome, it is possible to perform deep-sequencing and to identify the different and even rare mRNA isoforms. A targeted RNA-seq approach was used to analyse the naturally-occurring splicing events for a panel of 8 breast and/or ovarian cancer susceptibility genes (BRCA1, BRCA2, RAD51C, RAD51D, PTEN, STK11, CDH1, TP53), 3 Lynch syndrome genes (MLH1, MSH2, MSH6) and the fanconi anaemia SLX4 gene, in which monoallelic mutations were found in non-BRCA1 families. For BRCA1, BRCA2, RAD51C and RAD51D the results were validated by capillary electrophoresis and were compared to a non-targeted RNA-seq approach. We also compared splicing events from lymphoblastoid cell-lines with those from breast and ovarian fimbriae tissues. The potential of targeted RNA-seq to detect pathogenic changes in RNA-splicing was compared by the inclusion of samples with previously well characterized BRCA1/2 genetic variants. In our study, we update the catalogue of normal splicing events for BRCA1/2, provide an extensive

Key words: targeted RNA-seq, alternative splicing, inherited breast/ovarian cancer syndrome, Lynch syndrome, BRCA1/2

Additional Supporting Information may be found in the online version of this article.

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catalogue of normal RAD51C and RAD51D alternative splicing, and list splicing events found for eight other genes. Additionally, we show that our approach allowed the identification of aberrant splicing events due to the presence of BRCA1/2 genetic variants and distinguished between complete and partial splicing events. In conclusion, targeted-RNA-seq can be very useful to classify variants based on their putative pathogenic impact on splicing.

What’s new?
Hereditary familial breast/ovarian cancer (HBOC) syndrome involves numerous pathogenic variants, including variants of uncertain clinical significance (VUS). A subset of VUS, however, is suspected to influence RNA splicing, leading to the expression of potentially pathological transcript isoforms. Here, using a targeted RNA-seq approach, naturally occurring splice isoforms were described for BRCA1/2, RAD51C, RAD51D, and eight additional tumor-suppressor genes that are associated with HBOC and Lynch syndrome. The targeted RNA-seq approach also identified aberrant splicing events associated with the presence of BRCA1/2 genetic variants and successfully distinguished complete from incomplete splicing events, which is of major importance in determining pathogenicity.

Introduction
Pathogenic variants in the BRCA1/2 genes account for about 15–20% of the families with hereditary familial breast/ovarian cancer syndrome (HBOC). Recent studies have demonstrated that RAD51C and RAD51D should also be included in the genetic screening of ovarian cancer patients.1–5 Pathogenic variants in other known genes, such as PTEN, TP53, CDH1, STK11/LKB1, and NBS1 account for less than 10% of the non-BRCA HBOC families.6–9 Monoaclonal mutations in the Fanconi Anaemia SLX4 (or FANCP) gene were also found in non-BRCA families, although at an extremely low percentage.10,11 Cases of Lynch syndrome, account for 2–3% of all colorectal cancers and are associated with germline pathogenic variants in the MLH1, MSH2, MSH6 and PMS2 genes. Women are also at risk of endometrial and ovarian cancers.12

A large percentage of the sequence variants in the BRCA1 and BRCA2 genes that are detected by routine mutation screening are variants of uncertain (clinical) significance (VUS). A subset of VUS may affect splicing by disturbing the recognition of the donor and acceptor splice sites (DSS and ASS, respectively) or by disrupting intronic and exonic cis-elements necessary for the regulation of splicing.13,14 The effect of genetic variants in canonical DSS and ASS can be well predicted using in silico tools.15–17 For intronic or exonic splice enhancer/silencer elements the predictive power remains limited. To confirm or exclude an effect of variants on mRNA splicing experimental in vitro work is needed: usually, RT-PCR followed by Sanger sequencing. These experiments can be laborious and time-consuming. In addition, RT-PCR experiments are usually limited to the region containing the sequence variant of interest and thereby do not assess a putative effect of this variant on the overall splicing architecture of the mRNA. Moreover, these RT-PCR experiments often lead to the detection of non-canonical mRNA isoforms present in both HBOC patients and healthy controls.18 Genetic variants may lead to significantly changed expression of these transcripts’ isoforms and, when there is loss of the reference transcript, may be pathogenic. Known exceptions to this situation are in-frame deletion/insertion splicing events which lead to protein isoforms that retain tumour suppressor function, such as BRCA1Δ9,19,20 or BRCA2Δ12.20 When designing splicing assays, it is important to take all isoforms into account to either: a) consider them when analysing expression levels of the reference transcript, or b) target them more specifically to measure isoform-specific expression level changes. The ENIGMA consortium of investigators has recently published a comprehensive list of all naturally-occurring BRCA1/2 isoforms found by RT-PCR/capillary analysis.21,22 Such an extensive analysis remains lacking for many other human genes.

Nowadays, with the aid of RNA-seq, it is possible to analyse transcription events at an unprecedented depth.23,24 Through target-enrichment of a subset of the transcriptome, the different and even rare mRNA isoforms can be detected.25–27 Thousands of new isoforms and low abundant transcripts have been identified using this approach. Therefore, we sought to analyse in depth the naturally-occurring splicing events for a panel of tumour suppressor genes that are associated with HBOC and Lynch syndrome. We initially validated the targeted RNA-seq approach using previously published data for BRCA1 and BRCA2.21,22 Then a detailed analysis of RAD51C and RAD51D transcription was performed to obtain an extensivelist of naturally-occurring isoforms. These results were validated by capillary electrophoresis (CE) and compared to a (non-targeted) RNA-seq approach. Splicing events from lymphoblastoid cell-lines were compared to those from breast, ovarian, and ovarian fimbria tissues. Furthermore, we also assessed the diagnostic potential of targeted RNA-seq to detect pathogenic changes in RNA-splicing by the inclusion of samples with known effects on BRCA1/2-splicing.

