Insertion of an SVA-E retrotransposon into the CASP8 gene is associated with protection against prostate cancer

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Received: September 3, 2015. Revised: December 8, 2015. Accepted: December 21, 2015

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**Abstract**

Transcriptional and splicing anomalies have been observed in intron 8 of the CASP8 gene (encoding procaspase-8) in association with cutaneous basal-cell carcinoma (BCC) and linked to a germline SNP rs700635. Here, we show that the rs700635[C] allele, which is associated with increased risk of BCC and breast cancer, is protective against prostate cancer (odds ratio (OR) = 0.91, \(P = 1.0 \times 10^{-6}\)). rs700635[C] is also associated with failures to correctly splice out CASP8 intron 8 in breast and prostate tumours and in corresponding normal tissues. Investigation of rs700635[C] carriers revealed that they have a human-specific short interspersed element-variable number of tandem repeat-Alu (SINE-VNTR-Alu), subfamily-E retrotransposon (SVA-E) inserted into CASP8 intron 8. The SVA-E shows evidence of prior activity, because it has transduced some CASP8 sequences during subsequent retrotransposition events. Whole-genome sequence (WGS) data were used to tag the SVA-E with a surrogate SNP rs1035142[T] (\(r^2 = 0.999\)), which showed associations with both the splicing anomalies (\(P = 6.5 \times 10^{-32}\)) and with protection against prostate cancer (OR = 0.91, \(P = 3.8 \times 10^{-7}\)).

**Introduction**

CASP8 encodes an initiator caspase that functions in the extrinsic pathway of apoptosis, controlled by death receptors of the tumour necrosis factor receptor superfamily. Somatic deletions and point mutations of CASP8 have been observed in several cancer types including colorectal, head and neck, gastric cancer and neuroblastoma. Germline sequence variants in and near the gene (encoding procaspase-8) in association with cutaneous basal-cell carcinoma (BCC) and chronic lymphocytic leukaemia (3,4). Germline sequence variants in and near different cancer types including breast cancer, melanoma, basal-cell carcinoma (BCC) and chronic lymphocytic leukaemia (3–7).

We have sequenced the genomes of 8383 Icelanders and imputed the genotypes of 150 656 chip-typed, long-range phased Icelanders and their relatives. This allows us to search for associations between over 25 million sequence variants and a broad range of phenotypes (7,8). Recently, we reported an association between variants near CASP8 and BCC. One of the BCC risk variants, rs700635[C], is strongly associated with transcriptional and splicing anomalies involving CASP8 (7). Here, we show that rs700635[C] tags an SVA-E retrotransposon that is inserted into intron 8 of the CASP8 gene and is linked to the observed splicing anomalies. Surprisingly, we find that insertion of the SVA-E is also associated with protection against prostate cancer.

**Results**

We previously found that the BCC risk variant rs700635[C] (Icelandic population allele frequency: 0.299) is associated with impaired splicing of CASP8 intron 8. A highly correlated variant (\(r^2 = 1\) in Iceland), rs1830298 has been shown to be associated with breast cancer (9) and we confirmed this effect for rs700635[C] in 5534 breast cancer patients and 309 172 controls from Iceland (odds ratio (OR) = 1.08, \(P = 0.0035\), 95% confidence interval (CI): 1.02–1.13). Given the potential for multi-cancer susceptibility conferred by this variant, we tested its association with other forms of cancer, using Icelandic Cancer Registry (ICR) records. We saw a signal suggesting that rs700635 is associated with prostate cancer (Table 1) based on 5274 cases and 97 905 controls. However the OR indicated that, unlike for BCC and breast cancer, the rs700635[C] allele appeared to protect against prostate cancer (OR = 0.92, \(P = 0.0018\), 95% CI: 0.87–0.97). We sought to confirm this by examining prostate cancer genome-wide association study chip data from the Netherlands and by single-track genotyping rs700635 in replication samples from the USA, Spain, Romania and the UK, a total of 5245 non-Icelandic prostate cancer cases and 9971 controls (Table 1). The replication sample set corresponds to an effective sample size of 6441 cases and 6441 controls, providing an estimated power of 87% to reproduce the association. The initial finding was confirmed in the replication sample set (OR = 0.90, \(P = 1.4 \times 10^{-4}\), 95% CI: 0.85–0.94). Overall evidence for the association, including the Icelandic sample, was OR = 0.91, \(P = 1.0 \times 10^{-6}\), 95% CI: 0.88–0.95 and was without significant heterogeneity (Table 1). The published (9) breast cancer risk allele, rs1830298[C] showed similar protection against prostate cancer in Iceland (OR = 0.92, \(P = 0.0018\)) as would be expected from the strong correlation between the two variants.

In blood and sun-exposed skin tissues, rs700635[C] is associated with the presence of RNA isoforms of CASP8 in which intron 8 fails to splice out correctly and contain a variant exon unique to the NM_033358 transcript. These RNA isoforms are of interest because they conceivably could lead to attenuated caspase-8 signalling (7). We analysed RNA-sequencing (RNA-seq) data from 93 samples in the Genotype-Tissue Expression (GTEx) project and found that rs700635[C] is similarly associated with a significant retention of CASP8 sequences mapping to the 5′ half of intron 8 in both normal prostate and breast tissues (Fig. 1). A modest decrease in expression of the CASP8 major exons was also shown in breast although not in prostate tissue. We also noted the presence of RNA-seq reads in the 3′ end of intron 8, as we had previously seen in sun-exposed skin (7). In order to determine whether similar splicing anomalies are present in tumours of the breast and prostate, we examined the cancer genome atlas (TCGA) RNA-seq data. Indeed, both tumour types showed highly significant retention of intron 8 sequences in association with rs700635[C] (\(P = 8.3 \times 10^{-8}\) and 1.4 × 10⁻³⁴ for breast and prostate tumours, respectively; Supplementary Material, Fig. S1).

