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Hereditary Cancer in Clinical Practice

# RESEARCH Open Access



# Genetic variants of prospectively demonstrated phenocopies in *BRCA1/2* kindreds

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## **Abstract**

**Background:** In kindreds carrying *path\_BRCA1/2* variants, some women in these families will develop cancer despite testing negative for the family's pathogenic variant. These families may have additional genetic variants, which not only may increase the susceptibility of the families' *path\_BRCA1/2*, but also be capable of causing cancer in the absence of the *path\_BRCA1/2* variants. We aimed to identify novel genetic variants in prospectively detected breast cancer (BC) or gynecological cancer cases tested negative for their families' pathogenic *BRCA1/2* variant (*path\_BRCA1* or *path\_BRCA2*).

**Methods:** Women with BC or gynecological cancer who had tested negative for *path\_BRCA1* or *path\_BRCA2* variants were included. Forty-four cancer susceptibility genes were screened for genetic variation through a targeted ampliconbased sequencing assay. Protein- and RNA splicing-dedicated in silico analyses were performed for all variants of unknown significance (VUS). Variants predicted as the ones most likely affecting pre-mRNA splicing were experimentally analyzed in a minigene assay.

**Results:** We identified 48 women who were tested negative for their family's  $path\_BRCA1$  (n = 13) or  $path\_BRCA2$  (n = 35) variants. Pathogenic variants in the ATM, BRCA2, MSH6 and MUTYH genes were found in 10% (5/48) of the cases, of whom 15% (2/13) were from  $path\_BRCA1$  and 9% (3/35) from  $path\_BRCA2$  families. Out of the 26 unique VUS, 3 (12%) were predicted to affect RNA splicing (APC c.721G > A, MAP3K1 c.764A > G and MSH2 c.815C > T). However, by using a minigene, assay we here show that APC c.721G > A does not cause a splicing defect, similarly to what has been recently reported for the MAP3K1 c.764A > G. The MSH2 c.815C > T was previously described as causing partial exon skipping and it was identified in this work together with the  $path\_BRCA2$  c.9382C > T (p.R3128X).

**Conclusion:** All women in breast or breast/ovarian cancer kindreds would benefit from being offered genetic testing irrespective of which causative genetic variants have been demonstrated in their relatives.

Keywords: BRCA1, BRCA2, Breast cancer, Gene panel testing, RNA splicing

#### **Background**

Breast cancer (BC) is one of the most common human malignancies, accounting for 22% of all cancers in women worldwide [1]. A significant proportion of BC cases can be explained by hereditary predisposition and approximately 30% of this hereditary cancer risk is explained by the currently known high-penetrance

susceptibility genes [2–5]. Notably, carriers of pathogenic *BRCA1* or *BRCA2* variants (*path\_BRCA1* or *path\_BRCA2*) have an increased risk of developing BC (average lifetime risk of 35–85%) and ovarian cancer (average lifetime risk 11–39%). Further, carriers of pathogenic variants of *ATM*, *CHEK2*, *PALB2*, *NBS1* and *RAD50* have been found to confer two- to five-fold increased risk for developing BC [1, 6]. It is also known that pathogenic variants in *TP53*, *PTEN*, *STK11* and *CDH1*, resulting in Li-Fraumeni syndrome, Cowden syndrome, Peutz–Jeghers syndrome and hereditary diffuse

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gastric cancer, respectively, are associated with a high lifetime risk (>40%) of BC. Moreover, pathogenic variants in RAD51 paralogs, i.e., RAD51C, confer an increased risk of ovarian cancer [7]. The frequency of pathogenic variants in BC-associated genes varies significantly among different populations, as exemplified by the frequently studied founder pathogenic variant c.1100delC in CHEK2 [6].

The identification of path\_BRCA1 or path\_BRCA2 in an affected BC individual enables access to evidencebased screening for family members, and thus facilitates the implementation of appropriate cancer prevention in these families [1, 5, 6]. However, some women in families with an identified pathogenic variant will develop cancer despite testing negative for the family's pathogenic variant, often denoted as phenocopies [8]. In BC kindreds having a demonstrated path\_BRCA2 variant, the number of phenocopies is reportedly more frequent than expected by chance [8-10]. It has been proposed that these families may have additional genetic variants, which not only may increase the susceptibility of the families' path\_BRCA1/2, but also be capable of causing cancer in the absence of the path\_BRCA1/2 demonstrated in the families [5–7].

The current practice of genetic counselling for women who do not carry the *path\_BRCA1/2* variants of their relatives is challenging since their recognition is crucial for application of proper diagnostic and therapeutic approaches in these families. To discover additional inherited disease-causing variants in *path\_BRCA1/2* kindreds, we examined all prospectively detected BC or gynecological cancer cases in these kindreds by next-generation sequencing (NGS) using a panel of 44 cancer susceptibility genes. All detected variants were analyzed by RNA splicing- and protein-dedicated in silico methods. Variants predicted as the most likely to affect splicing were experimentally analyzed by using a cell-based minigene splicing assay.

#### Methods

#### Study population

For more than 20 years, we (the Hereditary Cancer Biobank from the Norwegian Radium Hospital, Norway; and the Department of Genomic Medicine from the University of Manchester, United Kingdom) have ascertained BC and breast/ovarian cancer kindreds by family history. The sisters and daughters of cancer patients were initially subjected to follow-up by annual mammography and gynecological examinations as appropriate at that time, and later they were all subjected to genetic testing [11].

Both collaborating outpatient genetic centers identified 48 women with prospective detected BC or gynecological cancer at follow-up, who were tested negative for their respective families' *path\_BRCA1/2* variants. Clinical data were obtained from pathology reports and clinical files.

Ethical approval for the prospective study was granted from the Norwegian Data Inspectorate and Ethical Review Board (ref 2015/2382). All examined patients had signed an informed consent for their participation in the study.

#### Targeted sequencing

Genomic DNA was isolated from peripheral blood samples and targeted sequencing was carried out using a TrueSeq amplicon based assay v.1.5 on a MiSeq apparatus, as previously described [12]. The 44-gene panel used in this work includes genes associated with cancer predisposition as described in a prior study [12].