Material and Methods
Bait design
We selected 12 genes, spanning a total region of 688,440 bp (Supporting Information Table S1). Double tiling SureSelect
baits (Agilent Technologies) were custom designed by Agilent for the regions of interest using two approaches: a) covering the known transcripts (which allows selecting baits for annotated splicing events from Agilent own data) and b) covering the genomic region (including introns and 1 Kb upstream and downstream). Duplicate baits were removed. A list of all baits is available on request.

**Cell cultures**
We used lymphoblastoid cell-lines (LCLs) from 2 BRCA1- and 2 BRCA2-mutation carriers (BRCA1:c.5467+5G>C, BRCA1: [c.594-2A+c.641A>G], BRCA2:c.8632+1G>A, BRCA2:c.9501 +3A>T) previously generated by the Kathleen Cunningham Consortium for Research into Breast Cancer (kConFab) as described elsewhere. Culture conditions and RNA isolation are described in the Supporting Information methods.

**Library preparation**
SureSelect RNA Target Enrichment for Illumina Paired-End Multiplexed Sequencing kit (Agilent; protocol version 2.2.1) was used. Briefly, 200 ng of mRNA were chemically fragmented and double-stranded cDNA was synthesized. After end-repair and dA-igation to the 3’-end of the cDNA fragments, paired-end adaptors were ligated. cDNA of about 250 bp was isolated with two rounds of clean-up with SPRI beads (AMPure XP, Agencourt) according to instructions. After amplifying the cDNA library for 11 cycles, the quality and quantity of each sample was determined with the 2100 Bioanalyzer (Agilent Technologies) and the Qubit 1.27 (Invitrogen), respectively. The prepared libraries were hybridized with the custom-designed SureSelect Oligo Capture library during 24 h at 65 °C. An amplification step of 12 cycles was used to add index tags. The quantity and quality of the samples were assessed as described above. The index-tagged sample libraries were pooled to an equimolar (4 nM) amount. 20pM were subject to cluster amplification and sequenced on a HiSeq2000 instrument (Illumina) using the TruSeq SBS kit-HS (2x100 cycles) on a single lane.

**Read alignment**
The STAR aligner (Version 2.4.1d) was used to map read pairs to *H. sapiens* reference genome HS.GRCh37 (iGenomes). The only set parameter for index construction was --sjdbOverhang 92. Explicitly adjusted parameters used in STAR include --outFilterMultimapNmax 2, --outFilterMismatchNmax 20 and --chimSegmentMin 0. Duplicate read pairs removal was performed with Picard tools (https://github.com/broadinstitute/picard). Alignment of the raw reads to specific events is described in the Supporting Information methods. Start and end positions from STAR output refer to the first nucleotide in the intron (AG|gu) and last nucleotide of the intron (ag|G), respectively.

**Non-targeted RNA-seq library preparation and mapping**
Described in the Supporting Information methods.

**Nomenclature**
The description of genetic variants follows the Human Genetic Variation Society (HGVS) approved guidelines, where c.1 (and r.1) is the A of the ATG translation initiation codon. Alternative splicing events are those incorporating splice junctions not present in the reference transcripts (BRCA1: NM_007294, lacking exon 4 as initially described, BRCA2: NM_000059, RAD51C: NM_058216, RAD51D: NM_002878). Splicing events in other genes were not annotated. The Supporting Information data provided shows the genomic positions (HS.GRCh37) of the splicing events detected. We described splicing events using the after symbols: Δ (skipping), ▼ (insertion), p (acceptor shift) and q (donor shift); see Supporting Information Figure S1. In case there is a new cassette exon we add a letter after the intron number, and we use A, B or C for the different cassette insertion events. For example, if, between exons 2 and 3 of the reference transcript, 2 cassette insertion events occur, these would be ▼2A and ▼2B. The letter designation was the same when events shared the acceptor splice site. A sub-index (skipping) or a super-index (insertion) indicates the number of nucleotides involved in the alternative event.

**Capillary electrophoresis**
Capillary electrophoresis (CE) was conducted for RAD51C and RAD51D as previously reported. CE analyses were performed in cDNAs obtained from control Lymphoblastoid cell-lines (LCLs) generated by kConFab, Tempus-stabilized (ThermoFisher) peripheral blood RNA from healthy control individuals, commercially available RNA from a non-malignant breast tissue (Clontech 636,576), and commercially available RNA from a pool of non-malignant ovarian tissues (Clontech 636,555). cDNA was amplified with various combinations of forward and FAM-labeled reverse primers spanning the full sequence of the reference transcripts (sequences are available upon request) and products were visualized with CE. In some cases, splicing isoforms were verified by automated Sanger sequencing.

**Quantitative evaluation of the targeted-enrichment RNA-seq**
Samples with known splicing events in BRCA1 (c.5467+5G>C and c.594-2A>G; 641A>G) and BRCA2 (c.8632+1G>A and c.9501+3A>T) were used for evaluation. The use of targeted RNA-seq to detect pathogenic changes in splicing was assessed taking into account the after: 1) detection of increased expression of splicing events in one sample compared to the other samples; 2) distinction of partial splicing events (variant allele still expresses the reference transcript) and complete splicing events (no residual expression of the reference transcript from the variant allele). The in-house developed QURNAS-tool (unpublished data), available at https://hdl.handle.net/10441/LY8ZQ4. A brief description of the tool is described in Supporting Information methods. Analysis of the read counts for reference exon-exon junctions, as described in the Supporting Information methods, was used to determine the expression of the reference transcript.
Results

In total, 425,665,943 reads were obtained for the 4 LCL samples. From these, 19% non-duplicate read pairs were uniquely mapped on the reference genome and about 7% of these were mapped onto the genes of interest (Supporting Information Table S2).

