

# Analysis of more than 400,000 women provides case-control evidence for *BRCA1* and *BRCA2* variant classification

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
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Clinical genetic testing identifies variants causal for hereditary cancer, information that is used for risk assessment and clinical management. Unfortunately, some variants identified are of uncertain clinical significance (VUS), complicating patient management. Case-control data is one evidence type used to classify VUS. As an initiative of the Evidence-based Network for the Interpretation of Germline Mutant Alleles (ENIGMA) Analytical Working Group we analyze germline sequencing data of *BRCA1* and *BRCA2* from 96,691 female breast cancer cases and 302,116 controls from three studies: the BRIDGES study of the Breast Cancer Association Consortium, the Cancer Risk Estimates Related to Susceptibility consortium, and the UK Biobank. We observe 11,207 *BRCA1* and *BRCA2* variants, with 6909 being coding, covering 23.4% of *BRCA1* and *BRCA2* VUS in ClinVar and 19.2% of ClinVar curated (likely) benign or pathogenic variants. Case-control likelihood ratio (ccLR) evidence is highly consistent with ClinVar assertions for (likely) benign or pathogenic variants; exhibiting 99.1% sensitivity and 95.3% specificity for *BRCA1* and 93.3% sensitivity and 86.6% specificity for *BRCA2*. This approach provides case-control evidence for 787 unclassified variants; these include 579 with strong or moderate benign evidence and 10 with strong pathogenic evidence for which ccLR evidence is sufficient to alter clinical classification.

Clinical genetic testing of disease-associated susceptibility genes is often complicated by the identification of variants of uncertain clinical significance (VUS)<sup>1</sup>. These individually rare variants may include missense changes, intronic and small in-frame insertions or deletions, as well as regulatory variants for which the association with disease is uncertain, complicating counselling and clinical management<sup>2</sup>. This is especially true for *BRCA1* and *BRCA2*, where pathogenic variants (PVs) confer a high and clinically significant risk of breast<sup>3,4</sup>, ovarian<sup>5,6</sup>, pancreatic<sup>7</sup> and (in the case of *BRCA2*) prostate cancer<sup>8,9</sup>; individuals with a VUS are generally not eligible to access well recognized clinical management options offered to individuals with a PV, such as risk-reducing prophylactic surgery to prevent cancer, magnetic resonance imaging (MRI) screening or PARP-inhibitor treatment for cancers that arise<sup>10–13</sup>.

To facilitate the classification of variants identified by genetic testing into (likely) benign or (likely) pathogenic, the American College of Medical Genetics and Genomics and the Association for Molecular Pathology (ACMG/AMP) groups<sup>14</sup> developed guidelines that incorporate a set of criteria representing different evidence types. In this framework, independent lines of evidence in favor of or against pathogenicity for a variant are weighted according to strength level, where possible based on likelihood ratios (LRs), and following recommendations based on Bayesian modeling of the ACMG/AMP criteria<sup>15</sup>. Specific guidelines for *BRCA1* and *BRCA2*, based on this framework, have been published by the ClinGen Evidence-based Network for the Interpretation of Germline Mutant Alleles (ENIGMA) *BRCA1* and *BRCA2* Variant Curation Expert Panel (VCEP)<sup>2</sup>. Classification of a variant as (likely) benign or (likely) pathogenic (or remaining as VUS) is

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determined by joint assessment of the evidence types, either according to the combination of criteria met<sup>4,15</sup> or following a points-scoring system recently developed<sup>16</sup>. Based on previous research by the ENIGMA consortium<sup>17</sup>, one evidence type adapted for use under the ACMG/AMP framework is that from case-control data. In the ACMG/AMP framework this was included in the PS4 criterion, defined as “the prevalence of the variant in affected individuals is significantly increased over controls, with advice on application that the relative risk (RR) or odds ratio (OR) from case-control studies should be  $> 5.0$ , and the confidence interval (CI) around the estimate should not include 1.0”<sup>14</sup>. ClinGen specifications of the ACMG/AMP framework for *BRCA1* and *BRCA2* suggest that an  $OR \geq 4.0$  is assigned at full strength for a statistically significant association with CI not including 2.0[2]. As an initiative of the ENIGMA Analytical Working Group, we have recently proposed a LR-based framework for the analysis of case-control data for variant classification, where derived LRs are applicable under the ACMG/AMP framework for variant classification<sup>18</sup>. The case-control LR method computes a likelihood ratio, based on the likelihood of the observed data (i.e., the distribution of the variant of interest in cases and controls) under the hypothesis that the variant confers similar age-specific risks as the “average” *BRCA1* or *BRCA2* PV, relative to the likelihood under hypothesis that the variant is not associated with an increased risk of the disease. By taking into account the known age-specific penetrance of PVs, this approach provides significantly greater power for providing evidence for clinical classification, as compared to traditional “agnostic” case-control analyses. Hence, compared to ORs derived by logistic regression analysis, the LR-based framework has vastly improved performance to provide evidence towards pathogenicity, and more importantly, it can also be used to derive evidence against pathogenicity. For this reason, the *BRCA1/2* VCEP specifications also state that cCLR estimates should be used in preference to case-control OR data for application of case-control information (<https://cspec.genome.network/cspec/ui/svi/>).

In this work, using the case-control LR (cCLR) method and logistic regression analysis, we perform a large-scale assessment of 4291 unique variants in *BRCA1* (2248 within the coding sequence (CDS)  $\pm 5$  bp) and 6916 unique variants in *BRCA2* (4661 CDS  $\pm 5$  bp) based on sequencing data from 96,691 female breast cancer cases and 302,116 unaffected female controls from three cohorts: the BRIDGES project of the Breast Cancer Association Consortium (BCAC)<sup>4</sup>, the Cancer Risk Estimates Related to Susceptibility (CARRIERS) consortium<sup>3</sup> and the UK Biobank (UKB)<sup>19</sup>. In summary, the obtained likelihood ratios strongly align with ClinVar pathogenicity assertions, and we provide case-control evidence towards or against pathogenicity for 787 unclassified variants.

## Results

Case-control LRs were aligned to evidence strength levels for or against pathogenicity based on the thresholds recommended under the ACMG/AMP framework<sup>15</sup>. To evaluate the effect of variant counts on the reliability of the results, we performed calibration analyses considering variants that were present in at least one, two or three individuals in each dataset separately (Supplementary Data 1-2) and in the combined dataset (Supplementary Data 3). Using ClinVar reference sets of (likely) benign and (likely) pathogenic variants, we determined the proportion of benign and pathogenic variants that fell into the different ACMG/AMP evidence strength categories based on variant-specific cCLR estimates<sup>15</sup>; this allowed us to confirm if variants reaching cCLR evidence at a given evidence strength were enriched in pathogenic variants at the expected rate for that evidence strength level e.g., for variants reaching cCLR pathogenic supporting evidence, we would expect at least 2.08-fold enrichment in the pathogenic reference set compared to the benign reference set.

To ensure low false discovery and false omission rates ( $< 0.05$ ) and reach strong evidence for or against pathogenicity when applying

cCLR estimates (Supplementary Data 1-3), the analyses indicated that a more conservative approach was necessary. Thus, we restricted results to variants present in at least three individuals in the combined dataset (*BRCA1* and *BRCA2*) and evidence from at least two datasets (only for *BRCA2*). By applying these criteria, the enrichment ratios for known pathogenic: benign variants within each cCLR evidence category were broadly consistent with the expectation for that category, with the exception of *BRCA2* for cCLR benign moderate evidence (Table 1, Supplementary Data 3).

The combined dataset of 96,691 female breast cancer cases and 302,116 unaffected female controls from BRIDGES, CARRIERS, and the UKB contained case-control counts for 11,207 variants. Specifically, 4291 variants were found in *BRCA1* and 6,916 in *BRCA2* (Supplementary Data 4). The dataset encompasses 21.2% (6495/30,609) of the ClinVar *BRCA1* and *BRCA2* curated variants, 19.2% (2982/15,565) of the ClinVar variants curated as (likely) benign or pathogenic, and 23.4% (3513/15,044) of the ClinVar variants annotated as VUS. Results for all 11,207 variants are detailed in Supplementary Data 4.

When focusing solely on exonic and proximal intronic sequences (CDS  $\pm 5$  bp), covered adequately by all three cohorts, the dataset represented 18.7% (5718/30,609) of the ClinVar database and 21.9% (3297/15,044) of the ClinVar VUS. Of the total CDS  $\pm 5$  bp variants, there are 2248 variants in *BRCA1* and 4661 in *BRCA2* (Supplementary Data 5). Further filtering for maximum credible allele frequency (AF) in the non-founder populations (gnomAD v4.1.0), denoted as filtering allele frequency (FAF)  $\leq 0.001$  (*BRCA1* and *BRCA2* VCEP “BA1” benign stand-alone cutoff), present in at least three individuals in the combined dataset, and with evidence from at least two datasets for *BRCA2*, led to a final set of 1710 variants (681 in *BRCA1* and 1029 in *BRCA2*). These included: 760 variants listed in ClinVar as (likely) pathogenic or (likely) benign; and 950 variants considered to be of uncertain significance because they were not reported in ClinVar or they were listed in ClinVar as VUS or with conflicting classifications of pathogenicity (Fig. 1a). Sequence ontology variant consequence was annotated as follows: 204 presumed loss-of-function (LOF) variants of which 126 were frameshift insertions or deletions, 23 canonical splice site ( $\pm 2$  bp), 55 nonsense variants, as well as 31 in-frame insertions or deletions, 367 synonymous, 1082 missense and 26 intronic ( $\pm 3$  to  $\pm 5$  bp) variants (Fig. 1b). Case-control LRs provided evidence in favor of pathogenicity for 264 variants and against pathogenicity for 1185 variants; of these 1449 variants, 787 were considered to be of uncertain significance (Fig. 1c). Case-control results for the final set of 1710 *BRCA1* and *BRCA2* variants, are summarized and displayed as genomic maps in Fig. 2 and Fig. 3.