#### Sequencing data analysis

Paired-end sequence reads were aligned to the human reference genome (build GRCh37) using the BWA-mem algorithm (v.0.7.8-r55) [13]. The initial sequence alignments were converted to BAM format and subsequently sorted and indexed with SAMtools (v.1.1) [13]. Genotyping of single nucleotide variants (SNV) and short indels was performed by GATK's HaplotypeCaller. Filtering of raw genotype calls and assessment of callable regions/loci were done according to GATK's best practice procedures, as described more detail previously [12].

Variants were annotated using ANNOVAR (version November 2015) [14] and were queried against a range of variant databases and protein resources (v29, December 2015), as previously described [12].

# Validation by cycling temperature capillary electrophoresis

The pathogenic variants identified in this study were validated by cycling temperature capillary electrophoresis. The method is based on allele separation by cooperative melting equilibrium while cycling the temperature surrounding capillaries [15]. This approach has previously been described and extensively used to detect somatic mutations and single nucleotide polymorphisms (SNPs) [16–19]. The amplicon design was performed by the variant melting profile tool (https://hyperbrowser.uio.no/hb/?tool\_id=hb\_variant\_melting\_profiles/) [20]. Primer sequences, PCR reaction conditions and electrophoresis settings are described in Additional file 1.

#### Genetic variants nomenclature and classification

The nomenclature guidelines of the Human Genome Variation Society (HGVS) were used to describe the detected genetic variants [21]. The recurrence of the identified variants was established by interrogating six databases (in their latest releases as of November 2016): Evidence-based Network for the Interpretation of Germline Mutant

Alleles (ENIGMA), Breast Cancer Information Core Database (BIC), the International Society of Gastrointestinal Hereditary Tumors (InSiGHT) Database, the Leiden Open Variation Database (LOVD), ClinVar, and the Human Gene Mutation Database (HGMD).

Novel variants were considered pathogenic if either one of the following criteria was met: a) introduced a premature stop codon in the protein sequence (nonsense or frameshift); b) occurred at positions +1/+2 or -1/-2 of donor or acceptor splice sites, respectively; and c) represented whole-exon deletions or duplications.

#### In silico analyses of VUS

Two types of bioinformatics methods were used to predict the impact of selected variants on RNA splicing. First, we used MaxEntScan (MES) and SSF-like (SSFL) to predict variant-induced alterations in 3' and 5' splice site strength, as described by Houdayer et al. [22], except that here both algorithms were interrogated by using the integrated software tool Alamut Batch version 1.5, (Interactive Biosoftware, http://www.interactive-biosoftware.com). For prediction of variant-induced impact on exonic splicing regulatory elements (ESR), we resorted to ΔtESRseq- [23], ΔHZei- [24], and SPANR-based [25] as described by Soukarieh et al. [26]. Score differences  $(\Delta)$  between variant and wild-type (WT) cases were taken as proxies for assessing the probability of a splicing defect. More precisely, we considered that a variant mapping at a splice site was susceptible of negatively impacting exon inclusion if ΔMES≥15% and ΔSSFL≥5% [22], whereas an exonic variant located outside the splice sites was considered as a probable inducer of exon skipping if negative  $\Delta$  scores (below the thresholds described below) were provided by all the 3 ESR-dedicated in silico tools. We chose the following thresholds: <- 0.5 for  $\Delta tESRseq$ -, <- 10 for  $\Delta HZei$ -, and < - 0.2 for SPANRbased scores. In addition, we evaluated the possibility of variant-induced de novo splice sites by taking into consideration local changes in MES and SSFL scores. In this case, we considered that variants located outside the splice sites were susceptible of creating a competing splice site if local MES scores were equal to or greater than those of the corresponding reference splice site for the same exon.

In silico protein impact predictions of VUS were performed with FATHMM (http://fathmm.biocompute.org.uk) (v2.3), PolyPhen2-HVAR (v 2.2.2), MutationTaster (data release Nov 2015), MutationAssessor (release 3), SIFT (Jan 2015) and PROVEAN (v1.1 Jan 2015) using dbNSFP v3.4.

#### Cell-based minigene splicing assays

In order to determine the impact of the APC c.721G > A on RNA splicing, we performed functional assays based

on the comparative analysis of the splicing pattern of WT and mutant reporter minigenes [27], as follows. First, the genomic region containing APC exon 7 and at least 150 nucleotides of the flanking introns (c.646-169 to c.729 + 247) were amplified by PCR using patient #12470 DNA as template and primers indicated in Additional file 2. Next, the PCR-amplified fragments were inserted into a previously linearized pCAS2 vector [26] to generate the pCAS2-APC exon 7 WT and c.721G > A minigenes. All constructs were sequenced to ensure that no unwanted mutations had been introduced into the inserted fragments during PCR or cloning. Then, WT and mutant minigenes were transfected in parallel into HeLa cells grown in 12-well plates (at ~ 70% confluence) using the FuGENE 6 transfection reagent (Roche Applied Science). Twenty-four hours later, total RNA was extracted using the NucleoSpin RNA II kit (Macherey Nagel) and, the minigene transcripts were analyzed by semi-quantitative RT-PCR using the One-Step RT-PCR kit (QIAGEN), as previously described [26]. The sequences of the RT-PCR primers are shown in Additional file 2. Then, RT-PCR products were separated by electrophoresis on 2.5% agarose gel containing EtBr and visualized by exposure to UV light under saturating conditions using the Gel Doc XR image acquisition system (Bio-Rad), followed by gel-purification and Sanger sequencing for proper identification of the minigenes' transcripts. Finally, splicing events were quantitated by performing equivalent fluorescent RT-PCR reactions followed by capillary electrophoresis on an automated sequencer (Applied Biosystems), and computational analysis by using the GeneMapper v5.0 software (Applied Biosystems).