Performance test

Initially, we established whether the read depth of the RNA-seq experiment was sufficient to achieve our objective to obtain an extensive list of splicing events for a given gene. Therefore, we compared the list of detected BRCA1/2 splicing events in our RNA-seq data to previously published naturally occurring events (Supporting Information Tables S3 and S4). Supporting Information Figures S2 and S3 depict the splice junctions that were identified in BRCA1 and BRCA2, as well as their relative expression. Compared to previously identified/reported BRCA1 splicing events, we detected 63 out of 67 events (94%), missing 3 multicassette and 1 mixed biotype event (Supporting Information Table S3). For BRCA2, we were initially able to detect 34 out of 36 known splicing events (Supporting Information Table S4), missing the identification of a cassette and one mixed biotype event. Overall, our method allowed to detect more known events than a previous targeted RNA-seq study. Nevertheless, we did not find BRCA1 and 1 BRCA2 junctions described in that study. So, we aligned the raw reads to the already known events not found by the STAR aligner and visually inspected the outcome. The BRCA1 events described by ENIGMA were indeed not present in our samples, but new events from Davy et al., ins 2A (donor splice site) and Δ15q were identified in all 4 samples with 282 and 144 reads, respectively. Also, the three BRCA2 remaining splicing events were detected. BRCA2Δ6q9p7 contains only 2 nucleotides of exon 6, which likely caused problems for the STAR aligner. BRCA2Δ18 was found in 3 out of 4 samples (138 reads). Introns 17 contains a rare GC donor splice site, but we were able to detect the normal 17–18 exon-exon junction, as well as the 18–19 junction, and other GC-donor splice sites. The donor site of exon 20C (previously described as 20B3) was also detected with 40 reads. It is unclear why the STAR aligner did not detect these events in our data.

In addition to the previously described events, a high number of new events was detected. This created the need to set a threshold: splice junctions must be present in at least one sample with a minimum of 25 reads, independently of the number of samples in which they were observed. Using this criterion, over 20 new events were found for each gene (not described in Gencode, Ensembl, or published), as described in Supporting Information Tables S3 and S4. Since CE was shown to be very sensitive for characterization and relative semi-quantitative analysis of splicing events, we reanalysed some of the unresolved CE peaks from the previous studies, taking into account the targeted RNA-seq data. PCR products with sizes consistent with some of the newly identified events could indeed be identified. More specifically, we observed CE evidence of 6 BRCA1 and 5 BRCA2 events not previously described. Most events that were not confirmed by CE were large retention/insertion events, which give technical limitations for CE. Moreover, events in the 3’ and 5’ ends of the genes could not be tested with CE or other PCR-based methods.

RAD51C and RAD51D splicing events

Once we established that our RNA-seq experiment performed well for BRCA1/2 genes, we analysed the data for RAD51C and RAD51D genes, using the above-mentioned threshold (at least one sample with a minimum of 25 reads). Splicing events and their relative expression are depicted in Supporting Information Figures S4 and S5. We detected 46 and 36 alternative splicing events (Tables 1 and 2) with expression levels ranging from 0.02–6% to 0.05–61% of the reference RAD51C and RAD51D exon-exon junctions, respectively (see Supporting Information Methods for details on the estimation strategy). Of the alternative splicing events, 14 and 11 events detected in RAD51C and RAD51D, respectively, were not previously described in Ensembl, Gencode or Davy et al. It is noteworthy that in 3 of 4 samples a frameshift isoform of RAD51D lacking exon 3 was more abundant than the reference transcript (isoform 1) and the isoform containing a downstream alternative exon 3 (Supporting Information Fig. S5).

CE was used both as a confirmation of the RNA-seq results and to help solve intricate events. Analysis with CE enables, at least to some extent, the identification of co-occurring events, which is not directly possible using solely RNA-seq data for events that are not captured in one read. For example, alternative cassette exons which result from the combination of two splicing events can be imputed from the exact CE-sizing data. Of the new splicing events, 23/27 RAD51C (88%, 1 event was not tested) and 13/20 RAD51D (76%, 3 events not tested) were confirmed by CE. Events not evaluated are located either at the 5’ and 3’ ends of the transcripts hindering an efficient, sensitive PCR-reaction. Furthermore, CE and PCR followed by sequencing also allowed identification of combinations of multicassette exons that are not adjacent to each other in each gene; RAD51CΔ1q103+Δ3, RAD51DΔ3+Δ3A179+Δ4,5 and RAD51DΔ3+Δ3A179+Δ4,6 (Δ4,6 was found below the threshold). Possibly, other event combinations exist, but they were not extensively tested.

Some CE peaks were difficult to be associated with splicing events. Four inferred events initially only found by CE were tested by mapping the raw data to them. RAD51C5D3 was found to be present in all 4 samples with 199 reads on average. Others are either not present in our samples or we missed the prediction of the event.

