ABSTRACT

Objective: Cancer ovary is one of the fatal gynecologic malignancies worldwide. Since breast cancer (BRCA) genes are considered tumor suppressor genes and play important roles in cancer by repairing of chromosomal damage with the error repair of DNA breaks. Therefore, breast cancer 1 (BRCA1) and breast cancer 2 (BRCA2) gene mutations strongly enhance the development of ovarian cancer risk among women. Here, we report that both genes are an essential mediator of progress ovarian cancer, to determine the influence of BRCA1 and BRCA2 mutations in the improvement of ovarian cancer.

Methods: A total of 25 subjects were chosen for the genetic studies, and three groups were recruited: fifteen ovarian cancer patients group, five healthy controls, and five first-degree relatives to a known case of ovarian cancer patients.

Results: A genetic analysis revealed that a strong correlation exists between both gene mutations’ status in ovarian cancer, and BRCA gene mutations (185delAG, 5382insC, and 4153delA in BRCA1 and 6174delT in BRCA2) remained to establish to have a relatively high frequency among people in this study among ovarian cancer patients. Furthermore, seven patients with ovarian cancer carried all of the four investigated mutations, and five had three mutations.

Conclusion: Otherwise, BRCA gene frequency showed low prevalence among first-degree relatives, and to a lesser extent among healthy controls, with only a few had all of the mutations combined. These data demonstrate for the first time a molecular link between BRCA1 and BRCA2 mutations in ovarian cancer progression in Iraq.

Keywords: Breast cancer 1, Breast cancer 2, Mutation, Ovarian cancer, Polymerase chain reaction.

INTRODUCTION

Cancer is one of the most important health problems of malignant diseases. This disease arises due to the abnormal division of cells without control. The condition that accompanies surrounding healthy tissue is due to tumor formation and abnormal growth of this tumor. This will result in an invasion of the neighboring tissues and metastasis in other organs through blood or lymphatic stream [1]. Ovarian cancer affects many women in the world. Therefore, the cancer of ovary has developed the greatest common malignancy and the subsequent main cause of cancer death in western women. The development and progression of ovarian cancer are controlled by complex mechanisms that are still not completely understood. Several aspects of this disease such as grade, genetics, and oncogene suppressor gene expression seem to be more heterogeneous, its related features involving tumor molecular events and hormonal and biological characteristics are particularly difficult. Ovarian cancer is a challenging health problem; there are two factors concerning its etiology and these are genetic and environmental issues. However, the pathogenesis of this cancer related with many hormonal and lifestyle aspects which demonstrated an essential role in this cancer, and these aspects include one of the largest risk factors of family history of such disease [2]. The first degree in a woman with a relative affected with ovarian cancer is correlated with improved risk of ovarian cancer in a three-fold ratio [3].

So far, there are several high-risk ovarian cancer susceptibility variants, two different genes named BRCA1 and BRCA2, both genes recognized to be best factors in the molecular events in the virulence of ovarian cancer. BRCA1 and BRCA2 have an important function as tumor suppressor genes and encode proteins involved in deoxyribonucleic acid (DNA) repair [2]. Several lines of evidence strongly suggest that the women carrying mutations in these 2 genes have an affectedly increased risk of developing ovarian cancer in their lifetime. Interestingly, more than 400 different mutations have been identified in these two genes, BRCA1 and a nuclear polypeptide of 220 kDa (1863 amino acids) encodes, and located on chromosome 17q12-21. In families with a high percentage of breast cancer (BRCA) either in early-onset BRCA or ovarian cancer happens, the majority of ovarian cancer incidence followed by the mutation of gene 1.