In both our RNA-seq data and our whole-genome DNA sequencing data, we observed very low read coverage over a ∼2.8 kb region at the 3′ end of intron 8. The region concerned is highly repetitive and GC rich. Using whole-genome sequence (WGS) data from 5253 individuals (from whom the majority of samples were prepared using Illumina’s polymerase chain reaction (PCR)-free or TruSeq Nano sample preparation methods, which are less susceptible to GC bias than older HiSeq methods) we detected a copy number variation at this location, with significant higher read counts being correlated with the presence of rs700635[C] (Fig. 2A). Further examination revealed that the region contains a polymorphic insertion/deletion of a 2792 bp short interspersed element-variable number of tandem repeat-Alu

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**Human Molecular Genetics, 2016, Vol. 25, No. 5 | 1009**

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Table 1. The association of rs700635 and rs1035142 with prostate cancer

<table>
<thead>
<tr>
<th>Sample Set</th>
<th>Number of controls</th>
<th>Number of cases</th>
<th>r1035142[C] Frequency in controls</th>
<th>OR (95%CI)</th>
<th>P</th>
<th>95%CI</th>
<th>Pp = a</th>
</tr>
</thead>
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<tr>
<td>Iceland</td>
<td>5274</td>
<td>9736</td>
<td>0.589</td>
<td>0.88 (0.68-1.13)</td>
<td>0.399</td>
<td>0.89 (0.76-1.06)</td>
<td>0.0038</td>
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<tr>
<td>Netherlands</td>
<td>1445</td>
<td>2936</td>
<td>0.394</td>
<td>0.84 (0.76-0.91)</td>
<td>0.027</td>
<td>0.81 (0.74-0.89)</td>
<td>0.0054</td>
</tr>
<tr>
<td>Sweden</td>
<td>759</td>
<td>2493</td>
<td>0.541</td>
<td>0.97 (0.88-1.08)</td>
<td>0.04</td>
<td>0.94 (0.75-1.18)</td>
<td>0.190</td>
</tr>
<tr>
<td>UK</td>
<td>543</td>
<td>1291</td>
<td>0.596</td>
<td>0.99 (0.87-1.12)</td>
<td>0.031</td>
<td>0.96 (0.83-1.11)</td>
<td>0.0001</td>
</tr>
<tr>
<td>USA</td>
<td>1547</td>
<td>1289</td>
<td>0.596</td>
<td>0.94 (0.85-1.04)</td>
<td>0.053</td>
<td>0.91 (0.80-1.04)</td>
<td>0.0001</td>
</tr>
<tr>
<td>Romania</td>
<td>981</td>
<td>1123</td>
<td>0.596</td>
<td>0.97 (0.88-1.07)</td>
<td>0.031</td>
<td>0.95 (0.82-1.12)</td>
<td>0.0001</td>
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<tr>
<td>All non-Icelandic</td>
<td>5245</td>
<td>9971</td>
<td>0.596</td>
<td>0.99 (0.88-1.09)</td>
<td>0.031</td>
<td>0.91 (0.80-1.04)</td>
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<td>All combined</td>
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<td>0.596</td>
<td>0.99 (0.88-1.09)</td>
<td>0.031</td>
<td>0.91 (0.80-1.04)</td>
<td>0.0001</td>
</tr>
</tbody>
</table>

*aLikelihood ratio test for heterogeneity. bEstimated effective sample size based on 8434 cases and 8444 controls, after adjusting for genotypic control. cEstimated effective sample size based on 14 678 cases and 14 578 controls, after adjusting for genotypic control. dEstimated effective sample size based on 150 656 chip-typed, long-range phased Icelanders and their relatives (see the ‘Materials and Methods’ section). We then searched for a SNP allele or haplotype that tags the presence of the SVA-E with high confidence. We estimated effective sample size based on 150 656 chip-typed, long-range phased Icelanders and their relatives (see the ‘Materials and Methods’ section). We then searched for a SNP allele or haplotype that tags the presence of the SVA-E with high confidence.

We used DNA sequence reads with paired ends near the SVA-E insertion site breakpoints to detect the presence of the retrotransposon and assess its association with rs700635 genotype. There was a strong association between reads that cross the (empty) insertion site and the rs700635[A] allele (Fig. 2C, P < 2 × 10^{-15}). Similarly, reads that cross from native CASP8 sequence into SVA-E retrotransposon sequence were strongly associated with the rs700635[C] allele (Supplementary Material, Fig. S2). Thus, rs700635[C] is associated with protection against prostate cancer, failure to correctly splice CASP8 intron 8, and with the presence of the SVA-E retrotransposon in the intron.