We also compared our initial results with non-targeted RNA-seq data from an immortalized lymphocyte cell line, and...
| Genomic coordinates start and end | Event description | HGVS nomenclature | Biotype | Functional Annotation | Read counts | Davy et al. | G/E |
|----------------------------------|------------------|-------------------|---------|----------------------|-------------|------------|-----|
| 56,769,994 56,772,291 | 1q15Δ6 r.-11_145del | r.11_145del | donor shift | Non-coding | 1,172 | 28 | 12 | 6 | Y | Y | GE |
| 56,769,994 56,770,711 | 1q15Δ6 r.-11_145del+145_146ins145 | +563_145+862 | mixed | PTC-NMD | 62 | 0 | 0 | 2 | Y | N | G |
| 56,770,047 56,772,291 | 1q15Δ6 r.43_145del | donor shift | PTC-NMD | 3,777 | 80 | 82 | 26 | Y | Y | GE |
| 56,770,047 56,770,711 | 1q15Δ6+ r.43_145del+r.145_146ins145 | +563_145+862 | mixed | PTC-NMD | 125 | 4 | 1 | 0 | Y | Y | G |
| 56,770,150 56,772,294 | 1q103 Δ r.43_148del | mixed | PTC-NMD | 19 | 0 | 0 | 0 | N | N | G |
| 56,770,150 56,770,711 | 1q103+ Δ r.43_148del+r.145_146ins145 | +563_145+862 | mixed | PTC-NMD | 59 | 0 | 0 | 0 | N | N | G |
| 56,771,173 56,772,291 | 1q171 Δ r.145_146ins145+1_145+171 | donor shift | PTC-NMD | 272 | 7 | 1 | 0 | Y | Y | GE |
| 56,770,150 56,772,294 | 2q175 Δ r.146_148del | acceptor shift | No FS | 187 | 1 | 0 | 0 | Y | Y | GE |
| 56,774,221 56,772,673 | 3A Δ r.571_572ins571+2334_571+3,395+r.572_1131del | Terminal modification | intronic STOP+polyA | 55 | 0 | 0 | 0 | N | N | G |
| 56,774,221 56,772,673 | 3B Δ r.572_577ins572+3016_572+3,394+r.571_1131del | Terminal modification | intronic STOP+polyA | 310 | 18 | 6 | Y | Y | GE |
| 56,774,221 56,772,673 | 4 Δ r.572_652del | acceptor shift | No FS | 179 | 5 | 1 | 0 | Y | Y | GE |
| 56,774,221 56,772,673 | 5 Δ r.572_837del | cassette | PTC-NMD | 161 | 54 | 22 | 10 | 0 | 0 | Y | N | G |
| 56,774,221 56,772,673 | 6 Δ r.572_705del | cassette | PTC-NMD | 25 | 34 | 1 | 1 | 0 | 0 | Y | N | G |
| 56,774,221 56,772,673 | 7 Δ r.572_837del | cassette | PTC-NMD | 36 | 4 | 0 | 0 | Y | N | G |
| 56,774,221 56,772,673 | 8 Δ r.572_837del+4_016_837+4_089 | cassette | PTC-NMD | 881 | 327 | 29 | 24 | 5 | 1 | 0 | Y | Y | GE |
| Genomic coordinates start and end | Event description | HGVS nomenclature | Biotype   | Functional Annotation | Read counts | Davy et al. | G/E |
|----------------------------------|-------------------|------------------|-----------|----------------------|-------------|-------------|-----|
| 56,787,352 56,797,928            | ▼5D             | r.837_838ins838-178_838-131 | cassette | PTC-NMD              | 199/327     | Y           | N   |
| 56,787,352 56,801,400            | △6              | r.838_904del       | cassette | PTC-NMD              | 200         | Y           | GE  |
| 56,787,352 56,809,844            | △6,7            | r.838_965del       | multicassette | PTC-NMD | 397     | 0           | Y   |
| 56,787,352 56,809,841            | △6,7+▼8p       | r.838_965del+r.965_966ins966-3_966–1 | mixed | PTC-NMD              | 153         | Y           | Y   |
| 56,787,352 56,811,478            | △6-8            | r.838_1026del      | multicassette | PTC-NMD | 81       | 0           | Y   |
| 56,798,174 56,801,378            | △7              | r.905_965del       | cassette | PTC-NMD              | 677         | Y           | GE  |
| 56,798,174 56,809,844            | △7+▼8p         | r.905_r.965del+r.965_966ins966-3_966–1 | mixed | PTC-NMD              | 227         | Y           | G   |
| 56,798,174 56,811,478            | △7,8            | r.905_1026del      | multicassette | PTC-NMD | 612/1091 | 0           | Y   |
| 56,801,462 56,803,052            | ▼7A            | r.965_966ins965+1592_965+1,663 | cassette | PTC-NMD              | 46/39       | Y           | N   |
| 56,801,462 56,807,546            | ▼7B            | r.965_966ins966-2298_966–2,177 | cassette | PTC-NMD              | 612/1091    | Y           | G   |
| 56,807,669 56,809,841            | ▼7B,▼7B        | r.965_966ins966-2298_966–2,177+r.965_966ins966-3_966–1 | mixed | PTC-NMD              | 48          | Y           | Y   |
| 56,801,462 56,809,841            | ▼7p            | r.965_966ins966-2298_966–2,177+r.965_966ins966-3_966–1 | mixed | PTC-NMD              | 48          | Y           | Y   |
| 56,801,462 56,811,478            | △8              | r.966_1026         | cassette | PTC-NMD              | 49          | Y           | Y   |
| 56,809,906 56,811,484            | △9p            | r.1027_1032del     | acceptor shift | PTC-NMD | 21        | 1           | Y   |

Combination of individual splicing events was inferred from CE-data.

