Germline Variations of Apurinic/Apyrimidinic Endonuclease 1 (APEX1) Detected in Female Breast Cancer Patients

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Abstract

Apurinic/apyrimidinic endonuclease 1 (APEX1) is a multifunctional protein which plays a central role in the BER pathway. APEX1 gene being highly polymorphic in cancer patients and has been indicated to have a contributive role in Apurinic/apyrimidinic (AP) site accumulation in DNA and consequently an increased risk of cancer development. In this case-control study, all exons of the APEX1 gene and its exon/intron boundaries were amplified in 530 breast cancer patients and 395 matched healthy controls and then analyzed by single-stranded conformational polymorphism followed by sequencing. Sequence analysis revealed fourteen heterozygous mutations, seven 5’UTR, one 3’UTR, two intronic and four missense. Among identified mutations one 5’UTR (rs41561214), one 3’UTR (rs17112002) and one missense mutation (Ser129Arg, Mahjabeen et al., 2013) had already been reported while the remaining eleven mutations. Six novel mutations (g.20923366T>G, g.20923435G>A, g.20923462G>A, g.20923516G>A, g.20923539G>C>T) were observed in 5’UTR region, two (g.20923585T>G, g.20923589T>G) in intron1 and three missense (Glu101Lys, Ala121Pro, Ser123Trp) in exon 4. Frequencies of 5’UTR mutations; g.20923366T>G, g.20923435G>A and 3’UTR (rs17112002) were calculated as 0.13, 0.1 and 0.1 respectively. Whereas, the frequency of missense mutations Glu101Lys, Ser123Trp and Ser129Arg was calculated as 0.05. A significant association was observed between APEX1 mutations and increased breast cancer by ~9 fold (OR=8.68, 95%CI=2.64 to 28.5) with g.20923435G>A (5’UTR), ~13 fold (OR= 12.6, 95%CI=3.01 to 53.0) with g.20923539G>A (5’UTR) and ~5 fold increase with three missense mutations [Glu101Lys (OR=4.82, 95% CI=1.97 to 11.80), Ser123Trp (OR=4.62, 95% CI=1.7 to 12.19), Ser129Arg (OR=4.86, 95% CI=1.43 to 16.53)]. The incidence of observed mutations was found higher in patients with family history and with early menopause. In conclusion, our study demonstrates a significant association between germ line APEX1 mutations and breast cancer patients in the Pakistani population.

Keywords: Germline variations - BER pathway - APEX1 - breast cancer - Pakistan

Introduction

DNA damage caused by exogenous carcinogens and endogenous oxygen species is the most important factor for carcinogenesis and genomic instability through activation of oncogenes and inactivation of tumor suppressor genes. Thus, DNA repair play a role in maintaining genomic stability and protecting the cells against carcinogenesis (Zhou et al., 2012). There are several distinct DNA repair pathways including mismatch repair, nucleotide excision repair (NER), base excision repair (BER), homologous recombination repair and non-homologous end joining repair pathways. BER pathway is the primary guardian against damage that results from cellular metabolism including reactive oxygen species, methylation, deamination and hydroxylation. Genetic variations in BER may result in reduced or defective repair capability and accumulated DNA damage consequently leading to permanent mutations in the genome, which may ultimately contribute to carcinogenesis. Therefore, base excision repair is a universal event in cells for preventing mutagenesis (Zhou et al., 2011; Alanazi et al., 2013; Li et al., 2013; Luo et al., 2014).

Apurinic/apyrimidinic Endonuclease 1 (APEX1) also known as REF-1 is a multifunctional protein which plays a central role in BER pathway. APEX1 initiates DNA repair activity by hydrolyzing the 5’-phosphodiester backbone at the apurinic/apyrimidinic (AP) sites produced by endogenous and exogenous sources thus protecting the cells from accumulation of AP sites in DNA. APE1 can also act as 3’-phosphodiesterase to initiate repair of DNA single strand breaks, which are produced either directly by reactive oxygen species or indirectly through enzymatic removal of damaged bases. In humans, APEX1 is located

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Materials and Methods

Collection of samples

Present study was conducted with a prior approval from ethical committees of both COMSATS Institute of Information Technology Islamabad (CIIT) and collaborating hospitals. A total of 530 female patients with histologically confirmed breast cancer were recruited from Nuclear Medicine, Oncology and Radiotherapy Institute (NORI) Islamabad Pakistan. Additionally 395 age and ethnicity matched cancer free healthy female individuals were also recruited as controls. The inclusion criterion for the controls was absence of prior history of cancer or any other familial disease (diabetes, blood pressure and cardiovascular impairment) were excluded from this study. After obtaining informed consent, all individuals were personally interviewed using the specifically designed questionnaire. Information about age, age at menarche, menopausal status, menopausal age, family history, ethnic group was collected from both patients and control individuals. Detailed histopathological reports of patients were also recorded for their clinical characterization.