Furthermore, in high risk families the second gene which encodes a 384 kDa (3418 amino acids) and located on chromosome 13q12.1 is similarly considered as an increased risk factor of ovarian and BRCA mutation [4]. Numerous reports demonstrated that the majority of breast and ovarian cancers within these high-risk families appeared to involve by these two genes, and molecular investigations for the percentage of breast and ovarian cancer within the people that might be address such ovarian susceptibility genes still undocumented. The observation demonstrated that BRCA1 protein has a significant role in the signaling of DNA damage and in DNA repair, although the mechanism of action has not yet been adequately addressed. For this reason, several genetic studies have suggested that intracellular mechanism pathway involving a response to DNA damage, transcription, and interaction with other proteins included in DNA repair and apoptosis was contributed to the major roles of both genes [5]. BRCA1 and BRCA2 mutations have
been investigated in numerous studies using breast and ovarian cancer. However, to the best of our knowledge, only two studies have reported a comparison of percentage of the disease related to the expression profiles of mutation, BRCA1 mutation in women assessed to have risk factor between 26% and 54% for developing ovarian cancer, while in BRCA2 mutation assessed the percentage between 10% and 23% [6,7].

El-Harith et al. [8] had further examined the prevalence of both genes mutation for tumorigenesis in ovarian and BRCA and assessment of the benefits and limitations of molecular testing for these cancers. Essentially, some authors have described that testing of molecular events for susceptibility to cancer appears mainly suitable for the family to have been documented at high-risk cancer. Expressing of such abnormal mutations related to the subgroup of this family, mutation of these genes is considered as a possible diagnostic tool with clinical findings. The early truncate of proteins induced non-sense mutation which is considered common pathological mutations [5]. Since the lack of general population documents and multiplicity of BRCA1 and BRCA2 mutations developed problematic to control the specific risk estimate for each mutation which may enhance these mutations that are significant raise the risk for breast and ovarian cancer. One study described two specific population mutations with BRCA1 185delAG and 5382insC while 6174delT mutation with BRCA2 among Ashkenazi women in percent of 60% in ovarian and 30% in primary BRCA respectively [9].

The aim of the current study was to classify BRCA1 and BRCA2 mutation in Iraqi women with ovarian cancer and considered as molecular tools detection. Here, we show that in ovarian cancer gene mutations, ranks are upregulated and the mechanism by which maintenance of these mutations is succeeded to improve progression in ovarian cancer. Finally, fast development is one guarantee of ovarian tumor and these data demonstrated that this gene may promise for specific gene therapy for this type of cancer.

**METHODS**

**Patient samples**

This is a reconsidering study of patients diagnosed and treated for ovarian malignancy during their life. Clinical samples (blood) were collected from three groups of women; and two of these groups are undergoing radical ovariectomy for the treatment of ovarian cancer. A total of 25 samples were selected for the genetic studies, and the primary set of samples were obtained from women (age 45–70 years) attending the clinic of detection of ovarian cancer who were admitted to Al-Ramadi Teaching Hospital in Ramadi city (Iraq). Again, three groups were recruited: Fifteen ovarian cancer patients group, five healthy as controls, and five first-degree relatives to a documented case of ovarian cancer. A total of 25 samples were selected for the genetic studies, and the primary set of samples were obtained from women (age 45–70 years) attending the clinic of detection of ovarian cancer who were admitted to Al-Ramadi Teaching Hospital in Ramadi city (Iraq). Again, three groups were recruited: Fifteen ovarian cancer patients group, five healthy as controls, and five first-degree relatives to a documented case of ovarian cancer patients. Patient’s history was obtained from patient giving blood samples. Patient approval to participate in scientific research was taken according to the approval certificate from Maternity and Child Teaching Hospital.

**Polymerase chain reaction (PCR)**

**Total DNA extraction from blood samples**

In brief, genomic DNA was obtained from blood samples by Quick Prep Total DNA Purification Kit (Geneaid, UK) according to the producer guidelines. The DNA isolated from the samples was separately labeled and kept for the following procedure.

**PCR primers**

The oligonucleotide primers were synthesized by SIGMA®. On arrival, they were suspended to the appropriate stock concentration with ultrapure water and stored at –20°C. For PCR applications, the primers were diluted to 10 pmol/µl as a working concentration.

