

# Mutational Analysis of *BRCA1* and *BRCA2* Genes in Breast Cancer Patients from Eastern Sicily

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**Purpose:** Germline mutations of *BRCA1* and *BRCA2* are associated with a defined lifetime risk of breast (BC), ovarian (OC) and other cancers. Testing *BRCA* genes is pivotal to assess individual risk, but also to pursue preventive approaches in healthy carriers and tailored treatments in tumor patients. The prevalence of *BRCA1* and *BRCA2* alterations varies broadly across different geographic regions and, despite data about *BRCA* pathogenic variants among Sicilian families exist, studies specifically addressing eastern Sicily population are lacking. The aim of our study was to investigate the incidence and distribution of *BRCA* pathogenic germline alterations in a cohort of BC patients from eastern Sicily and to evaluate their associations with specific BC features.

**Patients and Methods:** Mutational status was assessed in a cohort of 389 BC patients, using next generation sequencing. The presence of alterations was correlated with tumor grading and proliferation index.

**Results:** Overall, 35 patients (9%) harbored a *BRCA* pathogenic variant, 17 (49%) in *BRCA1* and 18 (51%) in *BRCA2*. *BRCA1* alterations were prevalent among triple negative BC patients, whereas *BRCA2* mutations were more common in subjects with luminal B BC. Tumor grading and proliferation index were both significantly higher among subjects with *BRCA1* variants compared to non-carriers.

**Conclusion:** Our findings provide an overview about *BRCA* mutational status among BC patients from eastern Sicily and confirm the role of NGS analysis to identify hereditary BC patients. Overall, these data are consistent with previous evidences supporting *BRCA* screening to properly prevent and treat cancer among mutation carriers.

**Keywords:** hereditary breast cancer, BRCA1/2, NGS, tumor grading, Ki-67

## Introduction

Breast cancer (BC) represents the most frequent malignancy worldwide and the most lethal in women.<sup>1</sup> Over time, the biological features determining BC prognosis and clinical behavior have been extensively investigated and partly elucidated. Indeed, several surrogate markers are currently employed to classify BC into different molecular subtypes. These are estrogen (ER) and/or progesterone receptors (PgR), Human Epidermal growth factor Receptor 2 (*HER2*) amplification, proliferation index Ki-67 and tumor grading (G).<sup>2</sup> The combination of these variables identifies the following BC categories: 1) luminal tumors, which present ER and/or PgR expression and account for up to 75% of BC. These tumors are further classified into luminal A, when Ki-67 is below 20% and *HER2* is negative, and luminal B, when Ki-67 is equal or higher than 20% and in case of *HER2* amplification, regardless of proliferation index; 2) *HER2*+ tumors, which are negative for ER and PgR but display *HER2* amplification. This group encompasses 10% of all breast tumors; 3) triple negative BC (TNBC), which do not display ER and PgR expression and *HER2* amplification and represent about 15% of breast cancer.<sup>2-4</sup>

Across these BC subtypes, tumor grading and proliferation index represent transversal biomarkers, directly and independently correlating with tumor aggressiveness and prognosis.<sup>5,6</sup>

Besides the abovementioned biological features, the role of inherited genetic alterations leading to BC development has gained an increasing importance during the last few years.<sup>7</sup> Approximately one out of ten breast tumors is hereditary due the presence of germline alterations in specific genes.<sup>8</sup> Two large epidemiological studies including more than 180.000 women recently identified a set of 8 genes (ie *ATM*, *BARD1*, *BRCA1*, *BRCA2*, *CHK2*, *PALB2*, *RAD51C* and *RAD51D*) mainly responsible for hereditary BC. Among these genes, *BRCA1* and *BRCA2* (hereafter abbreviated as *BRCA1/2*) exhibited the strongest correlation with the development of breast tumors.<sup>9–12</sup> Indeed, germline *BRCA1/2* mutations considerably increase the lifetime risk of BC as well as of other malignancies, including ovarian, prostate, pancreatic, colorectal cancer and melanoma.<sup>13</sup> The cumulative incidence of BC by 80 years of age is 72% for women carrying *BRCA1* pathogenic variant (PV) and 69% for those with *BRCA2* PV.<sup>14</sup>

Noteworthy, a recent publication showed that BC risk depends on the type of PVs. In fact, photogenic missense variants, especially in *BRCA1* gene, are associated with a reduced risk of BC compared with pathogenic truncated mutations, particularly for older women.<sup>15</sup>

The presence of *BRCA1* or *BRCA2* PV correlates with distinct biological and clinico-pathological characteristics.<sup>16,17</sup> *BRCA1*-related BC tend to be clinically aggressive, poorly differentiated and highly proliferative. These tumors are frequently triple negative and present an earlier age of onset. Tumors occurring in *BRCA2* mutated patients usually display an intermediate to high differentiation grade and a variable proliferation index. These tumors are more often luminal B and usually occur in older individuals.<sup>16–18</sup> Of note, mutations in either *BRCA1* and *BRCA2* confer an increased sensitivity towards specific treatments, including platinum salts and targeted agents, such as poly(ADP-ribose) polymerase inhibitors (PARPi).<sup>19,20</sup>

In the last few years, the implementation of next generation sequencing (NGS) in the clinical practice gave access to an increasing number of BC patients to molecular tests for cancer predisposition syndromes, including *BRCA1/2*.<sup>21</sup> At the same time, definition of precise criteria based on family history, demographic characteristics and clinicopathological features allowed to better identify individuals who deserve *BRCA1/2* testing.<sup>22,23</sup> In this context, evidences are accumulating about *BRCA1/2* screening in specific populations, highlighting the differences between distinct geographical regions.<sup>24–27</sup> While reports exist about BC cohorts from western Sicily, fewer data are available about *BRCA1/2* screening among eastern Sicily population.<sup>28,29</sup>

We describe here the results of germline *BRCA1/2* screening in BC patients from eastern Sicily, further correlating the presence of a *BRCA1* or *BRCA2* mutations with the main clinico-pathological features of these tumors.

