Detection of \textit{BRCA1}/2 Gene Mutation Rate Among Women in Hilla Province

Wala’a Nori Bearem* Mohammad Sabri A. Razzak** Mohammed Abbod Muhsin** Ifad Kerim Alshibly**
* Breast Cancer Center, Hilla General Teaching Hospital, Iraq.
** College of Medicine, University of Babylon, Hilla, Iraq.

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Abstract

\textbf{Background:} breast cancer is one of the leading causes of death among women. One-fourth of women will be affected with cancer at some point in their lives. Breast cancer is the most prevalent malignancy in women. Mutations in breast cancer susceptibility genes \textit{BRCA1} and \textit{BRCA2} account for the majority of breast cancer cases. \textit{BRCA1} and \textit{BRCA2} are tumor suppressor genes that repair and correct errors in DNA, act as sensors of DNA damage and participate in the DNA repair processes.

\textbf{Aim:} to illustrate the role of and analyzing the types and frequencies of the most common \textit{BRCA1}, \textit{BRCA2} mutations in different groups of local Iraqi women.

\textbf{Materials and methods:} This is a case-control study design. A total of forty women were chosen for the genetic study which was performed to detect \textit{BRCA1} and \textit{BRCA2} mutations, DNA was amplified by polymerase chain reaction (PCR), exon 2 and 20 of \textit{BRCA1} and exon 11 of \textit{BRCA2} genes, using a specific primer pairs and PCR conditions in four groups of Babylonian populations, including patients with breast cancer, healthy control people, first degree relatives, and patients with benign breast tumors.

\textbf{Results:} The overall frequency of \textit{BRCA} genes mutation was more often detected among breast cancer patients than other groups in the study ($P=0.05$).

\textbf{Conclusions:} There was a surprising high degree of \textit{BRCA} gene mutation carriage rate, \textit{BRCA} gene mutations were found to have a relatively high frequency (up to 75%) among breast cancer patients in the study.

Introduction

Breast cancer is the most common malignancy affecting women and its incidence is increasing worldwide, making it the most common type of non-skin cancer in women and the fifth most common cause of death due to cancer [1]. The majority of breast and ovarian cancers are sporadic or not inherited [2]. In addition to specific genetic changes, many personal and environmental factors have been identified that may influence a person's risk of developing breast cancer[3]. Mutation in some genes such as the breast cancer susceptibility genes \textit{BRCA1} and
BRCA2 predispose to familial risk of breast cancer [4]. BRCA1 and BRCA2 are human genes that belong to a class of genes known as tumor suppressors. In normal cells, BRCA1 and BRCA2 help ensure the stability of the cell’s genetic material (DNA) and help prevent uncontrolled cell growth. Mutation in these genes has been linked to the development of hereditary breast and ovarian cancer [5]. Over 200 individual BRCA mutations have been described, they are found throughout the length of the gene, some areas appear to be "hot spots" for mutation [6]. A mutated BRCA gene usually makes a protein that does not function properly because it is abnormally short [7]. Mutations in the BRCA genes are inherited in an autosomal dominant fashion [8]. A number of screening test have been employed for breast cancer including: clinical and self breast exams, mammography, genetic screening, ultrasound, and magnetic resonance imaging [9]. A breast cancer (BRCA) gene test is a blood test to check for specific mutations BRCA genes. This can help determine the chance of developing breast cancer and ovarian cancer. This test is only done for people with a strong family history of breast cancer or ovarian cancer [10]. Early diagnosis and screening, especially when combined with adequate therapy, offer the most immediate hope for a reduction in breast cancer mortality. This was the basis of the Iraqi national program for early detection of breast cancer, which was initiated in 2000 in an attempt to downstage this disease at the time of presentation. Since then specialized centers and clinics for early detection of breast cancer have been established in the major hospitals in all Iraqi provinces [11]. Genetic assays to test patients for specific mutations are commercially available, but their use remains somewhat controversial and they should not be applied to the population at large. When carriers are identified, it is appropriate to offer radiological studies at an increased frequency and beginning at an earlier age, as well as to discuss the option of prophylactic surgery and counseling/testing other family [6].