We characterized the insertion site of the CASP8 SVA-E on chromosome 2 (Fig. 3). The SVA-E is inserted 20 bp to the right of the NM_033358 variant exon and extends to within 112 bp of exon 9. The insertion is flanked by a 10 bp target site duplication (TSD). Incidentally, we noted some anomalies in the correlation between reads spanning the left and the right ends of the SVA-E insertion in certain individuals. Upon further investigation, it became evident that a copy of the SVA-E has transposed out of the CASP8 gene and inserted into chromosome 19 (±6 bp around chr19:46833489, Fig. 3). It appears that when this transposition event occurred, the SVA-E transduced with it a 288 bp stretch of 3’ flanking sequence from the CASP8 gene (which, due to the orientation of the SVA-E, is to the left of the insertion site, Fig. 3). SVA elements are known to occasionally transduce 3’ flanking sequences during transposition (10). For the CASP8 element, the probable directionality of transposition was established by the observations that (i) the transduced segment exists as native chr2 sequence on copies of chromosome 2 which do not contain the SVA-E insertion, (ii) the transduced segment does not occur on chr19 except in association with the SVA-E insertion, (iii) the chr2 TSD comprises the first 10 bp of the transduced segment, (iv) on chr19, the transduced segment is followed by a short polyA tract then by a TSD comprised of native chr19 sequence and (v) the polyA tract is preceded in the transduced segment by an AATAAA polyadenylation sequence comprised of native CASP8 intron 8 sequence. Taken together, these observations suggest that a transcript arising at the 5’ end of the chr2 SVA-E failed to terminate at the internal SVA-E polyadenylation site, but rather ran on into native chr2 sequence until it encountered a fortuitous polyA signal on the reverse stand of CASP8 intron 8 sequence. The resulting RNA gained a short polyA tail then retrotransposed into the site on chr19, creating a new TSD at the insertion site. This indicates that the SVA-E insertion in the CASP8 gene has been active and capable of retrotransposition. Ongoing activity of the SVA-E might interfere with the correct functioning of the CASP8 gene.

We wanted to impute CASP8 SVA-E insertion genotypes into our set of long-range phased Icelanders (11–13) and to genotype the SVA-E in non-Icelandic populations. Even though the association between the SVA-E retrotransposon and rs700635[C] is highly significant, the correlation is not perfect (we estimate the r^2 as 0.71 in the Icelandic population). Genotyping the SVA-E directly is complicated by the chromosome 19 retrotransposition/transduction event described above. Nevertheless, by restricting genotype calls to information derived from insertion site-spanning and right end read pairs, we were able to determine SVA-E genotype likelihoods for a set of 8434 whole-genome sequenced individuals and impute these SVA-E genotypes for 150 656 chip-typed, long-range phased Icelanders and their relatives (see the ‘Materials and Methods’ section). We then searched for a SNP allele or haplotype that tags the presence of the SVA-E with high confidence. We...
identified rs1035142[T] (MAF = 0.38), which has an \( r^2 \) of 0.999 with the SVA-E insertion in the Icelandic population. In order to validate this tagging SNP and see whether the linkage disequilibrium (LD) relation holds in non-Icelandic populations, we generated a Centaurus single-track assay for rs1035142 and a PCR assay to detect the SVA-E directly (see the 'Materials and Methods' section). The presence of the SVA-E showed perfect LD with rs1035142[T] in a set of 184 individuals of European ancestry from the USA (\( r^2 = 1 \), \( P = 2 \times 10^{-16} \)). Accordingly, we concluded that rs1035142[T] is a reasonable surrogate for the SVA-E-positive (SVA-E[ pos]) genotype in populations of European ancestry.

We then re-assessed the association with prostate cancer using knowledge of the SVA-E and its surrogate SNP. As expected, the SVA-E[ pos] and rs1035142[T] genotypes gave similar association results for prostate cancer in the Icelandic population sample (\( P = 0.0069 \) and 0.0064, respectively). Inclusion of non-Icelandic replication sample genotypes yielded an overall result of \( \text{OR} = 0.91, P = 3.8 \times 10^{-7} \) (cf. \( \text{OR} = 0.91, P = 1.0 \times 10^{-5} \) for rs700635[C], Table 1). In conditional analysis, the association with rs1035142[T] remained nominally significant when adjusted for the effect of rs700635 (\( \text{P}_{\text{adj}} = 0.044 \)).

Next, we examined the association of the SVA-E and its surrogate rs1035142[T] with the transcriptional and splicing anomalies that were observed in CASP8 intron 8. Using WGS imputed genotypes and Icelandic RNA-seq data from 628 blood samples, we scanned the region surrounding rs700635 to identify variants that correlate most strongly with the retention of intron 8 sequences. The SVA-E and a cluster of correlated variants, including rs1035142, were the most significantly associated with intron retention (\( P = 2.2 \times 10^{-32} \) for the SVA-E, \( P = 6.5 \times 10^{-32} \) for rs1035142, Supplementary Material, Fig. S3A). In conditional analysis, the signal from rs700635[C] was fully explained by either SVA-E[ pos] or rs1035142[T] whereas both SVA-E[ pos] and rs1035142[T] retained significant associations when adjusted for rs700635[C] (Supplementary Material, Table S1). SVA-E[ pos] and rs1035142[T] were also better able to explain the associated reduction in