### *BRCA1* Likelihood Ratios

Informative case-control LRs were obtained for 604 out of 681 CDS  $\pm 5$  bp *BRCA1* variants (88.7%) (Fig. 4). Among these, evidence in favor of pathogenicity was observed for 114 variants; of those, 27 were categorized as very strong, 31 as strong, 31 as moderate, and 25 as supporting. Conversely, evidence against pathogenicity was provided for 490 variants; 157 categorized as very strong, 128 as strong, 119 as moderate, and 86 as supporting. In comparison, using logistic regression analysis only 14 variants reached strong evidence in favor of pathogenicity according to the *BRCA1/2* ACMG/AMP classification criterion description for OR application (PS4 criterion,  $OR \geq 4.0$ ,  $P$  value  $< 0.05$  and CI not including 2.0)<sup>2</sup>, and all of these were also assigned pathogenic evidence using the cCLR method.

Variants having case-control LR evidence in favor of pathogenicity consisted of 64.9% (74/114) LOF (39 frameshift insertions or deletions, 10 canonical splice site and 25 nonsense variants). Missense and synonymous variants accounted for 33.3% (38/114) and 1.8% (2/114), respectively. In contrast, variants with evidence against pathogenicity included 0.8% (4/490) LOF (1 frameshift, and 3 canonical splice site variants), 1.8% (9/490) in-frame, 68.6% (336/490) missense, 2.7% (13/

**Table 1 | Likelihood ratios (LRs) towards pathogenicity for (likely) benign and (likely) pathogenic variants assigned cCLR ACMG/AMP evidence strengths**

|   | (Likely) Benign |       | (Likely) Pathogenic |       | LR towards pathogenicity | LCI    | HCI      | ACMG/AMP Evidence strength |
|---|-----------------|-------|---------------------|-------|--------------------------|--------|----------|----------------------------|
|   | N               | Prop  | N                   | Prop  |                          |        |          |                            |
| <b>BRCA1 reference set</b>                  |                 |       |                     |       |                          |        |          |                            |
| Pathogenic Strong                           | 0               | 0.000 | 56                  | 0.615 | 302.152                  | 18.862 | 4840.167 | Pathogenic Strong          |
| Pathogenic Moderate                         | 2               | 0.008 | 18                  | 0.198 | 24.231                   | 5.736  | 102.368  | Pathogenic Strong          |
| Pathogenic Supporting                       | 2               | 0.008 | 7                   | 0.077 | 9.423                    | 1.994  | 44.528   | Pathogenic Moderate        |
| No evidence                                 | 26              | 0.106 | 8                   | 0.088 | 0.828                    | 0.389  | 1.763    | No evidence                |
| Benign Supporting                           | 23              | 0.094 | 0                   | 0.000 | 0.057                    | 0.003  | 0.927    | Benign Moderate            |
| Benign Moderate                             | 43              | 0.176 | 1                   | 0.011 | 0.063                    | 0.009  | 0.448    | Benign Moderate            |
| Benign Strong                               | 149             | 0.608 | 1                   | 0.011 | 0.018                    | 0.003  | 0.127    | Benign Strong              |
| <b>Total variants identified in dataset</b> | <b>245</b>      |       | <b>91</b>           |       |                          |        |          |                            |
| <b>BRCA2 reference set</b>                  |                 |       |                     |       |                          |        |          |                            |
| Pathogenic Strong                           | 1               | 0.004 | 51                  | 0.405 | 112.119                  | 15.669 | 802.270  | Pathogenic Strong          |
| Pathogenic Moderate                         | 8               | 0.029 | 19                  | 0.151 | 5.221                    | 2.349  | 11.606   | Pathogenic Moderate        |
| Pathogenic Supporting                       | 4               | 0.014 | 14                  | 0.111 | 7.694                    | 2.584  | 22.910   | Pathogenic Moderate        |
| No evidence                                 | 34              | 0.123 | 25                  | 0.198 | 1.616                    | 1.009  | 2.590    | No evidence                |
| Benign Supporting                           | 23              | 0.083 | 5                   | 0.040 | 0.478                    | 0.186  | 1.228    | <i>Benign Supporting</i>   |
| Benign Moderate                             | 45              | 0.162 | 9                   | 0.071 | 0.440                    | 0.222  | 0.871    | Benign Supporting          |
| Benign Strong                               | 162             | 0.585 | 3                   | 0.024 | 0.041                    | 0.013  | 0.125    | Benign Strong              |
| <b>Total variants identified in dataset</b> | <b>277</b>      |       | <b>126</b>          |       |                          |        |          |                            |

Reference sets included (likely) pathogenic and (likely) benign variants located within coding sequence (CDS) or splice acceptor/donor site dinucleotide positions ( $\pm 2$  bp), classified as such by the ClinGen *BRCA1/2* historical expert panel or the ClinGen *BRCA1/2* VCEP following ACMG/AMP guidelines, or by multiple submitters without conflicts in the ClinVar database, and also with gnomAD (v4.1.0) maximum credible allele frequency in the non-founder populations  $\leq 0.001$  consistent with the *BRCA1* and *BRCA2* VCEP's proposed BA1 cutoff. For likelihood ratio estimation, the Haldane-Anscombe correction was applied for any category where the cell count for pathogenic or benign variant reference set was zero. Code weights based on LR with confidence intervals spanning 1 are shown in italics.

490) intronic and 26.1% (128/490) synonymous variants (Figs. 4 and 5a). Among the 14 variants reaching the PS4 criterion, 78.6% ( $n=11$ ) were LOF (8 frameshift and 3 nonsense variants) alongside 21.4% ( $n=3$ ) missense variants. For *BRCA1* LOF variants, pathogenicity was provided for the majority of the frameshift insertions or deletions (97.5%, 39/40), canonical splice site (76.9%, 10/13) and nonsense (100%, 25/25) variants with informative case-control evidence.

The proportion of variants with informative case-control evidence towards pathogenicity was 10.2% (38/374) for missense, and 1.5% (2/130) for synonymous variants. All the *BRCA1* in-frame ( $n=9$ ) and intronic ( $n=13$ ) variants with informative case-control evidence were assigned benign evidence (Fig. 4). The majority of the identified CDS  $\pm 5$  bp *BRCA1* variants were within exon 10 ( $n=419$ ), followed by exons 15 ( $n=30$ ) and 14 ( $n=21$ ). One splice acceptor (c.594-2 A > C) and two splice donor (c.301+1 G > A and c.4096+1 G > A) variants within or proximal to exons 8, 5 and 10 were assigned benign evidence. Furthermore, missense variants with evidence in favor of pathogenicity were enriched in the exons encoding the RING domain (nucleotides 4-294; exons 2-5) (18.4%, 7/38) and the *BRCA1*-C-Terminal (BRCT) repeats (nucleotides 4987-5577, exons 16-23) (26.3%, 10/38) (Fig. 6). None of the in-frame variants assigned benign case-control evidence ( $n=9$ ) fell within the RING or BRCT domains.

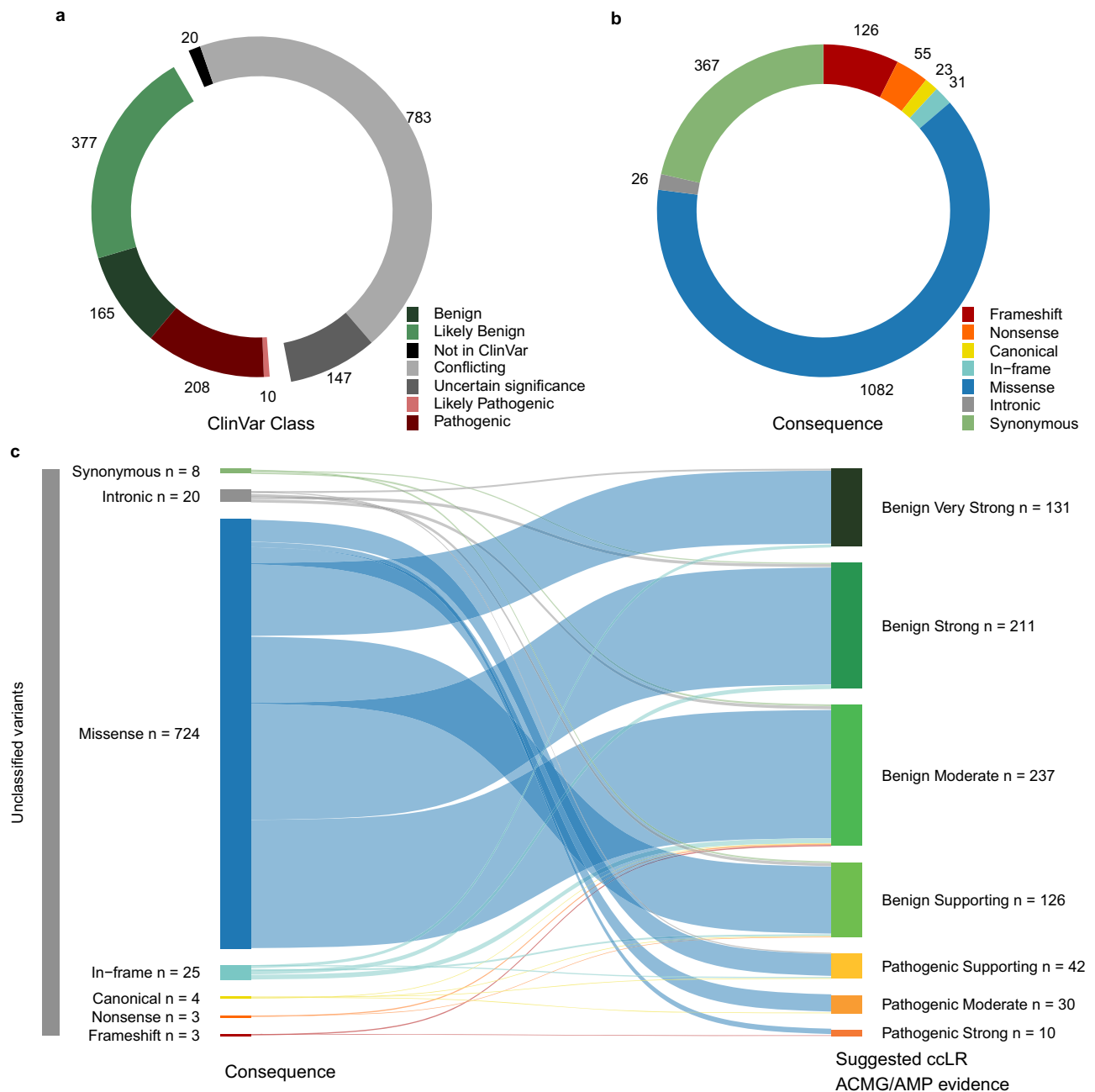
Overall, 25% (29/114) of the variants with evidence in favor of pathogenicity and 53.9% (264/490) of the variants with assigned evidence against pathogenicity, were considered to be of uncertain significance prior to evaluation (not reported in ClinVar or listed in ClinVar as VUS, with conflicting classifications of pathogenicity, or with classification "not provided"). For the remaining variants with an established ClinVar clinical classification as (likely) benign or (likely) pathogenic, we observed 99.1% (224/226) consistency for variants assigned benign evidence and 95.3% (81/85) for variants assigned pathogenic evidence (Figs. 4 and 5b). For the 14 variants that reached