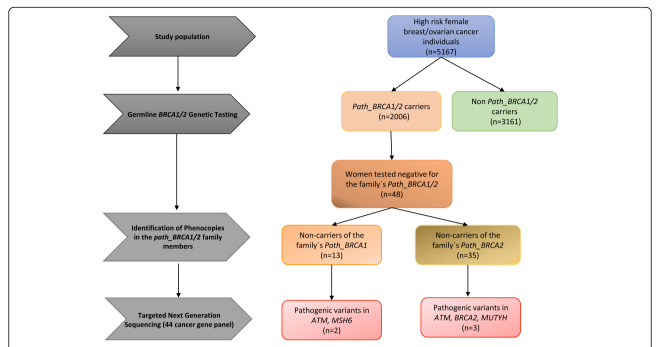
## Results

## Family history and clinical characteristics

In total, we identified 48 cases, of whom 18 BC or gynecological cancer patients who did not carry their respective families'  $path\_BRCA1$  or  $path\_BRCA2$  variants (n = 13 and n = 5, respectively) came from the Hereditary Cancer Biobank from the Norwegian Radium Hospital, while the Department of Genomic Medicine from the University of Manchester identified a total of 30 BC patients, all non-carriers of the family's  $path\_BRCA2$  variants (Fig. 1). The median age at first cancer diagnosis was 53.5 years (range 31–79 years). The incidence was higher for BC (92%), followed by ovarian cancer (4%) and endometrial and cervical cancer (2% each) (Table 1).

#### **Germline findings**

In the 48 cases, we identified five (10%) to carry pathogenic variants in *ATM* (c.468G > A, p.Trp156Ter and c.9139C > T, p.Arg3047Ter), *BRCA2* (c.9382C > T, p.Arg3128Ter), *MSH6* (c.2864delC, p.Thr955fs) and



**Fig. 1** Flow chart showing the study population selection from the Hereditary Cancer Biobank from the Norwegian Radium Hospital, Norway. It contains ascertained BC and breast/ovarian cancer kindreds by family history that were all subjected to genetic testing. The identification of phenocopies involved 48 women with prospective detected BC or gynecological cancer at follow-up, who were tested negative for their respective families' *path\_BRCA1/2* variants. Among these cases, 13 were identified in non-carriers of the family's *path\_BRCA1* variant and in 35 non-carriers of the family's *path\_BRCA2* variant (*n* = 30 from the Department of Genomic Medicine from the University of Manchester). Pathogenic variants were identified in 5/48 (10%) BC or gynecological cancer cases

MUTYH (c.1178G > A, p.Gly393Asp). Among these five cases, 2/13 were identified in non-carriers of the family's path\_BRCA1 variant and in 3/35 non-carriers of the family's path\_BRCA2 variant (Fig. 1). Disease type, familial path\_BRCA1/2 and pathogenic variants found in this study are shown in detail in Table 1.

Interestingly, one case with a familial *path\_BRCA2* (c.6591\_6592delTG) was found to carry another pathogenic variant in the same gene (*BRCA2* c.9382C > T, p.Arg3128Ter), which causes a premature stop in the codon 3128 and is known to be a high risk pathogenic variant (Table 1).

The pathogenic variants in BC-related genes (2 in *ATM* and 1 in *BRCA2*) were found in 3 women with BC or ovarian cancer, while the *MSH6* and the heterozygous *MUTYH* p.Gly393Asp pathogenic variant was found in a woman with endometrial cancer at 57 years and BC diagnosis at 56 years, respectively (Table 1).

#### Validation of the cancer gene panel output

The presence of the five pathogenic variants detected by targeted NGS was confirmed by cycling temperature capillary electrophoresis, showing 100% correspondence between both methods.

# Variants of unknown significance (VUS) and predicted protein alterations

In total, we found 26 unique VUS in 30 out of 48 patients (63%). Common polymorphisms (with an allele frequency  $\geq$  1% in the general population according to the ExAC database) and benign variants classified according to either ClinVar or the American College of Medical Genetics and Genomics (ACMG) guidelines were excluded from further analyses [41, 58].

The VUS were detected in 17 genes, namely: AXIN2, RAD51B (in 4 patients each), MAP3K1 (in 3 patients), APC, ATM, MSH2, NBN, POLE (in 2 patients each), BRCA1, CDH1, CDX2, DVL2, MRE11A, MUTYH, NOTCH3, PTEN and RAD51D (in 1 patient each) (Table 2). The minor allele frequencies (MAF) of these variants in public databases were very low or no frequency data have been reported (Table 2).

The VUS were furthermore analyzed by using 6 in silico protein prediction tools with different underlying algorithms (Fig. 2). The *MRE11A* c.1139G > A and the *MUTYH* c.881G > A variants were suggested to have a potentially damaging effect on protein level by all six predictions programs. For the variants in the *MSH2*, *NBN*, *POLE* and *BRCA1* genes (*MSH2* c.815C > T, *NBN* c.283G > A, *POLE* c.2459 T > C and *BRCA1* c.1927A > G,

 Table 1 Summary of the 48 prospective BC or gynecological cancer patients included in the study

