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Alternative mRNA splicing can attenuate the pathogenicity of presumed loss-of-function variants in *BRCA2*

Romy L. S. Mesman, MSc¹, Fabienne M. G. R. Calléja, MSc¹, Miguel de la Hoya, PhD², Peter Devilee, PhD^{1,3}, Christi J. van Asperen, MD, PhD⁴, Harry Vrieling, PhD¹ and Maaïke P. G. Vreeswijk, PhD¹

Purpose: Current interpretation guidelines for germline variants in high-risk cancer susceptibility genes consider predicted loss-of-function (LoF) variants, such as nonsense variants and variants in the canonical splice site sequences of *BRCA2*, to be associated with high cancer risk. However, some variant alleles produce alternative transcripts that encode (partially) functional protein isoforms leading to possible incorrect risk estimations. For accurate classification of variants it is therefore essential that alternative transcripts are identified and functionally characterized.

Methods: We systematically evaluated a large panel of human *BRCA2* variants for the production of alternative transcripts and assessed their capacity to exert *BRCA2* protein functionality. Evaluated variants included all single-exon deletions, various multiple-exon deletions, intronic variants at the canonical splice donor and acceptor sequences, and variants that previously have been shown to affect messenger RNA (mRNA) splicing in carriers.

Results: Multiple alternative transcripts encoding (partially) functional protein isoforms were identified (e.g., Δ [E4–E7], Δ [E6–E7], Δ E[6q39_E8], Δ [E10], Δ [E12], Δ E[12–14]). Expression of these transcripts did attenuate the impact of predicted LoF variants such as the canonical splice site variants c.631+2T>G, c.517-2A>G, c.6842-2A>G, c.6937+1G>A, and nonsense variants c.491T>A, c.581G>A, and c.6901G>T.

Conclusion: These results allow refinement of variant interpretation guidelines for *BRCA2* by providing insight into the functional consequences of naturally occurring and variant-related alternative splicing events.

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INTRODUCTION

Genetic testing of individuals with an enhanced risk of developing breast or ovarian cancer is routine clinical practice. Predicted loss-of-function (LoF) variants in *BRCA1* and *BRCA2*, such as nonsense variants, frame-shifting indels, and variants at the canonical splice sites, are considered to be associated with high cancer risk and carriers and their family members are managed accordingly.

Recently, however, it was established that some naturally occurring alternative transcripts of *BRCA1* and *BRCA2* encode protein isoforms with residual tumor suppressive activity.^{1–7} As a consequence, the pathogenic potential of predicted LoF variants located in an exon absent in these alternative transcripts may be substantially smaller than assumed.

Current gene-specific variant classification guidelines by ENIGMA (<https://enigmaconsortium.org/>) as well as the generic guidelines published by the American College of Medical Genetics and Genomics (ACMG) and Association for Molecular Pathology (AMP)⁸ have therefore included a

cautionary note. ENIGMA classification rules (<https://enigmaconsortium.org/>) state that variants found to produce messenger RNA (mRNA) transcript(s) predicted to encode isoforms that do not disrupt known clinically important functional domains should be considered class 3. The ACMG/AMP guidelines pose that the Pathogenic Very Strong (PVS1) code for predicted loss-of-function variants (nonsense, frameshift, canonical ± 1 or 2 splice sites, initiation codon, single or multiexon deletion) may no longer be valid if a variant induces an in-frame deletion or insertion that leaves the functional domains of the protein intact.⁹ Furthermore, caution is warranted for a variant allele that produces multiple mRNA transcripts as both transcript ratios and the functional integrity of the isoforms can affect its clinical relevance. Although alternative transcripts have been described for both *BRCA1* and *BRCA2*,^{10,11} a systematic analysis of the functionality of encoded protein isoforms has not been performed, which complicates the application of these variant classification guidelines.

¹Department of Human Genetics, Leiden University Medical Center, Leiden, The Netherlands; ²Molecular Oncology Laboratory, Instituto de Investigacion Sanitaria San Carlos, Hospital Clinico San Carlos, Madrid, Spain; ³Department of Pathology, Leiden University Medical Center, Leiden, The Netherlands; ⁴Department of Clinical Genetics, Leiden University Medical Center, Leiden, The Netherlands. Correspondence: Maaïke P. G. Vreeswijk (Vreeswijk@lumc.nl)

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For many *BRCA1* and *BRCA2* variants (both intronic and exonic) an effect on mRNA splicing has been reported using either patient RNA or minigene analysis.^{12–21} The analysis of patient RNA is however often hampered by the inability to determine allele-specific transcript expression. It then remains unclear if and to what extent wild type (WT) mRNA is still produced from the variant allele. To more directly assess the impact of an individual variant on both the nature and level of aberrant transcripts, minigene assays have been developed. These assays however lack the genomic context of the complete gene, limiting the detection of potential alternative transcripts. Jointly, the currently available approaches may provide evidence toward pathogenicity, but they all suffer from the same limitation: they do not provide insight into the in vivo functional consequences of variants that affect splicing, an important component of assessing variant pathogenicity. This shortcoming underscores the need for more detailed analyses per gene in which the presence and expression levels of alternative transcripts, either naturally occurring or induced by a variant, can be linked to protein function.

We recently validated a mouse embryonic stem cell (mESC)-based assay as a sensitive test for functional characterization of *BRCA2* missense variants.²² As sequence alterations are introduced in the full-length (FL) human *BRCA2* gene, the functional impact of all types of variants can be assessed including those that affect mRNA splicing. In addition, the presence of only a single human *BRCA2* allele makes the mESC system eminently suited for alternative mRNA transcript analysis.

In the present study, we show that the nature and relative contribution of naturally occurring transcripts to the overall expression of human *BRCA2* expressed in mESC is highly similar to those detected in various human tissues and cell lines. Furthermore, we systematically characterized a large panel of alternative transcripts for their ability to encode for (partially) functional *BRCA2* protein.

