In silico analysis of \( \text{BRCA1} \) and \( \text{BRCA2} \) missense variants and the relevance in molecular genetic testing

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Over the years since the genetic testing of \( \text{BRCA1} \) and \( \text{BRCA2} \) has been conducted for research and later introduced into clinical practice, a high number of missense variants have been reported in the literature and deposited in public databases. Polymorphism Phenotyping v2 (PolyPhen-2) and Sorting Intolerant from Tolerant (SIFT) are two widely applied bioinformatics tools used to assess the functional impacts of missense variants. A total of 2605 \( \text{BRCA1} \) and 4763 \( \text{BRCA2} \) variants from the ClinVar database were analysed with PolyPhen2 and SIFT. When SIFT was evaluated alongside PolyPhen-2 HumDiv and HumVar, it had shown top performance in terms of negative predictive value (NPV) (100%) and sensitivity (100%) for ClinVar classified benign and pathogenic \( \text{BRCA1} \) variants. Both SIFT and PolyPhen-2 HumDiv achieved 100% NPV and 100% sensitivity in prediction of pathogenicity of the \( \text{BRCA2} \) variants. The performances of PolyPhen-2 and SIFT in predicting functional impacts varied across the two genes. Due to lack of high concordance in prediction outcomes among the two tested algorithms, their usefulness in classifying the pathogenicity of VUS identified through molecular testing of \( \text{BRCA1} \) and \( \text{BRCA2} \) is hence limited in the clinical setting.

The \textit{Homo sapiens BReast CAnce 1, early onset (BRCA1)} and \textit{BReast CAnce 2, early onset (BRCA2)} genes were identified in 1994\cite{1} and 1995\cite{2}, respectively. They are the two main susceptibility genes which confer high risks in their pathogenic variant carriers for the hereditary breast and ovarian cancers (HBOC). In addition to the \( \text{BRCA1} \) and \( \text{BRCA2} \) genes, there are other susceptibility genes reported to date as high penetrance and moderate penetrance genes involving in the development of breast cancer. The high penetrance genes include the \( \text{TP53} \)\cite{3}, \( \text{PTEN} \)\cite{4}, \( \text{STK11} \)\cite{5} and \( \text{CDH1} \)\cite{6}. Not only conferring high risk to breast cancer, mutation carriers of the above-mentioned genes are also having higher chances for developing other types of cancerous syndrome with different tiers of risk. On the other hand, moderate penetrance genes such as \( \text{PALB2} \)\cite{7}, \( \text{CHEK2} \)\cite{8}, \( \text{ATM} \)\cite{9} and \( \text{RAD51C} \)\cite{10} are also well-studied for their conferred risks to breast cancer. Among of all the afore-mentioned susceptibility genes, \( \text{BRCA1} \) and \( \text{BRCA2} \) remain the two most frequently tested genes for the genetic testing of hereditary breast cancer in the current clinical settings.

The \textit{BRCA1} gene (NCBI RefSeq NG_005905; LRG_292) is located at the chromosome 17q21 and spanning a genomic region of 193 kilo bases. It has several isoforms of messenger RNA (mRNA) transcript. Current standard mRNA transcript, NM_007294 contains 22 coding exons resulting in a translated protein with 1863 amino acids (NP_009225). The \textit{BRCA2} gene (NCBI RefSeq NG_012772; LRG_293) with chromosomal position at 13q13 has a genomic sequence of 91 kilo bases. The standard mRNA transcript is NM_000059. This transcript encodes 3418 amino acids (NP_000050) in 26 coding exons. Both \textit{BRCA1} and \textit{BRCA2} play roles in homologous recombination pathway for repairing double-strand DNA. They are tumor-suppressor genes, hence the mutated alleles give predisposition of cancers to the carriers\cite{11}.