the PS4 criterion using logistic regression analysis, one was of uncertain significance and the remainder had an established ClinVar clinical classification consistent with PS4 evidence in favor of pathogenicity.

### BRCA2 Likelihood Ratios

Informative case-control LR were obtained for 82.1% of the *BRCA2* variants (845/1,029) (Fig. 4). Evidence in favor of pathogenicity was provided for 150 variants, with 24 being very strong, 36 strong, 46 moderate, and 44 supporting. Evidence against pathogenicity was provided for 695 variants, of which 180 reached very strong, 196 strong, 223 moderate, and 96 supporting. Using logistic regression analysis only 17 variants reached strong evidence in favor of pathogenicity following the *BRCA1/2* ACMG/AMP PS4 classification criterion ( $OR \geq 4.0$ ,  $P$  value  $< 0.05$  and CI not including 2.0); all of these were assigned pathogenic evidence using the cCLR method.

Variants assigned cCLR evidence in favor of pathogenicity comprised of 50.7% (76/150) presumed LOF (53 frameshift insertions or deletions, 18 nonsense and 5 canonical splice site variants), 38% (57/150) missense, 8.7% (13/150) synonymous, 1.3% (2/150) intronic and 1.3% (2/150) in-frame variants. In contrast, variants assigned cCLR evidence against pathogenicity comprised of 3% (21/695) presumed LOF (12 frameshift, 8 nonsense and 1 canonical splice site variants), 2.6% (18/695) in-frame, 69.6% (484/695) missense, 1.2% (8/695) intronic and 23.6% (164/695) synonymous variants. The 17 variants reaching the PS4 OR criterion were comprised of 14 presumed LOF (82.4%) (11 frameshift insertions or deletions, 2 nonsense and 1 canonical splice site variants) and 3 (17.6%) missense variants (Figs. 4 and 5c). For *BRCA2* LOF variants, we observed that pathogenicity was proposed for the majority of the frameshift insertions or deletions (81.5%, 53/65), canonical splice site (83.3%, 5/6) and nonsense (69.2%, 18/26) variants assigned informative case-control evidence. The proposed pathogenicity rate for non-LOF variants (in-frame, missense, synonymous



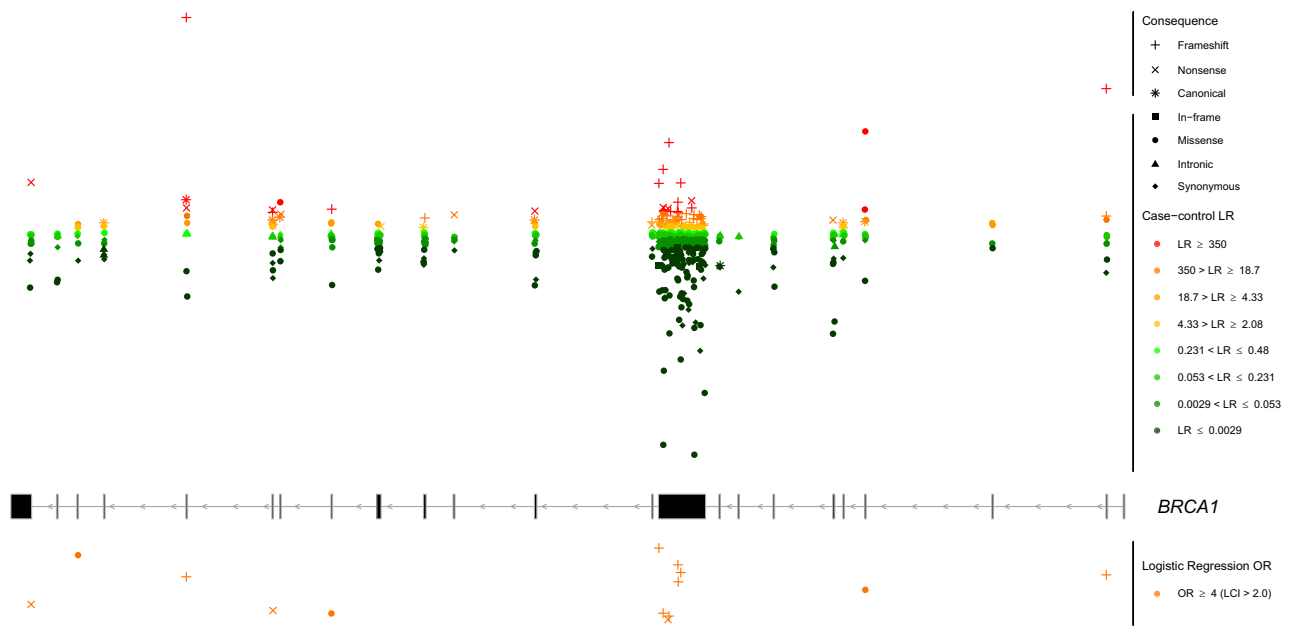
**Fig. 1 | Overview of *BRCA1* and *BRCA2* variants with case-control likelihood ratio evidence.** Donut plots showing the distribution of the **a** clinical classification (“ClinVar Class”) and **b** sequence ontology variant consequence (“Consequence”) for the 1717 CDS  $\pm$  5 bp variants with filtering allele frequency (FAF) > 0.001 (i.e., variants not meeting the *BRCA1* and *BRCA2* VCEP “BA1” benign stand-alone criterion), present in at least three individuals in the combined dataset, and with evidence from at least two datasets for *BRCA2*. The clinical classification status

(“ClinVar Class”) of variants was retrieved from the ClinVar database (last accessed on January 7, 2024). **c** Sankey plot depicting “suggested case-control likelihood ratio (cLR) ACMG/AMP evidence” (excluding variants with a cLR of “No evidence”) provided for unclassified variants (i.e., variants not reported in ClinVar or listed in ClinVar as VUS, variants of conflicting classifications of pathogenicity or variants with classification “not provided”), per sequence ontology variant consequence (“Consequence”).

and intronic variants) with informative case-control evidence, was 10% (2/20) for in-frame, 10.5% (57/541) for missense, 7.3% (13/177) for synonymous and 20% (2/10) for intronic variants (Fig. 4). The majority of the evaluated *BRCA2* variants were within exons 11 ( $n = 479$ ) and 10 ( $n = 111$ ) (Fig. 6). Two intronic variants (c.425+3 A > G, c.7435+5 T > C) were assigned evidence in favor of pathogenicity, falling within the splice donor sites of exons 4 and 14. Moreover, two in-frame variants within exons 11 and 20 were assigned evidence in favor of pathogenicity. One splice donor variant (c.8331+2 T > C) in the splice donor site of exon 18 was assigned evidence against pathogenicity. Finally,

missense variants assigned evidence in favor of pathogenicity were enriched in the exons encoding the DNA binding domain (DBD) (amino acids 2481-3186; exons 15-26) (38.6%, 22/57) (Fig. 6).

Overall, 35.3% (53/150) of the variants with evidence in favor of pathogenicity and 63.5% (441/695) with assigned evidence against pathogenicity, were considered to be of uncertain significance prior to evaluation (not reported in ClinVar or listed in ClinVar as VUS, with conflicting classifications of pathogenicity, or with classification “not provided”). For the remaining variants with an established ClinVar clinical classification as (likely) benign or (likely) pathogenic, we



**Fig. 2 | Genomic mapping of the case-control analysis for *BRCA1*.** Overlay of the case-control likelihood ratios (LRs) and the logistic regression odds ratio (OR) estimates is represented within each exon (middle panel). Exons are sequentially numbered from 1 to 23 and annotated from right to left to match the MANE Select transcripts. Although *BRCA1* was initially described with 24 exons (GenBank Accession ID U14680.1), exon 4 is missing following further assessment of the gene. We implement the most updated version of exon numbering (excluding legacy exon numbering). Case-control LRs (top panel) are represented on a continuous log<sub>2</sub>-transformed y axis with axis breaks. For the case-control LR analysis, the red color gradient represents LR reaching suggested ACMG/AMP evidence in favor of pathogenicity with strength levels ranging from very strong (dark red) to supporting (yellow). The green color gradient represents LR reaching ACMG/AMP evidence against pathogenicity with strength levels ranging from very strong (dark green) to supporting (light green). Variants with LR of “No evidence” are not

plotted. For the logistic regression analysis (bottom panel), orange color represents OR estimates reaching the strong PS4 criterion (OR ≥ 4.0, *P* value < 0.05, and confidence interval (CI) not including 2.0). Variants with OR estimates not reaching the PS4 criterion are not plotted. Associations were adjusted for age and study country (BCAC dataset), age and ethnic group (CARRIERS dataset), and age and genetic ancestry (UKB dataset). Only variants present in both cases and controls were analyzed. ORs with 95% CIs were estimated for each dataset and combined using a fixed-effects, inverse-variance meta-analysis in the ‘metafor’ R package to derive an overall test of association. A two-sided likelihood ratio test (LRT) was used to calculate *P*-values. No adjustments were made for multiple comparisons. For visualization purposes, the y-axis for logistic regression is represented in reverse order. LCI, lower confidence interval. Sequence ontology variant consequence (“Consequence”) is represented with different symbols.

observed 93.3% (237/254) consistency for variants assigned benign evidence and 86.6% (84/97) for variants assigned pathogenic evidence (Figs. 4 and 5b). For the 17 variants that met the PS4 OR criterion, one was of uncertain significance and the remainder had PS4 evidence consistent with an established ClinVar clinical classification.