## Patients and Methods

### Study Population

A retrospective study was carried out at the “Center of Experimental Oncology and Hematology” of the Hospital Policlinico “G. Rodolico - San Marco” of Catania. From January 2017 to March 2021, a total of 455 patients with breast and ovarian cancer, melanoma, pancreatic tumour or prostate carcinoma were referred to our Molecular Diagnostics Laboratory for *BRCA1/2* genetic testing. The present study was conducted in accordance with the Declaration of Helsinki and all participants provided written informed consent before undergoing molecular analysis.

Histological and biological features of BC (ER, PgR, HER2 status, Ki-67 and grading) were evaluated on core biopsies or surgical samples, considering only the invasive tumor component. Based on these features, BC were classified as follow: luminal A (ER+ and/or PgR+, HER2-, Ki-67<20%), luminal B (ER+ and/or PgR+, HER2-, Ki-67≥20%), luminal B-HER2+ (ER and/or PgR+, HER2+), HER2+ (ER and PgR-, HER2+) or triple negative (ER and PgR-, HER2-).

An oncogenetic counselling was performed for each patient by a multidisciplinary team including an oncologist, a geneticist and a psychologist before assessing *BRCA1* and *BRCA2* mutational status, in order to identify individuals at high risk of harboring a PV in *BRCA1* and/or *BRCA2* genes. Patients selection was performed according to the Italian Association of Medical Oncology (AIOM) guidelines and Sicily local recommendations.<sup>30,31</sup> These criteria included: (i) family history of known pathogenic variant in a predisposing gene (eg *BRCA1*, *BRCA2*, *TP53*, *PTEN*); (ii) male with BC;

(iii) women with BC and OC; (iv) woman with BC <36 years, with TNBC <60 years or with bilateral BC <50 years; (v) personal history of BC <50 years and at least one first-degree relative with: (a) BC <50 years; (b) non-mucinous and non-borderline OC at any age; (c) bilateral BC; (d) male BC; (e) pancreatic carcinoma; (f) prostate carcinoma; (vi) personal history of BC >50 years and family history of BC, OC or pancreatic carcinoma in two or more relatives who have a first-degree relationship with each other (including one who has a first-degree relationship with her); (vii) personal history of OC and at least one first-degree relative with: (a) BC <50 years; (b) OC; (c) bilateral BC; (d) male BC; (vii) woman with high-grade serous OC.

## Molecular Testing

### DNA Isolation and Next Generation Sequencing Analysis for *BRCA1/2* Genes

Samples of 20 mL of peripheral blood were obtained from each patient and collected into EDTA tubes (BD Biosciences). Genomic DNA was isolated from 0.7 mL of whole blood samples using the QIASymphony DSP DNA Midi kit Isolation Kit (QIAGEN, Hilden, Italy) and quantified by Qubit<sup>®</sup>3.0 fluorometer (Thermo Fisher Scientific, Waltham, MA, USA), according to manufacturer's instructions. Target enrichment and library preparation was performed by Oncomine<sup>™</sup> BRCA Research Assay Chef ready to load into an Ion AmpliSeq<sup>™</sup> Chef Reagents DL8 cartridge for automated libraries preparation, according to the manufacturer's instructions. The kit consists of two multiplex PCR primer pool and allowed to investigate all *BRCA1* (NM\_007300.3) and *BRCA2* (NM\_000059.3) genes. Briefly, 15 µL of each diluted sample DNA (10 ng) were added to the barcode plate for library preparation and loaded with all reagents and consumables on the Ion Chef<sup>™</sup> Instrument. An automated library preparation and pooling of barcoded sample libraries were then performed on the Ion Chef<sup>™</sup> Instrument. The quantity of prepared libraries was then evaluated by Qubit<sup>®</sup>3.0 fluorometer (Thermo Fisher Scientific, Waltham, MA, USA), according to manufacturer's instructions. Finally, libraries were combined in an equimolar ratio in the Ion Chef<sup>™</sup> Library Sample Tube (barcoded tube) and loaded onto the Ion Chef<sup>™</sup> Instrument. The sequencing was performed with Ion 510 Chip (Thermo Fisher Scientific) using Ion Torrent S5 (Thermo Fisher Scientific) instrument (Thermo Fisher Scientific). Data analysis was done by Amplicon Suite (SmartSeq s.r.l.) and Ion Reporter Software.

### Data Analysis, Genetic Classification and Sanger Sequencing

All variants nomenclature followed current guidelines of the Human Genome Variation Society available online (HGVS, <http://www.hgvs.org/mutnomen>). The clinical significance of *BRCA1/2* variants was defined using the classification of the international consortium ENIGMA (evidence-based network for the interpretation of germline mutant alleles, <https://enigmaconsortium.org/>) and consulting different databases, such as ARUP, BRCAEXCHANGE, ClinVar, IARC\_LOVD and UMD. The classification consists of five different classes of risk: benign (class I), likely benign (class II), variant of uncertain significance (VUS, class III), likely pathogenic (class IV) and pathogenic (class V). The impact of mutations on protein structure and function was also analyzed by VarSome, an informatic tool that allows the access to 30 databases.<sup>32</sup>

To assign a potential clinical significance to each VUS, the following in silico protein prediction algorithms were used: MUTATION TASTER,<sup>33</sup> PROVEAN-SIFT (<http://provean.jcvi.org/index.php>), POLYPHEN-2 (<http://genetics.bwh.harvard.edu/pph2/>) and Align-GVGD ([http://agvgd.hci.utah.edu/agvgd\\_input.php](http://agvgd.hci.utah.edu/agvgd_input.php)). Variants classified as class 1 and 2 were considered wild type.

The presence of each pathogenic variants was confirmed by Sanger sequencing. Briefly, a couple of specific primer was designed for each of the detected variants by using the *BRCA1* and *BRCA2* gene reference sequences (NG\_005905.2, NM\_007294.3 and NG\_012772.3, NM\_000059.3, respectively). Therefore, a target- PCR was performed, followed by Sanger sequencing.

