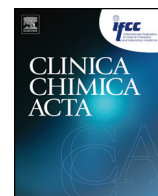




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## Q3 The molecular analysis of BRCA1 and BRCA2: Next-generation sequencing supersedes conventional approaches

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### A B S T R A C T

**Background:** Accurate and sensitive detection of *BRCA1/2* germ-line mutations is crucial for the clinical management of women affected by breast cancer, for prevention and, notably, also for the identification of at-risk healthy relatives. The most widely used methods for *BRCA1/2* molecular analysis are Sanger sequencing, and denaturing high performance liquid chromatography (dHPLC) followed by the Sanger method. However, recent findings suggest that next-generation sequencing (NGS)-based approaches may be an efficient tool for diagnostic purposes. In this context, we evaluated the effectiveness of NGS for *BRCA* gene analysis compared with dHPLC/Sanger sequencing.

**Methods:** Seventy women were screened for *BRCA1/2* mutations by both dHPLC/Sanger sequencing and NGS, and the data were analyzed using a bioinformatic pipeline.

**Results:** Sequence data analysis showed that NGS is more sensitive in detecting *BRCA1/2* variants than the conventional procedure, namely, dHPLC/Sanger.

**Conclusion:** Next-generation sequencing is more sensitive, faster, easier to use and less expensive than the conventional Sanger method. Consequently, it is a reliable procedure for the routine molecular screening of the *BRCA1/2* genes.

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## 1. Introduction

*BRCA1* and *BRCA2* are the most important genes predisposing to inherited breast and ovarian cancers [1]. Germline mutations in these two highly penetrant genes can increase the lifetime risk of developing these tumors by as much as 80%, and are also associated with an earlier onset of disease. Moreover, other kinds of cancers (i.e. prostate, pancreatic and gastric cancers) are more common in *BRCA1* and *BRCA2* carriers than in the general population [2]. Therefore, early identification of carriers among women affected by breast or ovarian cancer is crucial to enable patient stratification and to guide clinicians in deciding the most appropriate therapeutic strategy and follow-up program. In addition, accurate genetic counseling can identify at-risk healthy members of affected families, who can then be enrolled in appropriate surveillance programs.

**Abbreviations:** BC, breast cancer; dHPLC, denaturing high-performance liquid chromatography; FN, false negative; FP, false positive; NGS, next-generation sequencing; TN, true negative; TP, true positive.

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Direct Sanger sequencing continues to be the routine procedure for the molecular analysis of the *BRCA* genes [3,4]. However, given the large size of both *BRCA1* and *BRCA2* and the consequent cost of their direct sequencing, large-scale mutation scanning strategies such as denaturing high-performance liquid chromatography (dHPLC) and high-resolution melting, are also commonly used pre-sequencing methods [5,6]. By dramatically increasing the throughput of sequencing and reducing its costs, next-generation sequencing (NGS) has had a major impact on several fields of molecular research [7–9]. Furthermore, NGS is increasingly being applied in the field of diagnostics, including *BRCA* analysis [10–15].

Here, we report the outcome of an evaluation study conducted to assess the analytic performances of an NGS-based strategy for the molecular analysis of the *BRCA1* and *BRCA2* genes versus a well established, commonly used, dHPLC/Sanger sequencing strategy.

## 2. Materials and methods

### 2.1. Enrollment of patients and sample collection

Seventy women attending the Breast Unit, Istituto Nazionale dei Tumori, Fondazione G. Pascale in Naples, were consecutively enrolled