### Concordance with predicted or experimental impact on function

Concordance with evidence in favor or against pathogenicity using the cCLR method was higher for experimental evidence from higher-throughput functional assays compared to in silico prediction methods (Supplementary Data 6). We observed 72.4% sensitivity and 98.7% specificity for functionally evaluated variants located in 13 exons of *BRCA1* encoding the RING and BRCT domains (exons 2-5 and 15-23, respectively)<sup>20</sup> and 100% sensitivity and 93.3% specificity for functionally evaluated missense *BRCA2* variants<sup>21</sup>. We also observed a strong concordance between the cCLR method and the recent functional evaluation of variants located in the region encoding the DBD of *BRCA2*<sup>22–24</sup>, with maximum sensitivity reaching 66.7% and specificity at 92.1% across three assays. For in silico prediction methods evaluated, including BayesDel<sup>25</sup>, REVEL<sup>26</sup>, VEST4<sup>27</sup>, MutPred2<sup>28</sup> and AlphaMissense<sup>29</sup>, we observed a moderate concordance with sensitivity ranging from 29.7% to 62.5%, and specificity ranging from 69.1% to 96.3% for *BRCA1*, and sensitivity ranging from 13.2% to 30.3% and specificity ranging from 88.2% to 95.1% for *BRCA2* (Fig. 7 and Supplementary Data 6).

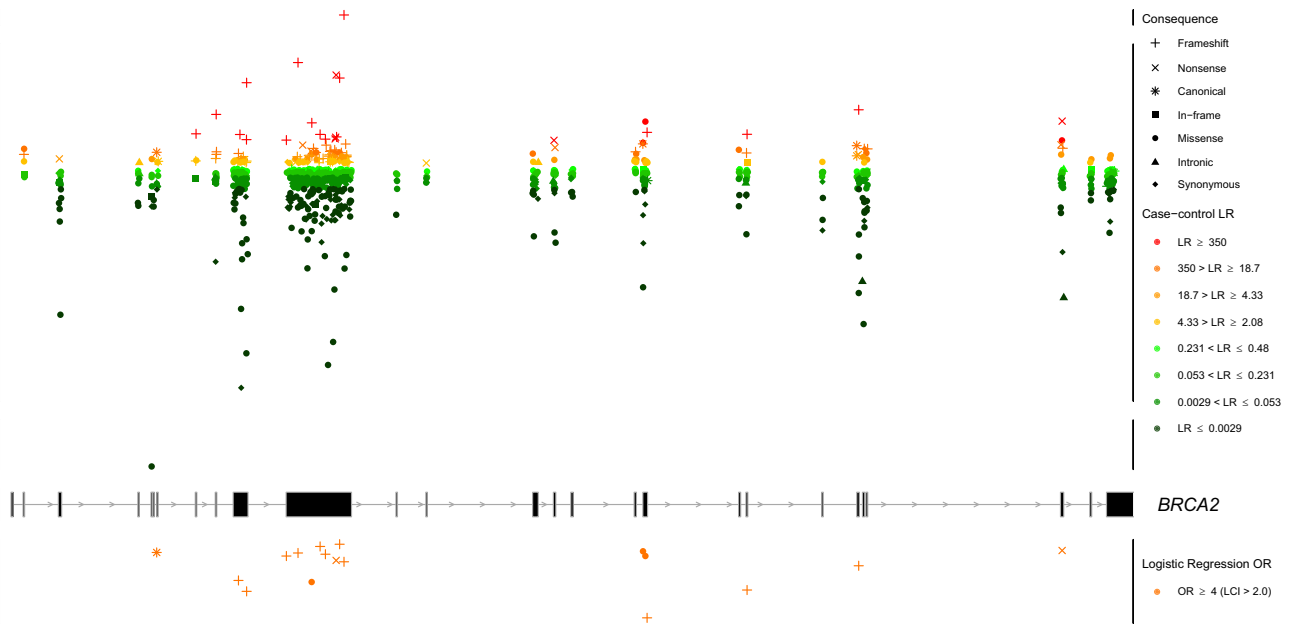
Taken together, high-throughput assay results align with cCLR evidence, offering a robust evaluation framework. Concordance between the cCLR method and predicted or experimental impact is

displayed as sequence-pathogenicity heatmaps (Fig. 7), which provide pathogenicity predictions for overlapping variants and functional evidence (binary categories) with cCLR.

### Discussion

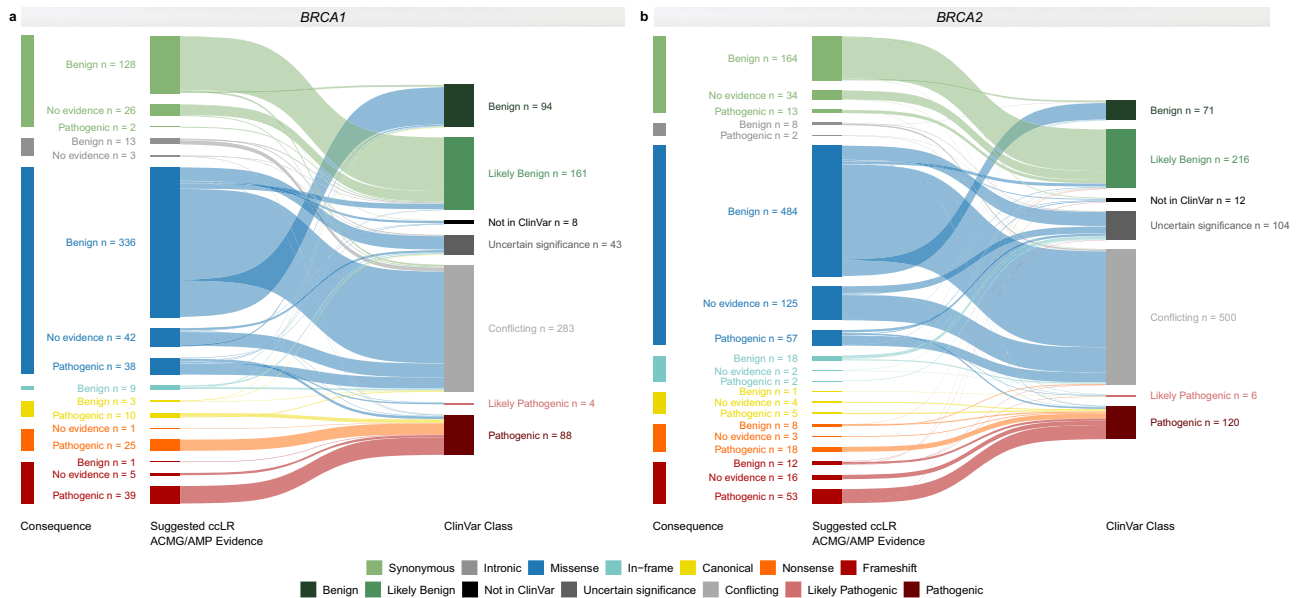
Significant efforts have been dedicated to the clinical classification of variants in *BRCA1* and *BRCA2*, owing to their elevated risk association with multiple cancer types. To date, only 111 *BRCA1* and *BRCA2* variants have been assigned case-control LR evidence<sup>18,30</sup>, and only 20 of these were previously used in clinical classification<sup>30</sup>. In this work, we present a large-scale multicenter case-control analysis of 11,207 *BRCA1* and *BRCA2* rare variants, of which 6909 are coding (within CDS ± 5 bp). Using sequencing data from 96,691 female breast cancer cases and 302,116 unaffected controls of the BRIDGES, CARRIERS and the UK Biobank datasets we utilized our recently developed cCLR method<sup>18</sup> and logistic regression analysis to leverage case-control data and provide evidence in favor or against variant pathogenicity. Our dataset comprehensively covers all exons, proximal intronic sequences and regulatory regions, accounting for 21.2% of the ClinVar curated variants and 23.4% of the ClinVar VUS, in these genes.