**PCR of ovarian cancer markers**

Here, we describe a simple and rapid method for the simultaneous detection of four common mutations: The primers were designed to amplify a 275 bp fragment of185delAG, 425 bp of 5382insC, and 134 bp of 4153delA in BRCA1 and 534 bp of 6174delT in BRCA2. For more description, to amplify genes encoding ovarian cancer with PCR technique, the primers and PCR conditions are indicated in Fig. 1. The sample of PCR reaction in the volume of 25 µl contains the following materials, 2.5 µl of each upstream and downstream primer, 2.5 µl of free nuclease water, 5 µl of DNA extraction, and 12.5 µl of master mix.

**DNA agarose gel electrophoresis**

The PCR products were electrophoresed for sizing and qualitative analysis using a horizontal submarine mini-gel apparatus, 1% agarose gels (Bioscience Services, UK), and electrophoresis power supply (Kodak UK). 15 µl of the PCR reaction was added to 2 ul of Gel Loading Buffer (Bioline, UK) and electrophoresed for 1 h at 60 V in a ×1 TAE (89 mM Tris-base, 2 mM Na2-EDTA, 89 mM Boric Acid, pH 8.3) buffered 2% w/v agarose gel (Bioline, UK) with ×1 TAE which is buffer for running. 2 µl of a 10 mg/ml solution was added of ethidium bromide (Sigma, UK) to the agarose before pouring to permit visualization of the DNA fragments electrophoresed on a 312 nm UV trans illuminator (Syngene, UK). The size marker used was appropriate to the size of the ampiclon and in this instance was an allelic ladder (Promega, USA).

**RESULTS**

In ovarian cancer, the expression of genes is varies, these genes contain several well-known ovarian cancer-associated gene transmutations, involving 185delAG, 5382insC and 4153delA (BRCA1) and 6174delT (BRCA2). PCR technique analysis showed a single band for all the mutations studied at the corresponding molecular weight in each gene. Therefore, to identify these mutations above in human ovarian cancer, we evaluated in this study major markers which have been screened for the main mutations overhead and are illustrated in (Figs. 2-5) respectively.

In the present study, the results were shown that the overall frequency of BRCA1 mutation was more often detected among ovarian cancer patients than others included in the study. Seven patients with ovarian cancer carried all of the four analyzed mutations, while five of them had three mutations. Otherwise, BRCA1 gene frequency showed low prevalence among first-degree relatives and to a lesser extent among healthy controls, with only a few had all of the mutations combined.

![Fig. 1: Overview of primers sequences and polymerase chain reaction condition to detect breast cancer 1 and breast cancer 2 genes expression](image-url)
The frequency of BRCA1 (185delAG) was the most frequently recorded mutation, it was detected in 66% (10/15) of patients of ovarian cancer, while only 20% (1/5) healthy control subjects carried this mutation. On the other side, both the first-degree relatives had the frequency of this type of BRCA mutation giving a value of 40% (2/5). Interestingly, the significant increase of mutation related with the patient no 8 presented that the assessed generation risk of ovarian cancer progress was higher in 2 times due to overexpressed of this c.6869delAG (185delAG) mutation, it was detected in 66% (10/15) of patients of ovarian cancer, and activity in all samples, we next examined the BRCA1 (5382insC). Next, we assessed the BRCA2 gene mutation (6174delT). The results showed that five ovarian cancer patient samples (33%) were carrier of mutation in the exon 11 of BRCA2 gene (mean±SEM 81.14±2.56, 60.08±3.82, 60.32±2.52, 53.00±4.36 and 85.85±5.09), and only one out of five (20%) of first-degree relatives have been showed appeared of this mutation (mean±SEM 51.9±5.24). In this population, BRCA1 mutations were common (185delAG and 5382insC) (Fig. 5).

Nevertheless, when compared to the incidence of both genes’ mutation between targeted carriers, the incidence of BRCA2 mutations is different about 50% that of BRCA1 mutations, giving a finding of carriers with BRCA2 persist unaffected. In this study, we established that, in which people were determined frequently depending on the sufficient family history, the BRCA2-6174delT mutation was approximately half as frequent as the BRCA1-185delAG and 5382insC mutations combined.