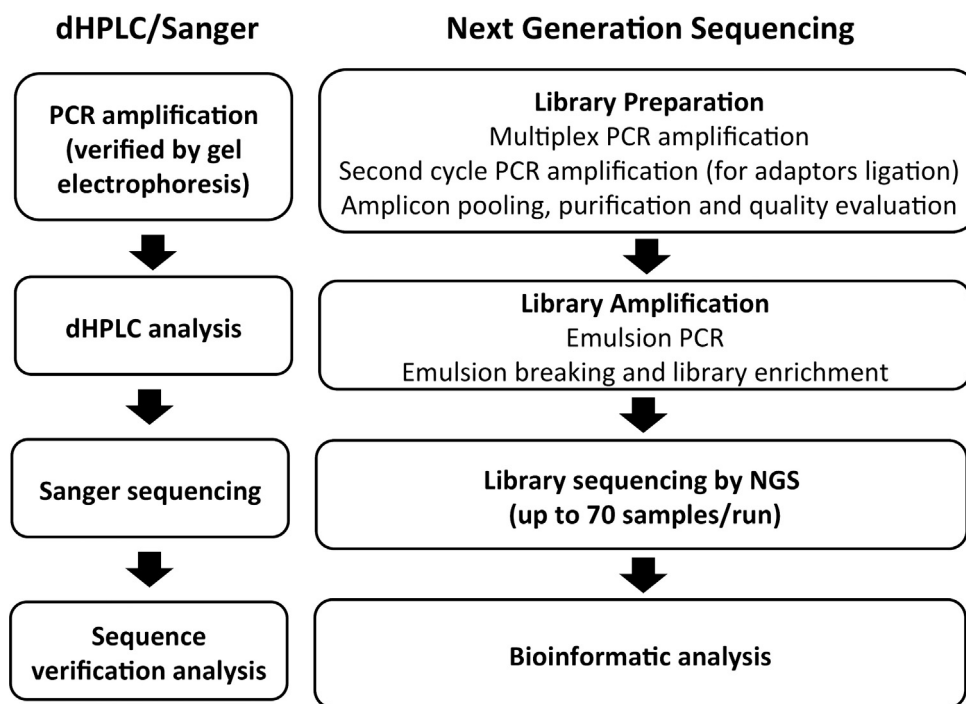


Fig. 1. Complete workflow of the two analytic strategies, dHPLC/Sanger and NGS, used to analyze the 70 at risk women.

in the study. All enrolled patients underwent pre-test counseling during which they were informed about the significance of molecular screening, provided information about their personal and familial history, and gave written informed consent to the study. The 70 enrolled women had at least one of the following conditions: early-onset breast cancer (BC) (diagnosed at 35 years or earlier); bilateral BC; multiple organ cancers, including BC; BC diagnosed at any age with at least one first- or two second-degree relatives with breast and/or ovarian cancer; BC diagnosed with advanced tumor staging; and patients with benign mammary alterations

with a BC-positive family history. The latter group consisted of 18 patients and can be considered a small control cohort.

A blood EDTA sample was collected from each subject. Genomic DNA was isolated from peripheral blood using the Nucleon BACC3 Genomic DNA Extraction Kit (GE Healthcare, Life Sciences, Little Chalfont, UK), according to the manufacturer's instructions. The quality of DNA samples was assessed by agarose gel electrophoresis and their quantity was evaluated with the NanoDrop 2000c Spectrophotometer (Thermo Fisher Scientific, Waltham, MA, USA).