For coding variants, case-control evidence is highly consistent with ClinVar pathogenicity data, exhibiting 99.1% sensitivity and 95.3% specificity in distinguishing (likely) pathogenic and (likely) benign variants for *BRCA1* and 93.3% and 86.6% for *BRCA2*. The somewhat lower sensitivity and specificity for *BRCA2* is consistent with the lower penetrance of pathogenic variants in *BRCA2* compared to *BRCA1*<sup>4</sup>, so that the power of the analysis to classify variants, for a given number of observations, is lower for *BRCA2*. Notably, the majority of presumed



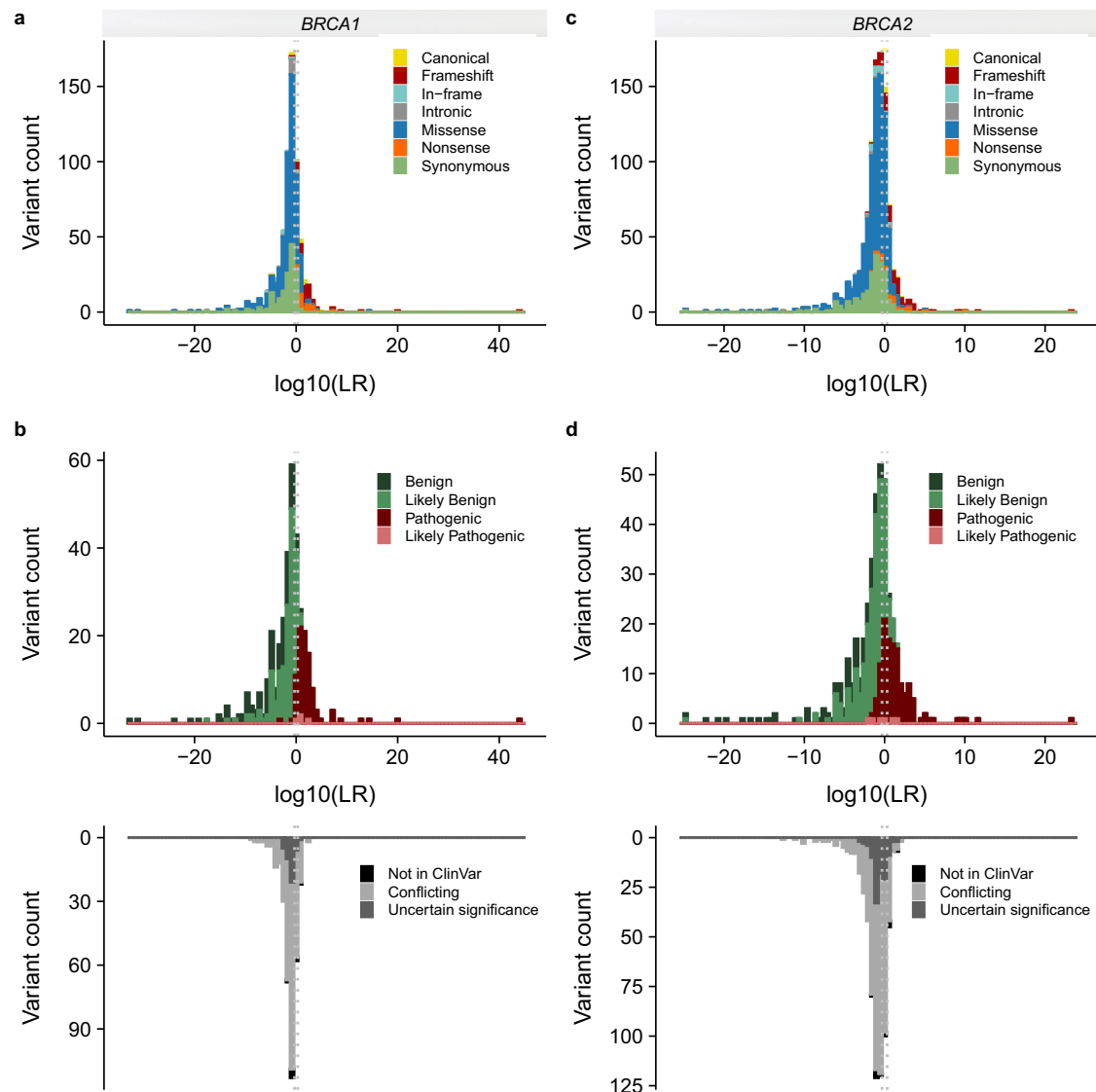
**Fig. 3 | Genomic mapping of the case-control analysis for *BRCA2*.** Overlay of the case-control likelihood ratios (LRs) and the logistic regression odds ratio (OR) estimates is represented within each exon (middle panel). Exons are sequentially numbered from 1 to 27 and annotated from left to right to match the MANE Select transcripts. Case-control LRs (top panel) are represented on a continuous log<sub>2</sub>-transformed y axis with axis breaks. For the case-control LR analysis, the red color gradient represents LR reaching suggested ACMG/AMP evidence in favor of pathogenicity with strength levels ranging from very strong (dark red) to supporting (yellow). The green color gradient represents LR reaching ACMG/AMP evidence against pathogenicity with strength levels ranging from very strong (dark green) to supporting (light green). Variants with an LR of “No evidence” are not plotted. For the logistic regression analysis (bottom panel), orange color represents

OR estimates reaching the strong PS4 criterion (OR ≥ 4.0, *P* value < 0.05, and confidence interval (CI) not including 2.0). Variants with OR estimates not reaching the PS4 criterion are not plotted. Associations were adjusted for age and study country (BCAC dataset), age and ethnic group (CARRIERS dataset), and age and genetic ancestry (UKB dataset). Only variants present in both cases and controls were analyzed. ORs with 95% CIs were estimated for each dataset and combined using a fixed-effects, inverse-variance meta-analysis in the ‘metafor’ R package to derive an overall test of association. A two-sided likelihood ratio test (LRT) was used to calculate *P*-values. No adjustments were made for multiple comparisons. For visualization purposes, the y-axis for logistic regression is represented in reverse order. LCI, lower confidence interval. Sequence ontology variant consequence (“Consequence”) is represented with different symbols.



**Fig. 4 | Case-control likelihood ratio evidence per sequence ontology variant consequence compared to ClinVar clinical classification.** Sankey plots for a *BRCA1* and b *BRCA2*. The variants assigned case-control likelihood ratio (LR) evidence in favor of or against pathogenicity (with suggested supporting, moderate, strong, or very strong evidence strength) are simplistically annotated as

“Pathogenic”, “Benign”, “Suggested case-control LR (ccLR) ACMG/AMP Evidence”, respectively. Variants with LRs between 0.48 and 2.08 are defined as “No evidence” in the “Suggested ccLR ACMG/AMP Evidence” panel. The clinical classification status (“ClinVar Class”) of variants was retrieved from the ClinVar database (last accessed on January 7, 2024).



**Fig. 5 | Distribution of the case-control likelihood ratios for *BRCA1* and *BRCA2*.** Histograms showing the distribution of case-control likelihood ratios (LRs) categorized by a sequence ontology variant consequence for *BRCA1*, **b** ClinVar classification for *BRCA1*, **c** sequence ontology variant consequence for *BRCA2*, **d** ClinVar classification for *BRCA2*. Histograms categorized by ClinVar classification are divided into two panels; the top panel represents variants curated in ClinVar as (likely)

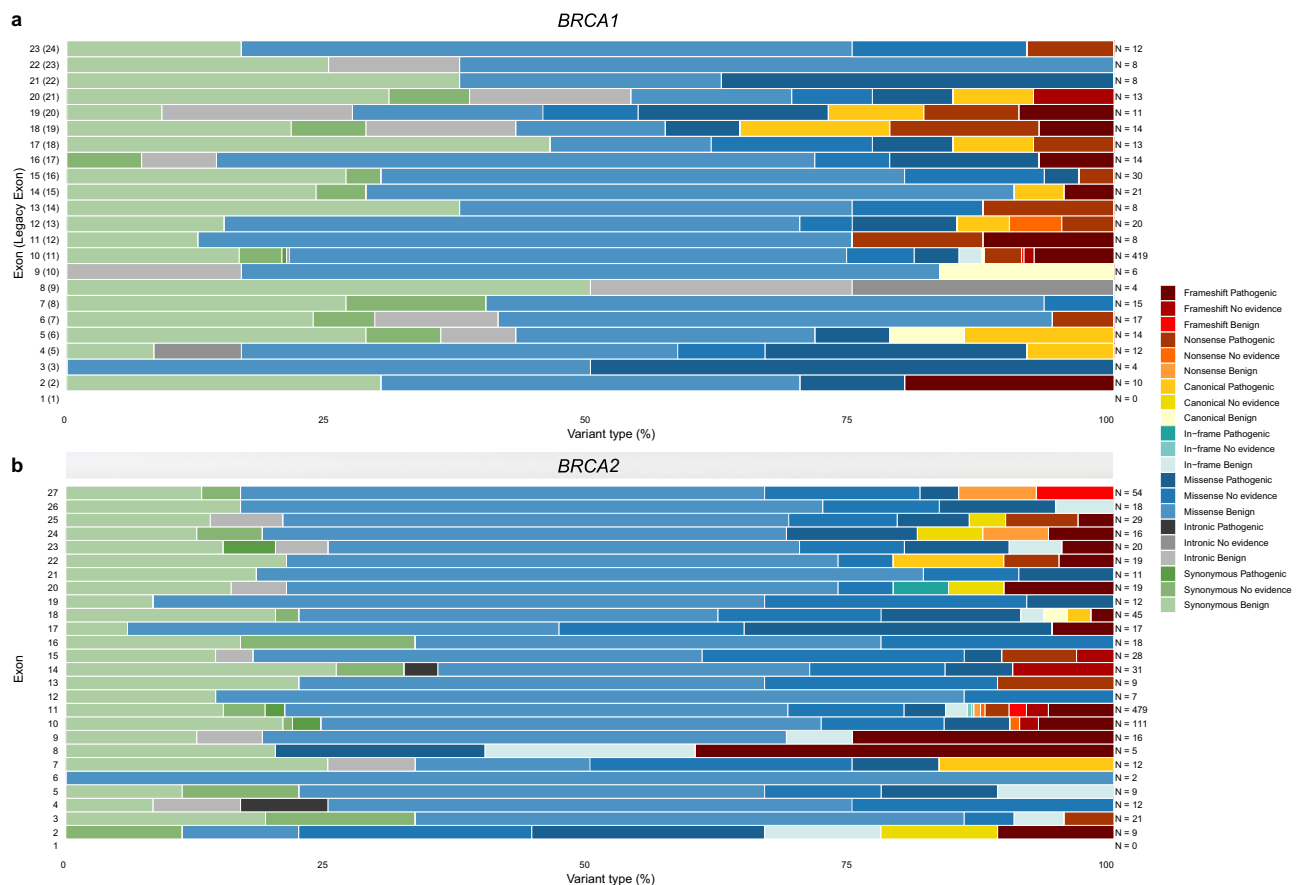
benign or (likely) pathogenic, while the bottom panel represents unclassified variants (variants not reported in ClinVar or listed in ClinVar as variants of uncertain significance or with conflicting classifications of pathogenicity). For visualization purposes the x axis represents  $\log_{10}(\text{LR})$  values. Dashed lines represent LRs between 0.48 and 2.08 considered as of “No evidence”.