Patient_ID	Institution	Familial path_ BRCA1 or path_BRCA2 variantFamilial path_BRCA1 or path_BRCA2 variant	ICD9 diagnosis (age)	Pathogenic variant identified in the current study		
17,161	HCBNRH	BRCA2 c.5217_5223delTTTAAGT (p.Tyr1739Terfs)BRCA2 c.5217_5223delTTTAAGT (p.Tyr1739Terfs)	OC (67)	ATM c.468G > A (p.Trp156Ter)*ATM c.468G > A (p.Trp156Ter)*		
6475	HCBNRH	BRCA1 c.1011dupA (p.Val340Glyfs)BRCA1 c.1011dupA (p.Val340Glyfs)	BC (52)	ATM c.9139C > T (p.Arg3047Ter)ATM c.9139C > T (p.Arg3047Ter)		
13,141	HCBNRH	BRCA1 c.1072delC (p.Leu358Cysfs)BRCA1 c.1072delC (p.Leu358Cysfs)	EC (57)	MSH6 c.2864delC (p.Thr955fs)*MSH6 c.2864delC (p.Thr955fs)*		
1873	HCBNRH	BRCA1 c.1556delA (p.Lys519Argfs)BRCA1 c.1556delA (p.Lys519Argfs)	MTHM (56), BC (70)	Not		
5378	HCBNRH	BRCA1 c.697_698delGT (p.Val233Asnfs)BRCA1 c.697_698delGT (p.Val233Asnfs)	BC (52)	Not		
5180	HCBNRH	BRCA1 c.5194-2A > CBRCA1 c.5194-2A > C	BC (39)	Not		
22	HCBNRH	BRCA2 c.3847_3848delGT (p.Val1283Lysfs)BRCA2 c.3847_3848delGT (p.Val1283Lysfs)	BC (63)	Not		
243	HCBNRH	BRCA2 c.3847_3848delGT (p.Val1283Lysfs)BRCA2 c.3847_3848delGT (p.Val1283Lysfs)	CVC (41)	Not		
5348	HCBNRH	BRCA1 c.1556delA (p.Lys519Argfs)BRCA1 c.1556delA (p.Lys519Argfs)	BC (68)	Not		
6031	HCBNRH	BRCA1 c.1556delA (p.Lys519Argfs)BRCA1 c.1556delA (p.Lys519Argfs)	BC (66)	Not		
6032	HCBNRH	BRCA1 c.3228_3229delAG (p.Gly1077Alafs)BRCA1 c.3228_3229delAG (p.Gly1077Alafs)	OC (55)	Not		
6207	HCBNRH	BRCA1 c.697_698delGT (p.Val233Asnfs)BRCA1 c.697_698delGT (p.Val233Asnfs)	BC (47)	Not		
8085	HCBNRH	BRCA1 c.3228_3229delAG (p.Gly1077Alafs)BRCA1 c.3228_3229delAG (p.Gly1077Alafs)	BC (55), CC (66)	Not		
11,717	HCBNRH	BRCA1 c.1556delA (p.Lys519Argfs)BRCA1 c.1556delA (p.Lys519Argfs)	BC(42,57)	Not		
12,470	HCBNRH	BRCA1 c.3178G > T (p.Glu1060Ter)	BC (39)	Not		
13,023	HCBNRH	BRCA2 c.5217_5223delTTTAAGT (p.Tyr1739Terfs)	BC (59)	Not		
15,529	HCBNRH	BRCA2 c.4821_4823delTGAins	BC (48)	Not		
22,325	HCBNRH	BRCA1 c.5047G > T (p.Glu1683Ter)	BC (45)	Not		
1,100,948	UM	BRCA2 c.6591_6592delTG (p.Glu2198Asnfs)	BC (44)	BRCA2 c.9382C > T (p.Arg3128Ter)		
12,010,643	UM	BRCA2 c.7360delA (p.lle2454Phefs)	BC (56)	MUTYH c.1178G > A (p.Gly393Asp)		
75,443	UM	BRCA2 c.5909C > A (p.Ser1970Ter)	BC (55)	Not		
88,295	UM	BRCA2 c.7977-1G > C	BC (44)	Not		
64,949	UM	BRCA2 c.5909C > A (p.Ser1970Ter)	BC (55)	Not		
67,723	UM		BC (46)	Not		

Table 1 Summary of the 48 prospective BC or gynecological cancer patients included in the study (Continued)

Patient_ID	Institution	Familial path_ BRCA1 or path_BRCA2 variantFamilial path_BRCA1 or path_BRCA2 variant	ICD9 diagnosis (age)	Pathogenic variant identified in the current study
		BRCA2 c.4866delA p.(Arg1622Serfs*14)		
84,510	UM	BRCA2 c.5946delT (p.Ser1982Argfs)	BC (67)	Not
13,007,862	UM	BRCA2 c.5909C > A (p.Ser1970Ter)	BC (31)	Not
9,009,462	UM	BRCA2 c.6535_6536insA (p.Val2179Aspfs)	BC (67)	Not
900,178	UM	BRCA2 c.1889delC (p.Thr630Asnfs)	BC (49,77)	Not
10,005,829	UM	BRCA2 c.9541_9554del p.(Met318CysfsTer13)	BC (38)	Not
10,007,016	UM	BRCA2 c.632-1G > A	BC (51)	Not
10,003,959	UM	BRCA2 c.6275_6276delTT (p.Leu2092Profs)	BC (55)	Not
12,852	UM	BRCA2 c.1929delG (p.Arg645Glufs)	BC (56)	Not
12,001,161	UM	BRCA2 c.7958 T > C (p.Leu2653Pro)	BC (67)	Not
13,017,067	UM	BRCA2 c.755_758delACAG (p.Asp252Valfs)	BC (74)	Not
688	UM	BRCA2 c.1929delG (p.Arg645Glufs)	BC (32)	Not
40,540	UM	BRCA2 c.8535_8538deIAGAG p.(Glu2846LysfsTer16)	BC (69)	Not
9,001,644	UM	BRCA2 c.4965C > G (p.Tyr1655Ter)	BC (39, 45)	Not
89,205	UM	BRCA2 c.5946deIT (p.Ser1982Argfs)	BC (77)	Not
10,002,068	UM	BRCA2 del exons 14–16	BC (37)	Not
10,004,590	UM	BRCA2 c.2672dupT	BC (67,67)	Not
40,286	UM	BRCA2 c.7069_7070delCT p.(Leu2357ValfsTer2)	BC (36,53)	Not
76,618	UM	BRCA2 c.4478_4481delAAAG (p.Glu1493Valfs)	BC (51)	Not
12,015,576	UM	BRCA2 c.9382C > T (p.Arg3128Ter)	BC (45)	Not
61,420	UM	BRCA2 c.5350_5351delAA p.(Asn1784HisfsTer2)	BC (59)	Not
960,579	UM	BRCA2 c.2808_2811del4 (p.Ala938Profs)	BC (39)	Not
14,965	UM	BRCA2 c.5682C > G p.(Tyr1894Ter)	BC (59)	Not
20,468	UM	BRCA2 c.6275_6276delTT (p.Leu2092Profs)	BC (38)	Not
56,193	UM	BRCA2 c.7884dupA (p.Trp2629Metfs)	BC (79)	Not