The prevalence of BRCA1 or BRCA2 mutations varies considerably among ethnic groups and geographical areas. In this study we tried to identify the carriage rate of some selected mutations among BRCA1/2 genes in a specified local Iraqi population.

Materials and Methods
Study design: This study is a pilot study design.

Patients:
A total of forty subjects were chosen for the genetic study, the primary set of subjects were drown from those attending the breast cancer center in Hilla General Teaching Hospital from March through October 2013. Four groups were recruited; 20 breast cancer patients, 10 healthy controls, 5 benign breast tumor patients and 5 first degree relatives to a known case of breast cancer. Both of the breast cancer and the benign breast tumor patients were clinically, radiologically and histopathologically confirmed as having breast cancer and benign breast tumor, respectively, while healthy control and first degree relatives to the breast cancer patients were clinically and radiologically confirmed as having no breast problems. All cases and controls were of Iraqi origin resident in Hilla city, and its peripheries. All subjects were screened for the presence of mutation in BRCA1 exon2, BRCA1 exon20, and BRCA2 exon11.

DNA extraction:
The screening of mutation in BRCA genes were carried out on genomic DNA. DNA was extracted from blood samples using the conventional methodologies supplied by the Promega company (www.promega.com), DNA was extracted from fresh whole blood collected in EDTA , heparin and citrate
anticoagulant tubes. The extracted DNA was stored in 2-8°C.

**Screening for BRCA1 and BRCA2 mutations:**
To detect BRCA1 and BRCA2 mutations, DNA was amplified by polymerase chain reaction (PCR), exon 2 and 20 of BRCA1 and exon 11 of BRCA2 genes, using the primer pairs and PCR conditions in table (1)[12].

**Ethical Issues:**
All patients and family members were counseled then they signed a written informed consent before testing. This study was performed with permissions from Babylon University; College of Medicine, Hilla General Teaching Hospital and from the Ministry of Health in Iraq.

**Results**
In the present study, the prevalence of BRCA1/2 gene mutation carriage rate was investigated among local Iraqi women, using conventional PCR technique and gel electrophoresis with required technical information illustrated in table(1).

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**Table 1** Primers sequences, product size and PCR conditions for BRCA genes mutation.

<table>
<thead>
<tr>
<th>Gene</th>
<th>Primer Sequence</th>
<th>Size /bp</th>
<th>PCR conditions</th>
<th>Temp.</th>
<th>time</th>
<th>cycles</th>
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<td>BRCA1</td>
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<tr>
<td>185delAG (exon 2)</td>
<td>F-5'-GAA GTT GTC ATT TTA TAA ACC TTT-3' R-5'-TGT CTT TTC TTC CCT AGT ATG T-3'</td>
<td>275</td>
<td>{95°C 5min 1x}</td>
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<td>5382insC (exon 20)</td>
<td>F-5'-ATA TGA CGT GTC TGC TCC AC-3' R-5'-GGG AAT CCA AAT TAC ACA GC-3'</td>
<td>425</td>
<td>{95°C 5min 1x}</td>
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<td>BRCA2</td>
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<tr>
<td>6174delT (exon 11)</td>
<td>F-5'-AAC GAA AAT TAT GGC AGG TTG TTA C-3' R-5'-GCT TTC CAC TTG CTG TAC ATC C-3'</td>
<td>534</td>
<td>{95°C 5min 1x}</td>
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In order to measure the frequency of BRCA mutation among local Iraqi population, a genetic analysis about the most common BRCA mutation was carried out, using conventional PCR technique and a highly specific primer sets. It was observed that there was a surprising high degree of BRCA gene mutation carriage rate. Figures (1) shows the electrophoretic imaging of BRCA1 exon-2, BRCA1 exon-20, and BRCA2 exon-11 gene mutation respectively. Figure (2) illustrates the results about the frequency of BRCA gene mutation among the study groups.
**Figure 1** The electrophoretic imaging of \( BRCA1 \) exon-2, \( BRCA1 \) exon-20, and \( BRCA2 \) exon-11 gene mutation.