LOF variants assigned evidence against pathogenicity are not assigned full or any pathogenic very strong (PVS1) code strength according to the recent ENIGMA classification criteria, since their predicted or known impact on splicing indicates that they are not associated with high risk of cancer<sup>2,31</sup>.

Using reference sets of (likely) pathogenic and (likely) benign variants drawn from ClinVar; using variants within the CDS  $\pm 2$  bp since this captures the regions more regularly tested in a clinical setting, we showed that the ccLR method maintains low false discovery and false omission rates ( $<0.05$ ) and can reach strong evidence both in favor and against pathogenicity, following the Bayesian adaptation of the ACMG/AMP framework<sup>15,16</sup>. Our results provide case-control LR evidence for 1449 variants with suggestive ACMG/AMP code strength levels; of these, 264 have evidence in favor of pathogenicity, and 1185 have evidence against pathogenicity.

This analysis provides evidence to inform clinical classification for 787 variants currently considered of uncertain clinical

significance. Specifically, strong or moderate benign evidence (equivalent to  $-4$  or  $-2$  points) observed for 579 unclassified variants should be sufficient to alter clinical classification with at least one other supporting benign evidence type, following standard ACMG/AMP classification protocols for combining evidence types<sup>14</sup> and requirement to reach  $-2$  points for a likely benign classification<sup>2,15,16</sup>. Likewise, strong pathogenic evidence ( $+4$  points) reached for 10 unclassified variants will provide valuable information towards pathogenicity in combination with other pathogenic evidence types, where  $+6$  points is sufficient to reach a likely pathogenic clinical classification<sup>2,14-16</sup>. This has important implications for clinical management. Individuals with a VUS are generally not eligible to access well-recognized clinical management options offered to individuals with a PV, such as risk-reducing prophylactic surgery to prevent cancer, MRI screening, or PARP-inhibitor treatment for cancers that arise<sup>10-13</sup>; thus, delaying effective interventions. Conversely, management of individuals with a VUS as if it were a PV that is actually



**Fig. 6 | Overview of the case-control likelihood ratio evidence assigned per exon and sequence ontology variant consequence for *BRCA1* and *BRCA2*.** Stacked bar plots of the suggested case-control likelihood ratio (LR) ACMG/AMP evidence per exon and sequence ontology variant consequence for **a** *BRCA1* and **b** *BRCA2*. Exons are sequentially numbered to match the MANE Select transcripts. Although *BRCA1* was initially described with 24 exons (GenBank Accession ID

U14680.1), exon 4 is missing following further assessment of the gene; legacy exon numbering for *BRCA1* is represented in brackets. Variants assigned case-control LR evidence in favor of or against pathogenicity (with suggested supporting, moderate, strong or very strong evidence strength) are simplistically annotated as “Pathogenic”, and “Benign” in the key, respectively. Variants with LRs between 0.48 and 2.08 are defined as “No evidence”.

benign can result in unnecessary anxiety and excessive medical intervention.

While our study provides valuable results that can be used in variant classification assuming the variant follows a high-risk penetrance as observed for “average” *BRCA1* and *BRCA2* PVs, however, it is important to acknowledge possible sources of bias such as potential sequencing artifacts, variant allele frequency differences between populations, case-control imbalances and possibility of ascertainment bias. We addressed this by applying stringent quality control measures and stratified analyses. It is also important to acknowledge the fact that when assessing numerous variants, particularly those with low counts, some may exhibit outcomes contrary to the expected, by chance. This issue is particularly relevant in the Supporting evidence category, where evidence is relatively weak. While the power to classify variants using case-control data alone can still be limiting, our likelihood-based approach lends itself naturally to combining with other evidence (e.g., co-segregation, tumor pathology, or in silico/functional scores). By integrating cCLR evidence with other lines of evidence, the likelihood of misclassification is minimized.

We further demonstrated that cCLR evidence aligns with functional predictors, offering a robust evaluation framework. These results can serve as a crucial resource for evaluating the consistency of other findings and refining existing methodologies.

Lastly, in order to accommodate the wider use of our case-control results, we provide the “cCLR browser” (<https://biostatuniting>.

[shinyapps.io/ccLRbrowser/](https://shinyapps.io/ccLRbrowser/)) that allows for easier exploration and visualization of our findings.

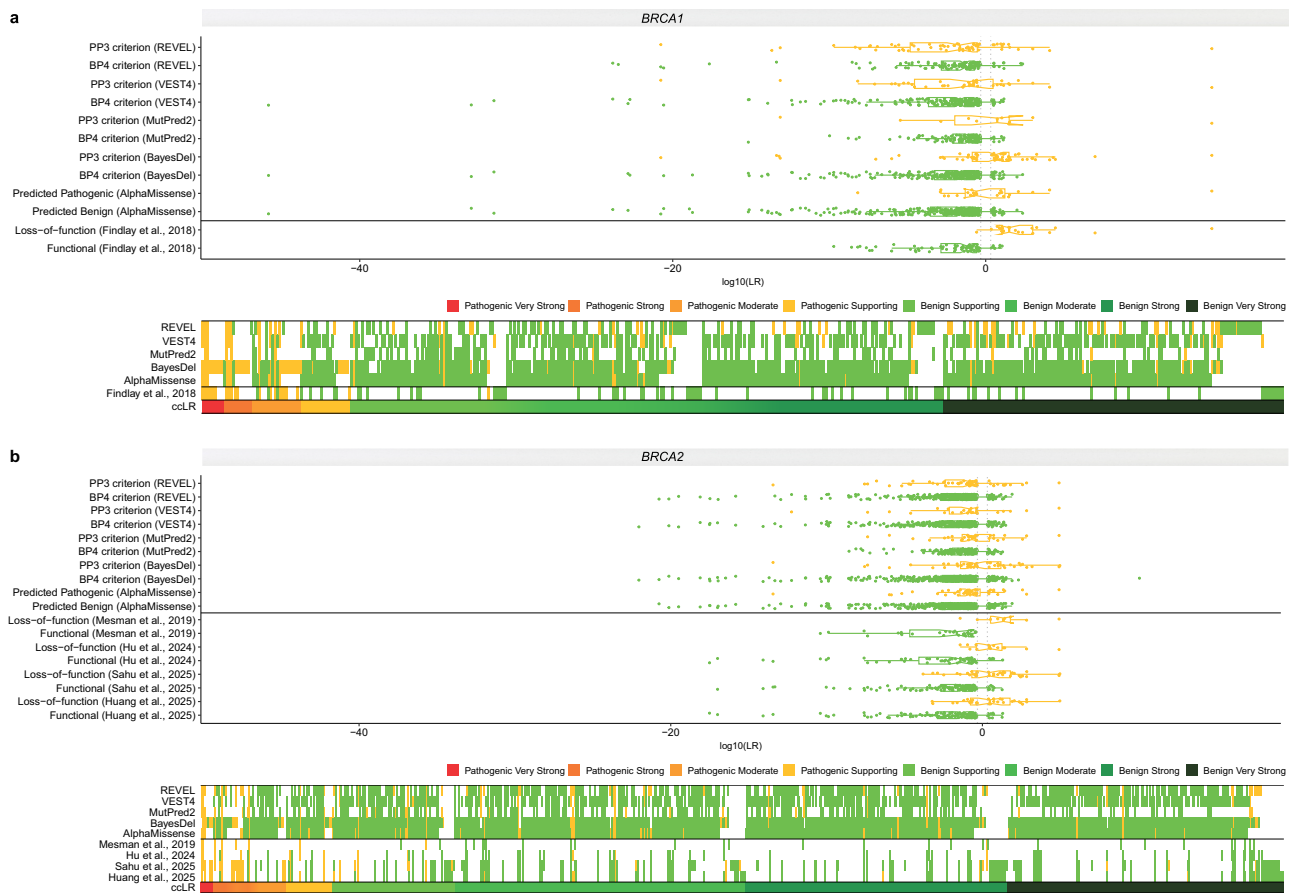
In summary, we present case-control evidence that strongly aligns with ClinVar pathogenicity assertions for non-VUS. We also provide case-control evidence towards or against pathogenicity for 787 unclassified variants. Our findings will be shared with the ClinGen ENIGMA *BRCA1* and *BRCA2* Variant Curation Expert Panel (VCEP), for expert panel curation activities that integrate multiple lines of evidence, including bioinformatic, clinical and experimental data, following with FDA-aligned variant classification processes. This can now be used in combination with other evidence for their final clinical classification, which is essential for accurate risk assessment and effective clinical decision-making, providing a larger number of patients and their relatives with clinically informative results.

## Methods

This research complies with all relevant ethical regulations and received approval by the Cyprus National Bioethics Committee (EEBK EP 2020.01.224). All the studies were approved by the relevant ethics review boards and used appropriate consent procedures.