HCBNRH Hereditary Cancer Biobank from the Norwegian Radium Hospital (Norway), UM University of Manchester (United Kingdom), ICD9 diagnosis International Classification of Diseases, 9th Revision, OC Ovary cancer, BC Breast cancer, EC Endometrial cancer, MTHM Malignant neoplasm of thymus, heart, and mediastinum, CC Colon cancer, CVC Cervical cancer, \*Considered pathogenic based in its nature (nonsense and frameshift), VUS Variants of unknown significance, NM for ATM NM\_000051, BRCA1 NM\_007294.3, BRCA2 NM\_000059.3, MSH6 NM\_001281492, MUTYH NM\_012222

five out of six predictions suggested a potentially damaging effect (Fig. 2).

Discrepancies in protein-related predictions were even more pronounced for the variants in *APC*, *AXIN2*,

*RAD51B, DVL2, RAD51D, CDH1* and *MSH2* c.2164G > A. In contrast, none of the six prediction tools showed deleterious effects for the detected variants in the *AXIN2, ATM, RAD51B* and *MAP3K1* genes (*AXIN2* 

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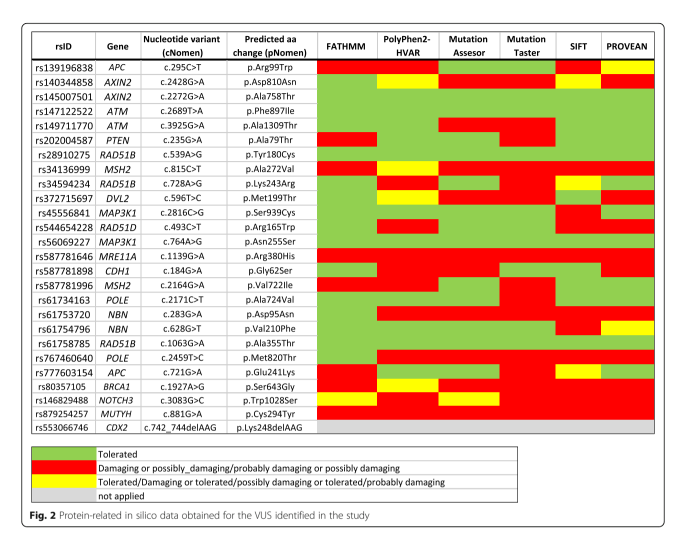
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Cryptic splice site- dedicated analyses	Potential local	splice effect					New Acceptor Site?	New Donor Site?						I		
		VAR vs	⊘(%)	0	0	0	0	-4.8	0	0	0	0	0	0	0	0
		Var		86. 5179	100	86. 8244	78. 4708	82. 5488	87. 3058	86. 1925	84. 8039	86. 1925	87. 0697	80.2	86. 8647	95. 7456
	SSFL	₩		86. 5179	100	86. 8244	78. 4708	86. 6769	87.	86. 1925	84. 8039	86. 1925	87. 0697	80.2	86. 8647	95. 7456
		VAR vs WT	(%) ∇	0	0	0	0	-11.7	0	0	0	0		0	0	0
		Var		8.17	10.8	6.19	7.5 2484	8.7	8.8 6265	6.3	7.4 9577	6.3	7.1 5277	11.8	9.6 515	9.9
	MES	$\mathbb{R}$		8.17	10.8	6.19	7.52	9.89	8.86 265	6.34	7.49	6.34	7.15	8.	9.6 515	941
ated		Туре	(3' or 5'ss)	m	ĩο	ñ	ìο	ìο	ñ	m	ñ	ñ	ìο	ώ	2ís	ñ
Reference splice site-dedicated analyses	Nearest reference	Distance	(nt)	21	-47	44	-71	8	12 57	35	75	35	6	27	-19	14
Non- Finnish European population frequency*				5.99 e-05	8.99 e-05	0.000	0.0269	0.00	N A	39861	0.00	39861	0.00	0.00 71658	0.000	4.5e-05
dbSNPrsID				587, 781, 898	587, 781, 996	61,754, 796	56,069, 227	61,734, 163	80,357, 105	145,007, 501	139, 196, 838	145, 007, 501	777, 603, 154	61,758, 785	202, 004, 587	587, 781, 646
Predicted protein change (pNomen)				p.Gly6 2Ser	p.Val 722lle	p.Val 210 Phe	p.Asn 255 Ser	p.Ala 724 Val	p.Ser 643 Gly	p.Ala 758 Thr	p.Arg 99Trp	p.Ala 758 Thr	p.Glu 241 Lys	p.Ala 355 Thr	p.Ala 79 Thr	p.Arg 380His
Nucleotide change (cNomen)				c.18 4G > A	c.216 4G> A	c.62 8G > T	c.76 4A > G	c.21 71C > T	c.19 27A > G	c.22 72G > A	c.29 5C > T	c.22 72G > A	c.72 1G > A	c.10 63G > A	c.23 5G > A	c.11 39G > A
Exon				m	13	9	т	19	10	01	4	10	_	=	4	11
Gene				CDH1	MSH2	NBN	MAP 3K1	POLE	BRCA1	AXIN2	APC	AXIN2	APC	RAD 518 RAD 518	PTEN PTEN	MRE 11A MRE 11A
Genomic position (GRCh37)				chr_16_68 835593 _G_A	chr2_47 703664 _G_A	chr_8_90 983475 _C_A	chr_5_56 155672 _A_G	chr 12_13 3244944 _G_A	chr17_4 1245621 _T_C		chr5_11 2102960 _C_T		chr5_11 2128218 _G_A	chr_14_6 9061228 _G_A	chr10_89 690828 _G_A	chr11_94 197365 _C_T
Patient ID				889			1873	5378	6031			12, 470		12, 852	88, 295	900,