1Genomic coordinates on chr 17, human genome built GRCh37.
2PTC-NMD, premature-stop codon; nonsense mRNA-mediated decay; FS, frameshift.
3Y, event was found; N, event was not found. -: not tested.
4Events described in Gencode or Ensemble are shown with a G or E, respectively. GE is used if an event is described in both databases.
5Read counts are shown as the average or read counts in the 4 samples. 2 numbers are shown for inserted exons.
Table 2. Events detected by STAR in RAD51D with RNA-seq compared to reference NM_002878. Combination of individual splicing events was inferred from CE-data

| Genomic coordinates start and end | Description | HGVS nomenclature | Biotype | Functional Annotation | Read counts | Targeted-LCLs | LCLs | breast | fimbria | CE | Davy et al. | G/E |
|----------------------------------|-------------|--------------------|---------|-----------------------|-------------|---------------|------|--------|--------|----|-------------|-----|
| 33,448,761 33,455,841           | Δ5'-gen-5'-UTR | r.2,256 -2127del    | terminal modification | unknown     | 27           | 0              | 0    | 0      | 0      | -  | N           |     |
| 33,448,756 33,455,841           | Δ5'-gen-5'-UTR | r.2,256 -2124del    | terminal modification | unknown     | 9            | 0              | 0    | 0      | 0      | -  | N           |     |
| 33,448,309 33,446,192           | Δ5’_1         | r.1678_82del        | Terminal modification | Non-Coding  | 25           | 0              | 0    | 0      | 0      | -  | N GE        |     |
| 33,446,192 33,448,309           | Δ5’_3         | r.1678_263del       | Terminal modification | Non-Coding  | 12           | 0              | 0    | 0      | 0      | -  | N G        |     |
| 33,448,309 33,446,501           | Δ5’_5         | r.1678_480del       | Terminal modification | Non-Coding  | 10           | 0              | 0    | 0      | 0      | -  | N GE        |     |
| 33,446,192 33,447,065           | Δ1q515_1      | r.433_82del         | Terminal modification | Non-Coding  | 23           | 0              | 2    | 0      | 0      | -  | N GE        |     |
| 33,448,718 33,446,192           | Δ1q168_1      | r.86_82del          | Terminal modification | Non-Coding  | 68           | 3              | 0    | 0      | 0      | -  | Y GE       |     |
| 33,446,67 33,446,550            | Δ2,3          | r.83_263del         | multicassette       | No Fs       | 70           | 1              | 0    | 0      | 0      | Y  | N           |     |
| 33,443,501 33,446,67            | Δ2,5          | r.83_480del         | multicassette       | PTC-NMD     | 41           | 1              | 0    | 0      | 0      | Y  | N           |     |
| 33,444,129 33,446,129           | Δ3            | r.145_263del        | cassette            | PTC-NMD     | 11,531       | 166            | 14   | 2      |        | Y  | Y GE       |     |
| 33,444,57 33,446,129            | Δ3+ ▼3A179    | r.145_263del+r.263_264ins263+1664_263+1642 | cassette + cassette | No Fs | 704 | 8 | 4 | 0 | Y | Y | GE |
| 33,443,812 33,446,129           | Δ3+ ▼3B98     | r.145_263del+r.263_264ins263+1709_263+1806 | cassette + cassette | PTC-NMD | 12 | 0 | 0 | 0 | Y | N |     |
| 33,443,142 33,446,129           | Δ3,4          | r.145_345del        | multicassette       | No Fs       | 191          | 4              | 1    | 0      | 0      | Y  | Y           |     |
| 33,443,501 33,446,129           | Δ3_5          | r.145_480del        | multicassette       | No Fs       | 7,372         | 54             | 46   | 10     |        | Y  | Y GE       |     |
| 33,443,497 33,446,129           | Δ3_6p4        | r.145_684del        | multicassette+ acceptor shift | PTC-NMD | 23 | 1 | 0 | 0 | Y | N |     |
| 33,430,64 33,446,129            | Δ3_6          | r.145_576del        | multicassette       | No Fs       | 55            | 2              | 1    | 0      | 0      | Y  | Y           |     |
| 33,444,057 33,445,519            | ▼3A179        | r.263_264ins263+1664_263+1642 | cassette | PTC-NMD | 788/1795 | 52/44 | 7/5 | 1/0 | Y | Y | GE |
| 33,443,812 33,445,519            | ▼3B98         | r.263_264ins263+1709_263+1806 | cassette | PTC-NMD | 14/62 | 11/2 | 0/0 | 0/0 | N | Y |     |