At this point, the purpose for the dominance of BRCA1 mutations among this population unclear; molecular explanation of this matter may be due to restricted to this gene in the founder effects. Even though the ovarian cancers related to inherited mutations in cancer susceptibility genes represent a small proportion of all cancers, it is of great importance for the clinician to detect the patients who are carrying these mutations. Consequently, the determination of potential risk among the family members of the mutation carrier can be estimated and prevention measures can be undertaken. Focusing on this target gene to reduce the ovarian cancer risk could develop a novel approach for early molecular diagnostic tools in ovarian malignancy.

The carriers of BRCA1 and BRCA2 gene mutations increase the risk factor and developing ovarian cancers during their lifetime. Particularly, the BRCA1 defect influences to early beginning of the hereditary ovarian cancer because both genetic and environmental converters are possible to influence low-penetrance mutations than high-penetrance mutations. These findings are dependable with earlier observations of lower penetrance of BRCA2 and more related to the BRCA2-6174delT mutation. As illustrated above, studies in a high-risk people found ovarian cancer risk which is higher in BRCA1 compared with BRCA2 families.

To our information, the different expression among BRCA1 and BRCA2 still needs more investigation in this present study. Consequently, women with BRCA1 mutations showed a high level of risk factor and developing the disease (40%) comparing with a low level of BRCA2 mutations group (20%). The diseases which develop the relation to ovarian cancers in both above gene mutations are commonly serous papillary carcinomas, though endometrial and clear cell carcinomas may be happening. In summary, this study provides evidence that BRCA1 plays a key role in ovarian cancer patients. It is important to highlight that there has been a significant interest in the development of these gene mutations as genetic therapy. During this study, samples investigated were limited number (25 samples for three groups), although these restrictions, our findings may be valued for additional studies on other genes related to this type of malignant organ.

DISCUSSION

In support of this discussion, a previous study by Fodor et al. revealed that almost of heritable ovarian carcinomas proposed due to the high prevalence of BRCA1 mutation comparing in small prevalence in the involvement of BRCA2 mutations [10]. Therefore, we found a significant difference in prevalence between BRCA1 and 2 which was a high percentage with BRCA1 mutations than in those with BRCA2 mutations among ovarian cancer women. In addition, there are some factors which play a critical role in enhancement the activate risk of
Fig. 4: The expression of breast cancer 1 (BRCA1) 4153delA amplicon product in ovarian cancer. Samples were prepared, separated by gel electrophoresis of polymerase chain reaction as outlined in methods, and then assessed for (a) BRCA1 4153delA amplicon product. Gels were quantified for (b) fold expression by scanning densitometry (mean±standard error of mean 83.46 ± 5.47); L: Ladder; NC: Negative control; 1, 2, 3, 4, 5, 6, 7, 8: Number of ovarian cancer patient; 9, 10: Number of the first-degree relatives; 11: Healthy control.

Fig. 5: The expression of breast cancer 2 (BRCA2) 6174delT amplicon product in ovarian cancer. Samples were prepared, separated by gel electrophoresis of polymerase chain reaction as outlined in methods, and then assessed for (a) BRCA2 6174delT amplicon product. Gels were quantified for (b) fold expression by scanning densitometry (mean±standard error of mean 66.12 ± 7.26); L: Ladder; NC: Negative control; 1, 2, 3, 4, 5, 6, 7, 8: Number of ovarian cancer patient; 9, 10: Number of the first-degree relatives; 11: Healthy control.
increasing ovarian cancer, such as parity, contraceptive oral drug, and oophorectomy. High levels of androgen in women indicated highly predisposed to a tumor of ovarian [11]. The recent study by Iyer et al. demonstrated that different population is established with a specific transcript of BRCA1 of the chromosome 17. These variations of nucleotide indicated risk issues in genetic ovarian and BRCA syndrome in the worldwide population [12]. In addition, another mutation in cytochrome b gene related to the mitochondrial genome indicated as expected to be possibly pathogenic in the event of ovarian cancer patients [13].