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**Figure 1.** Carriers of the rs700635[C] allele show preferential retention of CASP8 intron 8 sequences. RNA-seq data were obtained from normal prostate or breast tissue of donors who were genotyped for rs700635. (A) Median normalized read counts (normalized for each individual to the total number of aligned reads) from prostate RNA were stratified by rs700635 genotype and plotted over the genomic region covering the coding exons of CASP8. Data are from \( n = 6 \) C/C homozygotes (shown in red), \( n = 12 \) A/C heterozygotes (shown in green) and \( n = 19 \) A/A homozygotes (shown in blue). The X-axis is the genomic position in Mb (GRCh37/hg19). The structure of RefSeq CASP8 transcripts in the region is shown beneath the X-axis. (B) Zoom showing prostate RNA median normalized read counts by genotype over the region extending from the 3' end of intron 7, through the 5' region of intron 8 up to the NM_033358 variant exon. (C) Similarly plotted median normalized read counts from breast RNA. Data are from \( n = 7 \) C/C homozygotes, \( n = 27 \) A/C heterozygotes and \( n = 22 \) A/A homozygotes. (D) Zoom showing breast RNA median-normalized read counts over a region encompassing the 5' region of CASP8 intron 8. (E) Association between rs700635 genotype and prostate RNA-seq read count in intron 8. For each individual, the median of normalized RNA-seq read counts overlapping the 5' region of intron 8 [chr2:202141827-202146555 (GRCh37/hg19)] was determined. Association with genotype was then determined by linear regression against rs700635 variant allele count. \( P = 3.8 \times 10^{-8} \) and \( \beta = 0.33 \) (SD units). (F) Similarly determined association between rs700635 genotype and prostate RNA-seq read count over all major coding exons of CASP8. \( P = 0.678 \) and \( \beta = -0.07 \). (G) Association between rs700635 genotype and breast RNA-seq read count over all major coding exons. \( P = 0.0475 \) and \( \beta = -0.30 \).
expression of the major CASP8 isoforms observed previously (Supplementary Material, Table S1). However, when we carried out a similar analysis of blood RNA expression microarray data using a probe from the NM_033358 variant exon (14), the strongest expression quantitative trait locus mapped significantly better to rs700635 (Supplementary Material, Fig. S3B, Supplementary Material, Table S1). The reason for this discrepancy is not clear, but it suggests that genetic factors other than the SVA-E might affect expression of the NM_033358 variant exon as measured by the expression microarrays.

We investigated potential mechanisms by which the SVA-E might interfere with correct splicing of CASP8 intron 8. SVA-E element insertions have been observed previously to create anomalous splice junctions (15). However, in CASP8 RNA-seq data from blood, we did not detect any new splice sites associated with the presence of the SVA-E. No active splice donors or acceptors were detected within the body of the SVA-E.

We had previously shown that at least some of the RNA sequences from intron 8 have paired ends spliced from upstream CASP8 exons, which establishes that their transcriptional orientation is on the sense strand (7). Now, we examined more directly the possibility that the SVA-E might generate antisense transcripts reading through the 5’ region of intron 8. We carried out RT-PCR on blood RNA using orientation-specific primers in intron 8 for first-strand synthesis, followed by digital droplet PCR. In samples where the SVA-E is present, sense transcripts through the 5’ region of intron 8 were over 20-fold more abundant than antisense transcripts (Supplementary Material, Fig. S4). Moreover, normalized sense transcription through the region was significantly related to the presence of the SVA-E ($P = 8.8 \times 10^{-7}$). We also detected substantial sense-strand transcription in the 3’ end of intron 7 (Supplementary Material, Fig. S4, see also Fig. 1), raising the possibility of cryptic promoter activation. However, intron 7 transcripts did not vary in association with SVA-E genotype in blood RNA (Supplementary Material, Fig. S4F). Nevertheless, we can conclude that intron 8 RNAs whose abundance is increased in association with the SVA-E insertion are in the sense orientation with respect to the CASP8 gene. This is in agreement with our conclusions from the RNA-seq data.

**Discussion**

A strength of our association studies based on WGS is that they give a reasonably complete view of all SNPs and small indels present in Iceland down to a frequency of $\sim 0.05\%$ (8). Moreover, the sequence data make possible a detailed analysis of associated
structural variants, as we have done here. Using these methods, we have found that the insertion of an SVA-E retrotransposon into the CASP8 gene is associated with transcript anomalies and protects against prostate cancer. SVA retrotransposons are relatively young in evolutionary terms and the SVA-E family is found only in humans, where it is thought to be highly active.

Figure 2. rs700635[C] is associated with insertion of an SVA-E retrotransposon into the CASP8 gene. (A) The DNA sequence coverage of the CASP8 intron 8 region was normalized and the median coverage for each genotype was plotted at each position. A copy number variation was detected in the region chr2:201281936-201284718 (GRCh38/hg38). Linear regression of mean normalized coverage by SNP genotype showed a significantly higher copy number over this region in association with rs700635[C] \( (P < 2 \times 10^{-16}) \). (B) The general structure of an SVA element: these non-autonomous retrotransposons consist of a CCCTCT repeat region (which may have RNApolII promoter activity), followed by a series of Alu-like repeats, followed by a VNTR region, followed by a degenerate env gene fragment and long terminal repeat from an extinct endogenous human retrovirus (HERV-K). Transcription in the SVA may utilize a polyadenylation signal within the long terminal repeat, or it may continue into flanking chromosomal sequences (see text). [Image from (17), used with permission.] (C) Boxplot showing association of DNA-sequence read pairs spanning the SVA insertion site with rs700635 genotype: read pairs were selected such that the forward read was in CASP8 sequence within an interval of 1000 bp to the left of the SVA-E insertion site and the paired reverse read was in CASP8 sequence within an interval of 1000 bp to the right of the SVA-E insertion site. Such reads are expected to occur only if the SVA-E retrotransposon is absent. Counts of qualifying reads were normalized and plotted by genotype. Association was assessed by linear regression of read count versus rs700635 genotype \( (P < 2 \times 10^{-16}) \).

