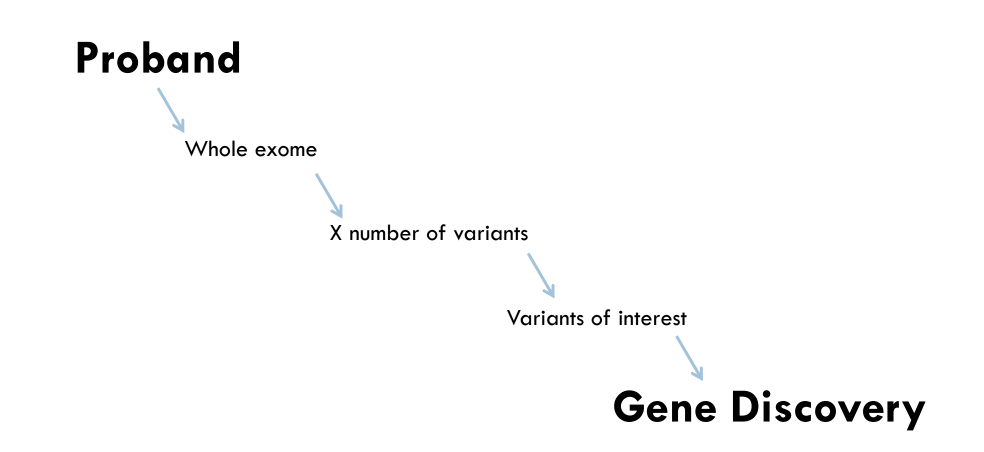
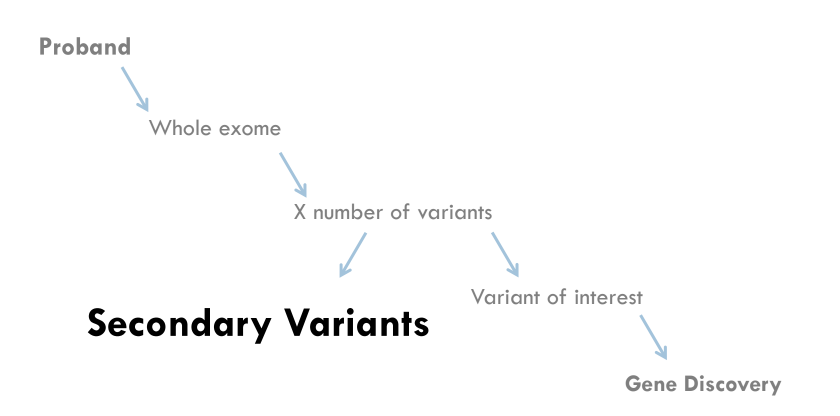
#### SECONDARY VARIANTS

Jennifer J. Johnston, PhD September 28, 2011

#### **Primary Variant**



#### Secondary Variant



#### What Do We Do With These Variants?

□ Ignore them

Analyze them and return useful results

#### Why return these variants?

Secondary line of research

Ethical obligation to research participant?

#### NHLBI Guidelines (should return)

- Important health implication of finding for participant, risk established and substantial
- Finding is actionable- therapy or prevention that could change course of disease
- Test analytically valid and disclosure complies with laws
- Participant has opted to receive results

#### NHLBI Guidelines (may return)

Benefit outweighs risk from participant's perspective

IRB approved disclosure plan

Test analytically valid and disclosure complies with laws

Participant has opted to receive results

Fabsitz et al. Circ Cardiovasc Genet (2010)

#### CLIA

Clinical Laboratory Improvement Amendments 1988

"Applies to research laboratories as well if they report patient-specific results for the diagnosis, prevention, or treatment of any disease or impairment of, or the assessment of the health of, individual patients."

#### ASHG Childhood Testing Guidelines

"If the medical or psychosocial benefits of a genetic test will not accrue until adulthood, as in the case of carrier status or adult-onset diseases, genetic testing generally should be deferred." - 1995

# So you have decided to return secondary variants....

# What do you return?

#### Diseases

Nature of disorder

- Severity/threat
- Actionability/treatability
- Alternative modes of diagnosis
- Proband vs. descendant risk

#### **Diseases to Consider**

- Cancer predisposition- Breast/Ovarian, Colorectal, other- BRCA1/2, APC, MLH1, MSH2, MSH6, PMS2
- Hypertrophic Cardiomyopathy- MYH7, MYBPC3, TNNT2, TNNI3, TPM1, MYL2, MYL3, ACTC1, CSRP3, TNN, ACTN2, MYH6, TCAP, TNNC1
- Long QT Syndrome- KCNQ1, KCNE1, KCNH2, KCNE2, SCN5A
- Malignant Hyperthermia- RYR1, CACNA1S

#### **Diseases to Consider**

Thrombophilia - F5 (Factor 5 Leiden, p.R506Q, 24/566), F2 (prothrombin, G20210A)

Hemochromatosis- HFE

Pharmacogenetics

Adult onset neurological disorders

Carrier variants that may affect future generations



#### Gene

□ HGMD, OMIM, GeneTests

- Variant
  - Return variants known to be causative
  - Can return *novel* variants highly likely to be causative
  - consider effect of telling versus not telling

http://www.ncbi.nlm.nih.gov/omim

http://www.ncbi.nlm.nih.gov/sites/GeneTests/



#### Gene

- □ HGMD, OMIM, GeneTests
- Variant
  - Return variants known to be causative
  - Can return *novel* variants highly likely to be causative
  - consider effect of telling versus not telling
    - CF BRCA1 CDH1

http://www.ncbi.nlm.nih.gov/omim

http://www.ncbi.nlm.nih.gov/sites/GeneTests/

At this point need to start filtering variants!

#### How to Work with Variant Data

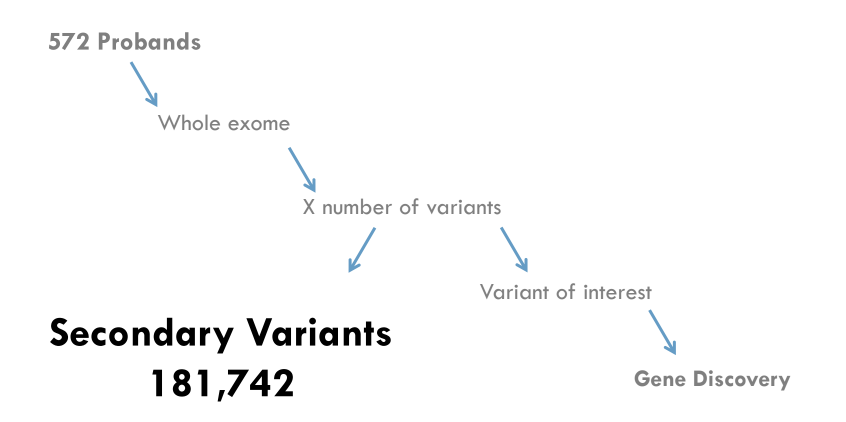
Annotation Source - VarSifter

#### Find variant n HGMD or LSDB

Analyze support for causation

Decide whether to return variant

#### **CS Secondary Variant**



# High Susceptibility Cancer Genes

APC	Familial adenomatous polyposis	FLCN	Birt-Hogg-Dubé syndrome	NF1	Neurofibromatosis type 1	RET	Multiple endocrine neoplasia Familial medullary thyroid cancer	TP53	Li-Fraumeni syndrome
BMPR1A	Familial juvenile polyposis	KIT	Gastrointestinal stromal tumor	NF2	Neurofibromatosis type 2	SDHAF2	Hereditary paraganglioma	TSC1	Tuberous sclerosis complex 1
BRCA1	Hereditary breast- ovarian cancer	MEN1	Multiple endocrine neoplasia type 1	PDGFRA	Gastrointestinal stromal tumor (GIST)	SDHB	Hereditary paraganglioma	TSC2	Tuberous sclerosis complex 2
BRCA2	Hereditary breast- ovarian cancer	MET	Hereditary papillary renal cell carcinoma	PMS2	Hereditary nonpolyposis colon cancer	SDHC	Hereditary paraganglioma	VHL	von Hippel-Lindau syndrome
CDC73 (HPRT2)	Hereditary hyperparathyroidism -jaw tumor syndrome	MLH1	Hereditary nonpolyposis colon cancer	PRKAR1A	Carney complex type 1	SDHD	Hereditary paraganglioma	WT1	Familial Wilms tumor 1
CDH1	Hereditary diffuse gastric cancer	MSH2	Hereditary nonpolyposis colon cancer	PTCH1	Nevoid basal cell carcinoma syndrome	SMAD4	Familial juvenile polyposis		
CDKN2A	Hereditary multiple melanoma	MSH6	Hereditary nonpolyposis colon cancer	PTEN	Cowden disease	SMARCB1	Schwannomatosis		
FH	Hereditary renal cell carcinoma	MUTYH	MYH-associated polyposis	RB1	Hereditary retinoblastoma	STK11	Peutz-Jeghers syndrome		