According to the ethnic groups, the evidence of 185delAG mutation was significantly different depending on the different population. The previous study demonstrated that the percentage of this mutation was the actual high frequency of 31.6% among non-Jewish Americans of Spanish ancestry from the San Luis Valley, Colorado [14]. In contrast to other studies, this mutation was variable low frequency (1.3–5.9 %) in a group of Americans Americans, the Spanish from Spain, Polish, Iranian, Pakistani, and Turkish women [15,16]. The evidence of BRCA1 185delAG mutation in “Jewish” was described for the non-Ashkenazi population through the Middle East, Greece, Turkey, and England (Yorkshire) in addition to two Indo-Pakistani families [17]. Taken together, this indicated that the mutation happens in a recombination segment of BRCA1, with non-Ashkenazi ethnic groups may explain how the different haplotypes depending on a high mutation rate in this repeat region [18]. Likewise, 40% (2/5) of the first-degree relative had the mutation. In addition, among Ashkenazi Jews, this mutation was common, and it was established in approximately 25% of mutation in high hereditary risk of ovarian and BRCA among Jewish women [19]. A previous study by Gosay et al. [20] has been demonstrated that the finding from limited cases proposed that the mutation of 5382insC also evident in women suffering breast and ovarian cancer from Hungary and Latvia.

Our results are agreement with another result, similar as previously determined by Gayther et al. [21], who also observed that the mutation of BRCA1 4153delA was the remarkable rate in percent of 2/177 (1.1%) in Russian cancer families related to ovarian cancer. Respectively, this mutation also indicated in Lithuania, Latvia, Belarus, and Poland [22,23]. Notably, other findings demonstrated that the prevalence of this mutation was more association in ovarian than breast type. On the another hand, several other reports have established an improved incidence of ovarian cancer patients with the mutation of 4153delA carriers [24,25]. While, other study indicated that the most common mutation among population level was BRCA2-6174delT in valued carrier incidence of 1.4%, compared with 1.1% with combined of BRCA1 mutations [26]. The correlation percentage between BRCA1 and BRCA2 from above studies was not observed at the same percentage in our study; however, the carriers of 6174delT mutation considered greater risk of ovarian cancer than the percent of BRCA2 risks (20% vs. 11%). These data in agreement with previous findings demonstrated that mutations in the ovarian cancer are related with high risk for this type of cancer [27]. The proportion of the high-risk ovarian cancer attributable to BRCA1 and BRCA2 has been shown to vary considerably between different studies. A study by Antoniou et al. [6] was shown that 75 carriers were BRCA1 185delAG, while 69 were BRCA1 5382insC carriers, and 52 carriers were BRCA2 6174delT. In contrast, the chances of ovarian cancer for BRCA1 and BRCA2 mutation carriers are 16-63% and 10-27%, respectively [7].

Since the syndromes of familial ovarian cancer linked to the gene mutations that happen on this gene, the people of ovarian cancers are affected by mutations on the BRCA1 or BRCA2 gene in approximately 10% [28]. Our findings in the present study advised that to consider the assessment of the common BRCA1 gene as a first target for screening in a high-risk population. Mutations with BRCA2 also confer a low risk of ovarian cancer. Interestingly, a family history of ovarian cancer is strongly predictive of a BRCA1 mutation and thus suggested that both genes retain highly candidate mutations in these genes substantially enhanced the developing risk of rising ovarian cancer.

CONCLUSION

The finding in this study has wide applications to deal with other types of tumor beyond ovarian cancer. Therefore, data in this study confirmed the identification of BRCA1 and BRCA2 among human ovarian cancer patients, which considered possible novel investigative or therapeutic targets in human ovarian cancer. Furthermore, these gene mutation expression sketches allow beneficial in convoluted gene appearance evaluates investing expression sketches from many various causes.

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CONFLICTS OF INTEREST

The authors that there were no conflicts of interest.

REFERENCES