Figure 3. The SVA-E insertion site in the CASP8 gene on chromosome 2 and the retrotransposition-transduction event on chromosome 19. Note that all structures are drawn to preserve the genomic orientation of the CASP8 gene and the chr2 SVA-E insertion. Thus, the 5\' end of the SVA-E is to the right in the diagram and chr2 sequences are from the genomic forward strand. (A) The chr2 SVA-E (pale blue bars) is inserted into CASP8 intron 8 between the NM_033358 variant exon and exon 9 (at chr2:201281936-201284718). A copy of the SVA-E is present on chr19 (green bars). The chr19 SVA-E is flanked by a transduced 288 bp segment, including the NM_033358 variant exon, from chr2 (darker green bar). (B) The structure of the chr2 breakpoints: the left panel shows the breakpoint between the transduced sequence and native CASP8 intron 8 sequence. The central panel shows the left end of the SVA-E insertion on chr2 (blue bars) with the TSD sequence indicated. The sequence of this junction is the same on the chr19 transduced copy (green bars). The right panel shows the breakpoint between the right end of the SVA-E and native CASP8 intron 8 sequence with the TSD indicated. (C) The structure of the breakpoint between the transduced sequence and native chr19 sequence: the transduced CASP8 intron 8 sequence contains a polyadenylation signal on the reverse strand (appearing here as TTTATT). Moving leftwards on the reverse strand, this is followed by a short polyA sequence (hatched green bar) then a TSD derived from native chr19 sequence. (D) The breakpoint between the 5\' end of the SVA-E and native chr19 sequence, with the TSD sequence indicated. (E) The structure of the chr19 insertion site when the SVA-E is not present. Note that here the TSD sequence is unduplicated.
SVA insertions have been seen to cause gene disruption and consequent disease (17). For example, the insertion of an SVA-E into the 3′ UTR of the FKTN gene disrupts the gene by creating an anomalous splice junction, leading to Fukuyama muscular dystrophy (15). SVA elements oriented on the sense strand of the genes into which they insert generally cause gene disruption by this mechanism, or by the insertion of new promoters or polyadenylation signals (17, 18). In CASP8, however, the SVA-E is inserted in the reverse orientation with respect to the gene and so it may have a less extreme effect. At least one case of an SVA element producing antisense RNA has been documented before. In the SYNE2 gene, an intronic SVA-D appears to utilize an internal promoter to produce antisense transcripts that terminate at a polyadenylation signal in the adjacent intron (19). It is not known whether this SYNE2 antisense RNA is pathogenic. Our study found that the CASP8 SVA-E does not produce substantial amounts of antisense RNA running through the 5′ region of intron 8. The transduction events we observed suggest that such antisense RNAs, if produced, are prone to terminate at the polyadenylation site in CASP8 intron 8 non-coding strand sequence. If, as seems likely, the CASP8 SVA-E causes the splicing and transcripational anomalies that we observe, then mechanisms other than antisense transcription from the inserted element are probable.

Variants that confer risk of some cancers and protect against others have been observed before, for example in the TERT-CLPTM1L locus (20). Satisfactory mechanistic explanations have not been forthcoming. In addition to its apoptotic functions, CASP8 has pleiotropic roles in necroptosis, inflammation, cell survival and motility (1,2,21). The effects of disturbance of CASP8 pathways might, therefore, vary between different tissue and tumour types. The CASP8 SVA-E retrotransposon described here is associated with increased risk of BCC and breast cancer, but decreased risk of prostate cancer. Yet the associated CASP8 transcriptional anomalies are similar in all three corresponding normal tissues and in breast and prostate cancers. This raises provocative questions about the mechanism through which CASP8 expression changes might be involved in the risk state for these three diseases.

Materials and Methods

Subjects

Approval for the Icelandic study was granted by the Icelandic National Bioethics Committee and the Icelandic Data Protection Authority. All participants provided informed consent. Affected individuals were identified through the ICR, which has maintained records of prostate cancer, breast cancer and melanoma diagnoses since 1955. Controls consisted of individuals selected from other ongoing association studies at deCODE who did not have a diagnosis of the cancer in question recorded in the ICR. In the Icelandic and non-Icelandic studies, it is not possible to rigorously exclude that some male controls might have had sub-clinical prostate cancer. Therefore, the effects we report could be somewhat underestimated.