The functional data presented here can be used to refine classification guidelines for variants in *BRCA2* and improve the validity of PVS1 assignments for this gene. Moreover, alternative splicing is a general feature of many multiexon coding genes, and should be considered as a mechanism by which the assumed pathogenic potential of predicted LoF variants may be attenuated or even circumvented.

MATERIALS AND METHODS

Generation of exon-deletion variants

Thirty different exon-deletion variants (i.e., 25 single-exon deletions as well as five multiple-exon deletions) were generated in the full-length human *BRCA2* gene located on a bacterial artificial chromosome (BAC) (clone RP11-777I19, BACPAC) as described previously²³ (Tables S1, S5). Once the deletion was confirmed by Sanger sequencing BAC DNA was isolated according to manufacturer protocol (NucleoBond® Xtra Midi, Macherey-Nagel).

Selection and generation of *BRCA2* variants

Single-nucleotide variants that are likely to affect *BRCA2* mRNA splicing were selected from the ClinVar database (<https://www.ncbi.nlm.nih.gov/clinvar>) consisting of variants in the canonical ± 1 or 2 splice sites (Table S2) or of the last nucleotide of an exon (Table S3). In addition, we included variants for which aberrant splicing patterns had been reported in the literature to assess whether *BRCA2* variants expressed in mESC yield similar patterns of alternative transcripts as human cells (Table S3). Furthermore, from the ClinVar database, we selected nonsense variants located within exons that are absent from naturally occurring alternative transcripts (Ex3–7, Ex12, Ex18, and Ex19) or other alternative in-frame transcripts comprising a single-exon deletion (Ex10 and Ex26) (Table S2). Variants were generated in the complete human *BRCA2* gene as described previously.²² Primer sequences are listed in Table S5.

mESC-based functional assay

The mESC-based functional assay involves the introduction of human *BRCA2* variants into a hemizygous *Brca2* mESC line as described previously (Fig. S1).²² For the cell viability assay, 6×10^4 cells were seeded in triplo on 60-mm cell culture dishes and subsequently treated for 16 hours with 1.0 μM 4-Hydroxytamoxifen (4-OHT) (Sigma Aldrich). The next day, cells were washed with phosphate-buffered saline (PBS) and cultured for six days in the presence of hypoxanthine-aminopterin-thymidine (HAT) and subsequently five days in the presence of hypoxanthine-thymidine (HT). Thirteen days after 4-OHT treatment, one culture dish was used to visualize clonal survival by methylene blue staining. For each variant, the number of clones was compared with WT *BRCA2* expressing cells and based on that categorized into one of three categories: full (similar numbers of clones as WT *BRCA2*), intermediate (fewer and smaller clones than WT *BRCA2*), and noncomplementing (absence of viable clones) variants (Fig. S2). Variants of the full and intermediate complementing categories were assessed in the homology directed repair (HDR) assay as described previously.²² A flowchart for the interpretation of functional data generated by the mESC assay is presented in Fig. S4.

Reverse Transcription-PCR (RT-PCR)

To study the effect of a variant on mRNA splicing, RNA was isolated using a trizol-based protocol and complementary DNA (cDNA) was synthesized using the ProtoScript II First Strand cDNA synthesis kit (NEB) according to manufacturer's instructions. For variants that failed functional complementation in the cell viability assay, RNA was isolated prior to removal of the conditional mouse *Brca2* (*mBrca2*) allele. Then, 2 μl of cDNA was amplified with GoTaq polymerase (Promega) and human *BRCA2* exon-specific primer pairs (Table S5) under the following polymerase chain reaction (PCR) conditions; 95 °C for 5 minutes, followed by

28 cycles of 95 °C for 30 seconds, 55 °C for 30 seconds, 72 °C for 2 minutes, and a final step at 72 °C for 10 minutes. RT-PCR products were separated on 0.8–1.5% agarose gels stained with ethidium bromide and visualized by exposure to ultraviolet (UV) light. Individual bands were reamplified by band-stab PCR²⁴ and purified PCR products were subjected to Sanger sequencing to identify which transcript they represented. Importantly, not every band on the gel reflected a unique mRNA transcript as some of the products represented single-stranded PCR products.

Quantitative analysis of naturally occurring alternative transcripts in mESC expressing WT *BRCA2*

Capillary electrophoresis (CE) analysis of alternative splicing has been extensively described previously.¹¹ In brief, we used a panel of overlapping RT-PCR assays (combinations of forward and fluorescent-labeled reverse primers located in different exons) that allowed a comprehensive screening of *BRCA2* splicing events by CE.

Analysis was performed on two technical replicas of RNA samples from mESC expressing WT *BRCA2*. RNA samples (approximately 1 µg) were subjected to cDNA synthesis using a PrimeScript RT reagent kit with random primers according to the manufacturer's protocol (Takara Biotechnology). We performed 13 different RT-PCR assays spanning exons 1–4, 1–6, 3–8, 4–9, 7–10, 11–14, 11–15, 14–16, 16–19, 16–22, 19–22, 20–24, and 22–27 (sequences of all primers are available upon request). PCR products were analyzed by CE (50-cm capillary arrays) in a 3130 Genetic Analyzer (Applied Biosystems) with GeneScan 500-LIZ/1200-LIZ size standards (Applied Biosystems) as internal markers. Size calling was performed with GeneMapper v4.0 Software (Applied Biosystems). For comparison, RNA samples from lymphoblastoid cell lines (LCLs) were analyzed in parallel. By comparing the relative contribution of the same alternative transcripts between samples from mESC and human cells, the quantification is not influenced by overestimation of the expression of shorter transcripts as previously shown to occur using RT-PCR in combination with CE.^{2,25,26} Only fragments over 50 relative fluorescent units (RFUs) were considered to represent distinct transcripts.

Western blot analysis

Western blot analysis was performed using NuPAGE™ Novex™ 3–8% Tris-Acetate Protein Gels (ThermoFisher Scientific). *BRCA2* protein was detected with the rabbit polyclonal antibody (BETHYL, A303–434A-T) directed against a region between amino acids 450–500 in exon 10 of *BRCA2*. Protein signal was detected by electrochemiluminescence (Amersham ECL RPN2235 Biocompare). It is important to note that most in-frame protein isoforms cannot be distinguished by western blot analysis due to the small difference in size between the full-length *BRCA2* protein (*BRCA2* FL protein isoform, 3418 aa) and *BRCA2* protein isoforms deleted for only one or a few small exons.