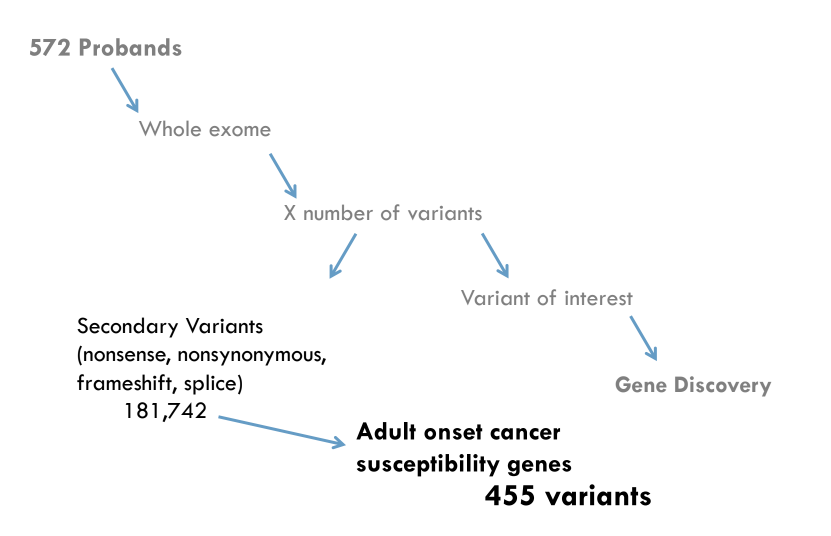
#### VarSifter – Gene Filter

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	934	1868191	chr1	864676	864678	Å	SNP	G	Non-synony	Include:
	935	485497	chr1	864679	864680	<u> </u>	INDEL	Т	DIV-fs	
	936	1579242		864679	864681	с	SNP	Ť	Non-synony	Hom. Recessive
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	951	1427808	chr1	868523	868532	GGAGGAGG	INDEL	GGAGG	Non-synony DIV-c	Inconsistent
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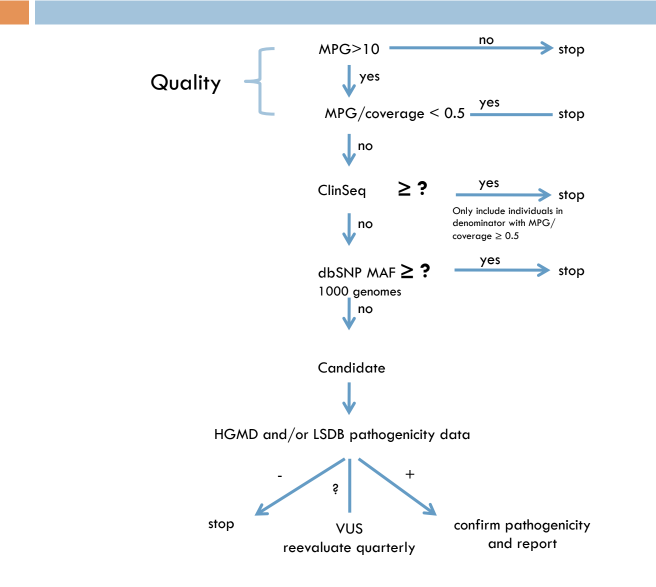
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chr9	97270930	97270932	G	SNP	A	Non-synonymous		Non-synonymous
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chr9	97308597	97308599	C	SNP	G	Non-synonymous		
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chr9	134770825	134770827	G	SNP	С	Non-synonymous		Dominant
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D100402.N D100416.N D100498.N D100569.N D100733.N D100818.N D100822.N D100840.N D100854.N	IA AA IA AA IA AA IA AA IA AA IA AA IA AA IA AA IA AA	38 46 11 7 12 8 16 11 4	48 52 63 13 7 14 8 20 13 2 2	riant Positio	ns: 455		X	Show Variants Show Genes Clear All Apply Filter /Users/jjohnsto/Desktop/VarSifter/ClinSeqOncogenes0104201
D100402.N D100416.N D100498.N D100569.N D100733.N D100818.N D100822.N D100840.N D100854.N	IA AA IA AA IA AA IA AA IA AA IA AA IA AA IA AA IA AA	38 46 11 7 12 8 16 11 4	48 52 63 13 7 14 8 20 13 2 2	riant Positio	ns: 455		×	Show Variants Show Genes      Clear All      Apply Filter      /Users/jjohnsto/Desktop/VarSifter/ClinSeqOncogenes0104201      Choose Gene File Filter      No Bed File Selected
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#### **CS Secondary Variant**



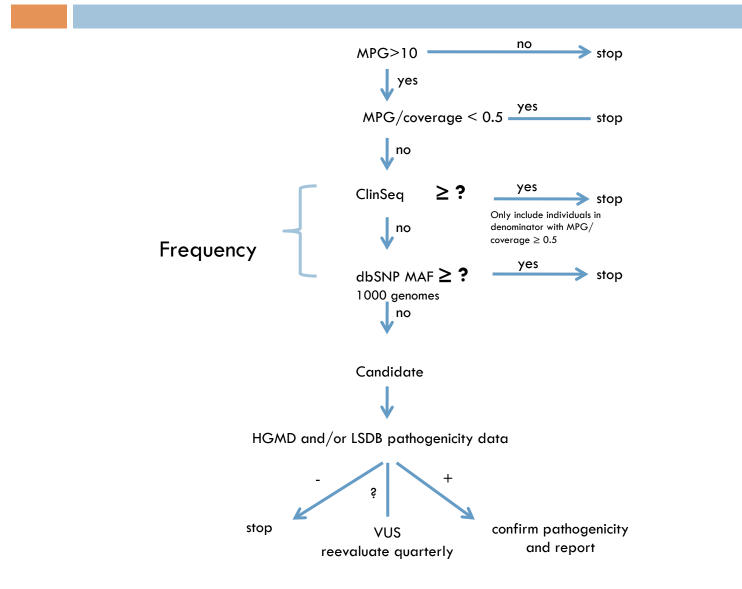
#### Framework for Variant Interpretation



# VarSifter – <u>Most Probable Genotype</u>/ Coverage

				VarSift	er – /Users/jjohnsto/De	sktop/VarSifte	r/VarSifter_1.0/572exc	omesnocontro	ols_cod.vs
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dex	VariantId	Chr	LeftFlank	RightFlank	ref_allele	muttype	var_allele	type	Include:
428194	989090	chr13	31804765	31804767	С	SNP	Т	N	DIV-c
42819	803617	chr13	31805128	31805130	т	SNP	С	N	_
42820	841738	chr13	31805178	31805180	G	SNP	С	N	DIV-fs
42820	1 1898438	chr13	31805406	31805408	A	SNP	G	N	Non-synonymous
428202	2 3182391	chr13	31805503	31805505	С	SNP	Т	N	
42820	7 1852661	chr13	31808455	31808457	С	SNP	G	N	Splice-site
42820			31808799	31808801	А	SNP	С	N	Stop
42821			31808841	31808843	A	SNP	G	N	
428214	\$ 562236	chr13	31809462	31809464	A	SNP	G	N	Exclude:
42821			31810006		С	SNP	Т	N	dbID
428220			31810053		A	SNP	G	N	
42822			31810072		G	SNP	A	N	Exclude Gene File
428224			31810749		G	SNP	Т	N	
42822			31811084		A	SNP	Т	N	Include:
42822			31811270		A	SNP	С	N	Hom. Recessive
42823	1534135	chr13	31811689	31811691	С	SNP	Т	N	
42823			31811803	31811805	G	SNP	A	N	🗌 Dominant
42823			31811970		ATTAAATT	INDEL	ATT		Inconsistent
42823			31812043		т	SNP	G	N	
42823	7 1551174	chr13	31812045	31812047	G	SNP	Т	N	Mend. Compound Het
42823	9 1578933	chr13	31812223	31812225	A	SNP	Т	N	🗹 Include Gene File
428240	862096	chr13	31812235	31812237	С	SNP	Т	N	🗌 Jasluda Dad Cila Dasiana
42824	1 1623569	chr13	31812437	31812439	т	INDEL	"		Include Bed File Regions
428242	2 718833	chr13	31812591	31812593	С	SNP	Т	N 🔺	Affected different from Norm
42824	3 1556721	chr13	31812813	31812815	С	SNP	Т	Nv	Diff. in at least: 0
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ample	Genotype	🗸 🗸 Geno	type score cov	erage					Case / Control
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180290.N 100199.N	Α ΑC 4 ΑΑ	17	36	)					Case / Control
180290.N 100199.N 100288.N	Α ΑC Α ΑΑ Α ΑΑ	17 112	36 160	)					Case / Control Var in cases (at least): 0 + Var in controls (this or fewer): 0 +
180290.N 100199.N 100288.N 100395.N	Α ΑC Α ΑΑ Α ΑΑ Α ΑΑ Α ΑΑ	17 112 85	36 160 120	)					Case / Control Var in cases (at least): 0
180290.N 100199.N 100288.N 100395.N 100402.N	Α ΑC Α ΑΑ Α ΑΑ Α ΑΑ Α ΑΑ Α ΑΑ	17 112 85 55	36 160 120 76						Case / Control Var in cases (at least): 0 + Var in controls (this or fewer): 0 +
130290 N 100199 N 100288 N 100395 N 100402 N 100416 N	AC           A         AA	17 112 85 55 92	16 160 120 76 130						Case / Control Var in cases (at least): 0 + Var in controls (this or fewer): 0 +
100199.N 100199.N 100288.N 100395.N 100402.N 100416.N 100498.N	AC           A         AA	17 112 85 55 92 83	10 160 122 76 130 117 82 43						Case / Control Var in cases (at least): 0 + Var in controls (this or fewer): 0 + Search gene names for:
100290.N 100288.N 100288.N 100395.N 100402.N 100416.N 100498.N 100569.N	AC         AC           A         AA	112 85 55 92 83 59	86 160 120 76 130 117 82						Case / Control Var in cases (at least): 0 + Var in controls (this or fewer): 0 +
ample 1100199.N. 100288.N. 100395.N. 100402.N. 100402.N. 100408.N. 1000569.N. 100733.N. 100818.N.	A         AC           A         AA	112 85 55 92 83 59 32	10 160 122 76 130 117 82 43						Case / Control Var in cases (at least): 0 + Var in controls (this or fewer): 0 + Search gene names for:
H 1230 N 0100199.N 0100288.N 0100395.N 0100402.N 0100416.N 0100498.N 0100569.N 0100733.N	A         AC           A         AA           A         AA	112 85 55 92 83 59 32 66	80 160 122 76 130 117 82 43 93	) ) ) 7				Ō	Case / Control Var in cases (at least): 0 + Var in controls (this or fewer): 0 + Search gene names for:
11.00241040 0100199.N. 0100288.N. 0100395.N. 0100402.N. 0100416.N. 01004468.N. 0100569.N. 0100733.N. 0100733.N. 0100818.N. 0100822.N. 0100840.N.	AC         AA           A         AA	112 85 55 92 83 59 32 66 33	16 120 76 130 117 82 43 93 45	) ) ) 7				Ō	Case / Control Var in cases (at least): 0 + Var in controls (this or fewer): 0 + Search gene names for:
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100199.N. 100288.N. 100395.N. 100402.N. 100402.N. 100416.N. 100569.N. 100569.N. 100733.N. 100818.N. 100822.N.	AC         AA           A         AA	112 85 55 92 83 59 32 66 33 81 46	1160 12( 76 13( 11) 82 43 93 45 114	) ) 7 4	riant Positions: 455			Ō	Case / Control Var in cases (at least): 0 + Var in controls (this or fewer): 0 + Search gene names for:
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H 100199.N. 0100199.N. 0100288.N. 0100395.N. 0100402.N. 01004416.N. 0100498.N. 0100569.N. 0100569.N. 0100733.N. 0100818.N.	AC         AA           A         AA	112 85 55 92 83 59 32 66 33 81 46	1160 12( 76 13( 11) 82 43 93 45 114	) ) 7 4	riant Positions: 455			Ō	Case / Control Var in cases (at least): 0 + Var in controls (this or fewer): 0 + Search gene names for: Show Variants Show Genes Clear All
11.00241040 0100199.N. 0100288.N. 0100395.N. 0100402.N. 0100416.N. 01004468.N. 0100569.N. 0100733.N. 0100733.N. 0100818.N. 0100822.N. 0100840.N.	AC         AA           A         AA	112 85 55 92 83 59 32 66 33 81 46	1160 12( 76 13( 11) 82 43 93 45 114	) ) 7 4	riant Positions: 455			Ō	Case / Control Var in cases (at least): 0 + Var in controls (this or fewer): 0 + Search gene names for: Show Variants Show Genes Clear All
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11.00241040 0100199.N. 0100288.N. 0100395.N. 0100402.N. 0100416.N. 01004468.N. 0100569.N. 0100733.N. 0100733.N. 0100818.N. 0100822.N. 0100840.N.	AC         AA           A         AA	112 85 55 92 83 59 32 66 33 81 46	1160 12( 76 13( 11) 82 43 93 45 114	) ) 7 4	riant Positions: 455			Ō	Case / Control Var in cases (at least): 0 ÷ Var in controls (this or fewer): 0 ÷ Search gene names for: Search gene names for: Clear All Clear All Vusers/jjohnsto/Desktop/VarSifter/ClinSeqOncogenes0104201
100199.N. 100288.N. 100395.N. 100402.N. 1004402.N. 100446.N. 100569.N. 100569.N. 100733.N. 100818.N. 100822.N.	AC         AA           A         AA	112 85 55 92 83 59 32 66 33 81 46	1160 12( 76 13( 11) 82 43 93 45 114	) ) 7 4	riant Positions: 455			Ō	Case / Control Var in cases (at least): 0 Var in controls (this or fewer): 0 Search gene names for: Search gene names for: Clear All Apply Filter /Users/jjohnsto/Desktop/VarSifter/ClinSeqOncogenes0104201 Choose Gene File Filter
100199.N. 100288.N. 100395.N. 100402.N. 100402.N. 100416.N. 100569.N. 100569.N. 100733.N. 100818.N. 100822.N.	AC         AA           A         AA	112 85 55 92 83 59 32 66 33 81 46	1160 12( 76 13( 11) 82 43 93 45 114	) ) 7 4	riant Positions: 455			Ō	Case / Control Var in cases (at least): 0 Var in controls (this or fewer): 0 Search gene names for: Search gene names for: Clear All Apply Filter /Users/jjohnsto/Desktop/VarSifter/ClinSeqOncogenes0104201 Choose Gene File Filter No Bed File Selected
100290, N 100199, N 100288, N 100395, N 100402, N 100402, N 100416, N 100498, N 100569, N 100569, N 100733, N 100818, N 100822, N	AC         AA           A         AA	112 85 55 92 83 59 32 66 33 81 46	1160 12( 76 13( 11) 82 43 93 45 114	) ) 7 4	riant Positions: 455			Ō	Case / Control Var in cases (at least): 0 Var in controls (this or fewer): 0 Search gene names for: Search gene names for: Clear All Apply Filter /Users/jjohnsto/Desktop/VarSifter/ClinSeqOncogenes0104201 Choose Gene File Filter