**Table 1**  
BRCA1 variants identified in the analyzed population by dHPLC/Sanger and NGS.

Exon/intron	HGVS <sup>1</sup> cDNA	HGVS <sup>1</sup> protein	Mutation type	Clinical relevance (BIC) <sup>2</sup>	Reference ID according to NCBI	Number of times a variant was observed (dHPLC + Sanger/NGS)
IVS7	c.442-3_442 – 3delT	–	Deletion	Unknown	n.r. <sup>3</sup>	1/1
9	c.591C > T	p.Cys197Cys	Synonymous	Polymorphism	rs1799965	1/1
IVS10	c.671-12_671 + 12delG	–	Deletion	Unknown	n.r. <sup>3</sup>	1/1
11	c.1067A > G	p.Gln356Arg	Missense	Unknown	rs1799950	15/20
11	c.1911 T > C	p.Thr637Thr	Synonymous	Unknown	n.r. <sup>3</sup>	1/1
11	c.2077G > A	p.Asp693Asn	Missense	Polymorphism	rs4986850	14/17
11	c.2082C > T	p.Ser694Ser	Synonymous	Unknown	rs1799949	31/39
11	c.2311 T > C	p.Leu771Leu	Synonymous	Unknown	rs16940	30/38
11	c.2612C > A	p.Pro871Gln	Missense	Polymorphism	rs799917	23/42
11	c.3113A > G	p.Glu1038Gly	Missense	Polymorphism	rs16941	36/36
11	c.3119G > A	p.Ser1040Asn	Missense	Unknown	rs4986852	3/3
11	c.3418A > G	p.Ser1140Gly	Missense	Unknown	rs2227945	0/1
11	c.3419G > T	p.Ser1164Ile	Missense	Deleterious	n.r. <sup>3</sup>	1/1
11	c.3548A > G	p.Lys1183Arg	Missense	Polymorphism	rs16942	33/37
11	c.3711A > G	p.Ile1237Met	Missense	Unknown	rs80357388	0/1
13	c.4308 T > C	p.Ser1436Ser	Synonymous	Polymorphism	rs1060915	11/37
14	c.4484G > T	p.Arg1495Met	Missense	Deleterious	rs80357389	0/1
16	c.4837A > T	p.Ser1613Cys	Missense	Unknown	rs1799966	1/1
16	c.4837A > G	p.Ser1613Gly	Missense	Polymorphism	rs1799966	32/38
16	c.4843G > A	p.Ala1615Thr	Missense	Unknown	rs80356987	1/1
16	c.4956G > A	p.Met1652Ile	Missense	Unknown	rs1799967	0/1
16	c.4964_4982del	p.Ser1655_Glu1661fs	Frameshift	Deleterious	rs80359876	1/1
IVS18	c.5153-1G > C	–	Substitution	Deleterious	rs80358137	1/1

<sup>1</sup> All identified variants are indicated both by cDNA base sequence (second column) and by protein sequence (third column) according to the HGVS (Human Genome Variation Society) nomenclature guidelines.

<sup>2</sup> Breast Cancer Information Core (<http://research.nhgri.nih.gov/bic/>).

<sup>3</sup> n.r., not reported in NCBI (National Center for Biotechnology Information) database.

t2.1 **Table 2**  
t2.2 **BRCA2 variants identified in the analyzed population by dHPLC/Sanger and NGS.**

t2.3	Exon/intron	HGVS <sup>1</sup> cDNA	HGVS <sup>1</sup> Protein	Mutation type	Clinical relevance (BIC) <sup>2</sup>	Reference ID according to NCBI	Number of times a variant was observed (dHPLC + Sanger/NGS)
t2.4	10	c.865A > C	p.Asn289His	Missense	Polymorphism	rs766173	6/6
t2.5	10	c.865A > G	p.Asn289Asp	Missense	Unknown	rs766173	1/1
t2.6	10	c.1114C > A	p.His372Asn	Missense	Polymorphism	rs144848	26/56
t2.7	10	c.1124C > T	p.Pro375Leu	Missense	Unknown	rs80358409	0/1
t2.8	10	c.1151C > T	p.Ser384Phe	Missense	Polymorphism	rs41293475	0/1
t2.9	10	c.1365A > G	p.Ser455Ser	Synonymous	Polymorphism	rs1801439	8/8
t2.10	IVS10	c.1909 + 12_1909 + 12delT	–	Deletion	Unknown	n.r. <sup>3</sup>	1/1
t2.11	11	c.2229 T > C	p.His743His	Synonymous	Polymorphism	rs1801499	5/5
t2.12	11	c.2971A > G	p.Asn991Asp	Missense	Polymorphism	rs1799944	4/6
t2.13	11	c.3396A > G	p.Lys1132Lys	Synonymous	Polymorphism	rs1801406	28/29
t2.14	11	c.3807 T > C	p.Val1269Val	Synonymous	Polymorphism	rs543304	19/31
t2.15	11	c.3824 T > C	p.Ile1275Thr	Missense	Unknown	rs80358625	0/1
t2.16	11	c.4131_4132insTGAGA	p.Asn1377_Thr1378	In Frame Insertion	Deleterious	rs80359429	1/2
t2.17	11	c.4563G > A	p.Lys1521Lys	Synonymous	Polymorphism	rs206075	27/69
t2.18	11	c.4585G > A	p.Gly1529Arg	Missense	Polymorphism	rs28897728	1/1
t2.19	11	c.5199C > T	p.Ser1733Ser	Synonymous	Polymorphism	rs28897734	1/2
t2.20	11	c.5312G > A	p.Gly1771Asp	Missense	Polymorphism	rs80358755	1/3
t2.21	11	c.3515C > G	p.Ser1172Trp	Missense	Unknown	rs80358600	0/2
t2.22	11	c.5744C > T	p.Thr1915Met	Missense	Polymorphism	rs4987117	3/1
t2.23	11	c.6037A > T	p.Lys2013Ter	Nonsense	Deleterious	rs80358840	1/1
t2.24	22	c.6486_6489delACAA	p.Lys2162_Gln2163fs	Frameshift	Deleterious	rs80359598	1/1
t2.25	11	c.6513C > G	p.Val2171Val	Synonymous	Polymorphism	rs206076	45/69
t2.26	14	c.7242A > G	p.Ser2414Ser	Synonymous	Polymorphism	rs1799955	16/23
t2.27	14	c.7354A > G	p.Asn2452Asp	Missense	Unknown	rs398122580	1/1
t2.28	IVS16	c.7806-14 T > C	–	Deletion	Unknown	rs9534262	1/1
t2.29	27	c.9976A > T	p.Lys3326Ter	Nonsense	Polymorphism	rs11571833	1/1