RESULTS

Naturally occurring alternative splicing of *BRCA2* mRNA

The mESC-based functional assay allows evaluation of any type of *BRCA2* variant in its natural genomic context. Variants are introduced in a human *BRCA2*-containing BAC, transfected into mESC containing a single, conditional *mBrca2* allele and assessed for their ability to rescue the cell lethality provoked by removal of endogenous *mBrca2* (Fig. S1). Three phenotypes can be distinguished for variants, i.e., fully complementing (similar number of clones as WT *BRCA2*), intermediate (fewer and smaller clones than WT *BRCA2*), and noncomplementing (absence of viable clones) (Fig. S2). Subsequently, variants of the full and intermediate complementing categories can be tested for their ability to perform HDR, the most prominent tumor suppressor function of *BRCA2*. The assay was previously validated by functional assessment of a large series of classified *BRCA2* missense variants and revealed a high sensitivity and specificity for variant classification.²² It is important to note that the complementation phenotype reflects the impact of variants on HDR as well as other *BRCA2*-associated cellular processes that play a role in the preservation of genome stability. Consequently, the correlation between complementation phenotype and HDR capacity is not absolute, but in general intermediate complementing variants display a severe reduction in repair capacity.

To determine whether processing of human *BRCA2* mRNA by the murine spliceosome accurately reflects the splicing process in human cells, we determined the presence and quantity of the major naturally occurring isoforms that are produced from a genomic copy of the human *BRCA2* gene in mESC. RNA analysis showed that the repertoire of the major naturally occurring mRNA transcripts (i.e., Δ[E3], Δ[E6q39_E7], Δ[E12], and Δ[E17–E18]) of mESC expressing WT *BRCA2* closely resembled that of human LCLs both qualitatively (all predominant splicing events are detected, novel splicing events are not observed) and quantitatively (similar expression ratios relative to FL transcript) (Fig. 1).¹¹ Up to this date, no tissue-specific transcripts have been observed in nonmalignant breast epithelia, ovarian epithelia, or ovarian fimbria.^{11,15}

Functional characterization of exon deletions in *BRCA2*

Although splicing is a highly coordinated process, it is currently impossible to predict which alternative transcripts will be produced when the splice recognition site of a particular exon is destroyed or when a complete exon has been deleted. Furthermore, it is unclear when in-frame transcripts are produced whether these encode for protein isoforms that retain tumor suppressor activity. To systematically bridge this knowledge gap, we generated 30 different exon-deletion (DelEx) variants in the human *BRCA2* gene, including all single-exon deletions and five in-frame multiple-exon deletions (Table S1) and analyzed the alternative transcripts these DelEx variants produced as well as their ability to preserve *BRCA2* functionality.

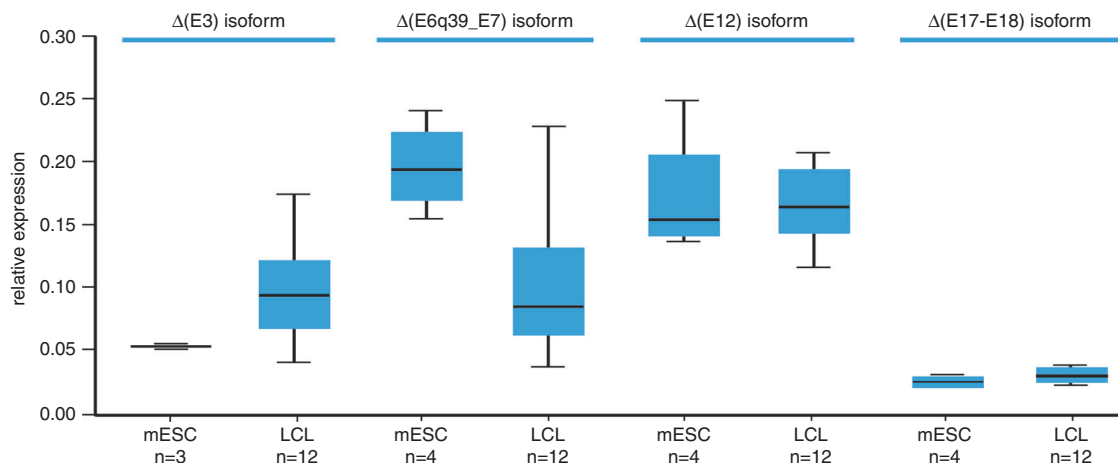


Fig. 1 Expression of the major naturally occurring alternative *BRCA2* transcripts relative to full-length transcript. Analysis by capillary electrophoresis of fluorescent RT-PCR products was performed to determine the relative expression level of isoforms in mouse embryonic stem cells (mESCs) transfected with wild type (WT) *BRCA2* and in lymphoblastoid cell lines (LCLs) from healthy blood donors.

After removal of the conditional *mBrca2* allele, 17 DelEx variants failed to complement the cell lethal phenotype induced by loss of *mBrca2*. Eight DelEx variants displayed full complementation, while complementation was intermediate for five other DelEx variants (Table S1, Fig. S2).