DNA extraction and polymerase chain reaction (PCR)

DNA was extracted from white blood cells, using standard phenol-chloroform extraction method (Masood et al., 2011) with minor alterations. Freshly extracted DNA was quantified by spectrophotometry and yield gel electrophoresis and stored at -20°C till further processing. Human APEX1 exon sequence was taken from Ensemble. Primers were designed using primer 3 software and checked for specific amplification using BLAST. Whole coding region including exon intron boundaries of approximately 60 bp sequence of APEX1 was investigated-identify novel and any splice site variation in addition-previous reported mutations. Each PCR reaction was performed in a 10 μl reaction mixture containing 1μl of genomic DNA (approximately 50 ng) templates, 1μl (10 mM) of each primer, 1μl nuclease free water and 5 μl PCR master mix (Thermo Scientific) containing 0.05U/μl Taq DNA polymerase, reaction buffer, 4mM MgCl2, 0.4 mM of each dNTP. PCR conditions were initial melting step at 94°C for 5 min, 35 cycles each comprised of 94°C for 45 sec, exon specific annealing temperature for 1 min and 72°C for 1 min. It was followed by a final extension step at 72°C for 10 min and finally held at 4°C. 2 μl of PCR products along with loading dye were electrophoresed on a 2 % agarose gel and stained with ethidium bromide. 100bp ladder was also loaded as standard for quantification of amount and confirmation of PCR product size.

Mutational screening and sequence analysis

Single stranded conformational polymorphism (SSCP) assay was used for mutational analysis of PCR products. Samples with altered electrophoretic mobility were re-amplified in a separate reaction and were analysed by direct sequencing-confirm and characterize the nature of mutations/ polymorphisms. Control (normal) samples were also sequenced along with cancerous samples-compare the sequencing results. Sequencing was carried out by MC lab (USA). Results after DNA sequencing were analysed using BioEdit (version 7.0.5) software.

Data analysis

χ²-test and Fisher exact test were used-evaluate the differences in selected demographic variables, family history, tumor types, tumor grades, ER/ PR and HER-2/nue status by using the Graph Pad Prism 5 Demo. Spearman correlation coefficient was used-assess the correlations among the observed mutations and clinical and histo-pathological parameters. Missense mutations were also analysed in-silico via Alamut biosoftware (version 2.2)-predict the pathogenicity of point mutations, PhyloP for conservation level of mutated nucleotides and amino acids along with Grantham distance for physico-chemical changes in amino acid structure.

Results

Mean age of breast cancer patients and controls was calculated as 46.4 (±11.59) and 42.80 (±12.96) years respectively. Patients and control individuals with age
>40 years were significantly (p <0.007) more afflicted in number than patients of ≤40 years age.

In the present study, all exons of APEX1 were screened extensively for germline mutations involving SSCP followed by sequence analysis in 530 patients and 395 control individuals. Fourteen different types of mutations were observed, which include seven 5'UTR, one 3'UTR, two Intronic and four missense mutations. All mutations observed in this study were heterozygous. Among identified mutations one 5'UTR (g.20923416C>T, rs41561214), one 3'UTR (20925669A>T, rs17112002) and one missense mutation (Ser129Arg, Mahjabeen et al., 2013) has already been reported while remaining elven mutations are novel. Six novel mutations (g.20923366T>G, g.20923435G>A, g.20923462G>A, g.20923516G>A, g.20923529C>T) were observed in 5'UTR region, two (g.20923585T>G, g.20923589T>G) in intron 1 and three (g.20924881G>A, Glu101Lys; g.20924941G>C, Ala121Pro; g.20924948C>G, Ser123Trp) were observed in exon 4 (Figure 1).