Over the years since the genetic testing of \( \text{BRCA1} \) and \( \text{BRCA2} \) has been conducted for research and clinical practice, a high number of sequence variants have been reported in the literature and deposited in public databases. The sequence variants identified from the studied subjects can be categorized into five classes\cite{12}. Common benign polymorphisms including variants without pathogenicity or of no clinical significance are grouped as Class 1. Class 2 includes variants that are likely benign or of little significance. Class 3 represents
variant of unknown significance (VUS) while classes 4 and 5 include likely pathogenic and definite pathogenic variants, respectively. The ENIGMA (Evidence-based Network for the Interpretation of Germline Mutant Alleles) consortium has been playing an important role to evaluate the clinical significance of sequence variants in high-risk breast cancer genes including BRCA1 and BRCA2 genes since its establishment in 2009. The BRCA Exchange portal also serves as a consolidated resource for aggregating curated BRCA1 and BRCA2 variants for public access.

The frequency of reporting VUS is up to 20% for individuals undergoing genetic testing of BRCA1 and BRCA2 but can be lower in other well-studied populations. Identification of VUS is considered as inconclusive and clinically not actionable. While creating uncertainty for treatment decision, reporting VUS has the potential to raise anxiety in its carriers. A large number of VUS have been listed in publicly available databases such as the Breast Cancer Information Core (BIC) database and ClinVar. The VUS usually consist of non-truncating single nucleotide changes resulting in missense variants, small in-frame insertions or deletions and variants in intronic, non-coding and untranslated regions. The VUS can be novel at the time it was reported due to the unavailability of prior clinical or functional evidence about its causality. Over the time, the status of a particular VUS can be shifted to either benign or pathogenic after sufficient findings have been accumulated.

With the advances of bioinformatics and computational biology, many in silico tools have been developed to help the biologists, scientists, geneticists and clinicians to predict the effects and potential significance of the missense coding variants. Polymorphism Phenotyping v2 (PolyPhen-2) and Sorting Intolerant from Tolerant (SIFT) are two widely applied bioinformatics tools used to assess functional effects of missense variants. The PolyPhen-2 gives predictions based on both sequence alignment and structural features characterizing the amino acid substitution while SIFT solely considers the sequence homology and protein conversation among species to help the biologists, scientists, geneticists and clinicians to predict the effects and potential significance of the missense variants.

### Results

#### Prediction results from PolyPhen-2 and SIFT on known benign and pathogenic variants.

There were 165 and 142 missense variants with known clinical significance (classified as benign or pathogenic), for BRCA1 and BRCA2 respectively which were subjected to both PolyPhen-2 and SIFT predictions. The prediction results for the benign and pathogenic entries were summarised in Supplementary Table S1 and Supplementary Table S2 for BRCA1 and BRCA2 respectively. For PolyPhen-2, the variants predicted as “possibly damaging” and “probably damaging” were grouped together as “damaging” variants. The positive predictive value (PPV), negative predictive value (NPV), sensitivity, specificity, and accuracy were summarised in Table 1.

#### Evaluation of prediction results of variants with known clinical significance.

Poor performances of PPV were noted for the BRCA1 and BRCA2 genes with the top-performers being PolyPhen-2 HumVar—33.85% for BRCA1 and 27.91% for BRCA2, respectively. In terms of PPV, all three approaches achieved good results (NPV > 85%) in the two genes. For BRCA1 (Fig. 1), SIFT and PolyPhen-2 HumVar showed highest sensitivity (100%) and highest specificity (67.18%), respectively. For BRCA2 (Fig. 2), both PolyPhen-2 HumDiv and SIFT showed 100% sensitivity while PolyPhen-2 HumVar achieved highest specificity (75.97%). PolyPhen-2 HumVar showed highest accuracies for BRCA1 (66.67%) and BRCA2 (77.46%) genes, respectively.