RNA analysis revealed that various DelEx variants expressed multiple alternative mRNA transcripts (e.g., DelEx4 variant did not only produce $\Delta[E4]$ but also $\Delta[E4-E7]$ transcript) (Table S1). Evaluation of the transcripts generated by the DelEx variants displaying full complementation allowed us to identify several potential rescue transcripts, i.e., encoding (partially) functional *BRCA2* protein isoforms (Table S1) that could be detected by western blot analysis (Fig. S3). The most potent in-frame rescue transcripts being $\Delta(E4-E7)$ (r.317_631del315), $\Delta(E6-E7)$ (r.476_631del156), $\Delta(E6q39_E8)$ (r.478_681del204), $\Delta(E10)$ (r.794_1909del1116), $\Delta(E12)$ (r.6842_6937del96) and $\Delta(E12-E14)$ (r.6842_7435del594) (summarized in Figs. 2b and 3, Table S1). *BRCA2* transcripts expressed by variants displaying intermediate complementation encoded protein isoforms that were either truncated and/or reduced in quantity (DelEx5, DelEx14, DelEx15, DelEx16, DelEx18). In some cases (DelEx15, DelEx16, DelEx18) the nature of the rescue transcript remains elusive. DelEx variants that failed to complement cell lethality either produced no detectable transcript (DelEx2) or (a mixture of) out-of-frame transcripts (DelEx6, 9, 13,, 20, 21, 22, 23, 24, 25) and nonfunctional in-frame transcripts (DelEx3, 3-7, 11, 14-16, 17, 19, 26) (summarized in Figs. 2b and 3, Table S1). Congruently with their complementation phenotype, the fully complementing DelEx variants displayed HDR levels above 50%. In contrast, HDR activity of the five variants that showed intermediate complementation was severely diminished to a level previously defined for variants associated with enhanced breast cancer risk (HDR < 30%) (Fig. 4a).²²

Based on the functionality of the transcripts lacking specific exons as summarized in Fig. 2b, it is concluded that exons 4,

5, 6, 7, 8, 10, 12, 13, and 14 do not encode essential parts of *BRCA2* protein and that protein isoforms encoded by naturally occurring or variant-induced in-frame alternative transcripts lacking one or multiple of these exons may (partially) retain *BRCA2*'s functionality in HDR.

Functional characterization of *BRCA2* PVS1 variants

Variant classification using ACMG/AMP guidelines involves several benign and pathogenic evidence criteria, including a pathogenic criterion (PVS1) for predicted LoF variants (nonsense, frameshift, canonical ± 1 or 2 splice sites, initiation codon, single- or multiexon deletion and duplications).⁹ The results from our DelEx variant analyses suggest that nonsense, out-of-frame indels, and spliceogenic variants either located in or affecting splicing of exons that encode nonessential domains of the *BRCA2* protein may not lead to complete LoF because of the production of rescue transcripts. To investigate this in more detail, we characterized a panel of 29 nonsense and spliceogenic PVS1 variants for their ability to produce *BRCA2* isoforms that retain residual protein activity (Table S2).

Of the ten nonsense variants that were evaluated, one variant (c.6901G>T [located in Ex12]) displayed full complementation while three nonsense variants (c.491T>A [Ex6], c.581G>A [Ex7], and c.9572G>A [Ex26]) showed intermediate complementation. Variant c.6901G>T almost exclusively produced the $\Delta(E12)$ transcript (Fig. 3b). In line with the previously demonstrated functionality of variant DelEx12 (51% HDR), variant c.6901G>T revealed a moderate functional impact and retained 43% HDR capacity (Fig. 4b). In cells expressing either variant c.491T>A or c.581G>A, the expression level of the naturally occurring $\Delta(E4-E7)$ transcript was slightly enhanced compared with cells expressing WT *BRCA2* (Fig. 3a). As shown for the DelEx4-7 variant this alternative transcript encodes a HDR-competent protein isoform. Nevertheless, the expression level of the $\Delta(E4-E7)$ transcript is apparently insufficient to retain full *BRCA2*