### CNV Analysis by Multiplex Ligation-Dependent Probe Amplification Analysis (MLPA)

Patients with a negative *BRCA1/2* genetic test were tested to evaluate the presence of Large Genomic Rearrangements (LGR) by Multiplex ligation-dependent probe amplification (MLPA), according to the manufacturer's instructions. Briefly, the DNA samples are denatured and linked up to 60 probes specific for *BRCA1* and *BRCA2* genes that each detect a specific DNA sequence of approximately 60 nucleotides in length. Probe amplification products, consisting of a

set of unique PCR amplicons, are then analyzed by capillary electrophoresis and analyzed by the Coffalyser.Net software, in combination with the appropriate lot-specific Coffalyser sheet ([www.mrcholland.com](http://www.mrcholland.com)).

## Statistical Analysis

Selected clinic-pathological variables (histological grade and Ki-67% proliferation index) were correlated with the presence of *BRCA1/2* PVs using the Fisher's Exact test assuming p-values <0.05 as significant, calculated using Prism software v. 8.4.

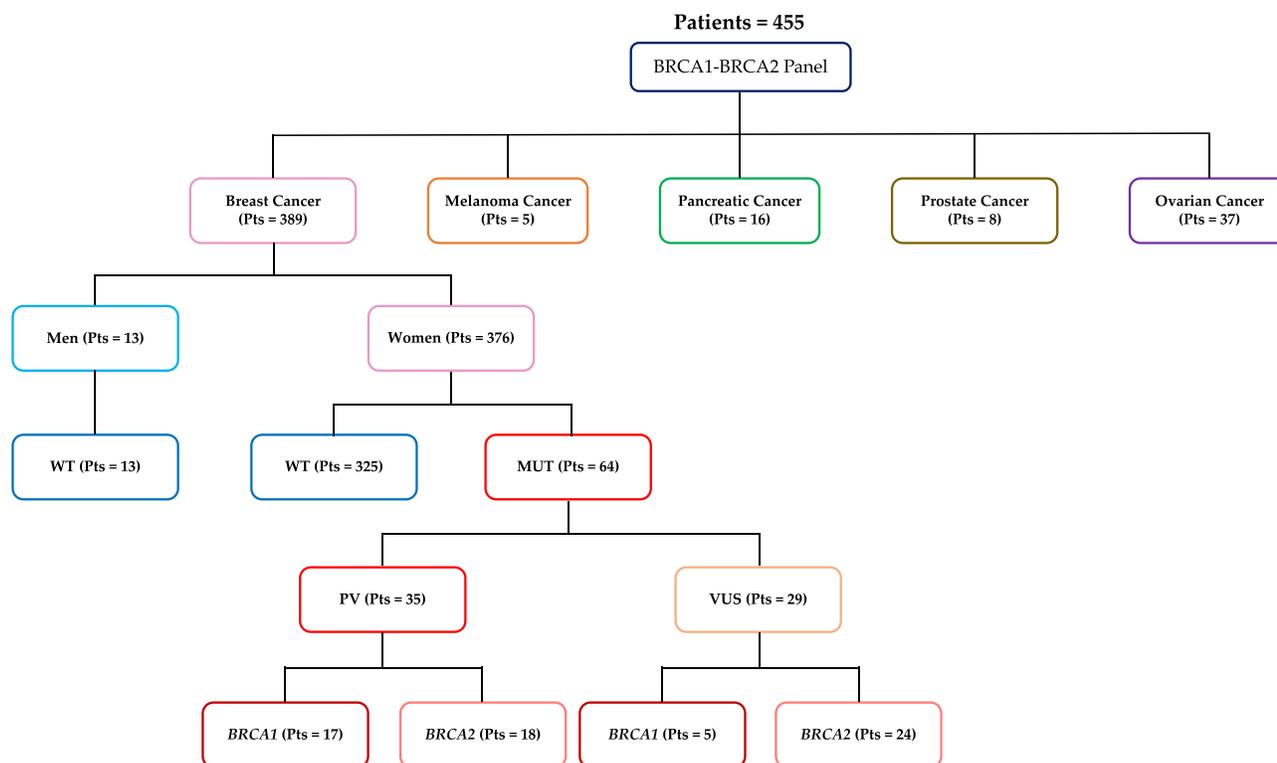
## Results

### Population Characteristics

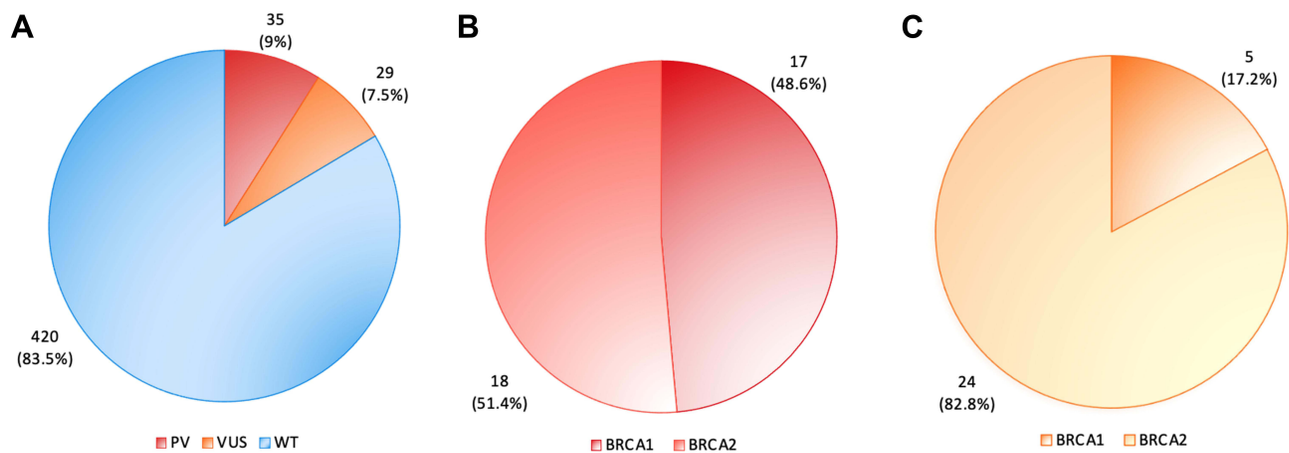
Between January 2017 and March 2021, 455 patients were screened for germline *BRCA1/2* mutations. Mutational tests were performed at the Center of Experimental Oncology and Hematology of the Hospital Policlinico "G. Rodolico - San Marco" of Catania, according to Sicily guidelines (<http://www.gurs.regione.sicilia.it/Indicep1.htm>, N. 02-Venerdì 10 Gennaio 2020) Overall, 389 patients had breast cancer, 37 ovarian cancer, 16 pancreatic cancer, 8 prostate cancer and 5 melanoma. Patients distribution according to cancer type and analysis results is shown in Figure 1.

