BRCA1/BRCA2: CanVIG-UK Gene-Specific Guidance



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A Garrett¹, L Loong¹, L King¹, S Allen¹, M Durkie², J. Drummond³, G.J. Burghel⁴, R. Robinson⁵, A Callaway^{6,7}, I. Berry⁵, A. Wallace⁴, S. Ellard⁸, E Baple⁸, H. Hanson^{1,9}, C.Turnbull^{1,10}

- 1) Division of Genetics and Epidemiology, The Institute of Cancer Research, London, UK.
- 2) Sheffield Diagnostic Genetics Service, Sheffield Children's NHS Foundation Trust
- 3) East Anglian Medical Genetics Service, Cambridge University Hospitals NHS Foundation Trust, Cambridge, UK
- 4) Manchester Centre for Genomic Medicine and NW Laboratory Genetics Hub, Manchester University Hospitals NHS Foundation Trust, Manchester, UK
- 5) Yorkshire Regional Genetics Service, Leeds Teaching Hospitals NHS Trust, Leeds, UK
- 6) Wessex Regional Genetics Laboratory, Salisbury NHS Foundation Trust, Salisbury, UK
- 7) Human Genetics and Genomic Medicine, Faculty of Medicine, University of Southampton, Southampton, UK
- Department of Molecular Genetics, Royal Devon & Exeter NHS Foundation Trust, Exeter, UK
- 9) St George's University Hospitals NHS Foundation Trust, Tooting, London, UK
- 10) The Royal Marsden NHS Foundation Trust, Fulham Road, London

For use in conjunction with CanVIG-UK Consensus Specification for Cancer susceptibility Genes of ACGS Best Practice Guidelines for Variant Classification. Evidence lines for which there are no gene-specific recommendations should be reviewed in context of CanVIG-UK Consensus Specification for Cancer Susceptibility Genes.

Evidence element and evide strengths allowed	ence	Thresholds/data-sources/applications specifically relevant to BRCA1/BRCA2
PS4: Case-control: The prevalence of the variant in affected individuals is significantly increased compared with the prevalence in controls	_VSTR _STR _MOD	 NHSD case control data can be used for case-control analysis: Controls should represent appropriate ethnicity and sex matching (i.e. female non-cancer NFE controls should be used if the case series consists predominantly of females, as with the current NHSD case series) and series denominator As this is an enriched series, OR≥10 is required Current data/denominator counts for base substitutions are available at <u>CanVar-UK</u> For non-base-substitutions i.e. deletions/duplications/insertions, NHSD counts can be accessed from <u>CanVIG-UK</u> If there are insufficient data to perform case-control analyses, PS4_sup can be applied: if there are observations of the variant in ≥5 different families with a pattern of diagnoses consistent with a hereditary breast and ovarian cancer syndrome and the variant is very rare or absent in control populations (i.e. PM2 has been applied)
PM2: Absent from controls (or at extremely low frequency if recessive) in ESP, 1000GP, or ExAC	_MOD _SUP	Cancer-free female controls (of any/all ethnicities) should be used (due to low penetrance in male pathogenic variant carriers). Otherwise, the main CanVIG-UK consensus guidance should be followed.