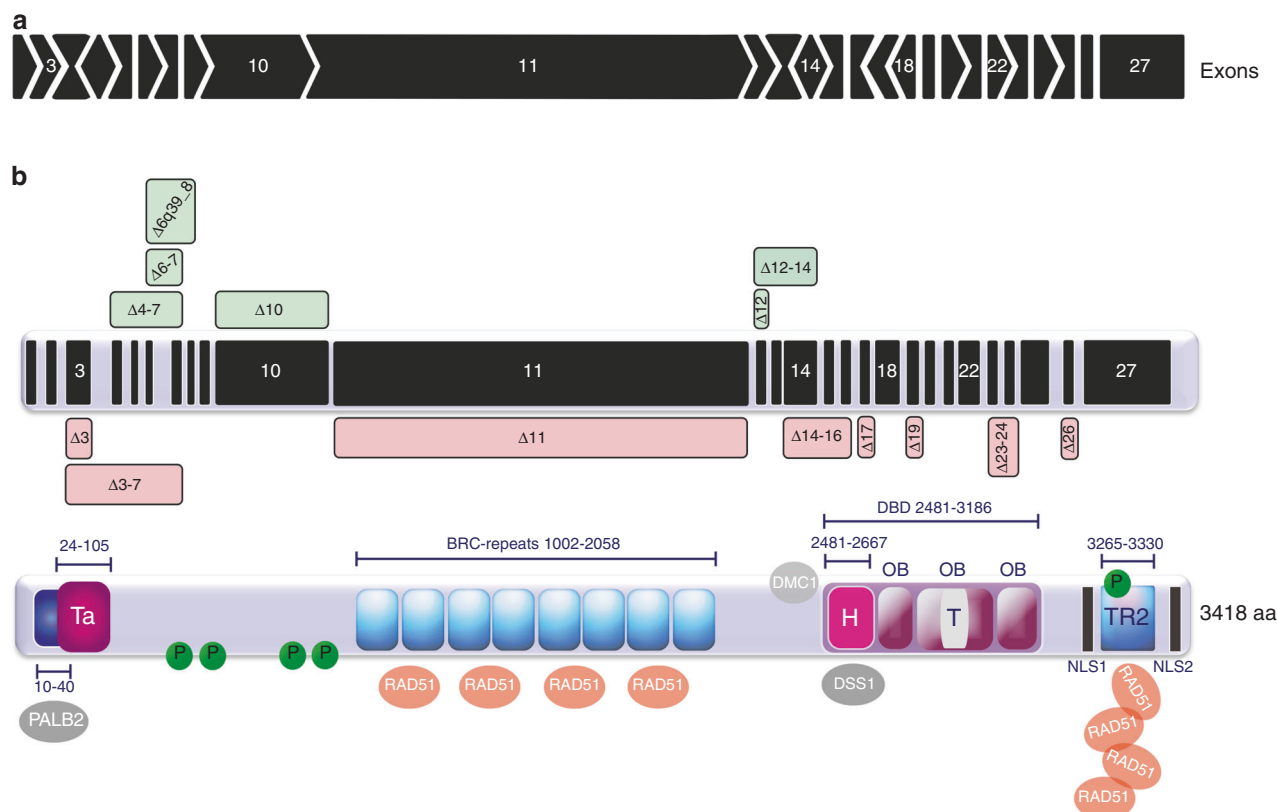


Fig. 2 Schematic overview of BRCA2 reading frame and functionality of alternative isoforms. (a) *BRCA2* reading frame and (b) the functionality conferred by alternative in-frame *BRCA2* isoforms including single- and multiple-exon deletions. Figure adapted from Mesman *et al.*²² Green box = homology directed repair (HDR) capacity >50%. Red box = no complementation or HDR capacity ≤30%.

functionality as both nonsense variants display a severe impact on HDR (Fig. 4b). Variant c.9572G>A produced two transcripts: the FL transcript containing the stop codon and a ΔEx26 transcript. As the DelEx26 variant failed to complement loss of endogenous *BRCA2*, the observed complementation of c.9572G>A is unlikely the consequence of increased Ex26 skipping. The severe reduction in HDR activity detected for c.9572G>A, only 25% activity compared with WT, most likely reflects some residual activity conferred by the truncated *BRCA2* protein isoform (Fig. 4b).

Remarkably, 5 of 19 canonical splice site variants tested were able to rescue cell lethality (Table S2). Enhanced expression of naturally occurring transcripts Δ(E4–E7) or Δ(E12) was detected for variants located in the canonical splice sites of Ex7 and Ex12 (i.e., c.517–2A>G [Ex7], c.631+2T>G ([Ex7], c.6842–2A>G [Ex12], c.6937+1G>A [Ex12]) and is likely responsible for their residual HDR activity (>50%, Fig. 4b). Variant c.7008–2A>T (Ex14) produced multiple alternative transcripts including three out-of-frame transcripts and one in-frame transcript containing a 246-bp (partial) deletion of Ex14 through an exon 14 cryptic acceptor site (Table S2, Fig. 3c). Although this variant was able to partially complement loss of endogenous *Brca2*, the level of HDR activity (35%) of this variant was severely impaired (Fig. 4b) and possibly results from the relatively low

expression level of the potential rescue transcript Δ(E14p246). It should be noted that this variant has been observed in *cis* with c.631G>A variant for which an effect on RNA splicing (i.e., exon 7 skipping) has also been reported.²⁷

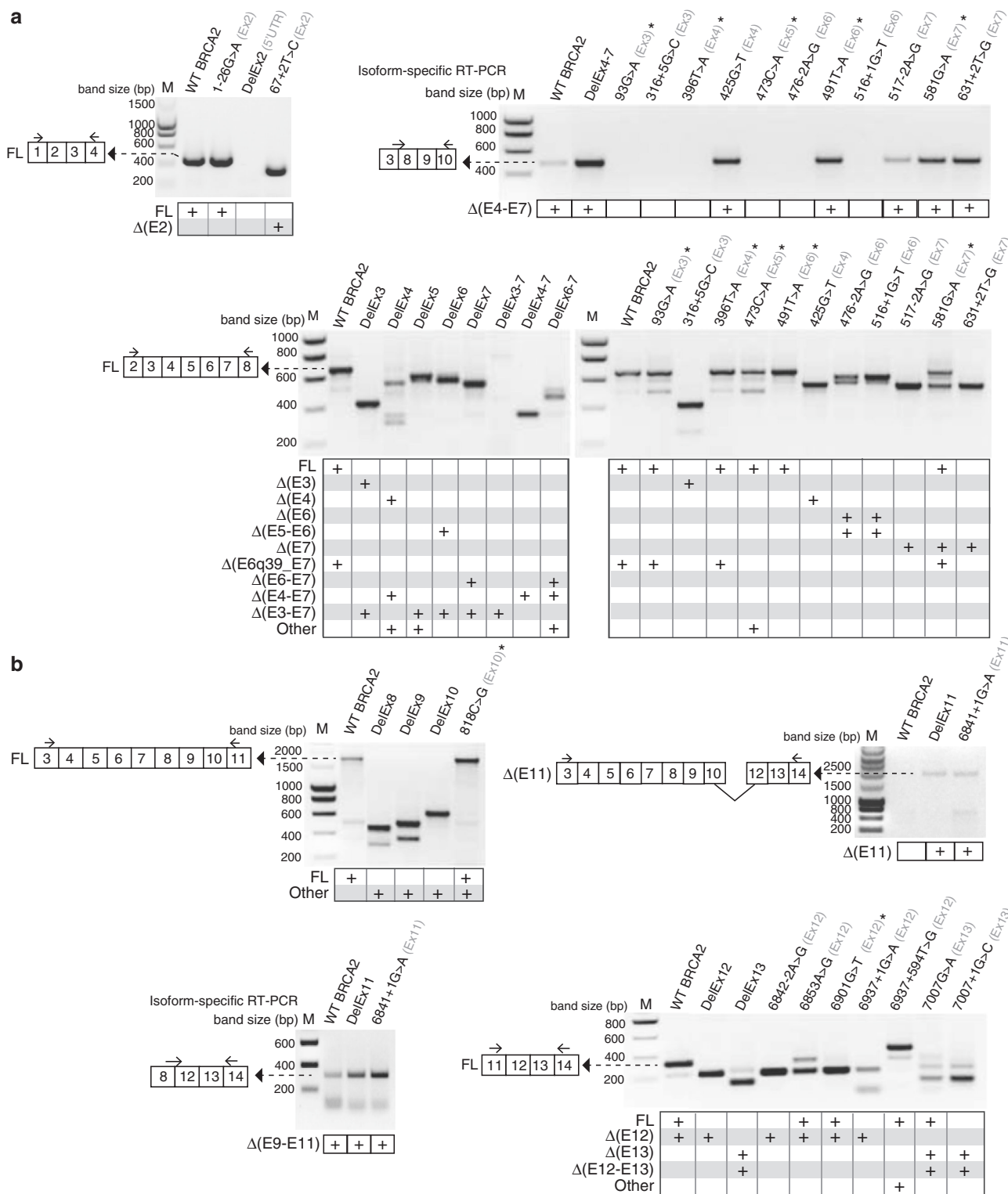
Functional characterization of potential spliceogenic *BRCA2* variants