We selectively focused our study on the breast cancer cohort. These patients displayed a median age of 49 years (range 23–89) and were mainly females (n=376, ie 97%).

Among these subjects, 64 (17%) carried a *BRCA1/2* mutation and all of them were women. Thirty-five (9%) had a PV, while 29 (7.5%) harbored a VUS. Seventeen of the 35 pathogenic variants (48.6%) occurred in *BRCA1* and 18 (51.4%) in *BRCA2*, whereas 5 VUS were in *BRCA1* (17.2%) and 24 (82.8%) in *BRCA2* (Figures 1 and 2). No LGR emerged from MLPA analysis.



**Figure 1** Flow chart showing an overview of the study. Patients with breast, melanoma, pancreatic, prostate or ovarian tumors tested for *BRCA1* and *BRCA2* gene mutation. **Abbreviations:** PVs, pathogenic variants; VUS, variants of uncertain significance; WT, wild-type *BRCA1/2* sequence.

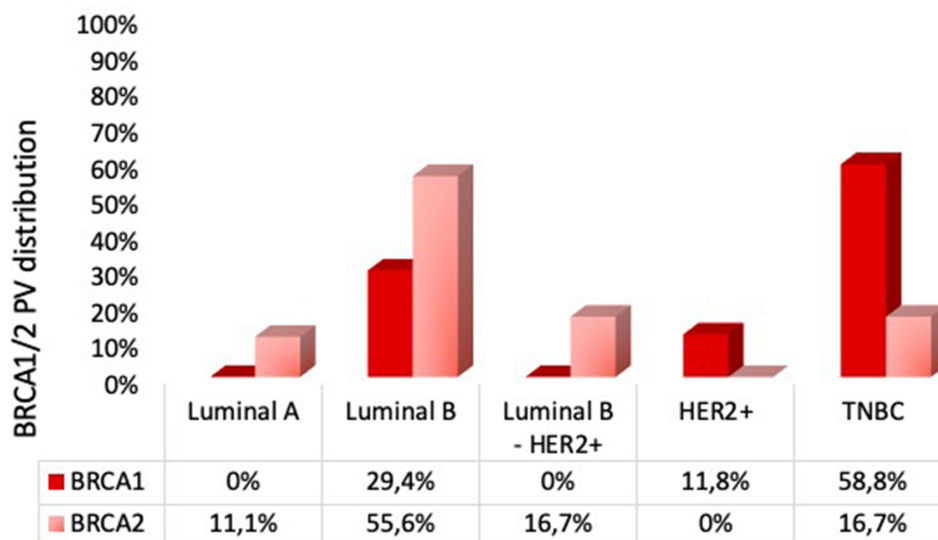


**Figure 2** *BRCA1* and *BRCA2* mutation analysis of 389 breast cancer patients. **(A)** Distribution of pathogenic variants (PVs) (red), variants of uncertain significance (VUS) (Orange) and WT (blue) in 389 patients with breast cancer; **(B)** 35 (9%) out of 389 breast cancer individuals showed *BRCA1/2* pathogenic variants (PVs). Among them, 17 (48.6%) were *BRCA1* PV carriers (dark red), while 18 (51.4%) were *BRCA2* carriers (light red); **(C)** 29 (7.5%) out 389 subjects harboured VUS, 5 (17.2%) in *BRCA1* gene (dark orange) and 24 (82.8%) in *BRCA2* gene (light orange).

**Abbreviations:** PVs, pathogenic variants; VUS, variants of uncertain significance; WT, wild-type *BRCA1/2* sequence.

### Distribution of Molecular Subtypes

We next investigated the prevalence of BC molecular subtypes in patients carrying *BRCA1/2* PVs. The distribution included 2 (5.7%) luminal A, 15 (42.9%) luminal B, 3 (8.6%) luminal B-HER2+, 2 (5.7%) HER2+ and 13 (37.1%) TNBC patients. Among *BRCA1* positive patients, 5 (29.4%) presented a luminal B BC, 2 (11.8%) a HER2+ disease and 10 (58.8%) a TNBC. No *BRCA1* mutated tumors were luminal A or luminal B-HER2+ (Figure 3). In the *BRCA2* positive subgroup, 10 (55.6%) tumors were luminal B, 3 (16.7%) luminal B-HER2+, 3 (16.7%) TNBC and 2 (11.1%) luminal A (Figure 3). No HER2+ tumors were present in this group. Hence, *BRCA1* mutations were prevalent in TNBC patients, whereas *BRCA2* alterations were predominant in luminal B individuals.



**Figure 3** Prevalence of breast cancer subtypes in patients carrying *BRCA1* and *BRCA2* pathogenic mutations. The histograms show the distribution of *BRCA1*- (dark red) and *BRCA2*- (light red) PVs in molecular subtypes of breast cancer patients. The numbers reported inside each box indicate the percentage of patients with *BRCA1* and *BRCA2* PVs for each breast cancer subtype.