#### Framework for Variant Interpretation



## VarSifter – MPG/Coverage

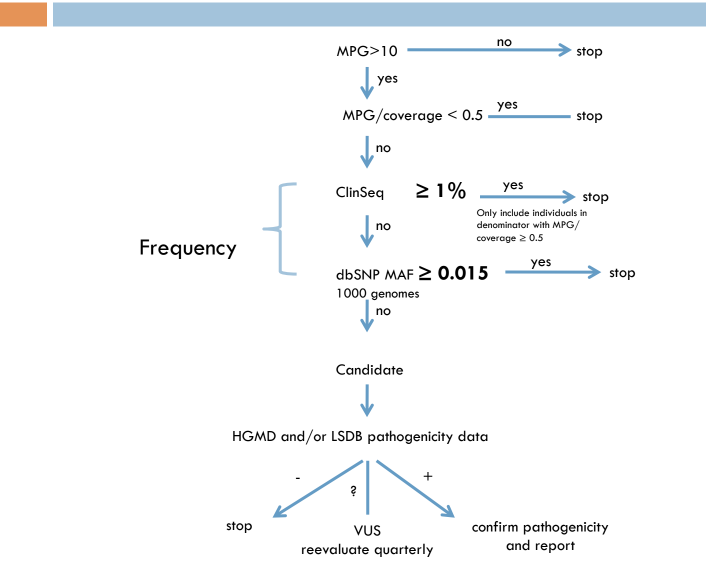
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VarSifter - /Users/jjohnsto/Desktop/VarSifter/VarSifter\_1.0/572exomesnocontrol

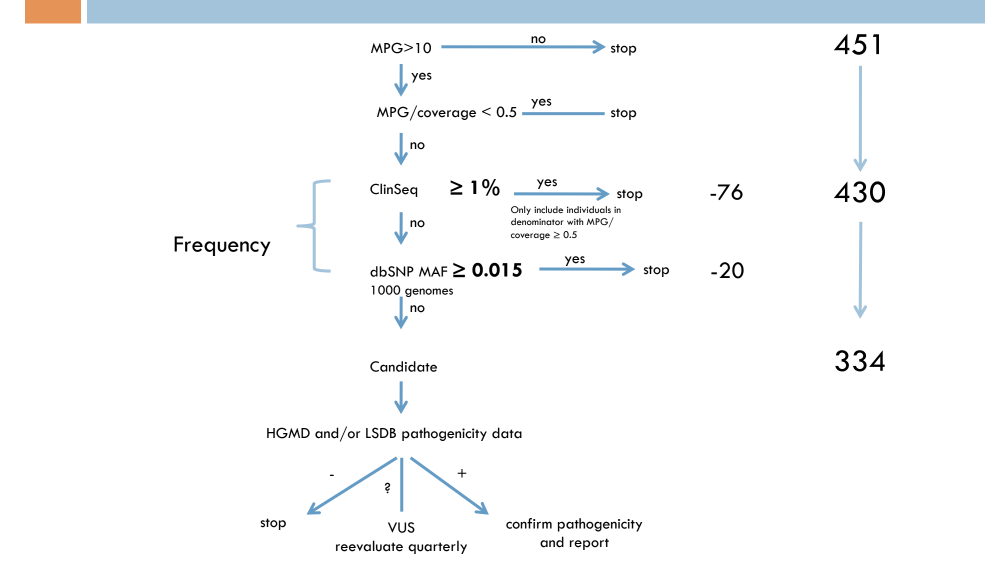
File View Help

Gene_name 🗚	ref_aa	aa_pos	var_aa	CSc_genotypes	CSc_homref	CSc_het	CSc_refallele	CSc_nonrefallele	CSmaf
BRCA2	S	1172	L	572	571	1	1143	1	0.000900
BRCA2	I	1188	v	572	571	1	1143	1	0.000900
BRCA2	G	1194	D	572	571	1	1143	1	0.000900
BRCA2	D	1420	Y	500	494	6	994	6	0.006000
BRCA2	к	1531	N	569	568	1	1137	1	0.000900
BRCA2	E	1593	D	572	571	1	1143	1	0.000900
BRCA2	S	1733	F	570	568	2	1138	2	0.001800
BRCA2	G	1771	D	562	561	1	1123	1	0.000900
BRCA2	NA	0	NA	551	550	1	1101	1	0.000900
BRCA2	I	1851	S	554	553	1	1107	1	0.000900
BRCA2	v	1852	F	555	554	1	1109	1	0.000900
BRCA2	D	1911	v	572	571	1	1143	1	0.000900
BRCA2	Т	1915	М	571	551	20	1122	20	0.017500
BRCA2	NA	0	NA	572	569	3	1141	3	0.002600
BRCA2	R	2034	С	572	568	4	1140	4	0.003500
BRCA2	R	2108	С	570	569	1	1139	1	0.000900
BRCA2	v	2109	I	570	569	1	1139	1	0.000900
BRCA2	N	2113	S	568	567	1	1135	1	0.000900
BRCA2	н	2116	R	565	564	1	1129	1	0.000900
BRCA2	I	2285	v	496	495	1	991	1	0.001000
BRCA2	н	2440	R	572	571	1	1143	1	0.000900
BRCA2	v	2466	А	566		0	0		-1.000
BRCA2	R	2502	С	572	571	1	1143	1	0.000900
BRCA2	Т	2515	I	572	571	1	1143	1	0.000900
BRCA2	А	2717	S	572	571	1	1143	1	0.000900
				(					) 4 1

#### **Cancer Variant Filtering**



#### **CS** Cancer Filtering



#### **Evaluation of Candidates**

- Human Gene Mutation Database (HGMD)
- Locus Specific Database (LSDB)
  - Controls
  - Multiple reports
  - Functional data
  - Presence with other causative mutations
  - Segregation with disease (LD & linkage caveat)
    - De novo (assuming parentage)
    - Penetrance
    - Phenocopies

#### VarSifter - HGMD

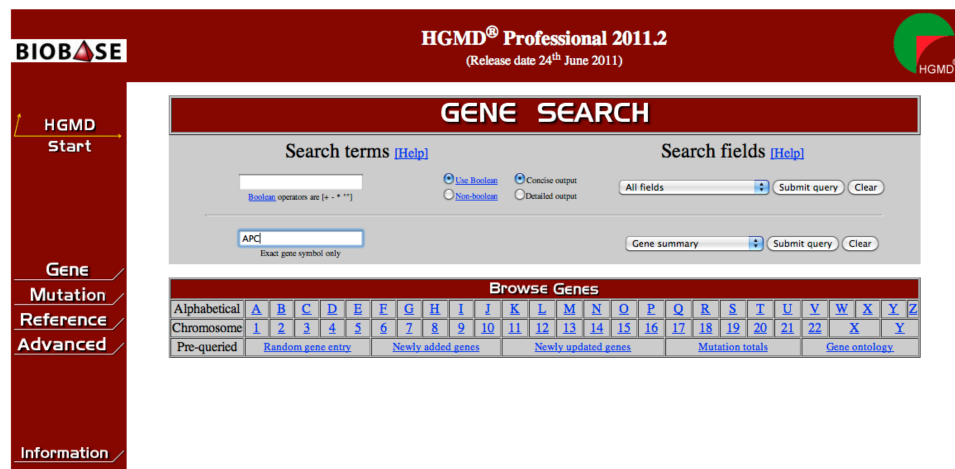
ile V	/iew Help							
Chr	LeftFlank	RightFlank	Gene_name 🗚	HGMDids	HGMDdisease	HGMDtags	HGMDinGene	transcript
chr13	31810006	31810008	BRCA2	CM050182	Breast cancer ?	DM	У	uc001uub.1
chr13	31810053	31810055	BRCA2	-	-	-	ý	uc001uub.1
chr13	31810072	31810074	BRCA2	-	-	-	y	uc001uub.1
chr13	31810749	31810751	BRCA2	CM003133	Breast and/or ovarian cancer ?	DM	y	uc001uub.1
chr13	31811084	31811086	BRCA2	-	-	-	У	uc001uub.1
chr13	31811270	31811272	BRCA2	-	-	-	У	uc001uub.1
chr13	31811689	31811691	BRCA2	-	-	-	У	uc001uub.1
chr13	31811803	31811805	BRCA2	CM041731	Breast and/or ovarian cancer ?	DM	У	uc001uub.1
chr13	31811970	31811979	BRCA2	-	-	-	У	uc001uub.1
chr13	31812043	31812045	BRCA2	-	-	-	У	uc001uub.1
chr13	31812045	31812047	BRCA2	-	-	-	У	uc001uub.1
chr13	31812223	31812225	BRCA2	-	-	-	У	uc001uub.1
chr13	31812235	31812237	BRCA2	CM010170	Breast cancer ?	DM	У	uc001uub.1
chr13	31812437	31812439	BRCA2	-	-	-	У	uc001uub.1
chr13	31812591	31812593	BRCA2	CM994286	Breast and/or ovarian cancer ?	DM	У	uc001uub.1
chr13	31812813	31812815	BRCA2	-	-	-	У	uc001uub.1
chr13	31812816	31812818	BRCA2	CM043917	Breast cancer ?	DM	У	uc001uub.1
chr13	31812829	31812831	BRCA2	-	-	-	У	uc001uub.1
chr13	31812838	31812840	BRCA2	CM022331	Breast and/or ovarian cancer	DM	У	uc001uub.1
chr13	31816705	31816707	BRCA2	-	-	-	У	uc001uub.1
chr13	31827308	31827310	BRCA2	-	-	-	У	uc001uub.1
chr13	31827386	31827388	BRCA2	CM960194	Breast cancer	DM	У	uc001uub.1
chr13	31828632	31828634		CM012590	Breast and/or ovarian cancer	DM	У	uc001uub.1
chr13	31828672	31828674	BRCA2	CM994287	Breast and/or ovarian cancer ?	DM	У	uc001uub.1
chr13	31835487	31835489	BRCA2	CM043984	Breast and/or ovarian cancer	DM	У	uc001uub.1
chr13	31835520	31835522	BRCA2	CM004715	Breast cancer	DM	v	uc001uub.1