Evidence towards Pathogenicity

PVS1: Predicted null	VSTR	It is pred	dicted that trur	ncating variants occurring at the 3' end of				
variant (in a gene where	STR	the gene	the gene will not undergo NMD. The residues below demarcate					
LOF is a known mechanism	MOD	the cons	sensus bound	ary, 3' of which protein truncating variants				
of disease)	SUP	are not	established to	result in NMD and/or impairment of				
	_001	function	of residual pr	otein.				
		BRCA1 (NM_007294.3): 1855 ¹						
		BRCA2 (NM_000059.3): 3309 ²						
		Based c	on ENIGMA re	commendations, as re-initiation sites have				
		also bee	en shown to re	esult in the loss of important functional				
		domains	s in BRCA1 ar	nd BRCA2, it is acceptable to use PVS1 at				
		a very s	strong level for	variants identified within the first 1000p of				
		both BRCA1 and BRCA2'.						
		known t	o lead to natu	rally occurring in-frame RNA isoforms that				
		may res	scue dene fun	ctionality ENIGMA has compiled the				
		below lis	st of splice val	riants for which the variant transcript may				
		be funct	tional and for s	some of which PVS1 may not be				
		applicat	ole.	·····				
		Gene	Region	Bases				
			intron 5	c.301+1				
			intron6	c.442-1				
			(exon 7 acceptor)	c.442-2				
				c.548-1				
				c.593+1				
			introns 8.9	c.593+2				
		BRCA1		c.594-1				
				c.670+1				
				c.670+2				
			intron 10 (oven 10 deper)	c.4096+1				
			intron 11	c.4186-1				
			(exon 12 acceptor)	c.4186-2				
			intron12	c.4358-1				
			(exon 15 accepor)	c.6842-1				
		BRCA2	intron12	c.6842-2				
		DITON		c.6937+1				
		Adapted	l from ENIGM	<u>10.0937+2</u> 10.2017 ¹				
PS1: Same amino acid	STR	Within f	orthcoming FI	NIGMA guidance it is anticipated that these				
change as an established	_0	element	ts will all be in	corporated within PP3 and only awarded to				
variant		variants	within key do	mains:				
PM4: Protein-length-	_MOD	In the ir	nterim, we re	commend:				
changing variant	_SUP	• Use	of PM1_sup a	and/or PM4_sup for any variant within				
PM5: Novel missense	_MOD	BRC	CA1 RING (aa	1-101), BRCT (aa1650-1863) COILED-				
change at an amino acid	_SUP	COI	L DOMAIN (a	a 1391-1424) and BRCA2 DNA-binding				
residue where a different		dom	ain (ad 2481-3	5180)				
missense change		• USE						
determined to be pathogenic			G. 10, 22, 31, T. 1695 1699	33, 41, 44, 47, 01, 04, 71 2 1607 1600 1706 1709 1715 1726				
seen before		1720	21.1000,1000 8 1720 1770	0, 1037, 1033, 1700, 1700, 1710, 1730, 1764 1766 1770 1775 1786 1837				
PP3: In silico: Multiple lines	_SUP	1839	0, 1700, 1740, 8 1830 1853	, 170, 1700, 1770, 1770, 1700, 1007,				
		DRF) 2607 2626	2627, 2663, 2722, 2723, 2748, 3052				
on the gene or gene product		3124	4	,,,, _, _, _, _, , , , ,				
	STD	 PM1 cannot be used where functional data are being used 						
Enrichment/constraint		for F	PS3, as per m	ain CanVIG-UK guidance				
PP2: Missense variant in a		• PP2	should not be	e used for BRCA1/BRCA2				
gene that has a low rate of	_30P	• Use	of PM5, PS1,	PP3 otherwise as per CanVIG-UK				
benign missense variation		Con	sensus Specil	fication				
and in which missense			•					
variants are a common								
mechanism of disease								

PM1: Located in a mutational hot spot and/or critical and well-established functional domain (e.g. active site of an enzyme) without benign variation PS3: Functional: Well- established in vitro or in vivo functional studies supportive of a damaging effect on the gene or gene product	_VSTR _STR _MOD _SUP	BRCA1: Findlay et al, 2018 ⁴ : Strong Bouwman et al, 2020 ⁵ : Strong Fernandes et al, 2019 ⁶ : Supporting Petitalot et al, 2019 ⁷ : Supporting BRCA2: Guidugli et al, 2018 ⁸ /Hart et al, 2019 ⁹ /Richardson et al, 2021 ¹⁰ : Strong See CanVIG Functional Assays Scores See the table at the bottom of this document for guidance on combining accounted
PP1: Co-segregation with disease in multiple affected family members in a gene definitively known to cause the disease	_VSTR _STR _MOD _SUP	Segregation evidence extracted from multifactorial analysis data can be used within PP1/BS4 using the thresholds specified in the PP5/BP6 guidance. Where combined with multiple evidence of other types, segregation evidence from multifactorial analysis data should be incorporated into the PP5/BP6 criteria Meiosis counting approaches may be used in addition if this evidence comes from families not already included in the multifactorial analyses. Evidence cannot exceed 'Very strong'
(maternity and paternity confirmed/unconfirmed) in a patient with the disease and no family history PM3: in trans with a pathogenic variant (recessive disorders)	_MOD _SUP _MOD _SUP	 Frequency data regarding co-occurrence in trans extracted from multifactorial analyses should be incorporated into PM3 or BP2 using the thresholds described in the PP5/BP6 guidance. Where combined with multiple evidence of other types, frequency data regarding co-occurrence in trans from multifactorial analyses should be incorporated into PP5/BP6 <i>In addition</i>, the <u>SVI recommendations for in trans Criterion</u> (PM3) can be used for either BRCA1 or BRCA2 for individuals with a Fanconi anaemia phenotype if this evidence comes from families not already included in the multifactorial analyses used for PP5. Evidence towards a Fanconi phenotype comprise: Clinical: diagnosis of childhood cancer or skeletal/structural/developmental abnormalities Molecular/Cellular: aberration on mitomycin-induced chromosomal breakage +/- depletion of BRCA2 in lymphocytes Both clinical and molecular/cellular aberrations must be present for a case to contribute to evidence Evidence cannot exceed 'Strong' Note: Caution is required in inferring the pathogenicity for the monoallelic phenotype, as variants may be hypomorphic (e.g. a variant contributing and causing a Fanconi anaemia phenotype
		variant contributing and causing a Fanconi anaemia phenotype may be low penetrance for breast cancer). Where the majority of evidence for variant pathogenicity comes from observations of the variant in cases of Fanconi Anaemia, it may be appropriate to comment on this in the clinical report