	φΔ			0	ı	0.54	03.	ı	ı	0.03	0.04	19.	0.1	0.04	0.01	90:00
	ΔHzei			-16.1	-100.08	-1.77	-46.5	-14.33	-50.64	24.6	32.96	-40.	-6.4	-7.06	87.95	-2.13
ESR- dedicated analyses	ΔtESRseq			-0.4 86881	-2.4 6964	0.0 509 416	-2.1 7832	-1.2 2987	-1.2 4035	0.31 8238	0.67 6556	-1.4 8785	0.3 00 115	0.4 %	0.5 54 269	1.2 8743
	Local MES scores	Var		1					ı			7.9				
	Local A	M		1					ı			6:0				
Cryptic splice site- dedicated analyses	Potential local	splice effect		ı					I			Cryptic 5'ss activation?				
		VAR vs	⊘(%)	0	0	0	0	0	0	0	0	0	0	0	0	0
		Var		100	87. 4307	80.	84. 3224	87. 3948	80.2	94.	84. 8076	78. 9497	82. 5954	72. 818	93. 4253	77. 9039
	SSFL scores	M M		100	87. 4307	80. 4452	84. 3224	87. 3948	80.2	94.	84.	78. 9497	82. 5954	72.	93. 4253	77. 9039
		VAR vs WT	√ (%)	0	0	0	0	0	0	0	0	0	0	0	0	0
		Var		12.	11.	6. 34 467	10. 3527	11.	11.8	10. 7663	9.9	9.0	11.	6. 31089	9.	6. 58677
	MES scores	M_		12.0	11. 7045	6.34	10. 3527	11. 6727	11.8	10. 7663	9.9 8517	9.09184	11.	6. 31089	9. 8979	6. 58677
ted		Туре	(3' or 5'ss)	ά	π´	2,	π́	3,	3,	2,		2,	2,	2,	π´	2,
Reference splice site-dedicated analyses	Nearest reference	Distance	(nt)	447	55	-61	23	23	27	-38		-29	09-	444	51	-10
Non- Finnish European population frequency*				0.0221	0.000	6.01e-05	3755	1.5e-05	0.007	0.00 30459	0.000	0.01	na	na	4.5e-05	0
dbSNPrsID				45,556, 841	553, 066, 746	372, 715, 697	34,136, 999	140, 344, 858	61,758, 785	61,753, 720	149, 711, 770	34,594, 234	rs14682 9488	rs8792 54257	147, 122,522	767, 460, 640
Predicted protein change (pNomen)				p.Ser 939 Cys	p.Lys 248de IAAG	p.Met 199 Thr	p.Ala 272 Val	p.Asp 810 Asn	p.Ala 355 Thr	p.Asp 95 Asn	p.Ala 1309 Thr	p.Lys 243 Arg	p.Trp 1028 Ser	p.Cys 294 Tyr	p.Phe 897Ile	p.Met 820 Thr
Nucleotide change (cNomen)				c.28 16C > G	c.742 _744 del	c.59 6 T > C	c.81 5C > T	c.24 28G > A	c.10 63G > A	c.28 3G > A	c.39 25G > A	c.72 8A > G	c.30 83G> C	c.88 1G > A	C.26 89 T > A	C.24 59 T > C
Exon				4	m	5	·5	=======================================	11	m	56	_	19	10	18	21
Gene				MAP 3K1	CDX2	DVL2 DVL2	MSH2	AXIN2	RAD 518	NBN	АТМ	RAD 518	NOT CH3	MUT YH	АТМ	POLE
Genomic position (GRCh37)				chr_5_56 177843 _C_G	chr13_28 537449 _ACTT_A	chr_17_ 7133187 _A_G	chr2_47 641430 _C_T	chr_17_6 3526198 _C_T	chr_14_6 9061228 _G_A	chr8_9 0993640 _C_T	chr_11_10 8155132 _G_A	chr_14_68 353893 _A_G	chr19_15 291551 _C_G	chr1_45 797881 _C_T	chr_11_10 8139187 _T_A	chr_12_133 241897 _A_G
Patient ID				960,	1,000,	1,100, 948		10,002, 068	10,005, 829			12,001, 161	12,015, 576	11,717	17, 161	22

Table 2 RNA splicing- dedicated in silico analyses for the VUS identified in our study (Continued)

	φΦ			19	-0.	60	1.42
	ΔHzei			7.23	6.7	0.12	-22. 32
ESR- dedicated analyses	ΔtESRseq ΔHzei			0.8 81 539	-1.1 8661	-0.9 42 617	–2.5 5724
	Local MES scores	Var			ω ω		
	Local M	M			1		
Cryptic splice site-dedicated analyses	Potential local	splice effect			New Acceptor Site?		
		VAR vs WT	⊲ %	0	0	0	0
		Var		83.	78. 4708	86. 1925	85. 1161
	SSFL	₩		83.	78. 4708	86. 1925	85. 1161
		VAR vs WT	√ (%)	0	0	0	0
		Var		9. 54919	7.52	6.34	8.20
	MES	M M		9. 54919	7. 52484	6. 34671	8. 20686
ted		Туре	(3' or 5'ss)	22	2,5	ω̈́	ώ
Reference splice site-dedicated analyses	Nearest reference	Distance	(nt)	-34	-71	35	13
Non- Finnish European population frequency*				0.00 45 906	0.0269	0.003 9861	6.94 e-05
dbSNPrsID				28, 910, 275	56,069, 227	145, 007, 501	544, 654, 228
Predicted protein change (pNomen)				p.Tyr1 80Cys	p.Asn 255Ser	p.Ala 758 Thr	p.Arg 165 Trp
Nucleotide Predicted change protein (cNomen) change (pNomen)				c.53 9A > G	c.76 4A > G	c.22 72G > A	c.49 3C > T
Exon				9	m	0	9
Gene				RAD 51B	MAP 3K1	AXIN2	RAD 51D
Patient Genomic ID position (GRCh37)				chr_14_68 352672 _A_G	chr_5_5 6155672 _A_G	chr_17_63 530163 _C_T	chr_17_33 433488 _G_A
Patient ID						6207	6475

not available, \*Non-Finnish European population based on ExAC database; NM for APC: NM\_000038; ATM: NM\_000551; AXIN2: NM\_00455; BRCA1: NM\_007300; CDH1: NM\_004360; CDX2: NM\_012: NM\_012: NM\_00422; MAPSK1: NM\_00521; MRH2: NM, 000551; MRH2 NA: NM\_00551; MRH2: NM, 000531; MRH2: NM, 0005