**Figure 2** Frequency of \( BRCA \) gene mutations
The overall frequency of \( BRCA \) genes mutation was more often detected among breast cancer patients than others in the study (figure 2). Eight patients with breast cancer carried all of the three analyzed mutations, and seven had two mutations. Otherwise, \( BRCA \) genes frequency showed low prevalence among first degree relatives and benign breast tumor patients, and to lesser extent among healthy controls, with only few, had all of the mutations combined.

The frequency of \( BRCA1 \) exon 2 mutation was found in 55% (11/20) patients of breast cancer, while only 20% (2/10) healthy control subjects carried this mutation. On the other side, both the first degree relatives and patients with benign tumors had the same frequency of this type of \( BRCA \) mutation giving a value of 40% (2/5).

In this study, \( BRCA1 \) exon 20 mutation was the most frequently recorded mutation, it was detected in (15/20) patient of breast cancer, forming 75% of this group. The healthy controls had the lowest frequency of this gene mutation, with 30% carriage rate (3/10). Likewise, 40% (2/5) of the first degree relative had the mutation, while 60% (3/5) of patients with benign tumors carried it.

\( BRCA2 \) gene mutation was also studied, the results found that 40% of breast cancer and benign tumor patients were carrier of mutation in the exon 11 of \( BRCA2 \) gene, 20% of first degree relatives had the mutation, and only 10% of the control were mutated.

**Discussion**

According to the Iraqi health ministry registrations of breast cancer, the incidence rate of breast cancer have been doubled from 3.26/100,000 in 1990 to 6.33/100,000 in 2000, with annual incidence rate of about 50,000/year [11]. Coming with accordance with the present study results, a study of \( BRCA \) mutations by Couch et al. (2007) [13], showed that the most commonly detected \( BRCA1 \) mutations are a deletion of adenine and guanine [\( BRCA1 \) 185delAG (exon 2)], and insertion of cytosine [\( BRCA1 \) 5382insC (exon 20)], and \( BRCA2 \) mutation is the deletion of thymine [\( BRCA2 \) 6174delT (exon 11)], which have a high frequency in the general population.

Likewise, Easton et al. (2009) [14] pointed that mutations in the breast cancer susceptibility genes \( BRCA1 \) and \( BRCA2 \) confer a high risk of developing breast cancer. Breast cancer is about twice as common in the first-degree relatives of women with the disease as in the general population, consistent with variation in genetic susceptibility to the disease.

There is an evidence that specific mutations in \( BRCA1 \) and \( BRCA2 \) more common in certain populations, for example, Friedenson, (2007) [15] had reported that three specific mutations, were the most common mutations found in these genes in the Ashkenazi Jewish population. Other ethnic and geographic populations around the world, such as the Norwegian, Dutch, and Icelandic peoples, also have higher frequencies of specific \( BRCA1 \) and \( BRCA2 \) mutations.

Ravi (2010) [16] has established that according to the cancer registries of almost all countries within the Eastern Mediterranean Region, as in Iraq, the breast cancer continues to rise. The Iraqi National Cancer Research Program has to better understand the underlying molecular and environmental causes in an effort to curb the incidence of cancer.

The results presented in this study illustrate the high rate of \( BRCA \) gene mutation carriers, this come in parallel with that of Al-Dujaily et al. (2008) and his colleagues who demonstrate that there were three-fold increase of all types of breast cancer in Iraq due to long-term exposure to radiation. Radiation levels were estimated to be around 320–800 tons in the aftermath of the first Gulf war in 1991 with further comparable levels occurring in 2003 [17]. The significance of this risk factor in women with a genetic susceptibility to breast cancer is
unclear. The possibility that genetic susceptibility to breast cancer occurs via a mechanism of radiation sensitivity raises questions about radiation exposure. However, they propose that despite the three-fold increase of all types of breast cancer in Iraq due to long-term exposure to depleted uranium, the increment in the incidence may not become evident with time until mutations are passed to future generations.