For the vast majority of potential spliceogenic variants that are located outside the canonical splice sites of *BRCA2* exons, it is unknown whether they truly affect splicing and if so, to what extent identified aberrant splicing events affect protein functionality. We selected 13 *BRCA2* variants for which RNA analysis has been reported in human cells and determined their impact on both mRNA splicing and protein function in mESCs (Figs. 3 and 4c, Table S3).

Overall, human *BRCA2* variants in mESCs rendered similar mRNA transcript profiles as previously detected in LCLs and minigene analysis with all major aberrant splicing events identified.^{12,28–31} The complementation phenotype of two variants, c.316+5G>C and c.7007G>A, resembled that of high-risk (class 4/5) variants with respect to their inability to rescue the cell lethality imposed by Cre-mediated loss of *mBrca2* (Fig. S2). Variant c.316+5G>C only produced transcripts lacking exon 3, which, as discussed above, encodes a stable but nonfunctional

protein isoform (Table S3, Figs. 3a and S3). Variant c.7007G>A (last nucleotide Ex13) expressed both two aberrant out-of-frame transcripts ($\Delta[E13]$ and $\Delta[E12-E13]$) and FL transcript (Fig. 3b). However, expression of the FL transcript was apparently too low to allow complementation (Fig. S3).

Variants c.8754+4A>G and c.9117G>A (last nucleotide Ex23) displayed full complementation of cell lethality but were severely impaired in their HDR capacity (Fig. 4c), in concordance with their recent classification as pathogenic variants.³² However, the nature of the transcript that is responsible for the rescue of cell viability remains elusive.



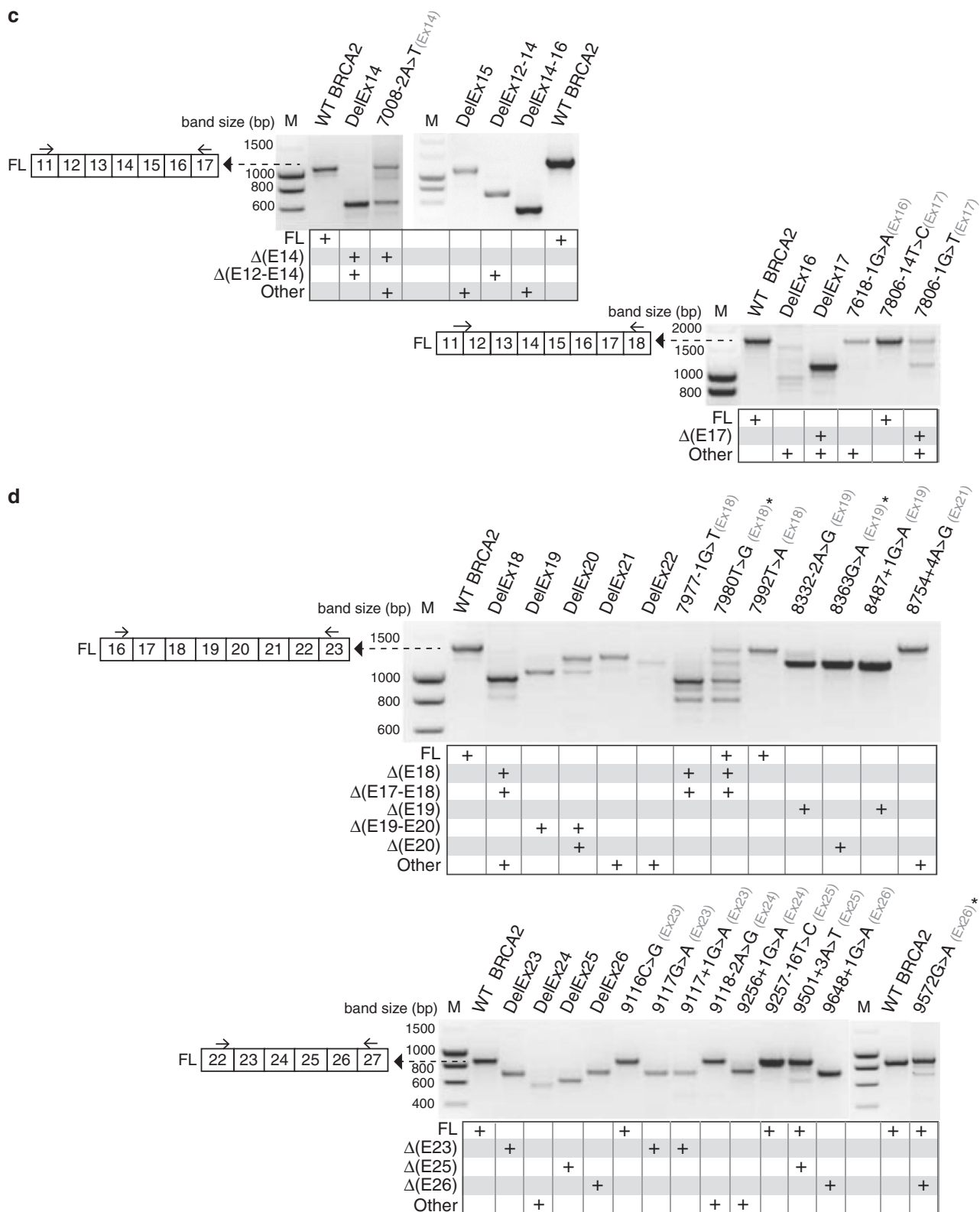


Fig. 3 RNA analysis of BRCA2 variants. Splicing patterns of BRCA2 variants in or surrounding (a) Ex2–7, (b) Ex8–13, (c) Ex14–17, (d) Ex18–26. The location of exonic primers used for RT-PCR is indicated by arrows above the schematic representation of the analyzed cDNA region. Unique mRNA transcripts indicated in this figure by “other” are defined in Tables S1–S3. Asterisks denote nonsense variants. + transcript detected. FL full length, M marker (Smartladder 200 to 10000 bp from Eurogentec) WT wild type.