t2.30 <sup>1</sup> All identified variants are indicated both by cDNA base sequence (second column) and by protein sequence (third column) according to the HGVS (Human Genome Variation Society) nomenclature guidelines.

t2.32 <sup>2</sup> Breast Cancer Information Core (<http://research.nhgri.nih.gov/bic/>).

t2.33 <sup>3</sup> n.r., not reported in NCBI (National Center for Biotechnology Information) database.

## 91 2.2. dHPLC/Surveyor nuclease and Sanger sequencing

92 The combined dHPLC/SURVEYOR® Nuclease approach  
93 (Transgenomic, Omaha, NE, USA) was used for molecular *BRCA1*  
94 and *BRCA2* screening, as previously described [5]. Briefly, PCR am-  
95 plifications were performed using specific primer pairs to amplify  
96 all the *BRCA1* and *BRCA2* coding exons and their flanking regions  
97 using a multiamplicon approach (Supplemental Table 1). After am-  
98 plification, each PCR reaction/sample was digested by SURVEYOR®  
99 Nuclease using 15 µL of a 1:1 mixture of DNA amplicons of each  
100 sample and wild-type DNA, 1.5 µL of 0.15 mM MgCl<sub>2</sub>, 1.5 µL of En-  
101 hancer Cofactor and 1.0 µL of SURVEYOR Nuclease (Transgenomic).  
102 The reactions were incubated at 42 °C for 60 min and stopped by  
103 adding 1.5 µL of stop buffer. For each amplicon pool, a gradient  
104 was chosen using Navigator software (Transgenomic) based on  
105 the size of the PCR fragments. Fragments were analyzed by HPLC  
106 on the Transgenomic WAVE Nucleic Acid High Sensitivity Fragment  
107 Analysis System (WAVE HS system; Transgenomic). Amplicon re-  
108 actions that showed a dHPLC profile suggestive of mutations were  
109 selected for Sanger sequencing confirmation. Direct sequencing  
110 was performed with an ABI 3100 capillary sequencer (Applied

111 Biosystems Inc., Foster City, CA, USA). Sanger electropherograms  
112 evaluation was carried out using the SeqMan tool (DNASTAR, Inc.,  
113 Madison, Wisconsin, USA).