**Abbreviations:** PVs, pathogenic variants; HER2 +, human epidermal growth factor receptor 2 positive; TNBC, triple negative breast cancer.

## Type and Localization of *BRCA1* and *BRCA2* Pathogenic Variants

Subsequently, we evaluated the type and gene location of *BRCA1* and *BRCA2* PVs. Among *BRCA1* PVs we observed 7 single nucleotide variants (SNVs), 6 deletions, 3 duplications and 1 insertion. Only one mutation (c.5522delG) represents a novel finding. The most frequent *BRCA1* PV, detected in two subjects, was c.5035\_5039delCTAAT. This alteration involves the deletion of five nucleotides (CTAAT) in *BRCA1* exon 15, resulting in the substitution of the amino acid leucine with tyrosine at codon 1679 and causing a premature truncation of the protein due to a translational frameshift with a predicted alternative stop codon. All the other alterations were detected in one case only. Of note, one of the reported PV is located in a splice site consensus region (c.4357+1G>T) (Table 1).

Concerning *BRCA2* PVs, we observed 6 deletions, 6 SNVs and 2 duplications. None of the identified alterations was novel. Three mutations were recurrent in our population, namely c.428dup and c.8487+1G>A observed in 3 subjects followed by c.5851\_5854delAGTT which has been retrieved in two cases. The c.428dup alteration involves the duplication of C in *BRCA2* exon 5 predicting to encode a truncated non-functional protein. The c.8487+1G>A mutation occurs in the intronic region ( $\pm$  1,2) of *BRCA2* intron 19 and affects the splice consensus sequence causing altered splicing leading to an abnormal or absent protein. The c.5851\_5854delAGTT pathogenic mutation, results from a deletion of 4 nucleotides at nucleotide positions 5851 to 5854 in the coding exon 10 of the *BRCA2* gene and results

**Table 1** *BRCA1/2* Pathogenic Variants Identified in Breast Cancer Patients

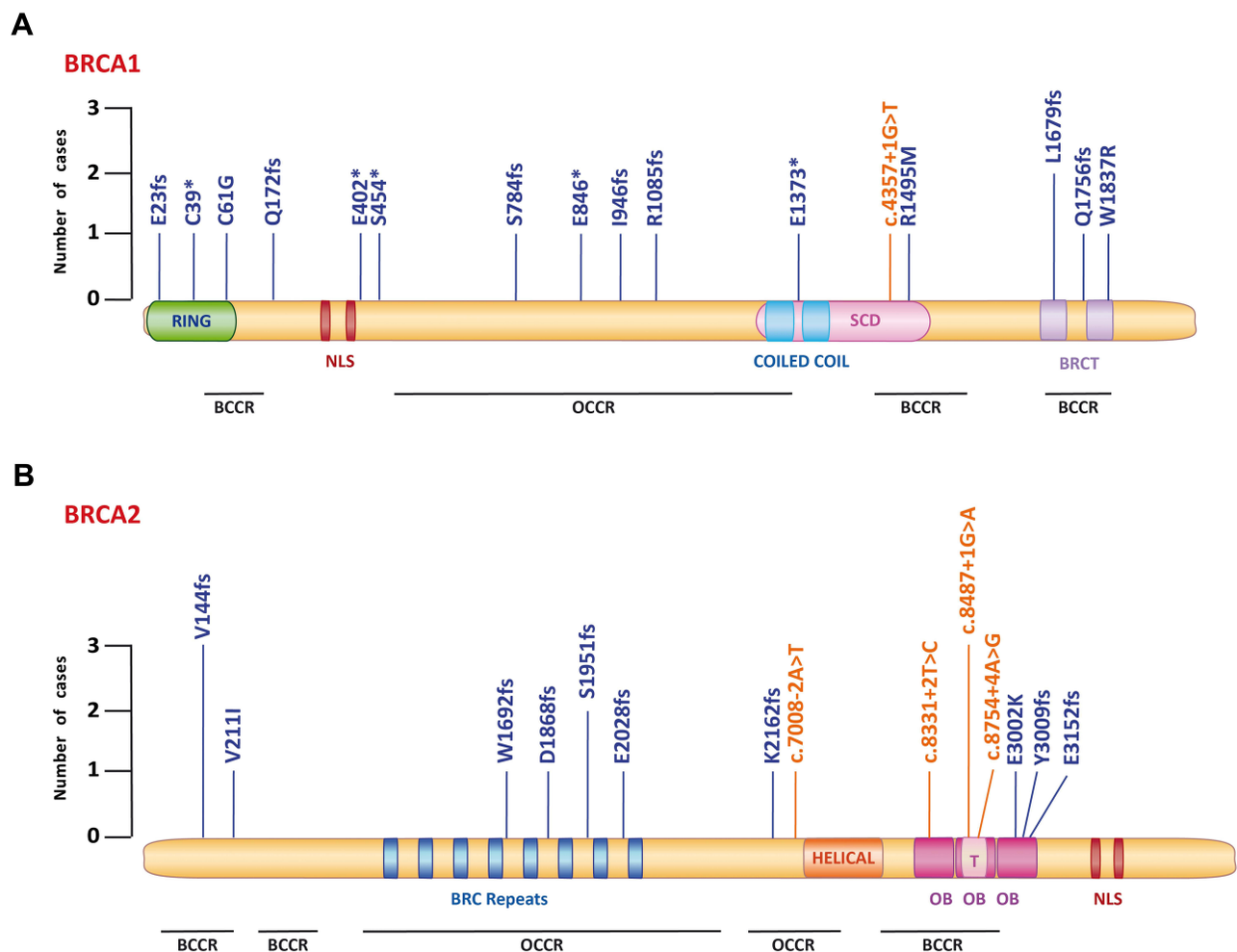
Gene	Type of PV	HGVS Nomenclature	Protein Change	No. Cases
<i>BRCA1</i>	Duplication	c.66_67insA	p.(Glu23Argfs)	1
<i>BRCA1</i>	Deletion	c.117_118delTG	p.(Cys39Ter)	1
<i>BRCA1</i>	SNV	c.181T>G	p.(Cys61Gly)	1
<i>BRCA1</i>	Deletion	c.514del	p.(Gln172Asnfs)	1
<i>BRCA1</i>	SNV	c.1204G>T	p.(Glu402Ter)	1
<i>BRCA1</i>	Deletion	c.1360_1361delAG	p.(Ser454Ter)	1
<i>BRCA1</i>	Deletion	c.2350_2351delTC	p.(Ser784Valfs)	1
<i>BRCA1</i>	SNV	c.2536G>T	p.(Glu846Ter)	1
<i>BRCA1</i>	Insertion	c.2835_2836insCC	p.(Ile946Profs)	1
<i>BRCA1</i>	Duplication	c.3253dup	p.(Arg1085Lysfs)	1
<i>BRCA1</i>	SNV	c.4117G>T	p.(Glu1373Ter)	1
<i>BRCA1</i>	SNV	c.4357+1G>T	/	1
<i>BRCA1</i>	SNV	c.4484G>T	p.(Arg1495Met)	1
<i>BRCA1</i>	Deletion	c.5035_5039delCTAAT	p.(Leu1679Tyrfs)	2
<i>BRCA1</i>	Duplication	c.5266dup	p.(Gln1756Profs)	1
<i>BRCA1</i>	SNV	c.5509T>C	p.(Trp1837Arg)	1
<i>BRCA1</i>	Deletion	c.5522delG	p.(Ser1841fs*)	1
<i>BRCA2</i>	Duplication	c.428dup	p.(Val144Cysfs)	3
<i>BRCA2</i>	SNV	c.631G>A	p.(Val211Ile)	1
<i>BRCA2</i>	SNV	c.7008-2A>T	/	1
<i>BRCA2</i>	Duplication	c.5073_5074insA	p.(Trp1692Metfs)	1
<i>BRCA2</i>	Deletion	c.5603_5606delACAG	p.(Asp1868Valfs)	1
<i>BRCA2</i>	Deletion	c.5851_5854delAGTT	p.(Ser1951TrpfsTer)	2
<i>BRCA2</i>	Deletion	c.6082_6086delGAAGA	p.(Glu2028Lysfs)	1
<i>BRCA2</i>	Deletion	c.6486_6489delACAA	p.(Lys2162Asnfs)	1
<i>BRCA2</i>	SNV	c.8331+2T>C	/	1
<i>BRCA2</i>	SNV	c.8487+1G>A	/	3
<i>BRCA2</i>	SNV	c.8754+4A>G	/	1
<i>BRCA2</i>	SNV	c.9004G>A	p.(Glu3002Lys)	1
<i>BRCA2</i>	Deletion	c.9026_9030delATCAT	p.(Tyr3009Serfs)	1
<i>BRCA2</i>	Deletion	c.9455_9456delAG	p.(Glu3152Glyfs)	1