#### Human Gene Mutation Database

http://nihlibrary.nih.gov/ResearchTools/Pages/Bioinformatics.aspx

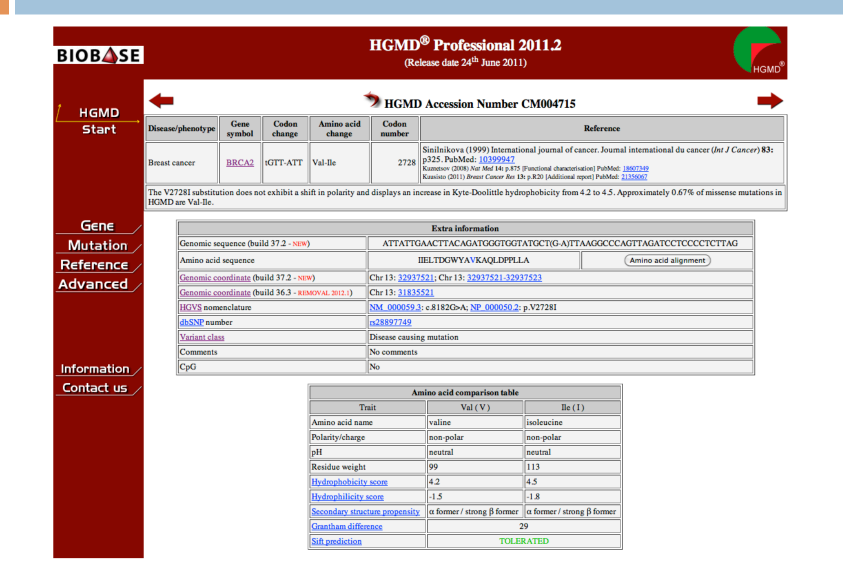
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D III POTS Main Page parkinson U	CSC HGMD OMIM Homence in Man nih Journals GeneClinics Home Page							
OBASE	HGMD <sup>®</sup> Professional 2011.2 (Release date 24 <sup>th</sup> June 2011)	Н						
	To start a search choose the search option in the menu to the left.							
Start	This release comprises the following tables:							
Data type:	Description:	Entries:						
Missense/nonse	Single base-pair substitutions in coding regions are presented in terms of a triplet change with an additional flanking base included if the mutated base lies in either the first or third position in the triplet.	63313						
Splicing	Mutations with consequences for mRNA splicing are presented in brief with information specifying the relative position of the lesion with respect to a numbered intron donor or acceptor splice site. Positions given as positive integers refer to a 3' (downstream) location, negative integers refer to a 5' (upstream) location.	10653						
Gene Autation Regulatory	Substitutions causing regulatory abnormalities are logged in with thirty nucleotides flanking the site of the mutation on both sides. The location of the mutation relative to the transcriptional initiation site, initiaton codon, polyadenylation site or termination codon is given.	2049						
Small deletions	Micro-deletions (20 bp or less) are presented in terms of the deleted bases in lower case plus, in upper case, 10 bp DNA sequence flanking both sides of the lesion. The numbered codon is preceded in the given sequence by the caret character (^).	17807						
Small insertion	Micro-insertions (20 bp or less) are presented in terms of the inserted bases in lower case plus, in upper case, 10 bp DNA sequence flanking both sides of the lesion. The numbered codon is preceded in the given sequence by the caret character (^).	7346						
Small indels	Micro-indels (20 bp or less) are presented in terms of the deleted/inserted bases in lower case plus, in upper case, 10 bp DNA sequence flanking both sides of the lesion. The numbered codon is preceded in the given sequence by the caret character (^).	1671						
Gross deletions	Information regarding the nature and location of each lesion is logged in narrative form because of the extremely variable quality of the original data reported.	7383						
Gross insertions/dupl	Information regarding the nature and location of each lesion is logged in narrative form because of the extremely variable quality cations of the original data reported.	1583						
Complex rearrangements	Information regarding the nature and location of each lesion is logged in narrative form because of the extremely variable quality of the original data reported.	1089						
Repeat variatio	Information regarding the nature and location of each lesion is logged in narrative form because of the extremely variable quality of the original data reported.	353						
	Mutation total	113247						
	With chromosomal coordinates (NCBI37.2/hg19)	90487						
	With chromosomal coordinates (NCB136.3/hg18 - REMOVAL 2012.1)	90480						

### HGMD - Search



Contact us

#### HGMD – Mutation Page



#### HGMD – Primary Literature

#### Display Settings: V Abstract

Send to: 🖂

Breast Cancer Res. 2011 Feb 28:13(1):R20. [Epub ahead of print]

Screening for BRCA1, BRCA2, CHEK2, PALB2, BRIP1, RAD50, and CDH1 mutations in high-risk Finnish BRCA1/2-founder mutation-negative breast and/or ovarian cancer individuals.

#### Kuusisto KM, Bebel A, Vihinen M, Schleutker J, Sallinen SL

Department of Pediatrics, Genetics Outpatient Clinic, Tampere University Hospital, Biokatu 8, Tampere, 33520, Finland. Satu-Leena.Sallinen@pshp.fi.

#### Abstract

ABSTRACT:

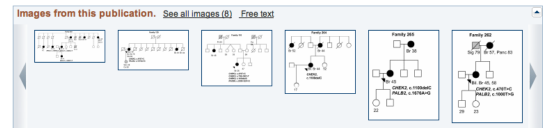
INTRODUCTION: Two major high-penetrance breast cancer genes, BRCA1 and BRCA2, are responsible for approximately 20% of hereditary breast cancer (HBC) cases in Finland. Additionally, rare mutations in several other genes that interact with BRCA1 and BRCA2 increase the risk of HBC. Still, a majority of HBC cases remain unexplained which is challenging for genetic counseling. We aimed to analyze additional mutations in HBC-associated genes and to define the sensitivity of our current BRCA1/2 mutation analysis protocol used in genetic counseling.

METHODS: Eighty-two well-characterized, high-risk hereditary breast and/or ovarian cancer (HBOC) BRCA1/2-founder mutation-negative Finnish individuals, were screened for germline alterations in seven breast cancer susceptibility genes, BRCA1, BRCA2, CHEK2, PALB2, BRIP1, RAD50, and CDH1. BRCA1/2 were analyzed by multiplex ligation-dependent probe amplification (MLPA) and direct sequencing. CHEK2 was analyzed by the high resolution melt (HRM) method and PALB2, RAD50, BRIP1 and CDH1 were analyzed by direct sequencing. Carrier frequencies between 82 (HBOC) BRCA1/2-founder mutation-negative Finnish individuals and 384 healthy Finnish population controls were compared by using Fisher's exact test. In silico prediction for novel missense variants effects was carried out by using Pathogenic-Or-Not -Pipeline (PON-P).

RESULTS: Three previously reported breast cancer-associated variants, BRCA1 c.5095C > T, CHEK2 c.470T > C, and CHEK2 c.1100delC, were observed in eleven (13.4%) individuals. Ten of these individuals (12.2%) had CHEK2 variants, c.470T > C and/or c.1100delC. Fourteen novel sequence alterations and nine individuals with more than one non-synonymous variant were identified. One of the novel variants, BRCA2 c.72A > T (Leu24Phe) was predicted to be likely pathogenic in silico. No large genomic rearrangements were detected in BRCA1/2 by multiplex ligation-dependent probe amplification (MLPA).

CONCLUSIONS: In this study, mutations in previously known breast cancer susceptibility genes can explain 13.4% of the analyzed high-risk BRCA1/2negative HBOC individuals, CHEK2 mutations, c.470T > C and c.1100delC, make a considerable contribution (12.2%) to these high-risk individuals but further segregation analysis is needed to evaluate the clinical significance of these mutations before applying them in clinical use. Additionally, we identified novel variants that warrant additional studies. Our current genetic testing protocol for 28 Finnish BRCA1/2-founder mutations and protein truncation test (PTT) of the largest exons is sensitive enough for clinical use as a primary screening tool.

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Review Pitfalls and caveats in BRCA sequencin [Ultrastruct Pathol. 200	
Review Breast cancer genetics in African Americans. [Cancer, 200	03]
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Related information	
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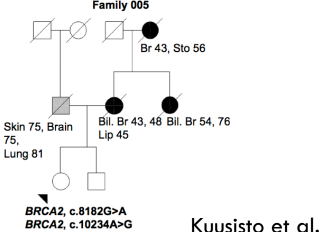
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#### **Primary Literature**

role of the three *BRCA2* missense variants, c.8182G > A, c.9976A > T, and c.10234A > G, in HBOC risk, is uncertain [<u>31-33</u>]. All three heterozygous variants were observed in two healthy women with a history of BrCa, one carrying the c.9976A > T variant and the other both the c.8182G > A and c.10234A > G variants (Tables <u>2</u> and <u>3</u>, Figure <u>8</u>, Family 005). At this stage, because we only have samples from the index individuals, no segregation analyses of the variants have been performed, but these families clearly warrant additional studies.



The

### Locus-Specific DataBases

Locus Specific Datab Based on various online resour	ase list ces and direct submissions of LSDBs	<u>LSDB list</u>   <u>Su</u>	<u>bmit new LSDB</u>   <u>Log i</u>
Locus Specific Mutation	Databases		
IMPORTANT NOTE: Genes ar be found under "CD40LG" ar	e in order of <u>HUGO APPROVED GENE DESIGNATION</u> , not alias. e.g. "p53" w d so on.	ill be found under "TP53" while "CD4	40L" or "TNFSF5" will
If you wish to add a gene yo	ou can <u>do so here</u> .		
397 public entries	A B C D E F G H I J K L M N O P Q R S T U V V Or, specify the HGNC Gene Symbol: Go to this gene »	ΧXZ	
• • • •	C Database	Curators	Software 🗘
A2M	Mendelian genes	Curator vacancy	LOVD 2.X
alpha-2-macroglobulin	http://grenada.lumc.nl/LOVD2/mendelian_genes/home.php?select_db=A2M	?	
A4GALT	Mendelian genes	Curator vacancy	LOVD 2.X
alpha 1,4-galactosyltransferase	http://grenada.lumc.nl/LOVD2/mendelian_genes/home.php?select_db=A4GALT Mendelian_genes	/ Curatar vacancy	LOVD 2.X
achalasia, adrenocortical insufficiency, alacrimia (Allgrove, triple-A)	http://grenada.lumc.nl/LOVD2/mendelian_genes/home.php?select_db=AAAS	Curator vacancy ?	LOVD 2.X
AANAT arylalkylamine N-acetyltransferase	Mendelian genes http://grenada.lumc.nl/LOVD2/mendelian_genes/home.php?select_db=AANAT	Curator vacancy ?	LOVD 2.X
AARS alanyl-tRNA synthetase	LOVD - Leiden Open Variation Database https://grenada.lumc.nl/LOVD2/shared1/home.php?select_db=AARS	Curator Vacancy Leiden University Medical Center	LOVD 2.X

http://www.hgvs.org/dblist/glsdb.html http://grenada.lumc.nl/LSDB\_list/lsdb.phpaction=view\_all&symbol\_start=M

### LSDB

\varTheta 🔿 🔿 Locus Specific Database list										
+ Shttp://grenada.lumc.nl/LSDB_list/lsdb.php?action=view_all&symbol_start=A       C Qr Google										
😔 💭 🇰 POTS Main Page parkinson UCSC HGMD OMIM Homence in Man nih Journals GeneClinics Home Page										
	<pre>nttp://risnmap2.igib.res.in/nome.pnp?select_db=APBB2</pre>									
APC Adenomatous Polyposis Coli	Colon cancer gene variant databases http://chromium.liacs.nl/LOVD2/colon_cancer/home.php?select_db=APC	Stefan Aretz & Waltraut Friedl & Kirsten Wöllner Institute of Human Genetics, Bonn Institute of Human Genetics Institute of Human Genetics, Bonn	LOVD 2.X							
APC	Zhejiang University Center for Genetic and Genomic Medicine http://www.genomed.org/LOVD/home.php?select_db=APC		LOVD 2.X							
APC	Zhejiang University Center for Genetic and Genomic Medicine (ZJU-CGGM) http://genomed.org/LOVD/HNPCC/home.php?select_db=APC		LOVD 2.X							
APC	LOVD - Leiden Open Variation Database https://australianhumanvariomedatabase.arcs.org.au/home.php?select_db=APC		LOVD 2.X							
APC adenomatous polyposis coli	The UMD APC mutations database http://www.umd.be/APC/	Christophe Beroud, Laboratoire de génétique Moléculaire et Chromosomique, Montpellier, France Thierry Soussi INSERM, Hopital Necker Enfants Malades, Paris	UMD							
APC adenomatous polyposis coli	Zhejiang University-Adinovo Center APC Database http://china-hvp.org/LOVD/?select_db=APC	Ming Qi, PhD, FACMG, Peikuan Cong and Yudong Gao	LOVD 2.X							
APC adenomatous polyposis coli	The APC Mutation Database http://fap.taenzer.me	Dr. Stefan Aretz and Dr. Waltraut Friedl	Unknown							