## 114 2.3. NGS sequencing

115 Multiple amplicon DNA libraries, covering all the *BRCA1* and *BRCA2*  
116 coding exons and their flanking regions, were obtained using the  
117 BRCA MASTR v2.1 Assay kit (Multiplicom, Niel, Belgium), following  
118 the manufacturer's instructions. In brief, for each patient, 250 ng of ge-  
119 nomic DNA was used to perform 5 multiplex PCR reactions able to am-  
120 plify the entire target region. Then, a 1:1000 dilution of each multiplex  
121 PCR was re-amplified using hybrid primers to univocally tag all the mul-  
122 tiplexes from the same patient with a barcode sequence (MID). These  
123 primers contained a universal adaptor sequence that is required for  
124 downstream sequencing reactions. The multiplexed tagged reactions  
125 were purified using Agencourt AMPure XP Beads (Beckman Coulter,  
126 Brea, CA, USA), quantified using the NanoDrop 2000c Spectrophotome-  
127 ter (Thermo Fisher Scientific, Waltham, MA, USA), and quality-assessed  
128 using the Experion DNA 1 kb Analysis kit (Bio-Rad, Hercules, CA, USA).  
129 Equimolar amounts of the 5 multiplexed tagged reactions from the

t3.1 **Table 3**  
t3.2 **Analytic performances of dHPLC/Sanger and NGS methods in detecting BRCA 1/2 sequence variants.**

t3.3	DHPLC/Sanger		NGS		
	BRCA1	BRCA2	BRCA1	BRCA2	
t3.5	N total variants	237	198	320	323
t3.6	N unique variants	19 (7 SNPs, 9 UCV, 3 causative mutations)	22 (15 SNPs, 3 UCV, 4 causative mutations)	23 (7 SNPs, 12 UCV, 4 causative mutations)	26 (16 SNPs, 6 UCV, 4 causative mutations)
t3.7	Analytic sensitivity % (95% CI)	67.6 (63.9–71.1)		100 (99.4–100)	
t3.8	Analytic specificity % (95% CI)	100 (100)		100 (100)	
t3.9	PPV %	100		100	
t3.10	NPV %	99.98		100	

t3.11 N, number; SNP, single nucleotide polymorphism; UCV, unknown significance variant; CI, confidence interval; PPV, positive predictive value; NPV, negative predictive value.

130 same sample were pooled to create a library/sample. Up to 35 different  
131 libraries were pooled and sequenced using the Genome Sequencer FLX  
132 System (454-Life Science and Roche, Branford, CO, USA), according to  
133 the manufacturer's instructions. Causative mutations or doubt variants  
134 were confirmed by Sanger sequencing.

#### 135 2.4. NGS data analysis

136 NGS sequence data were analyzed using the SeqPilot software (ver-  
137 sion 3.5.2) from JSI Medical Systems ([www.jsi-medsys.de](http://www.jsi-medsys.de)). The *BRCA1*  
138 (ENSG00000012048 for gene reference and ENST00000357654 for  
139 transcript) and *BRCA2* (ENSG00000139618 and ENST00000380152 re-  
140 spectively) sequences from the NCBI-database (<http://www.ncbi.nlm.nih.gov>)  
141 were used as reference sequences. Raw sequencing data were  
142 directly uploaded in the software and, for each MID, blasted against a  
143 target reference sequence. Thus, we obtained a report/patient contain-  
144 ing the sequence-coverage/exon and the list of high confidence variants.  
145 According to the manufacturer's protocol, a minimum absolute cover-  
146 age/exon of 40× is allowed and only variants present in both directions  
147 and with a minimum coverage of 10% default, excluding homopoly-  
148 mers, were contemplated in the genetic analysis. All sequence variants  
149 are named according to the nomenclature used by Human Genome Var-  
150 iation Society, HGVS (<http://www.hgvs.org>). The significance of vari-  
151 ants was attributed according to the Breast Cancer Mutation database,  
152 BIC (<http://research.nhgri.nih.gov/projects/bic/>), and the Ensemble Da-  
153 tabase (<http://www.ensembl.org>).

#### 154 2.5. Assessment of methodology performance

155 The sensitivity and specificity of the described procedures were  
156 assessed by evaluating their ability to correctly identify all the base  
157 changes identified in the same population by direct Sanger sequencing  
158 of all *BRCA1* and *BRCA2* exons. As specified above, missing Sanger data  
159 were obtained after the NGS analysis to determine if a specific variant  
160 was present or not, and were used as gold standard to identify each sub-  
161 stitution as true positive (TP), true negative (TN), false positive (FP) or  
162 false negative (FN), and to assess the analytic performances of the two  
163 approaches used. Sensitivity and specificity were calculated using the  
164 following formulas: sensitivity = TP / (TP + FN); specificity = TN /  
165 (TN + FP). Confidence intervals for sensitivity and specificity were esti-  
166 mated with the Pearson–Klopper method, and the R statistical software  
167 environment.