### The BCAC/BRIDGES dataset

The BCAC dataset included 47,201 women with in-situ or invasive breast cancer and 47,316 unaffected controls, from 29 BCAC studies defined as population-based<sup>4</sup>. All studies were approved by the



**Fig. 7 | Concordance between the case-control likelihood ratio method and functional predictors.** Concordance is shown separately for **a** *BRCA1* and **b** *BRCA2*. The top panels for each gene represent case-control likelihood ratios (LRs) compared to variants predicted as benign (“BP4 criterion”, “predicted benign” or “functional”) or pathogenic (“PP3 criterion”, “predicted pathogenic” or “loss-of-function”) by in silico prediction methods (AlphaMissense, BayesDel, MutPred2, VEST4 and REVEL) or through high-throughput functional assays (Findlay et al., 2018, Huang et al., 2025, Sahu et al., 2025, Hu et al., 2024, Mesman et al., 2019). Yellow and green colors represent variants predicted as pathogenic or benign by functional predictors, respectively. For visualization purposes the x axis represents log<sub>10</sub>(LR) values. Box plots with individual data points display the median value, with whiskers extending to a maximum of 1.5 × interquartile range (IQR) beyond the box. The notch in the box approximates the 95% confidence interval (CI) for the

median. Bottom panels for each gene represent sequence-pathogenicity heatmaps demonstrating the concordance between the case-control LR (ccLR) method and functional predictors. For the case-control LR (ccLR) evidence, red color gradient represents LR reaching suggested ACMG/AMP evidence in favor of pathogenicity with strength levels ranging from very strong (dark red) to supporting (yellow). The green color gradient represents LR reaching suggested ACMG/AMP evidence against pathogenicity with strength levels ranging from very strong (dark green) to supporting (light green). Variants with cclR of “No evidence” are not plotted. For the functional predictors, yellow and green colors represent evidence in favor and against pathogenicity, respectively (expressed as “pathogenic supporting”). The total number of variants included in the concordance analyses is depicted in Supplementary Data 6.

respective ethics review boards, adhering to appropriate consent procedures. Phenotype data were based on the BCAC database v14. Individuals from a minority ancestry for each study (i.e., non-Asian from the Asian studies and non-European ancestry from the other studies), based on genetic data or self-report, were excluded<sup>4</sup>. Women of unknown/uncertain ancestry/ethnicity were excluded<sup>4</sup>. All samples underwent panel sequencing for 34 known or suspected breast cancer susceptibility genes as part of the BRIDGES project<sup>4</sup>. Details on library preparation, sequencing, and bioinformatics analysis, including variant calling and quality control, are described elsewhere<sup>4</sup>. The final dataset used consisted of 46,306 women with breast cancer and 43,481 unaffected controls, diagnosed or interviewed at age 21 to 80 years (the age range for which penetrance estimates for *BRCA1* and *BRCA2* are available) (Fig. 8 and Supplementary Data 7).

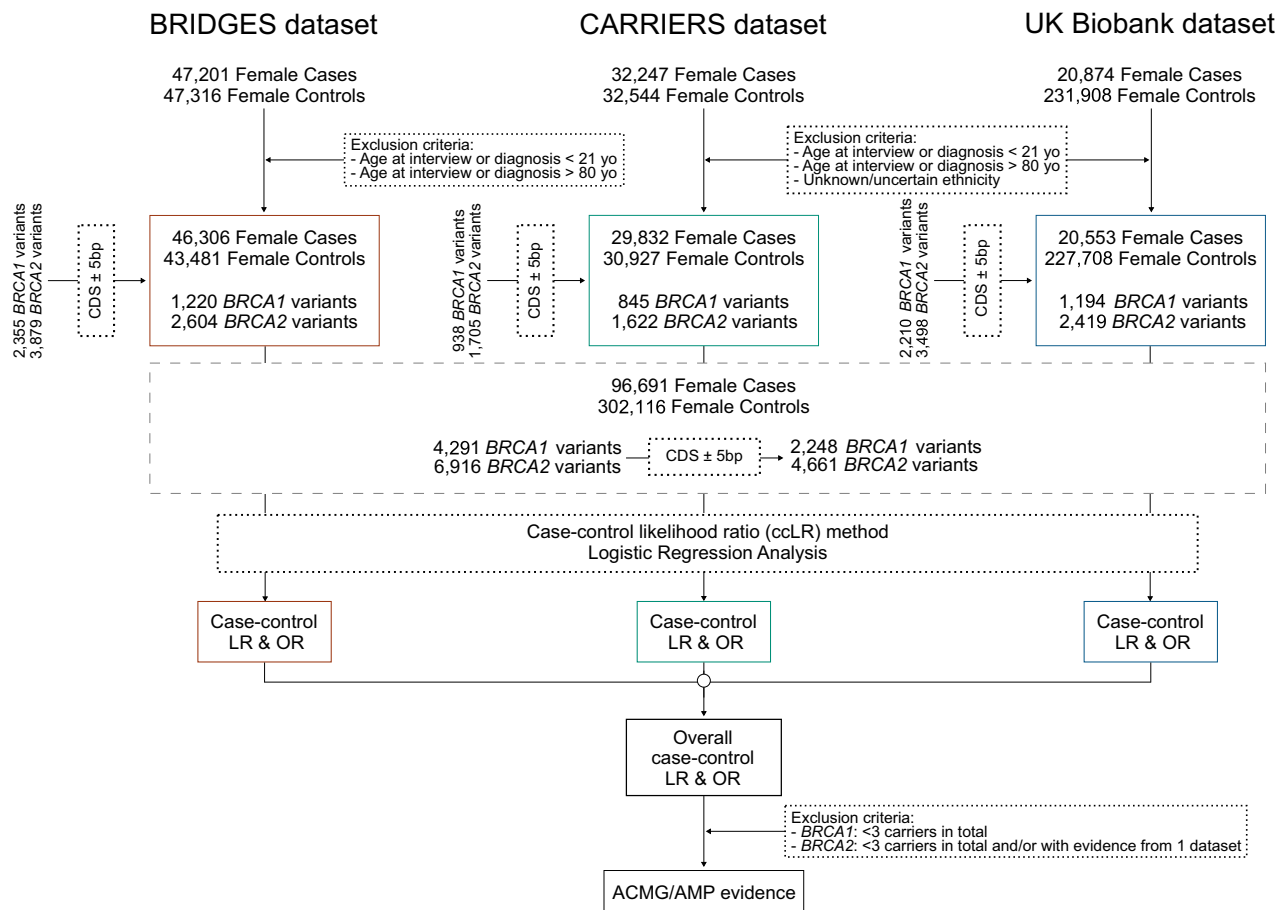
### The CARRIERS dataset

The CARRIERS consortium dataset included 32,247 women with in-situ or invasive breast cancer and 32,544 unaffected controls from 12 population-based studies<sup>3</sup>. The CARRIERS study was approved by the

institutional review board at the Mayo Clinic, and all participants provided informed consent for research. All samples were subjected to panel sequencing, targeting 37 cancer susceptibility genes. Details on library preparation, sequencing, and bioinformatics analysis, including variant calling and quality control, were previously documented<sup>3</sup>. Women of unknown/uncertain ethnicity were excluded. The final dataset consisted of 29,832 women with breast cancer and 30,927 unaffected controls diagnosed or interviewed, between the age range of 21 to 80 years (Fig. 8 and Supplementary Data 7).

### The UK Biobank dataset

UKB is a prospective cohort of more than 500,000 participants recruited in 22 assessment centers in the United Kingdom between 2006 and 2010<sup>32</sup>. Whole-exome sequencing (WES) data for 454,787 samples were released in October 2021 and were accessed via the UKB DNA Nexus platform<sup>19</sup>. Genetic ancestry was computed using a genetic principal components analysis from 2318 informative markers<sup>33</sup>. Women of unknown/uncertain ethnicity were excluded. Cases were defined as individuals with either invasive breast cancer (International



**Fig. 8 | Flowchart summarizing the study design.** Using sequencing data of 96,691 female breast cancer cases and 302,116 unaffected controls from the Breast Cancer Association Consortium (BCAC), the Cancer Risk Estimates Related to Susceptibility (CARRIERS) consortium and the UK Biobank (UKB) we calculated case-control likelihood ratios (LRs) and odds ratios (ORs) for 11,207 *BRCA1* and

*BRCA2* variants, of which 6909 are coding (coding sequence, CDS  $\pm$  5 bp). Derived LRs and ORs were further aligned to ACMG/AMP evidence strengths to provide evidence in favor of or against pathogenicity following sensitivity analyses-derived variant exclusion criteria.

Classification of Diseases (ICD)-10 code (C50) or carcinoma in situ (D05) based on linkage to the National Cancer Registration and Analysis Service (CRAS), or self-reported breast cancer incidence. Women diagnosed with ovarian cancer (International Classification of Diseases (ICD)-10 code C56) based on linkage to the National Cancer Registration and Analysis Service (NCRAS), or self-reported, were excluded from the controls. We accounted for both prevalent and incident cases, and only for cancers identified as an individual's first or second diagnosed cancer. Under these criteria, a total of 20,553 female breast cancer cases and 227,708 female controls were included, diagnosed or interviewed between the age range of 21 to 80 years (Fig. 8 and Supplementary Data 7). Access to the use of the UK Biobank data was granted under application number 102655.

### Data preparation

For all datasets, the Ensembl Variant Effect Predictor (VEP) v111 was used to annotate variants<sup>34</sup>. Annotations include the distance from the upstream or downstream gene, sequence ontology variant consequences, exon/intron number and Human Genome Variation Society (HGVS) nomenclature for the cDNA and protein level; the MANE Select transcript and protein were used (ENST00000357654.3 and ENSP00000350283.3 for *BRCA1*; ENST00000380152.3 and ENSP00000369497.3 for *BRCA2*)<sup>35</sup>. Exons and introns are sequentially numbered to match the MANE Select transcripts. Although *BRCA1* was initially described with 24 exons (GenBank Accession ID U14680.1), exon 4 is missing following further assessment of the gene. Herein, we

implement the most updated version of exon numbering (excluding legacy exon numbering). Allele frequency was retrieved from the gnomAD v4.1.0 release<sup>36</sup>. For filtering variants, we used the gnomAD maximum credible AF (the lower bound of the 95% CI) observed across the non-founder populations, including non-Finnish Europeans, African or African Americans, Admixed Americans, East Asians, South Asians, and Middle Easterners. Existing variant class was retrieved from the ClinVar database (<https://www.ncbi.nlm.nih.gov/clinvar/>, last accessed on January 7, 2024).