#### LOVD Gene homepage

General information	
Gene name	Adenomatous Polyposis Coli
Gene symbol	APC
Chromosome Location	5q22.2
Database location	chromium.liacs.nl
Curator	Kirsten Wöllner, Stefan Aretz and Waltraut Friedl
PubMed references	View all (unique) PubMed references in the APC database
Date of creation	September 09, 2009
Last update	September 21, 2011
Version	APC110921
Add sequence variant	Submit a sequence variant
First time submitters	Register here
Reference sequence file	coding DNA reference sequence for describing sequence variants
Genomic refseq ID	<u>NG_008481.1</u>
Transcript refseq ID	<u>NM_000038.4</u>
Exon/intron information	Exon/intron information table
Total number of unique DNA variants reported	1191
Total number of individuals with variant(s)	3782
Total number of variants reported	3792
Subscribe to updates of this gene	
NOTE	Aliases for APC are; BTPS2, DP2, DP2.5, DP3, GS

Graphical displays and utilities	
Summary tables	Summary of all sequence variants in the APC database, sorted by type of variant (with graphical displays and statistics)
Reading-frame checker	The Reading-frame checker generates a prediction of the effect of whole-exon changes
UCSC Genome Browser	Show variants in the UCSC Genome Browser (compact view)
Ensembl Genome Browser	Show variants in the Ensembl Genome Browser
NCBI Sequence Viewer	Show distribution histogram of variants in the NCBI Sequence Viewer

#### Sequence variant tables

Unique sequence variants	Listing of all unique sequence variants in the APC database, without patient data
Complete sequence variant listing	Listing of all sequence variants in the APC database
Variants with no known pathogenicity	Listing of all APC variants reported to have no noticeable phenotypic effect (note: excluding
	variants of unknown effect)

Search the database	
By type of variant	View all sequence variants of a certain type
Simple search	Query the database by selecting the most important variables (exon number, type of variant, disease phenotype)
Advanced search	Query the database by selecting a combination of variables
Based on patient origin	View all variants based on your patient origin search terms
Search through hidden entries	Find the number of variant entries in the database (including hidden entries) matching your search terms.

#### LSDB

View unique variants Search unique variants View all contents Full database search Variant listing based on patient origin Database statistics Switch gene

#### LOVD - Variant listings

Unhide all columns | Hide Specific Columns | Hide all columns

About this overview [Show]

3783 public entries

[ 100 🛊 ] entries per page	100	\$	entries	per	page
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Path. 😣 🗘	Exon 🕴 🛇	Codon_nr 😣 🗘	DNA change 🛛 🗘	DNA_reported 80	RNA change 😣 🗘	Protein 🛛 😂 📿	🗧 Туре 🛛 🚳 🗘	Cons_predicted 😣 🗘	DB-ID 🕹 🗘	Variant remarks 🛛 😂 📿	Origin 😂 🗘	Variant reference 😂
/?	00	-	c?C>G	-47306C>G (5' of ATG)	-	-	-	-	APC_00415	numbering 5' of ATG	-	-
/?	00	-	c.?C>T	-47287C>T	-	-	-	-	APC_00416	numbering 5' of ATG	-	-
/?	00	-	c.?insG	-47307insG	-	-		-	APC_00417	numbering 5' of ATG	-	-
/?	00	-	c.?T>G	-47408T>G	-	-		•	APC_00418	numbering 5' of ATG	-	-
+/?	01_15+promoter	del	cytogeneticdeletion	•		-	deletion, large	deletion, large	APC_00200	cytogenetic deletion	unknown	Raedle et al. 2001
+/?	01_15+promoter	del	cytogeneticdeletion			-	deletion, large	deletion, large	APC_00200	cytogenetic deletion	de novo	Aretz et al. 2005
+/?	01_15+promoter	del	cytogeneticdeletion		-	-	deletion, large	deletion, large	APC_00200	cytogenetic deletion	de novo	Aretz et al. 2005
+/?	01_15+promoter	del	cytogeneticdeletion		-	-	deletion, large	deletion, large	APC_00200	cytogenetic deletion	unknown	Aretz and Friedl (unpublished)
+/?	01	24	c.70C>T		-	p.Arg24X	substitution, base pair	nonsense	APC_00551	-	-	<u>Kanter-Smoler et al.</u> 2008
+/?	01_15	del	g.26940-? _133343+?del		-	-	deletion, large	deletion, large	APC_00587	-	familial	<u>Kanter-Smoler et al.</u> 2008
+/?	01+promoter	del	g.35041-?_52505+? del	-	-	-	deletion, large	deletion, large	APC_00526	-	familial	Aretz et al. 2005
+/?	01+promoter	del	g.35041-?_52505+? del			-	deletion, large	deletion, large	APC_00526		familial	Aretz et al. 2005
+/?	01+promoter	del	g.35041-?_52505+? del	-	-	-	deletion, large	deletion, large	APC_00526	-	familial	Aretz et al. 2005
+/?	01_05+promoter	del	g.35041-?_78383+? del	-	-	-	deletion, large	deletion, large	APC_00527		familial	Aretz et al. 2005

#### **LSDB**

View unique variants Search unique variants View all contents Full database search Variant listing based on patient origin Database statistics Switch gene

#### LOVD - Variant listings

Unhide all columns | Hide Specific Columns | Hide all columns

About this overview [Show]

#### 3783 public entries

100 \$	entries	per	pag
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Path. 🕴 🔇	Exon 😣 📿	Codon_nr 😣 🗘	DNA change 🛛 📿	DNA_reported 😣 🗘	RNA change 8	🗘 Protein 🛛 😣	🗘 Туре 🛛 🕴 🗘	Cons_predicted	DB-ID 😣 🗘	Variant remarks 😣 📿	Origin 😣 📿	Variant reference 😣 📿
/?	00	-	c?C>G	-47306C>G (5' of ATG)	-	-	-		APC_00415	numbering 5' of ATG	-	-
/?	00	-	c.?C>T	-47287C>T	-	-	-		APC_00416	numbering 5' of ATG	-	-
/?	00	-	c.?insG	-47307insG	-	-	-	-	APC_00417	numbering 5' of ATG	-	-
/?	00	-	c.?T>G	-47408T>G	-	-	-	•	APC_00418	numbering 5' of ATG	-	-
-/?	01_15+promoter	del	cytogeneticdeletion	•	-	-	deletion, large	deletion, large	APC_00200	cytogenetic deletion	unknown	Raedle et al. 2001
-/?	01_15+promoter	del	cytogeneticdeletion		-	-	deletion, large	deletion, large	APC_00200	cytogenetic deletion	de novo	Aretz et al. 2005
-/?	01_15+promoter	del	cytogeneticdeletion	-	-	-	deletion, large	deletion, large	APC_00200	cytogenetic deletion	de novo	Aretz et al. 2005
-/?	01_15+promoter	del	cytogeneticdeletion		-	-	deletion, large	deletion, large	APC_00200	cytogenetic deletion	unknown	Aretz and Friedl (unpublished)
-/?	01	24	c.70C>T		-	p.Arg24X	substitution, base pair	nonsense	APC_00551	-	-	Kanter-Smoler et al. 2008
-/?	01_15	del	g.26940-? _133343+?del	-	-	-	deletion, large	deletion, large	APC_00587	-	familial	<u>Kanter-Smoler et al.</u> 2008
-/?	01+promoter	del	g.35041-?_52505+? del	•	-	-	deletion, large	deletion, large	APC_00526	-	familial	Aretz et al. 2005
-/?	01+promoter	del	g.35041-?_52505+? del		-	-	deletion, large	deletion, large	APC_00526	-	familial	Aretz et al. 2005
-/?	01+promoter	del	g.35041-?_52505+? del		-	-	deletion, large	deletion, large	APC_00526	-	familial	<u>Aretz et al. 2005</u>
-/?	01_05+promoter	del	g.35041-?_78383+? del	•	-	-	deletion, large	deletion, large	APC_00527	-	familial	Aretz et al. 2005

#### Annotation Source - VarSifter

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File View Help

Chr 🗸 👘	LeftFlank 🗸	RightFlank	type	Gene_name	ref_aa	aa_pos	var_aa	dbID	HGMDids	HGMDdisease
chr5	112204359	112204361	Non-synonymous	APC	L	1724	V	-	-	-
chr5	112203573	112203575	Non-synonymous	APC	К	1179	E	-	-	-
chr5	112203562	112203568	DIV-c	APC	NA	0	NA	-	-	-
chr5	112203561	112203563	Non-synonymous	APC		1458	S	-	-	-
chr5	112203138	112203140	Non-synonymous	APC	E	1317	Q	rs1801166(C,G)	CM980089	"Colorectal cancer, predisp
chr5	112203109	112203111	Non-synonymous	APC	I	1024	К	rs1801155(A,T)	CM970090	"Colorectal cancer, predisp
chr5	112202773	112202775	Non-synonymous	APC	F	912	S	-	-	-
chr5	112202668	112202670	Non-synonymous	APC	Т	877	K	-	CM080043	Colorectal adenoma
chr5	112202649	112202661	DIV-c	APC	NA	0	NA	-	-	-
chr5	112202575	112202577	Non-synonymous	APC	L	1129	S	-	CM045407	Adenomatous polyposis coli
chr5	112202541	112202543	Non-synonymous	APC	N	1118	D	-	CM045405	Adenomatous polyposis coli
chr5	112202438	112202440	Non-synonymous	APC	D	1083	E	-	-	-
chr5	112202362	112202364	Non-synonymous	APC	D	1058	G	-	-	-
chr5	112201866	112201868	Non-synonymous	APC	E	893	К	-	CM013242	Adenomatous polyposis coli
chr5	112201797	112201799	Non-synonymous	APC	Р	870	S	rs33974176(C,T)	CM080070	Colorectal adenoma
chr5	112201627	112201629	Non-synonymous	APC	N	813	S	-	-	-
chr5	112201486	112201488	Non-synonymous	APC	A	766	V	-	-	-
chr5	112201393	112201395	Non-synonymous	APC	А	735	V	-	-	-
chr5	112192455	112192457	Non-synonymous	APC	I	544	Т	-	-	-
chr5	112191579	112191581	Non-synonymous	APC	S	535	F	-	-	-
chr5	112190789	112190791	Non-synonymous	APC	R	499	G	-	CM930023	Adenomatous polyposis coli
chr5	112182867	112182869	Non-synonymous	APC	R	414	С	-	CM910030	Adenomatous polyposis coli
chr5	112156122	112156124	Stop	APC	E	243	*	-	-	_
chr5	112156090	112156092	Non-synonymous	APC	R	232	Q	-	-	-
chr5	112144460	112144462	Non-synonymous	APC	Q	203	E	-	CM086466	Adenomatous polyposis coli
chr5	112130983	112130985	Non-synonymous	APC	E	140	D	-	-	_
chr5	112130951	112130953	Non-synonymous	APC	S	130	G	-	CM087822	"Colorectal cancer, severe
chr5	112130942	112130944	Non-synonymous	APC	S	127	G	-	CM024498	Adenomatous polyposis coli
chr5	112130880	112130882	Non-synonymous	APC	R	106	н	-	CM080058	Adenomatous polyposis coli
chr5	112118538	112118540	Non-synonymous	APC	М	18	К	-	-	_
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		previous page home about contact go to bottom
	Home	Mutalyzer 2.0 β-8
	Name Checker	released on 31 Jan 2011
•	Syntax Checker	HGVS nomenclature version 2.0
•	Position Converter	Position Conversion
•	SNP Converter	
•	Name Generator	Please supply the build which you want to use to convert your position, available builds at the moment are: hg18 (NCBI 36) and hg19 (GRCh37).
•	Batch Jobs	Example: NM_003002.2:c.274G>T
	Name Checker	or: chr11:g.111959693G>T
	Syntax Checker	or: NC_000011.9:g.111959693G>T
	Position Converter	Build hg18 🗘
	SNP Converter	Variant chr5:g.112202669C>A
	GenBank Uploader	
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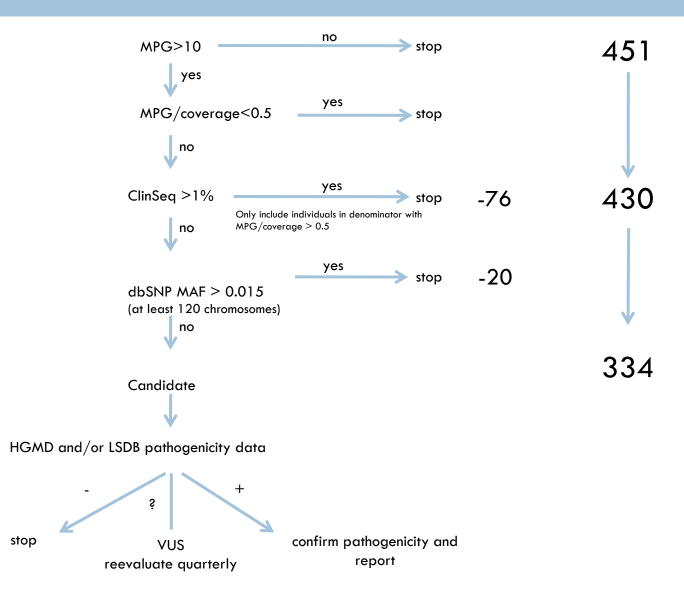
Mutalyzer http://www.mutalyzer.nl/2.0/