The Dutch prostate study population consisted of two recruitment-sets of prostate cancer patients; Group A consists of 360 hospital-based patients recruited between January 1999 and June 2006 at the urology outpatient clinic of the Radboud university medical center (Radboudumc); Group B consists of 1085 cases recruited between June 2006 and December 2006 through a population-based cancer registry held by the Netherlands Comprehensive Cancer Organization. The average age at diagnosis for patients in Group A was 63 years (median, 66 years; range, 43–83 years). The average age at diagnosis for patients in Group B was 65 years (median, 66 years; range, 43–75 years). The control individuals were cancer free at recruitment and matched for age. They were recruited from within a population-based survey in which 9371 individuals participated from a total of 22 500 age and sex stratified, randomly selected inhabitants of Nijmeegen initially contacted. All participants were of self-reported European descent. The study protocol was approved by the Institutional Review Board of Radboud University and all study subjects gave written informed consent. Samples were genotyped on Illumina HumanHap300, HumanHapCNV370, HumanHap610, 1 M or Omni-Quad bead chips at the deCODE genetics facility.

The Spanish study population consisted of prostate cancer cases recruited from the Oncology Department of Zaragoza Hospital in Zaragoza, Spain, from June 2005 to September 2008. All patients were of self-reported European descent. Clinical information including age at onset, grade and stage was obtained from medical records. The average age at diagnosis for the patients was 69 years (median, 70 years) and the range was from 44 to 83 years. The control individuals were recruited at the University Hospital in Zaragoza and were cancer free at the time of recruitment. Study protocols were approved by the Institutional Review Board of Zaragoza University Hospital. All subjects gave written informed consent.

In the UK, prostate cancer cases and controls were recruited within the ‘Prostate Testing for Cancer and Treatment’ trial (ProtecT) (22) and biological material collection was carried out within the ‘Prostate Cancer: Mechanisms of Progression and Treatment’ (ProMPT) study. Men aged 50–69 years were offered an appointment for counseling and prostate-specific antigen (PSA) testing. Recruitment took place at nine sites in the UK; 94,427 men agreed to participate in the study (50% of men contacted) and 8807 (−9%) had a raised PSA level. Of those with raised PSA levels, 2022 (23%) were diagnosed with prostate cancer; 229 men (−12%) had locally advanced (T3 or T4) or metastatic cancers, the rest having clinically localized (T1c or T2) disease. Men with a PSA level of ≥20 ng/ml were excluded from the trial. For the present study, 547 patients with PSA values >3 ng/ml and diagnosed with prostate cancer after undergoing a needle biopsy (average age at diagnosis is 63 years). Controls were 1889 men of average age 62 years with PSA values between 3 and 10 ng/ml, but not diagnosed with prostate cancer after undergoing a needle biopsy or with PSA values <3 ng/ml (and not biopsied). Ethical approvals for the ProtecT and ProMPT studies were obtained from Trent Multi-Centre Research Ethics Committee (MREC/01/4/025 and MREC/01/4/061). All subjects gave written informed consent prior to inclusion in the study.

In the UK, prostate cancer cases were recruited from the Pathology Core of Northwestern University’s Prostate Cancer Specialized Program of Research Excellence from May 2002 to May 2010. The average age at diagnosis for the patients was 60 years (median, 59 years) and the range was from 39 to 87 years. The controls were recruited as healthy control subjects for genetic studies at the University of Chicago and Northwestern University Medical School, Chicago. All patients and controls were of self-reported European descent. Study protocols were approved by the Institutional Review Boards of Northwestern University and the University of Chicago. All subjects gave written informed consent.

The Romanian study population consisted of prostate cancer cases recruited from the Urology Clinic ‘Theodor Burghel’ of The University of Medicine and Pharmacy ‘Carol Davila’ Bucharest, Romania, from May 2008 to November 2009. The average age at diagnosis for the cases was 70 years (median, 71 years) and the

Human Molecular Genetics, 2016, Vol. 25, No. 5
range was from 46 to 89 years. The controls were recruited at the General Surgery Clinic ‘St Mary’ and at the Urology Clinic ‘Theodor Burghéle’ of The University of Medicine and Pharmacy ‘Carol Davila’ Bucharest, Romania. The average age for controls was 60 years (median, 62 years) with a range from 19 to 87 years. The controls were cancer free at the time of recruitment. All subjects were of self-reported European descent. Study protocols were approved by the National Ethical Board of the Romanian Medical Doctors Association in Romania. All subjects gave written informed consent.

Genotyping and association analysis

Procedures for whole-genome sequencing, Illumina SNP chip genotyping, imputation and association testing have been described previously (7,8,23). In this study, we used WGS information from 8338 Icelanders to impute the genotypes of 150 656 Illumina chip-typed, long-range phased Icelanders and their ungenotyped relatives. Single-track genotyping was done using Centaurus assays (24). All genotyping was carried out at the Decode genetics facility with standardized quality control and allele calling protocols. We estimated the effective sample size for each cohort as $N_c$ cases and $N_c$ controls with $N_e = 2 \times N_c \times N_p/(N_p + N_c)$, which adjusts for the asymmetry in the number of cases, $N_c$, and controls, $N_p$. For Iceland we further adjusted $N_e$ for the relatedness of cases and controls by dividing by the genomic inflation factor $\lambda$ ($\sim 1.22$ for prostate cancer).