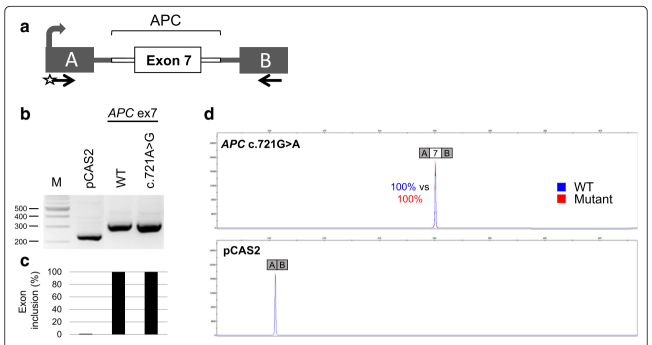


c.2272G > A, ATM c.2689 T > A, RAD51B c.539A > G and c.1063G > A and MAP3K1 c.764A > G) (Fig. 2).

# Splicing-dedicated in silico analysis and minigene splicing assays

Out of the 26 unique VUS, two (APC c.721G > A and MAP3K1 c.764A > G) were bioinformatically predicted as the most likely to affect RNA splicing, either by potentially creating a new splice site or by altering putative exonic splicing regulatory elements, respectively (Table 2). Given that RNA data was not available for APC c.721G > A, we set out to experimentally evaluate the impact on RNA splicing produced by this variant, by performing a cell-based minigene splicing assay. As shown in Fig. 3, we observed that c.721G > A did not affect the splicing pattern of APC exon 7 in our system. These results are reminiscent of those recently obtained for MAP3K1 c.764A > G by using a similar splicing assay, in which the variant did not cause an alteration in the minigene's splicing pattern (Dominguez-Valentin et al. under submission). It would be important in both cases to validate the minigene results by analyzing RNA from the variant carriers/patients as compared to those from healthy controls. However, we do not have such material in our biobank.

To our knowledge, the only other VUS from our list for which RNA data is available is MSH2 c.815C > T (p.Ala272Val). Previous results from different minigene assays revealed that, albeit located outside the splice sites, MSH2 c.815C > T induces partial skipping of exon 5 [28]. These results agree, at least in part, with those obtained by analyzing RNA from a LS patient carrying this same variant [29]. Indeed, the latter study revealed aberrantly spliced MSH2 transcripts associated with the presence of c.815C > T, but where the severity of the splicing defect was not addressed at the time. Of note, here we identified MSH2 c.815C > T together with another VUS (DVL2 c.596 T > C) and a path\_BRCA2 c.9382C > T (different from the familial path\_BRCA2) in a patient diagnosed with ductal carcinoma at 44 years of age (Patient 1,100,948) (Table 1).



**Fig. 3** Analysis of the impact on RNA splicing of *APC* c.721G > A by using a cell-based minigene splicing assay. **a** Structure of pCAS2-APC.ex7 minigene used in the assay. The bent arrow indicates the CMV promoter, boxes represent exons, lines in between indicate introns, and arrows below the exons represent primers used in RT-PCR reactions. The WT and c.721G > A minigenes were generated by inserting a genomic fragment containing the exon of interest and flanking intronic sequences into the intron of pCAS2, as described under Materials and Methods. **b** Analysis of the splicing pattern of pCAS2-APC.ex7 WT and c.721G > A minigenes. The two constructs were introduced into HeLa cells and the minigenes' transcripts were analyzed by RT-PCR 24 h post-transfection. The image shows the results of a representative experiment in which the RT-PCR products were separated on a 2.5% agarose gel stained with EtBr and visualized by exposure to ultraviolet light. M, 100 bp DNA ladder (New England Biolabs). **c** Quantification of splicing events observed in the minigene splicing assay. The relative levels of exon inclusion indicated under the gel are based on RT-PCR experiments equivalent to those shown in B but performed with a fluorescent forward primer and then separated on an automated sequencer under denaturing conditions. Quantification results were obtained by using the GeneMapper v5.0 software (Applied Biosystems) and correspond to the average of two independent fluorescent-RT-PCR experiments. **d** Representative fluorescent RT-PCR experiments. The panel shows superposed peaks corresponding to the WT and mutant products (in blue and red, respectively), as indicated

## Discussion

Among prospectively detected BC or gynecological cancer phenocopies in the path\_BRCA1/2 families, we found that 4/48 have pathogenic variants in highpenetrance cancer genes: two BC- and one CRCassociated gene (ATM, BRCA2 and MSH6, respectively). Our findings are in line with a previous study, which detected a likely pathogenic variant in a gene other than BRCA1/2 in a BC patient, i.e. MSH6 c.3848\_3862del (p.(Ile1283\_Tyr1287del) [30]. In addition, we found the MUTYH c.1178G > A (p.Gly393Asp) variant in a BC case, which is one of the most common path\_MUTYH variants. Pathogenic MUTYH variants may cause a recessively inherited colon cancer syndrome. Whether or not individuals who are heterozygous for MUTYH mutations may be at risk for cancer is debated [31]. Among the five cases found to carry pathogenic variants, 2/13 were identified from families with path\_BRCA1 and 3/ 35 with *path\_BRCA2* variants.