PP5: Reputable source	_VSTR	Published mul	tifactorial anal	ysis data providin	g likelihood ratios	
recently reports variant as	_STR	(LR) or log like	elihood ratios ((LLR) may be use	d as data sources	
pathogenic, but the	_MOD	encompassing	:	-		
the laboratory to perform an	_SUP	Segregation (PP1/BS4) Specificity of familial and/or tumour phonetype (PD4)				
independent evaluation		 Specificity of familial and/or tumour phenotype (PP4) Co. occurrence in trans. (PM2/PP2) 				
		• 00-000				
		Where individu	al likelihood r	atios for a particu	llar evidence type	
		do not line up with evidence (exponent) points required for a specific evidence strength, the combined LLR/LR				
		encompassing	multiple evide	ence types can be	e used instead to	
		represent the totality of evidence and applied within PP5				
		0 1 1				
		Suitable analys	Ses:			
			et al. 2007^{11}			
		 Elliuoi Parson 	etal 2020 ¹³	3		
		Whore multiple	notontially y	olid I D/I I Do oro	ovoilable for a	
		variant the va	lue from the m	allu LR/LLRS ale	available for a	
		used.				
		Where evidence	ce is supplied	as a LR (likelihoo	d ratio, e.g.	
		Parsons et al,	2020) this sho	build be converted	to Evidence	
		(Exponent) por	ints using the	lable below.		
		Where evidence	ce is supplied	as a natural LLR	(log likelihood	
		ratio, e.g. East	on et al, 2007), this should be a	converted to a LR	
		(for example u	el) before			
		conversion to l	Evidence (Exp	onent) Points us	ing the table below	
		(i.e. converted		0 a LLR base 2.0	0)	
		Likolibood	Evidence	Evidence		
		Ratio	(Exponent)	Strength		
			Points	ou ip		
		2.1	1	SUP		
		4.3	2	MOD		
		9	3			
		18.7	4	SIR		
		38.9	5			
		81	6			
		100.4	/	VOTD		
		350.4	8	VOIR	OR	
		PP5 can be ap	oplied at suppo	orting level on the	basis of any	
		• ≥2 acci	redited North	American comme	ercial diagnostic	
		laborat	ories OR		i olai alagnoolio	
		• ≥1 Nor	th American c	ommercial diagno	ostic laboratory	
		where	there is explic	it citation of utilisa	tion of otherwise	
		unavail		e from their data s		
		approv FNIGM	eu Ciingen Ex 1A	xpert Group (3 Sta	ai on Gillivar), le	
			17 1			
		This is an exc	eptional appli	cation, as per UK	-ACGS	
		specification.		less if is set a		
		For conflicts w		ciassifications, cor	TRACT ENIGMA.	

PP4: Phenotypic specificity (Patient's phenotype or family history is highly specific for a disease with a single genetic aetiology)	_STR _MOD _SUP	Tumour and family history phenotypic data extracted from multifactorial analyses should be incorporated into PP4 or BP5 using the thresholds described in the PP5/BP6 guidance. Where combined with multiple evidence of other types, tumour and family history phenotypic data from multifactorial analyses are incorporated into the PP5 evidence criterion. Patient phenotypic evidence whose strength cannot be quantified should not be used. Evidence cannot exceed 'Strong'
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Evidence towards Benignity