RNA-seq analysis

RNA sequencing reads for 628 Icelandic blood samples were aligned to Homo sapiens GRCh38/hg38 using TopHat (25) version 2.0.12 with a supplied set of known transcripts in GTF format (RefSeq hg38). TopHat was configured such that it attempts first to align reads to the provided transcriptome then, for reads that do not map fully to the transcriptome, it attempts to map them onto the genome. Read mapping statistics used for read count normalization were calculated using samtools version 1.1 (http://www.htslib.org). Mapped RNA-seq read data for 37 prostate and 56 breast tissues from the GTEx Release phs000424.v4. p1 from 17.01.2014 were downloaded from the GTEx consortium (26) (http://www.gtexportal.org/home/) via dbGAP. TCGA data from breast and prostate tumours were obtained from GHub (27). We used 1056 breast cancer and 374 prostate cancer samples from the same numbers of participants, after filtering for those who had genotype data for rs6743068 (a surrogate marker of rs700635) from corresponding samples.

Determination of association with DNA-seq coverage over the SVA insertion region

In order to generate a plot of the median coverage of DNA sequencing over the SVA-E insertion region, we counted the number of reads covering each position in the interval chr2:201230000–201290000 (GRCh38/hg38). We counted only reads with an average PHRED-scaled base calling quality of $\geq 20$, those that were not marked as a duplicate, not clipped by $\geq 10$ base pairs at both ends, and only the primary alignments. We computed the mean of local read counts over two intervals near-by the SVA-E insertion site chr2:201230000–201280000 and chr2:201285000–201290000 and used this mean for normalization within each individual. We then computed the median of the normalized read counts at every position in the interval chr2:201230000–201290000 for each set of individuals with the same SNP genotype. The resulting median normalized read counts are shown in the plot in Figure 2A. In order to assess the association between normalized read count and rs700635 genotype, the mean of the normalized coverage over the SVA-E insertion interval (chr2:201281936–201284718) was determined for each individual and then regressed against rs700635[C] allele count (with the sequencing platform included as a covariate). Analysis was based on 5253 whole-genome sequenced individuals taken from an earlier data freeze than the 8338 individuals on whom the disease-association testing was based.

Determination of association between SNP genotype and DNA sequence read pairs that detect the presence or absence of the SVA-E retrotransponson

We used three strategies to assess the presence or absence of the CASP8 SVA-E retrotransposon from whole-genome sequence data. Firstly, we considered read pairs that span the SVA-E insertion site, which should be observed only in the absence of the retrotransposon. Read pairs were counted if the forward read was in CASP8 sequence within an interval of 1000 bp to the left of the SVA-E insertion site and the reverse read was in CASP8 sequence within an interval of 1000 bp to the right of the SVA-E insertion site. Counts of qualifying reads were then normalized to the mean of local read counts as described above. The boxplot in Figure 2B shows these normalized read counts as a function of genotype. Secondly, we considered read pairs that cross from CASP8 sequence into the left end of an inserted SVA-E element. In this case, read pairs were counted if the forward read was in CASP8 sequence within an interval of 1000 bp to the left of the SVA-E insertion site and the reverse read was in the SVA-E element within 1000 bp of its left end. (Note here that we define the ‘left end’ of the SVA-E element as its 3' end, because the retrotransposon is inserted in an inverted orientation relative to the CASP8 gene.) Counts of qualifying reads were normalized and plotted by genotype as above. Thirdly, we considered read pairs that cross from the right end of an inserted SVA-E element into CASP8 sequence. In this case, read pairs were counted if the forward read was in the SVA-E element and within 1000 bp of its right end and the reverse read was in CASP8 sequences within an interval extending 1000 bp to the right of the insertion site. Counts of qualifying reads were normalized and plotted by genotype as above. We assessed the association between qualifying read count and genotype by linear regression of normalized qualifying reads against rs700635[C] allele count (with sequencing platform and insert size included as covariates).

The chromosome 19 transduction event was identified by reads crossing from the left end of the transduced chr2 sequence into native chr19 sequence (see Fig. 3). Subsequently, we identified a linked indel (at chr19: 46833095, GRCh38/hg38), which can be used reliably to detect the presence of the chr19 transduction event. We estimated a frequency of $0.213$ for the chr19 transduction event in the Icelandic population. We also noted the presence of an SVA-E carrying transduced CASP8 intron 8 sequence on an unmapped contig likely originating from chromosome 22 (chr22_K1270733v1_random). Due to its insertion into repetitive genomic DNA sequence, this transduction event could not be characterized further.

Genotyping the SVA-E

We determined genotypes of the SVA-E using methods outlined in (28). Briefly, each individual may have one of three genotypes: SS, a homozygous carrier of the SVA-E element; SD, a
heterozygous carrier of the SVA-E and the deletion or DD, a homozygous carrier of the deletion. For each individual i, the input to the genotyping consists of two counts of reads pairs, a count Bi of read pairs that span from the SVA-E across the right breakpoint and a count Ci of reads that span the (empty) insertion site. We compute the relative likelihood of the genotypes as:

\[ L(B_i, C_i|SS) = E^{B_i} \]
\[ L(B_i, C_i|SD) = 1/2^{(B_i+C_i)} \]
\[ L(B_i, C_i|DD) = E^{C_i} \]

Here, E is a constant, which we set to E = 0.001. Genotype likelihoods were determined for a set of 8434 whole-genome sequenced Icelanders. The genotypes were then imputed into a set of 150, 656 Icelanders using methods described previously (23).

PCR assay for the SVA-E

Two assays were employed, one which detects the presence of the SVA-E and a second which detects the empty insertion site. The forward primer was the same for both assays: AATGGGGGAG GGAAGAGAG. This primer is located to the left of the 288 bp transduced region in CASP8 intron 8. For the SVA-E-present assay, the reverse primer was CTGTGACCTGGCAATCC, located within the left end of the SVA-E element. For the SVA-E-absent assay, the reverse primer was TGTTGCCATGTTGTTG, located in CASP8 sequence to the right of the insertion site. PCR products were resolved on agarose gels and scored manually.