### Case-control analysis

The primary analysis involved the calculation of LRs for each individual variant using the cCLR method (<https://github.com/BiostatUnitCING/cCLR>)<sup>18,37</sup>. Under a survival analysis framework, the cCLR method compares the likelihood of the distribution of the variant of interest between cases and controls, under the hypothesis that the variant has similar age-specific relative risks as the "average" *BRCA1* or *BRCA2* PV, compared to the hypothesis that the variant is not associated with increased risk of the disease<sup>18</sup>. These risks are age-, sex-, and/or country-specific. Hence, the cCLR method requires specification of the age-specific risks in individuals with a PV and in the general population<sup>18</sup>. These were derived from the age-specific incidence rates for England and Wales (1998-2002) for ages 21-80 years, retrieved by the Cancer Incidence in Five Continents (CI5) Volume IX (<https://ci5.iarc.fr/>) and age-specific breast cancer odds ratio for individuals with *BRCA1* and *BRCA2* PVs estimated from the BRIDGES study<sup>4</sup>.

Case-control LRs were separately calculated for each dataset: BRIDGES, CARRIERS and UKB. To account for possible variant allele frequency differences by country or ethnicity, stratified LR calculations were performed within each dataset (BRIDGES stratified by country, and hence largely also by ethnicity; CARRIERS and UKB stratified by ethnicity) (Supplementary Data 8) and then multiplied across strata to provide a case-control LR for each independent dataset. Dataset-specific LRs were then used to obtain an overall LR for each variant. The LRs were aligned to evidence strength levels for or against pathogenicity based on the thresholds recommended under the ACMG/AMP framework<sup>15</sup>. Thus, in favor of pathogenicity was classified as very strong,  $LR \geq 350$ ; strong,  $350 > LR \geq 18.7$ ; moderate,  $18.7 > LR \geq 4.33$ ; or supporting,  $4.33 > LR \geq 2.08$ . Likelihood ratios against pathogenicity were classified as very strong,  $LR \leq 0.0029$ ; strong,  $0.0029 < LR \leq 0.053$ ; moderate,  $0.053 < LR \leq 0.231$ ; and supporting,  $0.231 < LR \leq 0.48$ . LRs between 0.48 and 2.08 were considered as “No evidence”.

Associations between variants and breast cancer risk were also assessed by logistic regression, adjusted for age and study country for the BCAC dataset, age and ethnic group for the CARRIERS dataset, and age and genetic ancestry for the UKB dataset. Logistic regression analysis was only performed for variants present in both cases and controls. Odds ratios and standard errors estimated from each dataset were combined in a fixed-effects, inverse-variance meta-analysis using the ‘metafor’ R package to derive an overall test of association. Using the *BRCA1/2* VCEP specification based on the PS4 ACMG/AMP classification criterion<sup>2</sup>, strong evidence in favor of pathogenicity was assigned to variants with  $OR \geq 4.0$ ,  $P$  value  $< 0.05$ , and CI not including 2.0.

## Reference sets

To assess the calibration of the cCLR method and perform sensitivity analyses, we selected reference sets of (likely) pathogenic and (likely) benign variants from the list of *BRCA1* and *BRCA2* variants identified in any of the three datasets. These reference sets included variants located within the CDS or at splice acceptor/donor site dinucleotide positions ( $\pm 2$  bp); since this captures the regions more regularly tested in a clinical setting, classified as such by the ClinGen *BRCA1/2* historical expert panel or the ClinGen *BRCA1/2* VCEP following ACMG/AMP guidelines<sup>2</sup>, or by multiple submitters without conflicts in the ClinVar database (last accessed on January 7, 2024). Variants were filtered to exclude any with gnomAD (v4.1.0) maximum credible AF in non-founder populations, denoted as FAF  $> 0.001$ , consistent with the *BRCA1* and *BRCA2* VCEP’s “BA1” benign stand-alone cutoff. Reference sets included 734 *BRCA1* variants (448, 369, and 379 with case-control LRs from BRIDGES, CARRIERS, and UKB, respectively) and 1240 *BRCA2* variants (762, 554, and 634 with case-control LRs from BRIDGES, CARRIERS, and UKB, respectively).

## Calibration of the cCLR method

To evaluate the calibration of the method, we used the case-control LRs for each of the variants in the reference sets and then determined the proportion of benign and pathogenic variants that would be classified in each ACMG/AMP evidence strength category<sup>15</sup>. We calculated enrichment for variants in each category  $j$  to be pathogenic as the ratio of proportions of variants in category  $j$  that are pathogenic or benign relative to the overall ratio of pathogenic and benign proportions<sup>38</sup>, that is:

$$\frac{n_{pj}n_B}{n_{Bj}n_P} \quad (1)$$

where  $n_{pj}$  and  $n_{Bj}$  are the number of pathogenic and benign variants in category  $j$ , and  $n_p$  and  $n_b$  are the total number of pathogenic and benign variants.

For calibration purposes, instances where variants were assigned either very strong or strong evidence according to the ACMG/AMP

criteria, either in favor or against pathogenicity, were collectively considered as instances of strong evidence in favor of or against pathogenicity, respectively. The Haldane-Anscombe correction was applied for any category where the cell count for the pathogenic or benign variant reference set was zero<sup>39</sup>.

## Concordance with predicted or experimental impact on function

To assess the concordance of the cCLR method with high-throughput functional assays and in silico prediction methods, we compared case-control LR values and suggestive ACMG/AMP code strength levels for variants located within the CDS or at splice acceptor/donor site dinucleotide positions ( $\pm 2$  bp); since this captures the regions more regularly tested in a clinical setting, present in at least three individuals in the combined dataset, and with evidence from at least two datasets for *BRCA2*.

Case-control evidence was compared with high-throughput functional assays performed for functionally critical *BRCA1* and *BRCA2* domains<sup>20–24</sup> and in silico prediction methods. In silico prediction methods included BayesDel<sup>25</sup> (without minor allele frequency) which is currently recommended (with optimal score cutpoints) by the ClinGen *BRCA1/2* VCEP for the curation of *BRCA1* and *BRCA2* variants (benign BP4 criterion “Multiple lines of computational evidence suggest no impact on gene or gene product”,  $\leq 0.15$  for *BRCA1* and  $\leq 0.18$  for *BRCA2*; pathogenic PP3 criterion “Multiple lines of computational evidence support a deleterious effect on the gene or gene product”,  $\geq 0.28$  for *BRCA1* and  $\geq 0.30$  for *BRCA2*)<sup>2</sup>, as well as other computational tools recommended (with appropriate score thresholds) by the ClinGen SVI Working Group<sup>40</sup>, including REVEL<sup>26</sup> (BP4,  $\leq 0.29$ ; PP3,  $\geq 0.644$ ), VEST4<sup>27</sup> (BP4,  $\leq 0.449$ ; PP3,  $\geq 0.764$ ) and MutPred2<sup>28</sup> (BP4,  $\leq 0.391$ ; PP3,  $\geq 0.737$ ). We also investigated the recent proteome-wide variant effect predictor AlphaMissense<sup>29</sup>.

Sensitivity and specificity metrics were used to assess the concordance of the functional predictors at predicting variants assigned evidence in favor or against pathogenicity using the cCLR method.

## Statistics & reproducibility

No statistical method was used to predetermine sample size. The experiments were not randomized and the Investigators were not blinded to allocation during experiments and outcome assessment.

## Web resources

Shiny app, <https://biostatunitcing.shinyapps.io/cCLRbrowser/>  
ClinVar database, <https://www.ncbi.nlm.nih.gov/clinvar/>  
gnomAD database, <https://gnomad.broadinstitute.org/>  
cCLR GitHub repository, <https://github.com/BiostatUnitCING/cCLR>  
Cancer Incidence in Five Continents (CI5) Volume IX, <https://ci5.iarc.fr/CI5I-X/Default.aspx>  
LiftOver tool, <https://genome.ucsc.edu/cgi-bin/hgLiftOver>

## Reporting summary

Further information on research design is available in the Nature Portfolio Reporting Summary linked to this article.

## Data availability

Individual level data for the BCAC data are not publicly available due to ethical review board constraints but are available on request through the BCAC Data Access Co-ordinating Committee (BCAC@medschl.cam.ac.uk). CARRIERS genotype data is available through dbGAP phs002820.v1. Phenotype data are not publicly available due to Institutional Review Board (IRB) constraints but are available on request through the CARRIERS Coordinating Committee (couch.fergus@mayo.edu). Requests for access to UK Biobank data should be made to the UK Biobank Access Management Team (access@ukbiobank.ac.uk). Summary statistics are published in Supplementary Data 4 and in the

shiny app (<https://biostatunitcing.shinyapps.io/cclRbrowser/>). According to the “UK Biobank best practice approaches on the publication of small numbers, Tables and/or results of any UK Biobank data there should have a minimum number of 5 reported participants within a cell”. Therefore, for variants present in less than 5 individuals, carrier count (Carriers, N) was replaced by “<5”.

### Code availability

Source code for the cclR method is publicly available in GitHub repository: <https://github.com/BiostatUnitCING/cclR><sup>37</sup>.

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## Competing interests

The following authors declare conflicts not directly relevant to this work as stated below. VK-MT and GDE have received consultancies from AstraZeneca and Everything Genetic Ltd. ACA and DFE are named creators of the BOADICEA model, which has been licensed by Cambridge Enterprise (University of Cambridge). SJH has received speaker fees from AstraZeneca and Pfizer Ltd. SY has received research funding from AstraZeneca and Repare Therapeutics and participates in the advisory board for AstraZeneca. JWW has received consulting fees from Natera, MyOme, Cancer IQ, equity from Natera, and speaker fees from AstraZeneca. All the other authors declare no competing interests.

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




























































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