(Continues)
| Genomic coordinates start and end | Description | HGVS nomenclature | Biotype | Functional Annotation | Read counts | CE | Davy et al. | G/E |
|----------------------------------|-------------|------------------|---------|----------------------|-------------|----|------------|-----|
| 33,434,142 33,443,877            | ▼3A179+Δ4  | r.263_264ins263 +1464_263 +1,642+264_365del | mixed | PTC-NMD | 20 | 0 | 0 | N | N | G |
| 33,433,501 33,443,877            | ▼3A179+Δ4,5 | r.263_264ins263 +1464_263+1,642+1.264_480del | mixed | PTC-NMD | 120 | 7 | 6 | 0 | Y | Y | GE |
| 33,434,142 33,445,519            | Δ4          | r.264_345del | cassette | PTC-NMD | 87 | 5 | 1 | 0 | Y | N | GE |
| 33,433,501 33,445,519            | Δ4,5        | r.264_480del | multicas- | PTC-NMD | 428 | 27 | 1 | 1 | Y | Y | GE |
|                                    | Δ4q17       | r.329_345del | donor shift | PTC-NMD | 33 | 3 | 0 | 0 | Y | N |
| 33,434,081 33,445,519            | Δ5          | r.346_480del | acceptor shift | PTC-NMD | 104 | 11 | 1 | 0 | Y | Y | GE |
| 33,433,301 33,434,401            | ▼6A163      | r.576_577ins576 +1,651_576+258 | cassette | PTC-NMD | 68/1278 | 0/54 | 2/2 | 0/0 | Y | N |
| 33,433,269 33,434,404            | ▼6B122      | r.576_577ins576 +137_576+258 | cassette | PTC-NMD | 1220/1278 | 45/54 | 5/2 | 0/0 | Y | Y | GE |
| 33,433,202 33,434,404            | ▼6C55       | r.576_577ins576 +204_576+258 | cassette | PTC-NMD | 89/1278 | 1/54 | 0/2 | 0/0 | Y | N |
| 33,428,385 33,434,404            | Δ7,8        | r.577_738del | multicas- | No Fs | 99 | 2 | 1 | 0 | Y | Y |
|                                    | Δ4q17       | r.329_345del | donor shift | PTC-NMD | 5/12 | 0/1 | 0/0 | 0/0 | N | N |
| 33,428,385 33,429,523            | ▼8A121      | r.738_739ins738 +629_738+749 | cassette | PTC-NMD | 5/12 | 0/1 | 0/0 | 0/0 | N | N |
|                                    | ▼8B117      | r.738_739ins738 +633_738+749 | cassette | PTC-NMD | 3/12 | 0/1 | 0/0 | 0/0 | Y | N |
|                                    | ▼9p16       | r.738_739ins739 -16_739−1 | acceptor shift | PTC-NMD | 31 | 0 | 0 | 0 | N | Y |
| 33,428,049 33,428,219            | Δ10p7       | r.904_910del | acceptor shift | FS-alternative stop | 108 | 1 | 0 | 0 | Y | Y |
| 33,427,911 33,428,219            | Δ10p15      | r.904_910del | acceptor shift | FS-alternative stop | 89 | 0 | 4 | 0 | N | Y |
| 33,353,581 33,434,219            | Δ10         | r.904_3161del | Terminal modification | To be defined | 139 | 0 | 0 | 0 | - | N | G |

1Genomic coordinates on chr 17, human genome built GRCh37.
2PTC-NMD, premature-stop codon- nonsense mRNA-mediated decay; FS, frameshift.
3Y, event was found; N, event was not found; -: not tested.
4Events described in Gencode or Ensemble are shown with a G or E, respectively. GE is used if an event is described in both databases.
5Read counts are shown as the average or read counts in the 4 samples. 2 numbers are shown for inserted exons.
It was 347 we obtained an average of 18,868 reads [9389–33,956], whereas exon-exon junctions of RAD51C and RAD51D varied among to note that the average number of reads for the reference and isoforms found in the normal breast and CE (data not shown), indicates that the lower number of splice geted RNA-seq were also found in breast or ovarian tissue by Int. J. Cancer: normal breast tissue and 10 [2-16] for normal breast tissue by the non-targeted approach (Table 3). Addi-
tions that are completely absent in the targeted RNA-seq data, but detected in non-targeted RNA-seq.

normal breast and fimbria tissues (Tables 1–3). It is important to note that the average number of reads for the reference exon-exon junctions of RAD51C and RAD51D varied among the different experiments. In the targeted RNA-seq on LCLs we obtained an average of 18,868 reads [9389-33,956], whereas it was 347 [118–484] for non-targeted LCLs, 134 [44–226] for normal breast tissue and 10 [2-16] for normal fimbria tissue. This, together with the fact that some events detected by tar-
targeted RNA-seq were also found in breast or ovarian tissue by CE (data not shown), indicates that the lower number of splice isoforms found in the normal breast and fimbria tissue is not related to tissue-specific transcription regulation, but due to lack of coverage in non-targeted RNA-seq experiments.

Interestingly, despite the lower coverage, RAD51CΔ8,9 +Δ10 and RAD51DΔ3,4+Δ5182 events were only observed in breast tissue by the non-targeted approach (Table 3). Additional 5 RAD51C events (Δ1A531; Δ1A461+Δ2p28; Δ5A57+Δ5C100; Δ5A57+Δ5D18; Δ931) were only observed in non-targeted RNA-seq of LCLs. None of these 7 events was observed after specific alignment of the raw targeted RNA-seq data for blood cells. These can be tissue-specific isoforms and/or a reflection of interindividual variability (events that are not present in one or more individuals). In CE tests, which were performed for multiple samples (average of 8 [2-32] samples), interindividual variability was observed for 54% of the splicing events. One particular event was only present in 16% of the samples. Interindividual variability was also observed among our 4 samples with targeted RNA-seq, although this was Amostly observed for lower expressed events. Yet, only the RAD51Dr.-2256..-2124del was observed in one single sample. It is noteworthy that also among the splicing events in other genes, the events observed in a single sample are a minority, i.e. 2 for CDH1 and 1 for MLH1.