Expression microarrays

Samples of total RNA from human peripheral blood (N = 1001) were hybridized to Agilent Technologies Human 25 K microarrays as described previously (14). We quantified expression changes between two samples as the mean logarithm (log_{10}) expression ratio (MLR) compared with a reference pool RNA sample. In comparing expression levels between groups of individuals with different genotypes, we denoted the expression level for each genotype as 10^{average MLR}, where the MLR is averaged over individuals with the particular genotype. We determined the significance by regressing the MLR values against the number of variant alleles carried. We took into account the effects of age, gender and differential cell-type count in blood as explanatory variables in the regression.

Strand-specific quantitative RT-PCR

Total RNA was isolated from whole blood using Qiagen RNA Maxi kits and the concentration and quality of the RNA was determined with Agilent 2100 Bioanalyzer (Agilent Technologies). Orientation-specific first-strand cDNA was synthesized using the High Capacity cDNA Reverse Transcriptase kits (Life Technologies, Inc.). Primers were intron 8 forward: CCAGCAACTA TTTTTATCTACTAC, intron 8 reverse: TGTAGGCTCTCTGG TTTAATTAT, intron 7 forward: CAGAGGCCAGGTATG, intron 7 reverse: CACACAGGACTCGACTCTCTGT, each used in separate reactions (500 nM final concentration). Quantitation was then carried out by digital PCR using both the forward and reverse primers and fluorescent labelled probes for intron 8: 6FAM-ACAGGGATTTGTCTTTC-MGB and for intron 7: 6FAM-ACACTGACCTCAAGATGG-MGB (Life Technologies) on a Bio-Rad QX200™ AutoDG™ Droplet Digital™ PCR System.

Supplementary Material

Supplementary Material is available at HMG online.

Authors’ Contributions


Acknowledgements

The authors thank those men with prostate cancer and the subjects who have donated their time and their samples that were used in this research. We thank G.H. Olafsdottir of the Icelandic Cancer Registry for assistance in the ascertainment of affected individuals, and Hjalti Sigurdsson for help with the graphics. We thank Haig H. Kazazian for permission to reproduce his drawing in Figure 2B.

Conflict of Interest statement. All authors from deCODE are employees of the biotechnology company deCODE genetics, a subsidiary of AMGEN.

Funding

The UK arm of the study acknowledges support from the National Cancer Research Institute (NCRI) for the ProMPT study [National Institute of Health Research (NIHR) collaborative study grant G0500966/75466]. The NCRI is formed by the Department of Health, Medical Research Council and Cancer Research UK. The collaborations are supported by the University of Cambridge, Cancer Research UK and the NIHR-funded Cambridge Bio-medical Research Centre, Cambridge, UK. We also thank the NIHR, Hutchison Whampoa Limited, the Human Research Tissue Bank (Addenbrooke’s Hospital) and Cancer Research UK. The Department of Health funded the ProtecT study through the NIHR Health Technology Assessment programme. We also acknowledge to support of the research staff in S4 who so carefully curated the samples and the follow-up data (Jo Burge, Marie Corcoran, Anne George and Sara Stearn). Professor Donovan, Professor Hamdy and Professor Neal are National Institute for Health Research (NIHR) Senior Investigators. The views and opinions expressed therein are those of the authors and do not necessarily reflect those of the Department of Health. The Austrian arm of the study acknowledges support from the Anniversary Fund of the Austrian National Bank (grant number 15079) and by the Medical and Scientific Fund of the Mayor of the City of Vienna (grant number 10077). We are grateful to the GTEx Consortium for early stage sharing of RNA-seq and genotypic data. The following acknowledgement refers to the GTEx data: The Genotype-Tissue Expression (GTEx) Project was supported by the Common Fund of the Office of the Director of the National Institutes of Health, through the National Human Genome Research Institute, NHGRI’s Extramural Research Award (U41HG008490).
Institutes of Health. Additional funds were provided by the NCI, NHGRI, NBLI, NIDA, NIMH and NINDS. Donors were enrolled at Biospecimen Source Sites funded by NCI/SAIC-Frederick, Inc. (SAIC-F) subcontracts to the National Disease Research Interchange (10X5170), Roswell Park Cancer Institute (10X5171) and Science Care, Inc. (X105172). The Laboratory, Data Analysis and Coordinating Center (LDACC) was funded through a contract (HHSN268201000029C) to The Broad Institute, Inc. Biorepository operations were funded through an SAIC-F subcontract to Van Andel Institute (10ST1035). Additional data repository and project management were provided by SAIC-F (HHSN262120080001E). The Brain Bank was supported by a supplement to University of Miami grants DA06227 and DA033684 and to contract N01MH000028. Statistical Methods development grants were made to the University of Geneva (MH090941 and MH101814), the University of Chicago (MH090951, MH090937, MH101820 and MH101825), the University of North Carolina (MH090948), Stanford University (MH01782), Washington University St Louis (MH101810) and the University of Pennsylvania (MH101822). Funding to pay the Open Access publication charges for this article was provided by deCODE genetics/AMGEN.

References