### 168 3. Results

169 The *BRCA1* and *BRCA2* genes were analyzed in the 70 enrolled sub-  
170 jects using the two analytic strategies described under Section 2  
171 (Fig. 1). The combined dHPLC/Surveyor® nuclease strategy revealed  
172 237 and 198 variants in *BRCA1* and *BRCA2*, respectively, and these  
173 were confirmed by Sanger sequencing. These variants correspond to  
174 19 *BRCA1* variants, i.e., 7 polymorphisms, 9 variants of unknown signif-  
175 icance (UCVs) and 3 causative mutations (Table 1), and to 22 *BRCA2* var-  
176 iants, i.e., 15 polymorphisms, 3 UCVs, and 4 causative mutations  
177 (Table 2). The NGS-based strategy revealed 320 and 323 variants in  
178 *BRCA1* and *BRCA2*, respectively. These variants correspond to 23 *BRCA1*  
179 variants, namely, 7 polymorphisms, 12 UCVs, and 4 causative mutations  
180 (Table 1), and 26 *BRCA2* variants, namely, 16 polymorphisms, 6 UCVs,  
181 and 4 causative mutations (Table 2).

182 Comparative analysis of the results obtained with the two analytic  
183 strategies showed that all the variants detected by the routinely used  
184 dHPLC/Sanger method were identified also by NGS. Interestingly, the  
185 NGS technique identified variants missed by conventional screening  
186 (Tables 1 and 2). Additional Sanger sequencing was carried out to assess  
187 all these discrepancies and confirmed the NGS data.

188 We assessed the reliability of the NGS-based approach by comparing  
189 the NGS results to the Sanger sequencing results (Table 3). While the

specificity was 100% for both methods, the NGS-based procedure had  
a higher sensitivity (100% versus 67.6%), which is in line with a previous  
report [16]. In addition, in our study the NGS-based method had also a  
higher diagnostic sensitivity since it was able to detect a causative  
*BRCA1* mutation missed by traditional screening (Table 3).

In terms of analytical time, it took about three months to analyze all  
the 70 enrolled subjects using the dHPLC/Sanger approach, and only  
10 days using NGS. In addition, the cost of NGS reagents was half that  
of the reagents required for the dHPLC/Sanger procedure.

### 4. Discussion

Thanks to the recent development of NGS technologies, and the con-  
sequent decreased cost of DNA sequencing, a wide range of technical  
possibilities are now available for studies aimed at elucidating the mo-  
lecular basis of human diseases. In addition, these technologies are  
now beginning to be exploited for diagnostic purposes. Therefore, it is  
conceivable that in the near future NGS may become an invaluable  
tool for both clinical biochemists and clinicians. In fact, increasing evi-  
dence indicates that information about the *BRCA1* and *BRCA2* mutation  
status, especially if available in a timely fashion, will enable a patient  
and her/his health-care provider to make informed decisions about can-  
cer prevention, screening and treatment [21].

In this optics, we evaluated the efficacy of an NGS-based method for  
the molecular analysis of *BRCA1* and *BRCA2* in 70 patients using both  
dHPLC/Sanger and NGS-based strategies. Here, we demonstrate that  
NGS is more sensitive than the conventional approaches in detecting se-  
quence variants. In addition, since we analyzed up to 70 samples in the  
same sequencing run, also the time and cost of the analysis were greatly  
reduced. Given these results, this NGS-based approach to the detection  
of *BRCA1/BRCA2* mutations is suitable for use in a routine diagnostic  
workflow.

Although several NGS-based procedures have been proposed for the  
molecular diagnosis of the predisposition for breast cancer, our study  
demonstrates that NGS-based tests are more sensitive, faster, easier to  
use, and less expensive than the conventional Sanger method, which  
is the most widely test currently used and is considered the gold stan-  
dard test.

Supplementary data to this article can be found online at <http://dx.doi.org/10.1016/j.cca.2015.03.045>.

### 5. Uncited reference

[17]

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