The proportion of the high risk breast cancer attributable to *BRCA1* and *BRCA2* has been shown to vary considerably between different studies. The observed frequency is not always the same in any two countries surveyed so far. The mutation frequencies reported in the present study came in parallel with another study undertaken in the north of Iraq by Majid *et al.* (2009)[18], who found that the age specific annual incidence rates of breast cancer were higher among Iraqi Kurds than for Israeli Arabs and Jordanians and were similar to Egyptians. Age specific breast cancer incidence rates for Jordanian and Israeli Arab women lag behind those in the United States.

The frequency of the *BRCA1*/*BRCA2* mutations in the Iraqi population has been compared with other populations including those of Ashkenazi. Lahad *et al.* (2007)[12] have studied the frequency of *BRCA1*/2 mutations in Ashkenazi Jews in Israel including cases of Iraqi and Iranian origin, they found that the frequency of *BRCA1* 185delAG and *BRCA1* 5382insC mutations was significantly higher than that of *BRCA2* 6174delT mutation. Analysis of these three markers in four Iranian/Iraqi *BRCA* gene mutation carriers revealed the same genotype that previous reports have described in Ashkenazi Jews, suggesting a common origin for the mutations in Ashkenazi and Iraqi/Iranian Jews. These results confirmed the present study reports, in which the three mutations accounted for most of high-risk breast cancer cases.

Identification of *BRCA* mutations in a substantial proportion of our Iraqi patients indicates that these genes play an important role in the incidence of breast cancer in the general population. These results were supported by Turkish study accomplished by Yazici *et al.* (2000)[19], who demonstrated the presence of *BRCA* mutations in patients with a personal and family history of breast cancer below age 50. Their results suggest that *BRCA1* and *BRCA2* mutations are observed in a significant proportion of Turkish families with breast cancer, and those with early onset of disease. This study demonstrates the importance of the consideration of inherited predisposition to breast cancer in the clinical management of breast cancer risk.

The data considering the frequency of *BRCA* gene mutation in the current study came in line with the fact that there is a significant association between the presence of *BRCA1* or *BRCA2* mutations and the occurrence of cancer. Al-Mulla *et al.* (2009)[2] have studied different mutations in the *BRCA* gene in Britain population, they reveal that more than half of individuals carried a mutated *BRCA1* gene, and less had a mutated *BRCA2* gene, while about one-third of the screened individuals carried wild-type genes. Exon 2 mutations were the most frequent with 185delAG being the most common mutation within exon 2, followed by mutation of exon 20 (5382insC). For *BRCA2* gene, mutations in exon 11 (6174delT) was the most frequent. Mutations in the *BRCA1* gene accounted for the majority of breast cancer. It was noted that carriers of the exon 2 of *BRCA1* gene mutation had significantly lower cancer incidence compared with carriers of other exons mutation. These findings are consistent with the present work, suggesting to that either the 185delAG mutation is of low penetrance or, more likely, carriers of this mutation would develop cancer at an older age compared with other *BRCA1* gene mutations.
It is well accepted that mutation frequencies of the BRCA1 and BRCA2 genes in high-risk people vary widely among different populations, the present study is consistent with this fact. The contributions of BRCA1 and BRCA2 to breast cancer in Italian patients appear to be less significant than in patients from Iraq and other communities with these mutations. In one Italian study, Ottini et al. (2000) [20] have found that BRCA1 mutations were detected in only one out of 10 cases from breast cancer families. This is a low proportion compared with other studies which suggested that mutation in BRCA1 and BRCA2 are responsible for the large majority of breast cancer families, with the greater proportion due to BRCA1. In this respect, the limits of mutation detection techniques and the small number of breast cancer cases tested should be taken into account.

Finally, it became clear that ethnicity affects frequency of BRCA gene mutation among populations. Showing that particular mutations in genes associated with breast cancer are more common among certain geographic or ethnic groups. The genetic changes occur more frequently in these groups because they have a shared ancestry over many generations. Different ethnic groups may have different mutations. The prevalence of these genes varies depending on ethnicity.

Conclusions

In conclusion, There was a surprising high degree of BRCA gene mutation carriage rate, BRCA gene mutations were found to have a relatively high frequency (up to 75%) among breast cancer patients in the study.

References