Home Name Checker	Mutalyzer 2.0 β-8 released on 31 Jan 2011
Syntax Checker	HGVS nomenclature version 2.0
Position Converter	Position Conversion
SNP Converter	Blasse supply the build which you want to use to convert your position, queilable builds at the memort and ha19 (NCRI 26) and ha10 (CRCh27)
Name Generator	Please supply the build which you want to use to convert your position, available builds at the moment are: hg18 (NCBI 36) and hg19 (GRCh37).
Batch Jobs	Example: NM_003002.2:c.274G>T
Name Checker	or: chr11:g.111959693G>T or: NC_000011.9:g.111959693G>T
Syntax Checker	
Position Converter	Build hg18 💠
SNP Converter	Variant chr5:g.112202669C>A
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Webservices	Submit Clear held
Help	
FAQ	Output:
Exercise	
Disclaimer	Chromosomal Variant:
Feedback	NC_000005.8:g.112202669C>A
External Links	
Human Gene	Found transprints in mutation region.
Nomenclature	Found transcripts in mutation region:
HGVS Variation	APC NM_001127510.2:c.3479C>A
Nomenclature HGVS Nomenclature	NM_001127511.2:c.3425C>A NM_000038.5:c.3479C>A
Extension Proposal	M_000038.4:c.3479C>A
= LOVD	NM_001127511.1:c.3479C>A NM_001127510.1:c.3479C>A
Mutalyzer 1.0.4	

HomeMutalyzer 2.0 β-8Name Checkerreleased on 31 Jan 2011Syntax CheckerHGVS nomenclature version 2.0	
= Syntax Checker HGVS nomenclature version 2.0	
Position Converter     Name Generator	
= SNP Converter	
Name Generator     Reference	
Batch Jobs Reference NM_000038.4:c.34790 Reference incorrect: should be of the format "NM_002001.2"	
Name Checker     Sequence Type Coding DNA	
= Syntax Checker Gene Symbol	
= Position Converter	
= SNP Converter Variant 1	
GenBank Uploader Mutation Type Substitution	
Webservices         Start Position         Start Position required. Start Position incorrect: position notation help	
Help     Deleted Sequence     Deleted Sequence     Deleted Sequence incorrect: substitution must consist of a single nucleotide / amino acid     Deleted Sequence incorrect: must consist of nucleotides [ACTG]	
FAQ     Inserted Sequence Inserted Sequence incorrect: substitution must consist of a single nucleotide / amino acid     Inserted Sequence incorrect: must consist of nucleotides [ACTG]	
Exercise	
Disclaimer	
Feedback     This field is optional	
External Links	
Human Gene     Constructed HGVS Name - Please click the link to check with the Name Checker	
Nomenclature         NM_000038.4:c.3479C>A:c.>	
= HGVS Variation	

<ul> <li>Home</li> <li>Name Checker</li> </ul>	Mutalyzer 2.0 β-8 released on 31 Jan 2011
Syntax Checker	HGVS nomenclature version 2.0
Position Converter	
SNP Converter	Name checker
Name Generator	Please insert the mutation name using the HGVS format:
Batch Jobs	<accession number="">.<version number="">(<gene symbol="">):<sequence type="">.<mutation></mutation></sequence></gene></version></accession>
Name Checker	Example: AB026906.1:c.274G>T
Syntax Checker	
Position Converter	NM_000038.4:c.3479C>A
SNP Converter	Submit Clear field
GenBank Uploader	
Webservices	Mutalyzer output:
Help	0 Errors, 0 Warnings.
FAQ	o Enois, o Warnings.
Exercise	Overview of the raw variants:
Disclaimer	
Feedback	Raw variant 1: substitution at 3564 CAGCATGAAGAAGAAGAAGAAGAAGAACAA C AAATTATAGCATAAAATATAATGAA
External Links	CAGCATGAAGAAGAAGAAGAAGAAGAACACAA A AAATTATAAGCATAAAATATAATGAA
Human Gene	
Nomenclature	Description relative to transcription start:
<ul> <li>HGVS Variation</li> <li>Nomenclature</li> </ul>	(Not for use in LSDBs in case of protein-coding transcripts).
= HGVS Nomenclature	<u>NM_000038.4:n.3564C&gt;A</u>
Extension Proposal	
= LOVD	Affected transcripts:
Mutalyzer 1.0.4	NM 000038.4(APC v001):c.3479C>A
	Affected proteins:
	NM_000038.4(APC_i001):p.(Thr1160Lys)

### **CS** Cancer Filtering



# International Association for Research on Cancer (IARC) Pathogenicity Scale

#### Proposed Classification System for Sequence Variants Identified by Genetic Testing

Class	Description	Probability of being pathogenic
5	Definitely pathogenic	>0.99
4	Likely pathogenic	0.95-0.99
3	Uncertain	0.05-0.949
2	Likely not pathogenic or of little clinical signi¢cance	0.001-0.049
1	Not pathogenic or of no clinical significance	<0.001

0 Insufficient information i.e. did not pass quality filter

Plon et al. Human Mut. (2008)

#### Variant Decision Examples

<i>APC</i> chr5	112,202,668- 112,202,670	NM_000038.4 c.3479C>A p.Thr1160Lys	3	1 in 258	-	CM080043 DM	Not in LSDB	two patients with CRA; rare variant hypothesis <sup>4</sup>
<i>APC</i> chr5	112,201,627- 112,201,629	NM_000038.4 c.2438A>G p.Asn813Ser	3	1 in 258	-	-	Not in LSDB	-
<i>BRCA2</i> chr13	31,812,437- 31,812,439	NM_000059.3 c.5946del p.Ser1982ArgfsX22	5	1 in 258	-	-	In LSDB <sup>1</sup> (7X): (?) BIC <sup>2</sup> (>1000X): clinically important	↑frameshift;↑cosegregation <sup>11</sup>
FLCN chr17	17,059,322- 17,059,324	NM_144997.5 c.1333G>A p.Ala445Thr	1L	1 in 255	rs41419545(C,T) no frequency data	-	In LSDB (1X): (-) https://grenada.lumc.nl/LSDB2/sh ared1/home.php?select_db=FLCN	-
<i>MSH6</i> chr2	47,879,811- 47,879,813	NM_000179.2 c.1186C>G p.Leu396Val	2	1 in 258	rs2020908(C,G) MAF 0.010 in 192 chr	CM101608 probable FP	In LSDB (19X): (?)	<ul> <li>◆3/200 control individuals;</li> <li>◆no significant mismatch repair defect<sup>24</sup></li> </ul>
<i>MUTYH</i> chr1	45,570,704- 45,570,706	NM_001048171.1 c.691C>T p.Arg231Cys	4	1 in 225	-	CM055444 DM	in LSDB (5X): (?)	↑biallelic; ↑MSH6 binding domain; 0/80 control individuals <sup>30</sup>
<i>RET</i> chr10	42,933,913- 42,933,915	NM_020630.4 c.2372A>T p.Tyr791Phe	2	2 in 258	-	CM971306 DM	In LSDB (7X)	

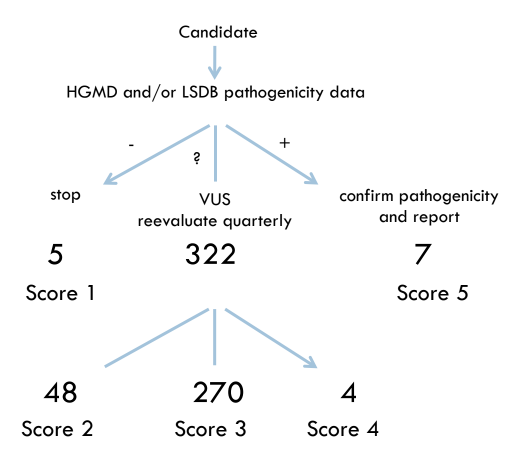
#### Variant Decision Examples

<i>APC</i> chr5	112,202,668- 112,202,670	NM_000038.4 c.3479C>A p.Thr1160Lvs	3	1 in 258	-	CM080043 DM	Not in LSDB	two patients with CRA; rare variant hypothesis⁴
<i>APC</i> chr5	112,201,627- 112,201,629	NM_000038.4 c.2438A>G p.Asn813Ser	3	1 in 258	-	-	Not in LSDB	-
<i>BRCA2</i> chr13	31,812,437- 31,812,439	NM_000059.3 c.5946del p.Ser1982ArgfsX22	5	1 in 258	-	-	In LSDB <sup>1</sup> (7X): (?) BIC <sup>2</sup> (>1000X): clinically important	↑frameshift;↑cosegregation <sup>11</sup>
FLCN chr17	17,059,322- 17,059,324	NM_144997.5 c.1333G>A p.Ala445Thr	1L	1 in 255	rs41419545(C,T) no frequency data	-	In LSDB (1X): (-) https://grenada.lumc.nl/LSDB2/sh ared1/home.php?select_db=FLCN	-
<i>MSH6</i> chr2	47,879,811- 47,879,813	NM_000179.2 c.1186C>G p.Leu396Val	2	1 in 258	rs2020908(C,G) MAF 0.010 in 192 chr	CM101608 probable FP	In LSDB (19X): (?)	↓3/200 control individuals; ↓no significant mismatch repair defect <sup>24</sup>
<i>MUTYH</i> chr1	45,570,704- 45,570,706	NM_001048171.1 c.691C>T p.Arg231Cys	4	1 in 225	-	CM055444 DM	in LSDB (5X): (?)	↑biallelic; ↑MSH6 binding domain; 0/80 control individuals <sup>30</sup>
<i>RET</i> chr10	42,933,913- 42,933,915	NM_020630.4 c.2372A>T p.Tyr791Phe	2	2 in 258	-	CM971306 DM	In LSDB (7X)	