#### Prediction results from PolyPhen-2 and SIFT on BRCA1 and BRCA2 missense VUS.

There were 2440 BRCA1 and 4621 BRCA2 missense VUS analysed by both PolyPhen-2 and SIFT. For BRCA1 (Fig. 3), PolyPhen-2 HumDiv predicted benign effects in 48.07% (N = 1173) and damaging effects in 51.93% tested VUS (N = 1267). PolyPhen-2 HumVar predicted benign effects in 63.07% (N = 1539) and damaging effects in 36.93% tested VUS (N = 901). SIFT predicted tolerated effects in 36.48% VUS (N = 890) and damaging effects in 63.52% VUS (N = 1550). Agreement was achieved in prediction outcomes from the three tested approaches in 55.04% (N = 1343) of the tested VUS with 620 and 723 VUS being predicted as benign and damaging, respectively (Fig. 4). On the other hand, PolyPhen-2 HumDiv achieved highest consensus with PolyPhen-2 HumVar in predicting benign VUS. However, for the pathogenic predictions, highest consensus was achieved between PolyPhen-2 HumVar and SIFT.
SIFT and PolyPhen-2 HumDiv (Fig. 5). Notably, about 26.32% (N = 452) and 35.44% (N = 645) of predictions observed were unique (outstanding predictions given by only one among three algorithm) for benign and damaging categories, respectively. This implied that higher concordance (consensus in at least two out of three tested approaches) was achieved for benign prediction in BRCA1 VUS.

On the other hand, for BRCA2 (Fig. 6), PolyPhen-2 HumDiv predicted damaging effects in 44.99% (N = 2079) and benign effects in 55.01% (N = 2542) tested VUS. PolyPhen-2 HumVar predicted damaging effects in 31.16% (N = 1440) and benign effects in 68.84% (N = 3181) tested VUS. SIFT predicted damaging effects in 51.48% (N = 2379) and tolerated effects in 48.52% (N = 2242) of tested VUS. Agreement was achieved in prediction outcomes from the three tested approaches in 68.97% (N = 3187) of the tested VUS with 1872 and 1315 VUS being predicted as benign and damaging, respectively (Fig. 7). On the other hand, PolyPhen-2 HumDiv achieved highest consensus with PolyPhen-2 HumVar in predicting benign VUS. However, for the damaging predictions, highest consensus was achieved between SIFT and PolyPhen-2 HumDiv (Fig. 8). Notably, about 15.70% (N = 519) and 33.28% (N = 915) of predictions observed were unique (outstanding predictions given by only one among three algorithm) for benign and damaging categories, respectively. Similar to the observations on BRCA1 VUS,
Agreement in prediction outcomes for BRCA1 missense variants

**Figure 4.** Summary of agreement in prediction outcomes for BRCA1 missense VUS.

Consensus of predictions between SIFT, PolyPhen-2 HumVar and HumDiv in BRCA1 missense VUS

**Figure 5.** Summary of consensus in prediction outcomes for BRCA1 missense VUS.

Distribution of prediction outcomes for BRCA2 missense VUS

**Figure 6.** Summary of distribution of prediction outcomes on BRCA2 missense VUS.
higher concordance (consensus in at least two out of three tested approaches) was achieved for benign predic-
tion in BRCA2 VUS.

Discussions

The current study compared the prediction performance of PolyPhen-2 HumDiv, PolyPhen-2 HumVar and SIFT using benchmark datasets for BRCA1 and BRCA2 genes. The study was further extended to evaluate their practical use to predict the pathogenicity of reported VUS in the context of the BRCA1 and BRCA2 genes. Analysis was based on the curated variants with expert review from the ClinVar18 which is widely referred by genetic laboratory personnel, clinicians and scientists nowadays for interpreting the molecular genetic testing results.