Our results are in concordance with the recently published NGS panel studies, which have demonstrated that

besides high-risk genes, like *BRCA1/2* and MMR genes, other genes may also contribute to familial cancer predisposition, thus providing a broader picture on the genetic heterogeneity of cancer syndromes [25, 32, 33]. In this regard, a molecular diagnosis yield of approximately 9% to identify a pathogenic or likely pathogenic variant in BC has been reported, and with yields of 13% in ovarian and 15% in colon/stomach cancer cases [25]. On the other hand, family history is currently used to identify high risk patients. However, the use of family history fails to identify women without close female relatives who are carriers of pathogenic variants [9].

Despite the potential of NGS to identify genetic causes among families that tested negative for pathogenic variants in high-risk genes using traditional methods [25, 32, 33], a high number of VUS are also detected and constitute a major challenge in oncogenetics [34]. In this study, we subjected 26 VUS to RNA splicing and protein in silico evaluations, and the bioinformatics predictions indicated that two VUS (*APC* c.721G > A and *MAP3K1* c.764A > G) were likely to affect RNA splicing. Our

results from minigene splicing assays suggest, however, that this is not the case. Complementary analysis of patients' RNA will be important to verify the impact on splicing of these variants in vivo. Of note, none of the six protein in silico prediction tools showed a deleterious effect for the MAP3K1 c.764A > G missense variant and inconsistences were found for the APC c.721G > A variant.

Bioinformatics prediction tools are widely used to aid the biological and clinical interpretation of sequence variants, although it is well recognized that they have their limitations. Co-segregation studies for further evaluation will be key for understanding whether some of the VUS detected in this work may have a causal effect. Some of the VUS may in the future be reclassified as deleterious or benign, but in the meantime, they cannot be used to make clinical decisions [30].

A polygenic model involving a combination of multiple genomic risk factors, including the effect of low- or moderate- penetrance susceptibility alleles may explain the increased BC risk in women who tested negative for family's *path\_BRCA1/2* variants [5]. In addition, heterozygous whole gene deletions (WGD) and intragenic microdeletions have been reported to account for a significant proportion of pathogenic variants underlying cancer predisposition syndromes, although WGD were not a common mechanism in any of the three high-risk BC genes, *BRCA1*, *BRCA2* and *TP53* [35].

The clinical utility of gene panels such as the one used in this study is not yet fully established and the appropriate routes for clinical deployment of such tests remain under discussion [36]. So far, the large patient datasets generated by NGS panels may be used to explore the specific penetrance of the genes included in these panels, and to assess the performance and implications of the use of NGS in clinical diagnostics [34].

#### **Conclusions**

In kindreds carrying *path\_BRCA1/2* variants, testing only for the already known *path\_BRCA1/2* variants in the family may not be sufficient to exclude increased risk neither for BC nor for ovarian cancer or other cancers in the healthy female relatives. Our findings suggest that all women in BC or breast/ovarian cancer kindreds would benefit from being offered genetic testing irrespective of which causative genetic variants have been demonstrated in their relatives. In addition, we found a number of VUS in genes other than *BRCA1/2* i.e. *AXIN2*, *APC*, *DVL2*, *MAP3K1*, *RAD51B*, *NBN*, *POLE*, *CDH1*, *CDX2*, *MRE11A*, *MUTYH*, *NOTCH3*, *PTEN* and *RAD51D*. All these may be suspected of being associated with cancer in the families studied and may be considered as candidates for being included in future gene

panel testing to better understand why some families present aggregation of cancer cases.

#### **Additional files**

**Additional file 1:** The concentration in a 10 ml PCR was 1xThermopol Reaction Buffer with 2 mM MgS04, 0.3 μM "reverse" primers, 0.15 μM "forward" primer, 0.1 μM, 6-Carboxyfluorescein-GC clamp primer, 600 μM dNTP, 100 μg Bovine Serum Albumine (Sigma-Aldrich, Oslo, Norway) and 0.75 U Taq DNA polymerase. Plates were sealed with two strips of electrical tape (Clas Ohlson, Oslo, Norway). The temperature cycling was repeated 35 times; 94 °C for 30 s, annealing temperature held for 30 s and extension at 72 °C for 60 s (Eppendorf Mastercycler ep gradient S (Eppendorf, Hamburg, Germany)). **Table S1.** primers used to amplify PCR product to be analysed by cycling temperature capillary electrophoresis. (DOCX 16 kb)

Additional file 2: Primers used in the pCAS2 minigene splicing assay. (DOCX 14 kb)

#### Abbreviations

ACMG: American College of Medical Genetics and Genomics; BC: Breast cancer; BIC: Breast Cancer Information Core Database; CRC: Colorectal cancer; ENIGMA: Evidence-based Network for the Interpretation of Germline Mutant Alleles; ESR: Exonic splicing regulatory elements; HGMD: Human Gene Mutation Database; InSiGHT: International Society of Gastrointestinal Hereditary Tumors Database; LOVD: Leiden Open Variation Database; LS: Lynch syndrome; MAF: Minor allele frequency; MES: MaxEntScan; NGS: Next generation sequencing; path\_BRCA1/2: Pathogenic (disease-causing) variant of the BRCA1 or the BRCA2 genes; SNPs: Single nucleotide polymorphisms; SNV: Single-nucleotide variants; SSFL: SSF-like; VUS: Variants of unknown significance; WGD: Whole gene deletions; WT: Wild type

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#### Availability of data and materials

All data generated or analyzed during this study are included in the manuscript.

#### Authors' contributions

All authors have taken part in the different steps of the study: MDV, DGRE, PM and EH designed the study, AM, HT performed in silico splicing predictions and the minigene assays, POE performed validation experiments, MM, AN and EHF performed in silico protein predictions, SN, DV performed the sequence analysis. MDV drafted the manuscript and all have read, revised and approved the manuscript.

#### Ethics approval and consent to participate

Ethical approval for the prospective study was granted from the Norwegian Data Inspectorate and Ethical Review Board (ref 2015/2382). All examined patients had signed an informed consent for their participation in the study.

#### Consent for publication

Not Applicable.

#### Competing interests

The authors declare that they have no competing interests.

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