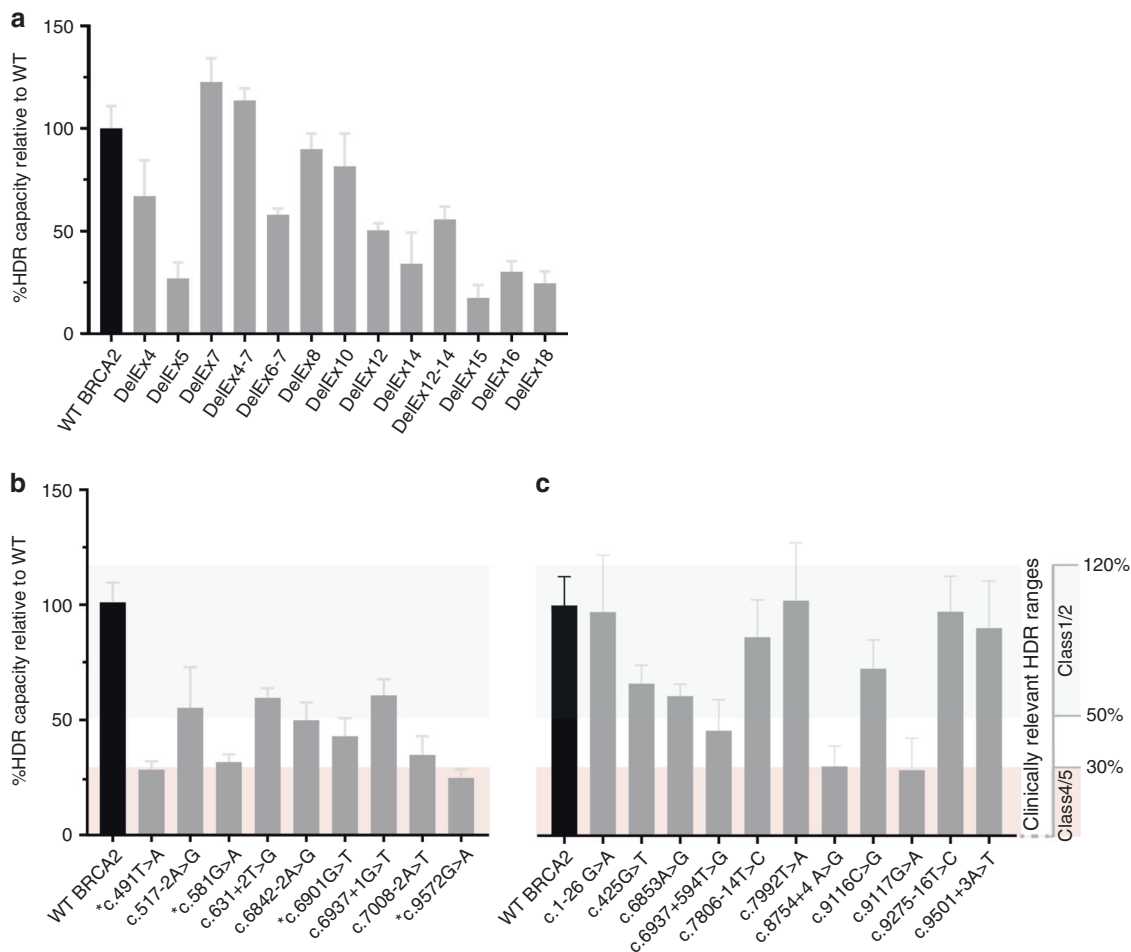


Fig. 4 Homology directed repair (HDR) capacity as measured in the DR-GFP reporter assay. Homology directed repair (HDR) capacity measured in the Direct Repeat - Green Fluorescent Protein (DR-GFP) reporter assay for (a) DelEx variants, (b) PVS1 variants, and (c) potential spliceogenic *BRCA2* variants outside the canonical splice sites. HDR capacity is expressed as the percentage GFP positive cells relative to the GFP positive population in wild type (WT) *BRCA2* samples. Error bars indicate the SD of six independent GFP measurements per variant. The HDR capacity was measured for all variants that were able to complement the loss of cell viability following Cre-mediated deletion of the conditional *mBrca2* allele. The HDR range of classified nonpathogenic (class 1/2) and pathogenic (class 4/5) *BRCA2* missense variants is plotted at the right y-axis.²² Asterisks denote nonsense variants.

Variant c.425G>T (last nucleotide Ex4) produced an out-of-frame transcript ($\Delta[E4]$) and a transcript ($\Delta[E4-E7]$) that preserves the reading frame, which is likely responsible for the residual 66% HDR capacity (Figs. 3a and 4c).

Also, deep intronic variants can impose aberrant splicing as previously reported for c.6937+594T>G.^{28,31} Due to the activation of a cryptic splice site an intronic fragment of 95 bases is inserted between exons 12 and 13 leading to an out-of-frame transcript. Molecular analysis revealed that although intron retention seems to be the predominant splicing event (Fig. 3b) for c.6937+594T>G, the variant allele produced sufficient FL transcript to rescue cell lethality and to retain residual HDR activity (46% compared with WT) (Fig. 4c). For the remaining seven potential spliceogenic variants mRNA splicing in five variants appeared not to be affected while in c.6853A>G and c.9501+3A>T sufficient quantities of FL transcript were produced to prevent substantial loss of *BRCA2* function (Fig. 4c).