Results from this study show differences between the performance of PolyPhen-2 HumDiv, PolyPhen-2 HumVar and SIFT. Of the missense variants with known clinical significance (classified benign or pathogenic), there was concordant agreement achieved in the predictions in 96 (58.18%) of 165 BRCA1 and in 95 (66.90%) of 142 BRCA2 variants for all three approaches used, respectively. The discordant predictions are attributed to the inherent characteristics of different systems used by different algorithms. The PolyPhen-2 combines information from protein sequence and structure to analyse missense variants. However, two different datasets had been used to train the HumDiv and HumVar, hence resulting in different analytical approaches to a given missense variant20. HumDiv data is mainly used to discriminate the rare alleles in complex disease phenotypes and in natural selection. On the other hand, HumVar data is more relevant in analysing variants of Mendelian disorders, which more stringently recognise highly disease-causing mutations from benign polymorphism existing in the normal population. The hereditary breast cancers do not clearly fit into either category of complex or Mendelian disorder because of the incomplete penetrance imposed by the two pre-disposing genes. The HumVar had shown higher overall accuracy than HumDiv i.e. 66.67% versus 60.00% and 77.46% versus 61.27% for BRCA1 and BRCA2 missense variants, respectively. Notably, both HumDiv and HumVar had higher NPV (> 85%) than PPV (< 35%) indicating the PolyPhen-2 had given more confident results in predicting benign variants. When SIFT was evaluated along with PolyPhen-2 HumDiv and HumVar, it had shown top performance in terms of NPV (100%) and sensitivity (100%) for BRCA1 and BRCA2. Notably, HumDiv had also shown 100% NPV and
100% sensitivity for BRCA2. Hence, in this study although some similar performances were observed, these algorithms were noted to have varied predictive performances for BRCA1 and BRCA2 missense variants, respectively. Notably, the NPV of the three tested approaches were all above 85% (Table 1). However, the other performances including PPV, specificity and accuracy were generally low (<80%), hence, rendering clinical applicability of these approaches unsuitable. A limitation of the current study is the small size of datasets of tested missense variants with known clinical significance for BRCA1 (Benign, N = 131; Pathogenic, N = 34) and BRCA2 (Benign, N = 129; Pathogenic, N = 13), respectively.

Genetic testing for BRCA1 and BRCA2 genes can be important to provide actionable results to the clinicians and genetic counsellors who are directly dealing with the patients and their relatives. The utility of such testing is strongly dependent on the interpretation of variants identified. The current laboratory practice in variant interpretation include searching the published literature and online curated databases for evidence of pathogenicity. Frequently the identified missense variants are novel and are not catalogued in the existing publicly available databases. In vitro assays are performed by research laboratories to study the functional impact of the missense variant to the protein, however, this is beyond the ability of most clinical laboratories. The functional tests for BRCA1 and BRCA2 are generally time-consuming, laborious and costly. Studies of familial co-segregation of variant with cancer is feasible by providing carrier testing but such extended testing involves genotyping of many relatives collectively before conclusion can be firmly drawn. Again, such strategy is more applicable in the research setting.

Although lacking evidence from functional analysis and familial co-segregation with cancer, a rare missense variant is not always considered as VUS. Apart from very frequent polymorphic variants, according to ENIGMA BRCA1/2 gene variant classification criteria, a missense variant could be classified as class 1 if it has a prior probability of pathogenicity ≤0.02 from clinically calibrated bioinformatic analyses and an allele frequency ≥ 0.001 and <0.01 in large outbred control reference groups. Reporting VUS in the clinical report could lead to problematic interpretation since such results cannot be readily utilised to identify the at-risk family members and provide indication for increased surveillances. Such uninformative results could also possibly raise the misperception in the patients and family members that they are at higher risk of developing cancers.

In the current study, low consensus was noted when the three approaches were used to predict the pathogenicity of the VUS. Agreement was achieved in prediction outcomes from the three tested approaches in 55.04% and 68.97% of the VUS for BRCA1 and BRCA2, respectively. The lack of concordance in prediction outcomes can cause confusion in interpreting the significance of the missense variants. Previous comparative studies which evaluated other in silico algorithms suggested that combining several prediction tools could improve the prediction performance, however, the study by Walters-Sen et al. had reported the opposite.

There had been several other studies conducted in silico analysis on BRCA1 and BRCA2 missense variants. Based on the datasets from a commercial testing laboratory, Kerr et al. reported that SIFT presented 100% sensitivity and NPV in predicting both BRCA1 and BRCA2 variants, a similar finding as ours. Ernst et al. also showed 100% sensitivity for SIFT predictions on both BRCA1 and BRCA2 variants, which was also in congruent with the findings from our study. However, for accuracy we have shown that PolyPhen-2 had better performance than SIFT as compared to the studies by Kerr et al. and Ernst et al. Another multifactorial likelihood study by Parsons et al. had reported quantitative analysis of BRCA1 and BRCA2 variants in which bioinformatic prediction of variant effect was part of the analysis. Parsons et al. showed that the incorporation of clinical information can improve variant curation over purely bioinformatic approaches. This again highlights that bioinformatic approaches can at most serve as adjunct evidence in variant classification, and should not be solely considered in calling variant pathogenicity.