DSS and ASS that gave rise to the new events detected by targeted RNA-seq in RAD51C and RAD51D were tested for in silico prediction (data not shown). Most events used a com-

Table 3. Number of individual splicing events detected by STAR for RAD51C and RAD51D per tissue type and sequencing approach

| Tissue Type | Targeted RNA-seq LCLs | Non-targeted RNA-seq |
|-------------|-----------------------|----------------------|
|             | LCLs                  | normal breast | normal fimbria |
| RAD51C      | ≥ 25 reads\(^1\) | 55            | 39         | 23        | 9 |
|             | < 25 reads\(^2\) | 13            | 10         | 1         | 0 |
|             | Not in the targeted\(^3\) | 5            | 1         | 1         | 0 |
| RAD51D      | ≥ 25 reads\(^1\) | 40            | 25         | 18        | 5 |
|             | < 25 reads\(^2\) | 13            | 6         | 2         | 0 |
|             | Not in the targeted\(^3\) | 0            | 1         | 0         | 0 |

There is a difference between the number of events described here and those shown in Tables 1 and 2, because here we count all separate splicing events as listed in the STAR output, whereas in the previous tables part of the separate splicing events were combined, e.g., to describe a cassette insertion, as imputed from CE data.

\(^1\)The events in the targeted RNA-seq are used as reference.

\(^2\)Due to the large amount of data, only events that are found by other method/tissue are taken into account.

\(^3\)Events that are completely absent in the targeted RNA-seq data, but detected in non-targeted RNA-seq.

Quantitative analysis

We also sought to investigate whether targeted-enriched RNA-seq could be used in a clinical diagnostic setting, i.e. to find clinically relevant aberrations in splicing caused by genetic variants in individual samples. For this reason, samples with previously well-characterized splicing events in either one of the BRCA1/2 genes were used. To identify putative pathogenic splicing events in RNA-seq data, it is important to be able to: 1) detect de novo or increased expression of splicing events in one sample compared to other samples using QURNAS (unpublished data); 2) know if the expression of reference transcript is decreased, by inferring loss of the reference exon-exon junctions. The latter will give an indication about partial or complete aberrant splicing events. In general, for tumour suppressor genes like BRCA1/2, complete splicing, which is characterized by the absence of reference transcript expression from the variant allele, is more likely to be pathogenic.\(^4\),\(^35\) Table 4 and Figure 1 show our results. In brief, these are in agreement with previous results obtained with conventional RT-PCR.

Sample 1, carrying variant BRCA1c.5467+5G>C, showed a strong enrichment for out-of-frame exon 23 skipping.
time of this study. The reads for the mutation-carrier and high enrichment scores are highlighted in bold.

1 The enrichment score shown is for the carrier of the mutation described in the second column.
2 Yes—the variant was previously described as pathogenic; No—the variant was previously described as non-pathogenic; Uncertain—the variant was classified as being a variant of uncertain clinical significance.
3 Event previously not detected in controls [Whiley et al, Clin Chem, 2014].
4 There are 3 other transcripts that include Δ11q (Δ9,11q; Δ9,10,11q; Δ10,11q), but RNA-seq results do not allow to distinguish them, since they are a combination of splice events, i.e., Δ9, Δ10 or Δ9,10 with Δ11q.
5 Newly described event.

(Table 4) and is accompanied by a decrease of the local reference exon-exon junctions (Fig. 1), indicating loss of the reference transcript. The deletion of this exon, which codes for the second BRCT domain, leads to a premature stop codon within the last exon. This information suggests that c.5467+5G>C could be pathogenic like other variants leading to BRCA1 frame exons 9 and 10 skipping. The latter was present in all samples (22 average reads). A recent study, which used saturation genome editing to predict the functional effects of thousands of BRCA1 variants, reports this variant as having loss of function.38 However this was based on few data and the most recent classification for this variant is that it is a class 3 (unclassified).37 A new study, which used saturation genome editing to predict the functional effects of thousands of BRCA1 variants, reports this variant as having loss of function.38 There are no other studies that confirm complete loss of the reference transcript from the variant allele. So, future studies are required to improve the classification of this variant.

Additional splicing events, previously described in this sample, did not seem to be enriched according to QURNAS. Yet, there was a slight increase in reads for BRCA1Δ22,23, accompanied by a decrease of the normal isoform BRCA1Δ22 (Table 4). QURNAS might be missing enrichment of BRCA1Δ22,23 because it is a minor event compared to BRCA1Δ23—97 reads and 8798 reads, respectively—and it was also found in the other samples (22 average reads).

Sample 2, carrying BRCA1:c.594-2A>G in cis with c.641A>G, showed two enriched events: a strongly enriched event (enrichment score = 9) leading to out-of-frame exon 10 skipping and a weakly enriched event (enrichment score = 2) leading to in-frame exons 9 and 10 skipping. The latter was present in all samples, already at a relatively high expression level. In fact, this is a major naturally-occurring alternative splicing event as
The QURNAS’ results for sample 4, carrier of BRCA2: c.9501+3A>T, indicated that out-of-frame deletion of exon 25 was the most prominent splicing event caused by the variant, with an enrichment score of 10. The intron 23 retention, previously described as a minor event occurring in this carrier, \(^{18}\) could not be confirmed. The raw read counts for the exons 24/25 junction were 16,890 (ranging from 15,966 to 23,858 in the other samples) and 28,849 (ranging from 27,518 to 33,774 in the other samples) for exons 25/26 junction. In contrast, the read counts for the aberrant boundary between exons 24/26 were only 1923 reads. Subsequent analysis of the relative expression levels of the reference exon-exon junctions confirmed that BRCA2Δ25 is incomplete, as previously demonstrated.\(^ {40,41}\) These results are compatible with the fact that c.9501+3A>T is not pathogenic.\(^ {36}\)