**Note:** \*Novel alteration.

**Abbreviations:** HGVS, Human Genome Variation Society; PV, pathogenic variant; SNV, single nucleotide variant; c, chromosome site; p, protein site; del, deletion; fs, frame-shift mutation.

in a translational frameshift with a predicted alternative stop codon (p.S1951WfsTer). Of note, the two alterations, c.631G>A and c.7008-2A>T, were detected in the same patient, as previously reported.<sup>34</sup> The first mutation involves the substitution of one nucleotide containing a guanine (G) with an adenosine (A) in *BRCA2* exon 7 causing the change of valine amino acid with an isoleucine, an amino acid with highly similar properties, at codon 211. This change affects the normal mRNA splicing. The second variant is located in an intronic region and causes a double A to thymine (T) substitution before coding exon 13 of the *BRCA2* gene. The c.7008-2A>T alteration may produce multiple transcripts of different lengths. Moreover, in the *BRCA2* PVs group, 4 (22.2%) of 18 alterations were intronic.

Then, we mapped the *BRCA1/2* deleterious mutations across the functional domain and proteins binding regions (Figure 4). In *BRCA1* gene, 50% PVs were located in the breast cancer cluster regions (BCCR), while 22% mutations in the ovarian cancer cluster regions (OCCR) (Figure 4A). Among *BRCA2* PVs, 35.7% variants were in the BCCR region and 42.8% of mutations in the OCCR (Figure 4B). Next, we evaluated the location of PVs in *BRCA1* and *BRCA2* protein domains. For *BRCA1* protein, we found three PVs placed in both the ring domain and the coiled coil domain, and two mutations in the BRCT domain (Figure 4A). For *BRCA2* protein, 4 PVs mapped in the BRC repeats domains, while



**Figure 4** Schematic representations of *BRCA1* and *BRCA2* proteins and localization of pathogenic variants. The diagrams show the distribution of *BRCA1* (A) and *BRCA2* (B) pathogenic variants in breast cancer patients. The exonic mutations are depicted in blue, while the intronic variants in orange. The bar heights indicate the number of cases. *BRCA1* and *BRCA2* proteins are reported with their functional domains. (A) The *BRCA1* protein contains a RING domain and a nuclear localization sequence (NLS), a coiled-coil domain, a SQ/TQ cluster domain (SCD) and *BRCA1* C terminus domains (BRCT). (B) The *BRCA2* protein contains eight BRC repeats, a DNA-binding domain with a helical domain (Helical), three oligonucleotide/oligosaccharide binding (OB) fold, a Tower domain (T) and a NLS at the C-terminus. Regions referred to be breast cancer cluster regions (BCCRs) and ovarian cancer cluster regions (OCCRs) are shown at the bottom. \*Represents mutations determining a stop codon.

3 intronic and 3 exonic alterations were detected in the Oligonucleotide/Oligosaccharide Binding (OB) and Tower (T) domains (Figure 4B).