DISCUSSION

In classification guidelines documented by ACMG and AMP⁸ and ENIGMA (<https://enigmaconsortium.org/>), cautionary notes are included for variants that produce in-frame alternative gene transcripts that retain clinically important functional protein domains. Now that we have revealed various functionally redundant regions in the *BRCA2* protein, it is possible to propose *BRCA2*-specific rules. Our results indicate that the majority of the presumed LoF variants will lead to inactivation of the *BRCA2* protein, and hence, be associated with high cancer risk (Fig. 5). However, for a number of variants additional analyses will be required before they can be considered to represent pathogenic variants associated with high cancer risk. In particular, for variants in the canonical splice site regions of exons 4, 7, 8, 10, 12, and 14 caution is warranted since LoF may be prevented through elevated expression of in-frame rescue transcripts. Furthermore, nonsense variants, out-of-frame indels, and complete

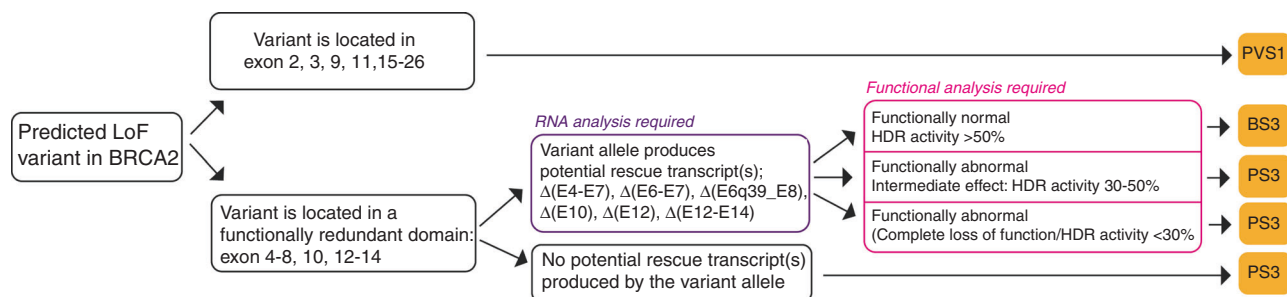


Fig. 5 Decision tree to assist in the interpretation of predicted loss-of-function variants according to American College of Medical Genetics and Genomics (ACMG) guidelines based on functional data presented in this paper. Additional data and considerations are needed to determine the appropriate strength of the PS3/BS3 criteria as stated in Brnich *et al.*³³ HDR homology directed repair.

deletion of functionally redundant exons may for the same reasons retain (partial) functionality. Expression of $\Delta(E4-E7)$ or $\Delta(E12)$ transcripts in mESCs with *BRCA2* nonsense variants in exons 6, 7, or 12⁷ was sufficient to retain substantial *BRCA2* protein functionality. These findings put into question whether the investigated splice site variants, nonsense variants, and complete exon deletions are associated with high cancer risk. As this model system provides an RNA splicing assay with a direct measure of protein function, the experimental data generated by this functional assay is eminently suited to be applied in variant interpretation. We would like to propose a refined provisional framework for functional evidence application in ACMG/AMP clinical variant interpretation guidelines.^{9,33,34} The decision tree shown in Fig. 5 may serve as a means to indicate those presumed LoF variants for which the PVS1 code might not be warranted.

In the current multifactorial likelihood model (MLM), a prior probability of pathogenicity is combined with likelihood ratios estimated from clinical data resulting in a final posterior probability that assigns the variant to one of the five classes of the International Agency for Research on Cancer (IARC) classification system.^{21,35,36} The prior probability is an in silico prediction of the functional impact based on variant location and bioinformatic prediction of variant effect.^{37,38} Due to the high prior probability assigned to nonsense (0.99) and canonical splice site (0.97) variants the prior heavily impacts the final classification of a variant. However, a high prior might not be justified for presumed LoF variants in functionally redundant exons. For this reason, a reduced prior probability of 0.5 was proposed for variants in *BRCA1* exons 9–10 or their proximal splice junction regions.³⁸ Likewise, the prior probability of pathogenicity was set at 0.5 for variants in the splice acceptor and donor site of *BRCA2* exon 12. Our results indicate that adjustment of the prior should be extended to other regions of *BRCA2* in which presumed LoF variants still display considerable *BRCA2* protein activity such as nonsense and splice site variants in exon 7 (Fig. S5). Furthermore, the design of the multifactorial likelihood model restricts its use to discrimination of variants that confer high cancer risk from those that do not. Recent data show that variants associated with reduced penetrance do exist in

BRCA1 and *BRCA2* and functional analysis might be required to identify these variants.^{39,40} Recently, Parsons *et al.*³² have performed multifactorial likelihood analyses for a large number of *BRCA1* and *BRCA2* variants, including 13 variants that were functionally characterized in this study (Table S4). For most variants, the IARC classification is in agreement with our functional data. However, two variants in respectively the splice acceptor site (c.517–2A>G) and donor site (c.631+2T>G) of exon 7 were classified as pathogenic based on multifactorial likelihood quantitative analysis, while in our analyses these variants show residual HDR capacity in the lower range of class 1/2 variants (Table S4). At this moment, the exact quantitative relationship between *BRCA2* protein functionality and cancer risk is still unclear. Although HDR activity around 50% of WT activity was shown to correlate with an odds ratio of 2.5 for breast cancer,⁴⁰ additional studies are required to define HDR activity ranges that allow assignment of variants to clinically relevant cancer risk categories (i.e., high, moderate, and low increased risk). The observation that presumed LoF variant alleles may retain (partial) functionality through the expression of alternative protein isoforms incites a shift in genetic diagnostics. These findings emphasize the need for inclusion of quantitative functional data to the MLM (as done in a qualitative way in ACMG/AMP guidelines) and specification of gene-specific classification guidelines.

SUPPLEMENTARY INFORMATION

The online version of this article (<https://doi.org/10.1038/s41436-020-0814-5>) contains supplementary material, which is available to authorized users.

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DISCLOSURE

The authors declare no conflicts of interest.

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