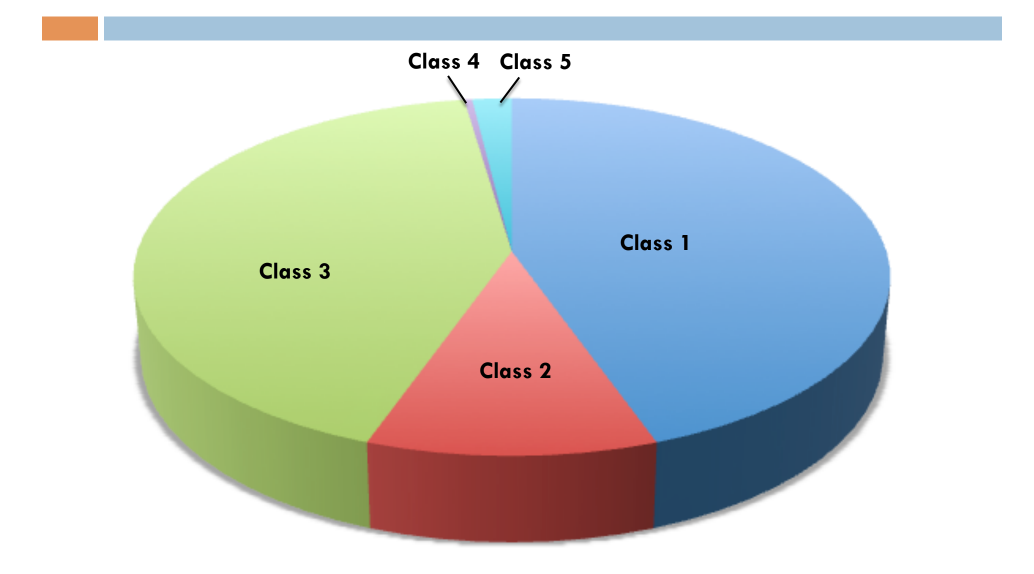
HGMD and LSDB often have conflicting information
 References cited do not always support causation

#### **MUST READ PRIMARY LITERATURE!!!!**

### **CS** Cancer Filtering



#### Summary of Variant Scores



#### **CS** Cancer Variants of Interest

- □ Three BRCA2 variants, both score 5
- □ Two BRCA1 variants, both score 5
- One SDHC variant, score 4- p.Arg15X, LOVD ?/+, Paraganglioma
- One FLCN variant, score 4- p.Lys508Arg, LOVD +/+?, Birt-Hogg-Dube syndrome
- Four variants in MUTYH (two 4, two 5s; AR; none biallelic)

#### Seven is a Big Number

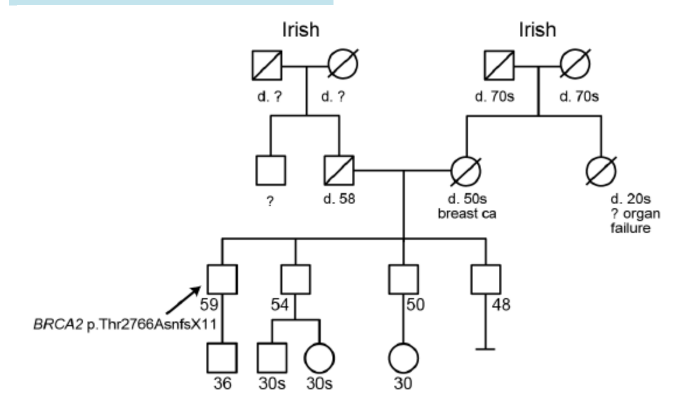
- Seven probands with BRCA1/2 variants in 572 ClinSeq cohort
  - □ All previously described
  - All associated with familial high-penetrance cancers
  - Only four had pedigrees that would lead to testing
  - Potentially life-saving results

#### Family History May not be Informative

#### BRCA2 c.8297delC Thr2766AsnfsX11 Classification: 5

#### **Evidence of Pathogenicity:**

Reported 41 times, no debate about pathogenicity



### Pathogenicity Score Criteria

Database Designation	Novel	Novel	Pathogenic	Pathogenic	VUS	Benign
Mutation Type	Missense	Nonsense Frameshift Splice	Missense	Nonsense Frameshift Splice	Any	Any
Score 5		Similar mutation type Consistent family history	Multiple reports, no evidence against	No evidence against		
Score 4		Similar mutation type Equivocal family history	Multiple reports, evidence against <b>OR</b> Single report, evidence for	Multiple reports, single evidence against	Multiple primary reports as pathogenic	
Score 3	All novel missense	Dissimilar mutation type Inconsistent family history	Single report, no supporting evidence	Multiple reports, multiple evidence against <b>OR</b> Single report, single evidence against	Primary reports as VUS	Single report OR primary reports as pathogenic
Score 2			Single report, multiple evidence against	Single report, multiple evidence against	Multiple evidence against	Multiple reports, no supporting evidence <b>OR</b> Single report, evidence against

# A Cautionary Tale

Gene symbol	Disease / phenotype	Location	HGMD accession
CDH1	Gastric cancer	16q22.1	CM041745

Disease/phenotype	Gene symbol	Codon change	Amino acid change	Codon number	Reference
Gastric cancer	CDH1	tGCC-ACC	Ala-Thr	298	Brooks-Wilson (2004) Journal of medical genetics ( <i>J Med Genet</i> ) <b>41</b> : p508. PubMed: <u>15235021</u> Mateus (2009) <i>Exp Cell Res</i> <b>315</b> : p.1393 [Functional characterisation] PubMed: <u>19268661</u>

The A298T substitution exhibits a shift in polarity from non-polar to polar and displays a decrease in Kyte-Doolittle hydrophobicity from 1.8 to -0.7. Approximately 1.77% of missense mutations in HGMD are Ala-Thr.

Extra information					
Genomic sequence (build 37.2 - NEW)	GACGCGGACGATGATGTGAACACCTACAAT(G-A)CCGCCATCGCTTACACCATCCTCAGCCAAG				
Amino acid sequence	DADDDVNTYNAAIAYTILSQD (Amino acid alignment)				
Genomic coordinate (build 37.2 - NEW)	Chr 16: <u>68845646;</u> Chr 16: <u>68845646-68845648</u>				
Genomic coordinate (build 36.3 - REMOVAL 2012.1)	Chr 16: <u>67403147</u>				
HGVS nomenclature	<u>NM_004360.3</u> : c.892G>A; <u>NP_004351.1</u> : p.A298T				
dbSNP number	No dbSNP ID				
Variant class	Disease causing mutation				
Comments	No comments				
CpG	No				

An	nino acid comparison table				
Trait	Ala(A)	Thr(T)			
Amino acid name	alanine	threonine			
Polarity/charge	non-polar	polar			
pH	neutral	neutral			
Residue weight	71	101			
Hydrophobicity score	1.8	-0.7			
Hydrophilicity score	-0.5	-0.4			
Secondary structure propensity	strong $\alpha$ former / $\beta$ indifferent	$\alpha$ indifferent / $\beta$ former			
Grantham difference	58				
Sift prediction	TOLERATED				

# **Cautionary Tale**

J Med Genet. 2004 Jul;41(7):508-17.

#### Germline E-cadherin mutations in hereditary diffuse gastric cancer: assessment of 42 new families and review of genetic screening criteria.

Brooks-Wilson AR, Kaurah P, Suriano G, Leach S, Senz J, Grehan N, Butterfield YS, Jeyes J, Schinas J, Bacani J, Kelsey M, Ferreira P, MacGillivray B, MacLeod P, Micek M, Ford J, Foulkes W, Australie K, Greenberg C, LaPointe M, Gilpin C, Nikkel S, Gilchrist D, Hughes R, Jackson CE, Monaghan KG, Oliveira MJ, Seruca R, Gallinger S, Caldas C, Huntsman D.

Table 2 Details of the gastric cancer families in the study and mutations detected

Family no	Cancer type, age	Study criteria met	Other family members with gastric cancers, n (ages)	Family members with breast cancer, n (confirmed lobular breast cancer)	CDH1 mutation: exon, nucleotide (amino acid)	Type of mutation
F26	DGC, 36	1	2 (32†, 33)	0	Exon 7, G892A (A298T)	Missense

#### The mutations found

include small insertions and deletions, splice site mutations, and three non-conservative amino acid substitutions (A298T, W409R, and R732Q). All three missense mutations conferred loss of E-cadherin function in vitro assays.

# What should we consider when returning carrier variants?

#### **Disease-Gene-Variant**

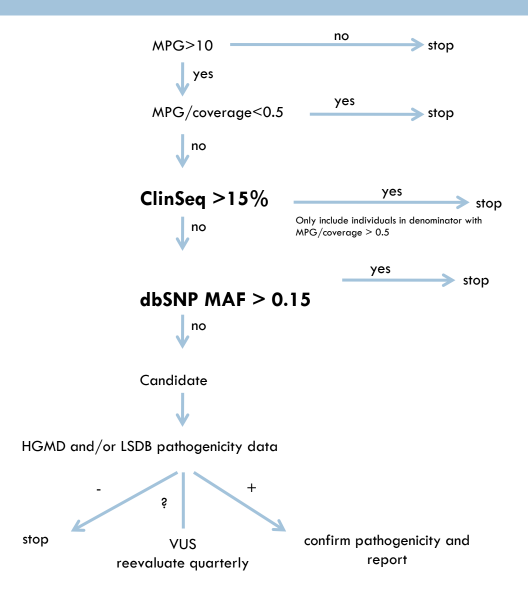
Severity of disease

Genes proven to cause disease

Variants with known pathogenicity

Threshold for disease incidence?