**Discussion**

Every gene undergoes alternative splicing, which is crucial in shaping transcriptome variation and proteome diversity. Changes in alternative splicing are also often associated with cancer. In order to recognize pathogenic splicing events, it is important to have a thorough understanding of the natural variation in splicing of expressed transcripts under healthy conditions.
conditions. Therefore, the aim of the study was to identify naturally occurring alternative splicing in transcripts from 12 tumour suppressor genes. The first task was to evaluate whether the targeted RNA-seq approach was able to identify splice junctions across the whole gene simultaneously at high-sensitivity. To accomplish this, BRCA1/2 genes were used as controls since extensive analysis of the splice isoforms repertoire of these genes was previously conducted using PCR-based techniques \(^{21,22}\) and recently also by targeted RNA-seq. \(^{31}\) Our results show that the approach used in our study was able to identify almost all previously described BRCA1/2 splicing events, i.e. 93% of the splicing events were detected with our standard analysis. Additional events, missed with STAR, were found after specific mapping. Five previously reported naturally occurring BRCA1 splicing events were not found in our four LCL samples, which seems to be due to the interindividual variability. Transcripts resulting from the combination of different splice events were also not always possible to detect. This is because the sequencing read-length often does not allow to know which events co-occur. This can be overcome with synthetic long-read sequencing (10x Genomics technology, www.10xgenomics.com), single molecule sequencing using PacBio sequencer, or sequencing of long-range PCR products using MinION nanopore sequencing, as previously reported for BRCA1. \(^{45}\) Nevertheless, it is noteworthy that the sequencing coverage used was high enough to detect additional new events occurring at low expression levels. Most genes had a sufficiently high expression (reference exon-exon junctions over 10,000 reads), except CDH1 and SLX4. As most of the events at low expression levels are probably due to stochastic effect of the splicing machinery, resulting from random combinations of splice sites and usage of weak splice and are assumed to have no biological significance, \(^{46}\) we set a threshold for the events to be described. The list of new events would otherwise be too extensive.

Once it was established that the approach used had sufficient sensitivity to detect virtually all previously known and even new BRCA1/2 alternative splicing events, RAD51C and RAD51D data was analysed. Using the above-mentioned read threshold (at least one sample with a minimum of 25x coverage), 24% and 30% of the detected events are described for the first time for RAD51C and RAD51D, respectively. The majority of these were confirmed by CE. In-frame events are of particular interest, since they do not lead to NMD and may lead to (partially) functional proteins. Within the BRCA1/2 transcripts, examples of functional isoforms (having tumour suppressor activity) are BRCA1Δ9,\(^{10,19}\) and BRCA2Δ12.\(^{20}\) In the absence of reference transcript and increased expression of these isoforms, there remains tumour-suppressor function. In contrast, BRCA1Δ16,17, BRCA2Δ3 and BRCA2Δ17 are examples of pathogenic in-frame deletions since these proteins lack important functional domains and tumour-suppressor activity.\(^{33,47,48}\) For RAD51C and RAD51D, practically all exons code for functional domains (UniprotKB, InterPro and Nextprot databases). No in-frame deletions that could lead to a functional protein were identified. Our findings for the additional eight genes analysed were similar.

In general, the frequency of alternative splicing depends on species complexity and cell type. It changes also during development and upon cellular differentiation, indicating that alternative splicing is an important cellular mechanism for the fine-tuning of gene expression both temporally and spatially.\(^{49,50}\) Therefore, RNA-seq data collected from healthy breast and fimbria tissues was analysed and compared to blood with the aim of finding different splice patterns between the tissues. The number of isoforms found in breast and fimbria was smaller than that found in LCLs. However, since we did not perform targeted RNA-seq in these tissue samples, it is not possible to make a good comparison. The mean coverage of the reference exon-exon junctions is more than 50 times larger in the targeted sequencing compared to non-targeted sequencing of LCLs. Compared to the data from breast and fimbria tissues, it is 214 and 1380 times higher, respectively. This coverage difference seen for the reference exon-exon junctions limits our conclusions about the number and type of alternative isoforms in these tissues. Similarly, publicly available data on the GTEx portal (www.gtexportal.org; version 4.1, build # 201) shows very low read numbers over reference exon-exon junctions and even lower for several known splicing events. Only sequencing at very high coverage, such as can be achieved with targeted RNA-seq, will provide sufficient insight into the different isoforms in the breast and fimbria tissues.

The samples used in our study contain BRCA1/2 variants leading to well defined aberrant splicing events which were all detected in the targeted RNA-seq data. Importantly, we could also correctly assess whether the events were complete or partial, which is crucial information to infer their pathogenicity. Therefore, targeted RNA-seq can be used to map RNA splicing for a complete locus with one test and can detect potential pathogenic splicing events in a gene, provided that the gene of interest is expressed in the available tissue. This technique can make a major contribution in the classification of variantic variants as either neutral or pathogenic, based on their effect on splicing, reducing the burden of VUS in genetic counselling.

In summary, here we describe an updated overview of the normal splicing events of BRCA1/2, and provide for the first time an extensive catalogue of normal RAD51C and RAD51D alternative splicing. We also provide an overview of normal alternative splicing for eight additional tumour suppressor genes. In-frame exon deletions that could potentially rescue protein function were not identified. The data can be further used in the design and interpretation of RNA-experiments to assess the effect of variants with a putative effect on splicing based on RNA-seq and conventional RT-PCR. Without targeted enrichment of the genes of interest, we would have not been able to detect splicing events that occur in these genes to the extent and depth that was achieved. Furthermore, we
validated our RNA-seq protocol in combination with the in-house developed QURNAS software for the identification of significant changes in splicing and developed a method to distinguish complete from partial loss of reference transcript. This is crucial information in finding aberrant splicing events caused by genetic variants and determining their clinical relevance.

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