In conclusion, the performances of PolyPhen-2 and SIFT in predicting functional impacts varied across a clinical dataset of BRCA1 and BRCA2 missense variants. Lack of high concordance in prediction outcomes highlighted their limited clinical application in classifying the pathogenicity of VUS identified through molecular testing of BRCA1 and BRCA2.

Methodology

Mining of BRCA1 and BRCA2 missense variants. ClinVar is a public depository of genetic variants allowing submissions of curations with clinical significance. ClinVar is a freely accessible database (https://www.ncbi.nlm.nih.gov/clinvar/). The variants of the BRCA1 and BRCA2 genes as of March 16, 2021 were downloaded after applying filters including “missense” for “molecular consequence” and “expert panel” for “review status”. The datasets were imported into a spreadsheet software, Microsoft Excel for Mac, visualized and analysed. Entries of different DNA alterations representing the identical missense protein change were removed. For BRCA1, a missense variant, c.5074G > C (p.Asp1692His) was excluded from analysis due to its proven effects on mRNA splicing instead of amino acid substitution. A total of 307 expert reviewed entries consisting of 34 pathogenic and 131 benign BRCA1, and 13 pathogenic and 129 benign BRCA2 variants were subject to in silico analysis for functional predictions.

The missense variants of unknown significance (VUS) were also sought for the BRCA1 and BRCA2 genes in ClinVar by applying filters including “missense” for “molecular consequence” and “uncertain significance” for “clinical significance”. Duplicate entries with identical missense protein change were removed. There were 2440 BRCA1 and 4621 BRCA2 VUS included in this study.

In silico analysis using PolyPhen-2. PolyPhen-2 is an online tool for prediction of the functional consequences of an amino acid substitution on a human protein. Polyphen-2 web interface was accessed at http://genetics.bwh.harvard.edu/pph2/index.shtml. Batch query mode was used for analysis. The query line of missense variant was outlined according to the format:
In silico analysis using SIFT. SIFT is an online tool that uses sequence homology to judge the functional impact of missense variants. SIFT web interface was accessed at http://sift.jcvi.org/. SIFT Human Protein was selected. The query line of missense was outlined by the Protein Ensembl ENSP ID followed by the specified substitution: “ENSP ID, substitution”. The Ensembl ENSP ID used for BRCA1 and BRCA2 were ENSP00000350283 and ENSP00000439902, respectively. The batch protein tool was used to multiple queries. SIFT give prediction outcomes for missense variants as damaging (score < 0.05) and tolerated (score ≥ 0.05).

Evaluation of prediction results of variants with known clinical significance. The performance of the PolyPhen-2 and SIFT was analyzed by comparing the statistical calculations:

- Positive Predictive Value (PPV) = TP / (TP + FP)
- Negative Predictive Value (NPV) = TN / (TN + FN)
- Sensitivity = TP / (TP + FN)
- Specificity = TN / (TN + FP)
- Accuracy = (TP + TN) / (TP + FP + TN + FN)

For these calculations, TP or True Positives are pathogenic variants with CLINVAR expert review called as possibly damaging or probably damaging by PolyPhen-2, and damaging by SIFT, respectively. FP or False Positives are benign variants called as possibly damaging or probably damaging by PolyPhen-2, and damaging by SIFT, respectively. TN or True Negatives are benign variants called as benign by PolyPhen-2, and tolerated by SIFT, respectively. FN or False Negatives are pathogenic variants called as benign by PolyPhen-2, and tolerated by SIFT, respectively. Calculations were performed using MedCal diagnostic test evaluator calculator.

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**Competing interests**

The author declares no competing interests.

**Additional information**

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