#### **Framework for Carrier Variants**



# Ambrygen Gene List

78 genes offered in prenatal panel

□ 75 AR, 3 X-linked

1:2,500 for CF to 1:1,000,000 for Beta ketothiolase deficiency

http://www.ambrygen.com/

#### **Common Recessive disease**

ETHNICITY	DISEASE	
Ashkenazi Jewish:	Tay-Sachs	1/30
	Canavan	1/40
	Cystic fibrosis	1/29
	Familial Dysautonomia	1/30
Mediterranean:	Thalassemia	1/20-1/50
	Sickle cell anemia	1/30-1/50
European Caucasian:	Cystic fibrosis	1/29
African American:	Sickle cell anemia	1/10
	Thalassemia	1/30-1/75
	Cystic fibrosis	1/65
Asian:	Thalassemia	1/20-1/50
	Cystic fibrosis	1/90
Hispanic:	Cystic fibrosis	1/46
French Canadian:	Tay-Sachs	1/15
	Cystic fibrosis	1/29

Population Risk: 1/30 \* 1/30 \* 1/4 = 1/3600

Known Carrier Risk: 1/30 \* 1/4 = 1/120

#### 30 X population risk

http://www.hopkinsmedicine.org/fertility/resources/genetic\_screening.html

#### Extremely Rare Recessive disease

Beta ketothiolase deficiency Population Risk (1 in a million): 1/500 \* 1/500 \* 1/4 = 1/1,000,000

Known Carrier Risk: 1/500 \* 1/4 = 1/2,000

500 X population risk

#### What did we find?

- 10 stops in HGMD
- □ 216 nonsynonymous in HGMD
- 11 novel stops
- 25 frame shifts
- □ 5 in frame deletions
- □ 14 splice not in HGMD

□ 623 were nonsynonymous changes not present in HGMD

#### CS Carrier Variants - 78 Genes

CFTR – Cystic Fibrosis - p.  $\Delta$  F508, 7/571

BBS10 – Bardet Biedl - c.271 dup, common mutation, 2/401

ASPA – Canavan disease - p.Glu285Ala, founder AJ, 1/564

IDUA - Hurler - p.Ala327Pro, common mutation, 1/522

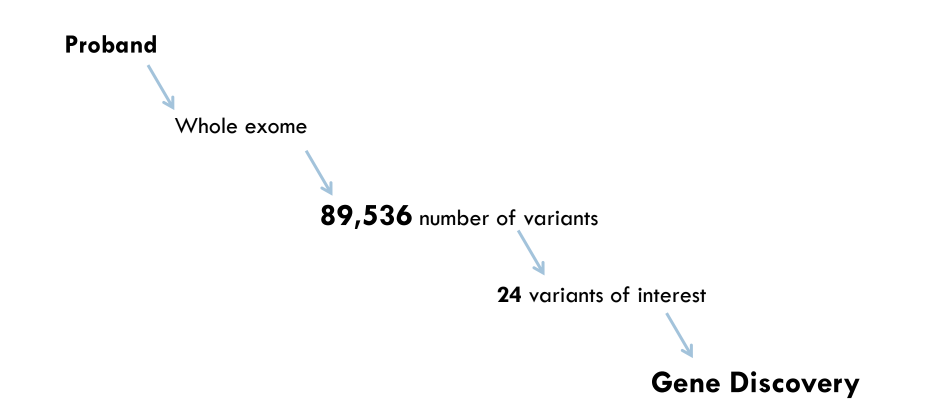
GALT – Galactosaemia, p.Gln188Arg, common mutation, 3/574

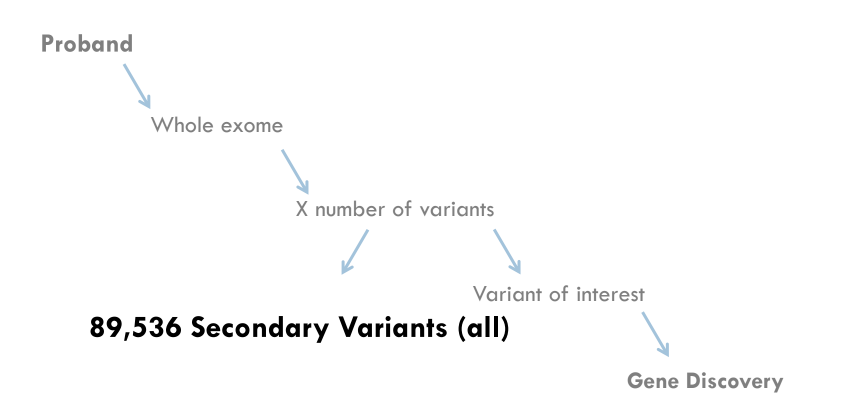
G6PC – Glycogen Storage 1a, p.Arg83Cys, founder AJ, 4/572

MUT - p.Asn219Tyr, common methylmalonic aciduria mutation 1/572

# How might we think of things differently for a trio?

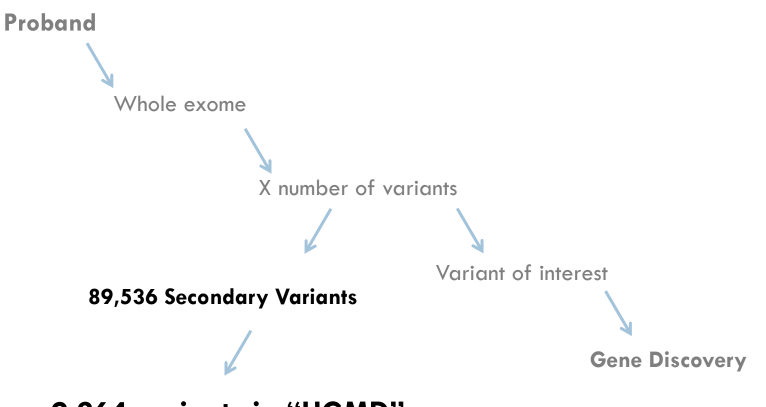




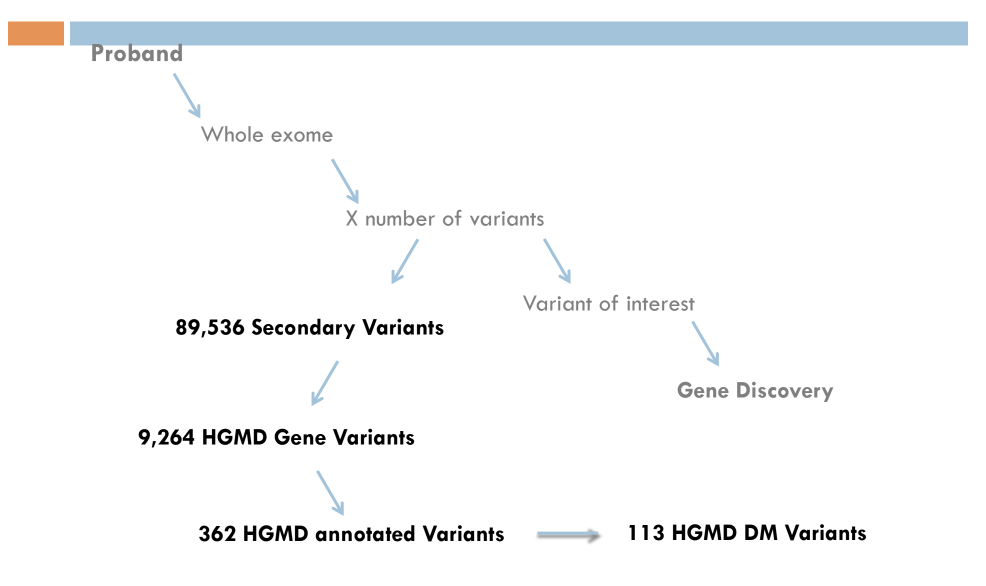


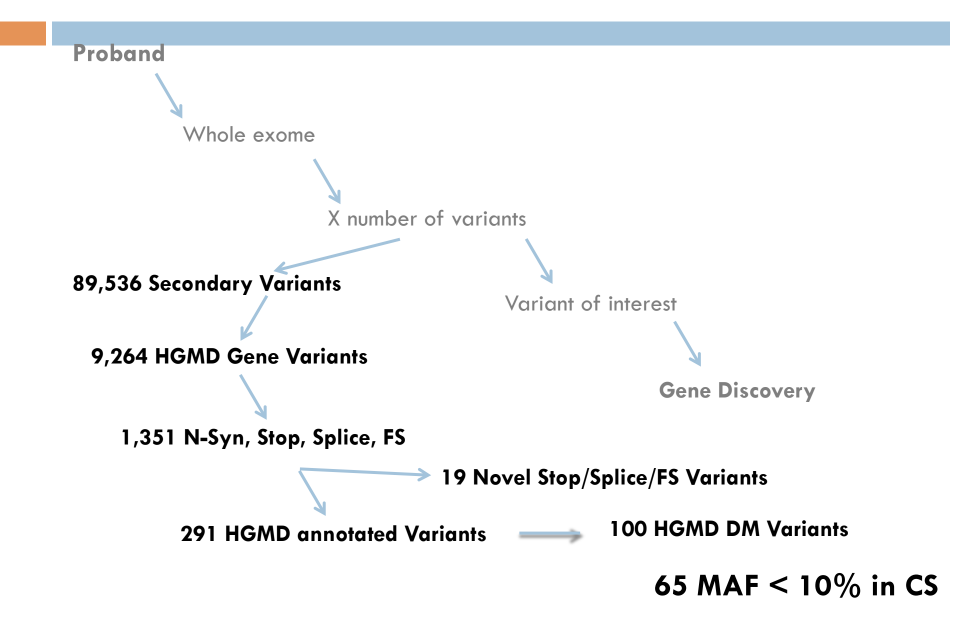
### VarSifter - HGMD

ile \	/iew Help							
Chr	LeftFlank	RightFlank	Gene_name 🗚	HGMDids	HGMDdisease	HGMDtags	HGMDinGene	transcript
chr13	31810006	31810008	BRCA2	CM050182	Breast cancer ?	DM	у	uc001uub.1
chr13	31810053	31810055	BRCA2	-	-	-	y	uc001uub.1
chr13	31810072	31810074	BRCA2	-	-	-	y	uc001uub.1
chr13	31810749	31810751	BRCA2	CM003133	Breast and/or ovarian cancer ?	DM	y	uc001uub.1
chr13	31811084	31811086	BRCA2	-	-	-	у	uc001uub.1
chr13	31811270	31811272	BRCA2	-	-	-	У	uc001uub.1
chr13	31811689	31811691	BRCA2	-	-	-	y	uc001uub.1
chr13	31811803	31811805	BRCA2	CM041731	Breast and/or ovarian cancer ?	DM	y	uc001uub.1
chr13	31811970	31811979	BRCA2	-	-	-	y	uc001uub.1
chr13	31812043	31812045	BRCA2	-	-	-	y	uc001uub.1
chr13	31812045	31812047	BRCA2	-	-	-	y	uc001uub.1
chr13	31812223	31812225	BRCA2	-	-	-	y	uc001uub.1
chr13	31812235	31812237	BRCA2	CM010170	Breast cancer ?	DM	ý	uc001uub.1
chr13	31812437	31812439	BRCA2	-	-	-	ý	uc001uub.1
chr13	31812591	31812593	BRCA2	CM994286	Breast and/or ovarian cancer ?	DM	ý	uc001uub.1
chr13	31812813	31812815	BRCA2	-	-	-	ý	uc001uub.1
chr13	31812816	31812818	BRCA2	CM043917	Breast cancer ?	DM	ý	uc001uub.1
chr13	31812829	31812831	BRCA2	-	-	-	ý	uc001uub.1
chr13	31812838	31812840	BRCA2	CM022331	Breast and/or ovarian cancer	DM	ý	uc001uub.1
chr13	31816705	31816707	BRCA2	-	-	-	y	uc001uub.1
chr13	31827308	31827310	BRCA2	-	-	-	y	uc001uub.1
chr13	31827386	31827388	BRCA2	CM960194	Breast cancer	DM	ý	uc001uub.1
chr13	31828632	31828634	BRCA2	CM012590	Breast and/or ovarian cancer	DM	ý	uc001uub.1
chr13	31828672	31828674	BRCA2	CM994287	Breast and/or ovarian cancer ?	DM	y	uc001uub.1
chr13	31835487	31835489	BRCA2	CM043984	Breast and/or ovarian cancer	DM	y	uc001uub.1
chr13	31835520	31835522	BRCA2	CM004715	Breast cancer	DM	v	uc001uub.1



9,264 variants in "HGMD" genes





#### Secondary Variant Paradox

- Exome/WGS sequencing can uncover life altering predictive information
- Value can only be appreciated if research practitioners annotate exomes for secondary variants

#### What is the future?

- Secondary annotation is a burden (for researchers)
- It is important
- NIH and others need to improve resources for this
  - Databases
  - Interpretation tools & services