Figure 1. Sequencing electropherogram of polymorphisms of APEX1. A, B, C, D, E, F & G are substitutions in 5'UTR region { (A) g.20923366 T>G substitution in 5'UTR region. (B) g.20923435 G>A substitution in 5'UTR region. (C) g.20923462 G>A substitution in 5'UTR region. (D) g.20923516 G>A substitution in 5'UTR region. (E) g.20923539 G>A substitution in 5'UTR region. (F) g.20923529 C>T substitution in 5'UTR region. (G) g.20923416 C>T (rs41561214) substitution in 5'UTR region.} (H) g.20925669 A>T (rs17112002) substitution in 3'UTR region. (I) g.20923585 T>G & g.20923589 T>G substitutions in intronic region. J, K & L are Missense substitutions in Exon 4 { (J) Missense mutation Glu101Lys showing g.20924881G>A substitution resulting in change of codon from GAG to AAG encoding the amino acid Glutamic acid instead of Lysine. (K) Missense mutation Ala121Pro showing g.20924941G>C substitution resulting in change codon from GCT to CCT encoding the amino acid Proline instead of Alanine and Missense mutation Ser123Trp showing g.20924948 C>G substitution resulting in change of codon from TCG to TGG encoding the amino acid Serine instead of Tryptophan. (L) g.20924967 T>G substitutions resulting in change of codon from AGT to AGG encoding the amino acid Serine instead of Arginine (Ser129Arg).}
Increased breast cancer risk was found associated with different mutations when compared with controls (Table 1). In case of 5'UTR mutations, ~4-fold increase in breast cancer risk was associated with g.20923667G>T (OR=4.08, 95% CI=1.97-8.43), g.20923516G>A (OR=3.86, 95% CI=1.46-10.18), g.20923529C>T (OR=4.26, 95% CI=1.41-12.88) mutations, whereas ~5 fold increased breast cancer risk was associated with g.20923416C>T (OR=5.07, 95% CI=2.27-11.32), ~6 fold increase with g.20923462G>A (OR=5.66, 95% CI=1.97-16.23), ~9 fold increase with g.20923435G>A (OR=8.68, 95% CI=2.64-28.50) and ~13 fold increase with g.20923539G>A (OR=12.63, 95% CI=3.01-53.01) mutations. In similar fashion ~5 fold increase in breast cancer risk was observed in 3'UTR mutations g.20925669A>T (OR=5.42, 95% CI=2.74-10.73) and both intronic mutations g.20923585T>G (OR=5.11, 95% CI=3.23-11.25) and g.20923589T>G (OR=5.19, 95% CI=2.43-11.10). While analyzing missense mutations, a significantly increased breast cancer risk was also found associated with missense mutations when compared with controls showing ~5 fold increase with three mutations (Glu101Lys (OR=4.92, 95% CI=1.97-11.80), Ser123Trp (OR=4.62, 95% CI=1.7-12.19), Ser129Arg (OR=4.86, 95% CI=1.43-16.53)) and ~4 fold increase with Ala121Pro (OR=3.68, 95% CI=0.98-13.86).

Missense mutations were also analysed via Alamut biosoftware-check the conservation levels of mutated nucleotides and amino acids along with in-silico predictions about Algin GVGD score, Grantham distance; SIFT score, Mutation Taster (Table 2). Mutation taster predicted three missense mutations (Glu101Lys, Ser123Trp, Ser129Arg) as potentially disease causing (p<1.0). Greater physicochemical difference was predicted between Serine and Tryptophan amino acids (Ser123Trp; Grantham distance=177) and Serine and Arginine amino acids (Ser129Arg; Grantham distance=110).

Association of mutations observed in this study were also correlated with different clinico-pathological parameters including family history, menopausal age, HER-2/neu and ER/PR status. Frequency of patients with family history was observed significantly higher (p<0.0001) compared-frequency of control individuals. As presented in Table 4, significantly higher frequency of 3'UTR mutation, g.20925669A>T (p<0.009) and two missense mutations Glu101Lys (p<0.009) and Ser129Arg (p<0.05) was observed in patients with family history when compared with controls. Correlations between frequency of APEX1 mutations and menopausal age of breast cancer patients (Table 4) showed that the frequency level PhyloP score its conservation level, dist. Taster
Score, Grantham Distance and Mutation Taster Prediction

| Change in Nucleotide with its conservation level PhyloP score | Amino acid change with its conservation level (considering 13 species) | SIFT score (median) | Align GVGD score (C0-C65) | Grantham dist. (0-215) | Mutation Taster (p-value) |
|-------------------------------------------------------------|-------------------------------------------------------------|--------------------|------------------------|----------------------|-------------------------|
| g.20924881G>A                                               | Highly conserved nucleotide phyloP: 4.32 Highly conserved amino acid, up to Baker’s yeast | Glu101Lys          | Tolerated 0.18          | CO (GV: 95.14-GD: 0.00) | 56 Disease causing (p-1.0) |
| Weakly conserved nucleotide phyloP: 1.66                     | Moderately conserved amino acid                             | Tolerated 0.22      | CO (GV: 227.66-GD: 0.00) | 27 Disease causing (p-1.0) |
| g.20924984G>G                                               | Weakly conserved nucleotide phyloP: 2.55                    | Ser123Trp          | Deleterious 0.01        | C15 (GV: 120.14-GD: 97.50) | 177 Disease causing (p-1.0) |
| g.20924967T>G                                               | Weakly conserved nucleotide phyloP: 0.12                    | Ser129Arg          | Tolerated 0.16          | CO (GV: 141.06-GD: 0.00) | 110 Disease causing (p-1.0) |