BA1/BS1: Allele frequency is "too high" in ExAC or gnomAD for disorder	_SA _STR	BA1: MTAF = 0.001 (0.1%) BS1: MTAF = 0.0001 (0.01%) The U95%CI should be used as the filtering allele count for the
		MTAF. This can be calculated using <u>cardiodb</u> or within gnomAD (see <u>training resources</u> from Miranda Durkie for methodology) Cancer-free female controls should be used (due to low penetrance in male pathogenic variant carriers)
BS2: Observation in	_STR	
controls inconsistent with	_SUP	
Observed in a healthy adult		
individual for a recessive		
(homozygous), dominant		
(heterozygous), or X-linked		
(hemizygous) disorder, with		
full penetrance expected at		
an early age		
BP4: In silico: Multiple lines	_SUP	
of computational evidence		
or gene product		
(conservation, evolutionary,		
splicing impact, etc.)		
BP1: Missense variant in a	_SUP	Can be used for missense variants with no predicted splicing
gene for which primarily		effect (as per main CanVIG-UK consensus specification) at non-
truncating variants are		conserved residues outside of BRCA1 RING (aa 1-101), BRC1
known to cause disease		(du 1050-1805) COILED-COIL DOMAIN (du 1591-1424) and BRCA2 DNA-binding domain (gg 2481-3186)
BP7: Synonymous (silent)	SUP	
variant for which splicing		
prediction algorithms predict		
no impact to the splice		
consensus sequence	0115	
BP3: In-frame	_SUP	
repetitive region		
BS3: Well-established in	STR	
vitro or in vivo functional	MOD	
studies show no damaging	SUP	
effect on protein function or	_	
splicing		* DD 4
BS4: Non segregation with	_STR	See PP1
uisease	_SOP	
BP2: Observed in trans	STR	*see PM3
with a pathogenic variant	SUP	
for a fully penetrant	_001	
dominant gene/disorder or		
observed in cis		

BP6: Reputable source	_STR	*see PP5			
benign, but the evidence is not available to the laboratory to perform an	_50P	Likelihood Ratio	Evidence (Exponent) Points	Evidence Strength	
independent evaluation		0.48	-1	SUP	
		0.23	-2		
		0.11	-3		
		0.05	-4	STR	
BP5: Alternate molecular basis for disease	_SUP	*see PP4			

Recommendations for the management of conflicting functional assay results See table below for management of discrepancy for BRCA1 variants between Findlay et al, 2018⁴ and Bouwman et al, 2020⁵ discordant assay results. For more general guidance regarding conflicting results from other functional assays, refer to the table in the main CanVIG-UK consensus specification.

Findlay Class	Findlay Score	Bouwman Platinum	Bouwman Olaparib	Bouwman DR-GFP	PS3_STR	BS3_STR
LOF	<-1.328	All deleterious/ likely deleterious (1 intermediate allowed)			~	×
LOF	<-1.328	Any are neutral/likely neutral			×	×
INT (towards LOF)	-1.328 to -1.038	All deleterious/ likely deleterious			~	×
INT (towards FUNC)	-1.038 to -0.748	All neutral/likely neutral		×	\checkmark	
INT	-1.328 to -0.78	Conflicting results or any intermediate			×	×
FUNC	>-0.748	All neutral/likely neutral (1 intermediate allowed)			×	~
FUNC	>-0.748	Any are o	deleterious/likel	y deleterious	×	×

N.B: Bouwman et al, 2020 "not clear" refers to opposite categorisation ± the standard deviation of repeat experiments and should be treated as conflicting assay results. Where a variant is LOF on the Findlay et al assay and has an RNA score of <-2, this indicates that LOF is due to interference with splicing and therefore should not be treated as conflicting evidence if the variant is neutral on the Bouwman et al assay.

Version History/Amendments

Revised	Date	Section	Update	Amended	Approved
1.12	01/09/2021	PP4	Guidance on use of LLRs from published epidemiological studies amended to account for the use of natural logs in the statistics presented	Garrett	Turnbull
1.12	01/09/2021	PM1	Addition of critical residues in the DNA binding domain of BRCA2. Critical residues in all listed functional domains updated to mirror draft 2021 ENIGMA guidance	Garrett	Turnbull
1.12	01/09/2021	BP1	Resolution of typo in BRCT region specification	Garrett	Turnbull
1.13	15/10/2021	PVS1	Clarification that PVS1 may not be applicable for some of the variants at ENGIMA specified positions	Garrett	Turnbull
1.14	02/12/2021	PS4	Terminology change to reflect transition of PHE to NHSD	Garrett	Turnbull

1.14	02/12/2021	PVS1	Addition of recommendations for variants within the first 100bp	Callaway	CStAG
1.15	28/04/2022	PM1/PM4	Clarification that PM1_sup and PM4_sup may be used in combination but PM4 not to be used at moderate. Removal of mention that CanVIG-UK provide non- white ethnicity counts under PS4	Garrett	CStAG

References

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