## Correlation Between Clinico-Pathological Characteristics and *BRCA* Pathogenic Variants

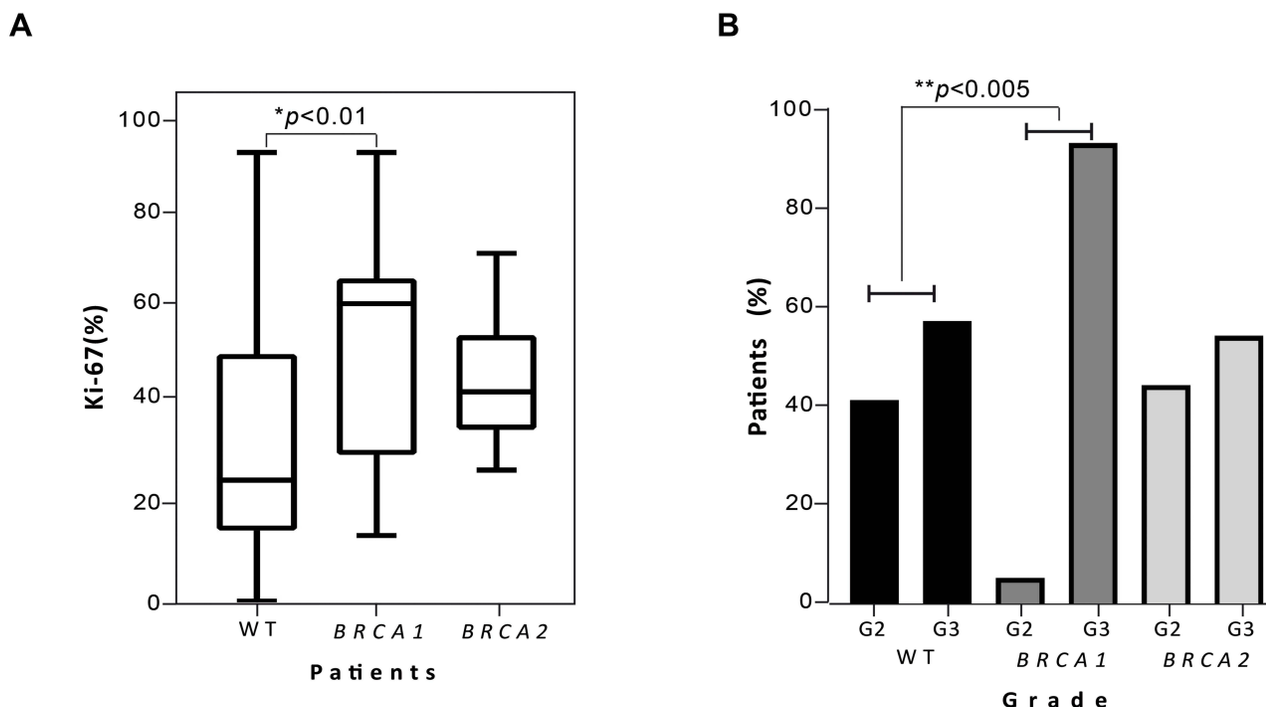
Then we looked at BC clinico-pathological features, which can correlate with the presence of *BRCA1/2* PVs. Complete clinical records were available for 181 patients with a negative *BRCA1/2* test (non-carriers) and for all the carriers (n=35). A correlation emerged for tumor proliferation rate and grade.

We calculated the distribution of Ki-67 according to median value of our cohort (25%, range <10–90%). Subjects with a Ki-67 <25% were defined as “low Ki-67” while individuals with value  $\geq 25\%$  were considered “high Ki-67”. A significant Ki-67 difference was found between non-carriers and *BRCA1* PV carriers ( $p < 0.01$ ) (Figure 5A).

Similarly, we examined whether tumor grading correlates with the presence of *BRCA1/2* PVs. Since no G1 BC were present in our population, we stratified patients in two groups (G2 or G3). Consistently with the Ki-67 results, the analysis revealed a statistically significant correlation between tumor grade and *BRCA1* mutations, with a higher proportion of G3 tumors among *BRCA1* carriers compared to non-carriers ( $p < 0.005$ ) (Figure 5B).

## Discussion

The development of DNA sequencing technologies determined an unprecedented progress in *BRCA1/2* genetic testing, with crucial consequences for patients presenting a family cancer history. To date, about 20.000 *BRCA1/2* variants have been identified and classified according to American College of Medical Genetics<sup>35</sup> and ENIGMA system.<sup>35,36</sup> It is known that the spectrum of *BRCA1/2* mutations varies greatly across different geographic regions.<sup>37</sup> Indeed, the rate of *BRCA1/2* PVs ranges from 8% to 37% throughout Italian territory, showing a broad intra-national variability.<sup>38,39</sup> With a population of almost 5 million people, Sicily represents the fifth Italian region in terms of number of inhabitants. While



**Figure 5** Correlation of Ki-67 and grading distribution in breast cancer women with or without *BRCA1* and *BRCA2* PVs. **(A)** Boxplots show comparison of median Ki-67 among 181 non-carriers BC patients and those with *BRCA1* (18) or *BRCA2* (17) PVs.  $P$  value below 0.5 was considered statistically significant. **(B)** The histograms represent the distribution of BC cancer patients into the histological grade group (G2 and G3) according to the *BRCA1* and *BRCA2* mutational status (WT subjects, *BRCA1* and *BRCA2* PVs carriers).

data exist about *BRCA1/2* distribution in western Sicily, no extensive evidences are available for the eastern part of the island.

Our study represents one of the first reports about the incidence of *BRCA1/2* PVs in BC patients from eastern Sicily.<sup>28</sup> We focused our analysis on BC, since this was by far the most frequent disease in our cohort.

Upon 389 BC patients tested, 9% harbored a *BRCA1/2* PV, evenly distributed between *BRCA1* and *BRCA2*. These results are in line with those previously reported in Italian population.<sup>28</sup> Interestingly, our cohort comprised 3% of men (13/389). This rate is higher than expected for male breast cancer (1% of all BC),<sup>40</sup> mirroring the selection of our population according to the risk of *BRCA1/2* mutations. However, none of the men presented a *BRCA1/2* PV, and they are therefore candidate to further molecular analyses to rule out the presence of less frequent mutations (eg *PALB2*, *RAD51C* and *D*, etc.). Variants of uncertain significance were retrieved in 7% of the subjects, with a clear prevalence of *BRCA2* VUS. Even this result is consistent with pre-existing evidences.<sup>28,41,42</sup>

When we analyzed the distribution of BC molecular subtypes among *BRCA1/2* mutated women, we confirmed the known association between TNBC and *BRCA1* PVs (58.8%) and between luminal B BC and *BRCA2* PVs (55.6%).<sup>16,43</sup> The scarce number of luminal A and HER2+ tumors among both *BRCA1* and *BRCA2* PV carriers is in line with the existing literature data.<sup>16,43</sup>

We then focused on the type and location of *BRCA1/2* PVs. In our cohort, the most frequent *BRCA1* PV was the c.5035\_5039delCTAAT. Although this variant has not been described by Incorvaia et al in their Sicilian cohort,<sup>28</sup> it has already been reported as germline *BRCA1* PV by other authors.<sup>34</sup> Several *BRCA1* PVs found in our cohort - such as c.181T>G, c.514del, c.3253dupA and c.5266dupC - have been observed in Sicily.<sup>28</sup> Among them, two *BRCA1* founder mutations (c.181T>G and c.5266dupC), typically retrieved among Ashkenazi Jews, in Eastern and Central Europe (Polish, Czech, Slovenian, Austrian, Hungarian, Belarusian and German),<sup>44,45</sup> but also in United States and Argentina, were recently defined as “recurring germline variants” among Italian BC and OC patients.<sup>34</sup> The c.514del variant was previously described in 8 breast cancer patients from Northern Sicily from Palermo and Messina. Interestingly, the c.3253dupA alteration has been found in some families from Catania even by Incorvaia et al.<sup>28</sup> The most representative *BRCA2* PVs were c.428dup, c.5851\_5854delAGTT and the intronic variant c.8487+1G>A, which were already described in a Sicilian population.<sup>28</sup> More in detail, the c.428dup was reported in patients from Palermo, the c.5851\_5854delAGTT PV was observed among families from north-western Sicily, mainly in the area of Trapani and Palermo, while the c.8487+1G>A variant was more frequently detected in subjects from Messina, Palermo and Caltanissetta.<sup>28</sup> The c.5851\_5854delAGTT alteration has been previously described by Rebbeck et al in Colombia.<sup>37</sup> Another *BRCA2* PV, c.631+1G>A, has been identified in BC and OC patients from Sicily (Agrigento, Siracusa and Ragusa).<sup>28</sup> Of note, we observed the coexistence of two *BRCA2* variants in the same patient (*BRCA2* c.631G>A and c.7008-2A>T) that we assume are segregated in *cis* modality, as previously reported.<sup>34,46</sup> These *BRCA2* double mutations were frequently observed in Italian regions and were found to introduce a premature stop codons, affecting messenger RNA splicing and leading to a nonfunctioning *BRCA2* protein.<sup>47,48</sup>

We also mapped the *BRCA1* and *BRCA2* PVs across the protein functional domains and the putative OCCR and BCCR regions of the genes. These regions were described by Rebbeck et al as risk regions for developing ovarian and breast cancer, respectively.<sup>49</sup> However, evidences about the correlation between the location of germline variants and the risk of breast or ovarian cancer remain controversial.<sup>28,50-52</sup> In our population, *BRCA1* PVs were mainly distributed in the BCCR domain, while *BRCA2* PVs were mostly located in the OCCR region. However, we were not able to find any association between the putative OCCR and BCCR regions and BC features. This may be due to the limited number *BRCA1/2* mutated patients. Looking at the protein domains, *BRCA1* PVs were distributed along the entire protein, *BRCA2* alterations were preferentially found in the BRC repeats domain.

Lastly, we correlated the BC clinico-pathological features with *BRCA1/2* PVs. Because of the limited number of patients included, we only found a significant correlation for Ki-67 and tumor grade. Even though Ki-67 assessment and interpretation still present some controversies, it is certain that a high proliferation rate is associated with an increased risk of disease relapse and worse survival. To date, the cut-off to distinguish between “high” and “low” Ki-67 is 20%. However, this threshold did not apply to our population of *BRCA1/2* mutated patients, which presented a Ki-67 median value of 25%. This trend towards a high Ki-67 rate can be explained by the prevalence in our cohort of luminal B and

TNBC, with few luminal A tumors. However, several evidences seem to suggest that a higher Ki-67 cut-off (25–30%) could better stratify patients according to their prognosis.<sup>53,54</sup> Looking at the results of our analysis, it is not surprising that a significant correlation emerged between high Ki-67 and grade and the presence of *BRCA1* PVs. Indeed, *BRCA1*-associated tumors are classically TNBC and display more aggressive features.<sup>16,17</sup>

## Conclusion

In conclusion, the present study provides a report about *BRCA1/2* mutational status in a cohort of BC from eastern Sicily. Overall, our findings are consistent with pre-existing evidences, either in terms of mutations prevalence and clinico-pathological features of BC. Additional studies on a larger population of *BRCA1/2* mutated BC patients are warranted, such as the extension of mutational analysis using multigenic panels to assess the presence of PVs different and less frequent than *BRCA1/2*. This would allow identifying and properly managing a growing number of subjects at increased risk of cancer due to inherited mutations.

## Ethics Statement

We confirm that the patients signed an informed consent to release anonymously their tumor samples for research purposes. All patients gave written informed consent in accordance with the Declaration of Helsinki. This study was exempt from ethical review based on the policy of the A.O.U. Policlinico “G.Rodolico - S.Marco” since *BRCA1/2* analysis was performed as per clinical practice and all the patients gave their written informed consent. Patients also gave consent for the use of their data for research purposes.

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## Disclosure

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