PhyloP was used as a conservation score rating the nucleotides from “not conserved” (-14.1) to “highly conserved” (6.4) (Pollard et al., 2010). Align GV GD score: most likely deleterious (C65)

Table 3. Correlation between APEX1 Mutations and Tumor Types, ER/PR Status and HER-2/neu Status in Breast Cancer Patients

| Mutations                              | Type of Tumor | ER Status | PR Status | HER-2/neu Status |
|----------------------------------------|---------------|-----------|-----------|-------------------|
| DCIS No (%)                            | IDC No (%)    | ILC No (%)| -ve No (%) | +ve No (%) |
| g.20923366G>T 5'UTR                    | 8 (10.25)     | 29 (13.87) | 19 (9.09)  | 8 (10.25)       |
| g.20924345G>A 5'UTR                    | 7 (8.97)      | 24 (11.11)| 17 (8.61)  | 24 (11.11)      |
| g.20924622G>T 5'UTR                    | 10 (2.05)     | 16 (7.44)| 8 (6.50)   | 14 (7.44)       |
| g.20925161G>T 5'UTR                    | 4 (5.13)      | 16 (7.87)| 8 (6.50)   | 14 (7.44)       |
| g.20925394G>T 5'UTR                    | 9 (11.54)     | 11 (7.92)| 9 (6.50)   | 14 (7.44)       |
| g.20925928T>C 3'UTR (rs41561214)       | 5 (6.41)      | 25 (2.77)| 6 (3.50)   | 25 (2.77)       |
| g.20925669A>T 3'UTR (rs1712002)        | 6 (7.69)      | 11 (5.09)| 15 (7.18)| 8 (6.48)        |
| g.20925855T>G Inttron 1               | 6 (7.69)      | 15 (6.94)| 17 (8.13) | 8 (6.48)        |
| g.20925857T>G Inttron 1               | 6 (7.69)      | 15 (6.94)| 16 (7.55)| 8 (6.48)        |
| g.20924881G>A Exon 4 g.2091101Lys     | 2 (2.56)      | 14 (6.48)| 10 (4.78)| 7 (6.48)        |
| g.20924914G>C Exon 4 p.1211Pro       | 2 (2.56)      | 5 (2.31) | 5 (2.39)  | 1 (0.56)        |
| g.20924984C>G Exon 4 p.Ser123Trp      | 4 (5.13)      | 10 (4.63)| 10 (4.78)| 7 (6.48)        |
| g.20924967T>G Exon 4 p.Ser129Arg      | 3 (3.85)      | 8 (3.70) | 9 (4.30)  | 8 (3.70)        |
| Correlation coefficient                 | 0.710**       | 0.756**  | 0.720**   | 0.782**         |
| p-value                                | 0.0004        | 0.002    | 0.004     | 0.001           |

Pearson correlation coefficient; *p value for ** test; p<0.05 is considered statistically significant; IDC: invasive ductal carcinoma; DCE: Ductal Carcinoma in Site; LCT: Lobular Carcinoma in Situ; ER: Estrogen Receptor; PR: Progesterone Receptor; HER-2/neu: Human Epidermal Growth Factor Receptor 2

Table 4. Association of mutations observed in this study with different clinico-pathological parameters including family history, menopausal age, HER-2/neu and ER/PR status. Frequency of patients with family history was observed significantly higher (p<0.0001) compared-frequency of control individuals.
Germline Variations of Apurinic/Apyrimidinic Endonuclease 1 Detected in Female Breast Cancer Patients

Epidemiologic studies have linked single nucleotide polymorphisms (SNPs) in BER pathway genes—increased human cancer risk including breast cancer which have been evaluated in different populations in past by focusing on three key genes: X-ray repair cross-complementing protein 1 (XRCC1), 8-oxoguanine DNA glycosylase (OGG1) and apurinic/apyrimidinic endonuclease 1 (APEX1) (Hu et al., 2001; Zhang et al., 2006; Lo et al., 2009; Lu et al., 2009; Li et al., 2011; Karahalil et al., 2012; Mahjabeen et al., 2013; Kim et al., 2013; Popanda et al., 2013). APEX1 is a multifunctional protein playing an important role in DNA repair process through AP-endonucleolytic 3’ phosphodiesterase, 3’-5’ exonuclease and 3’ phosphatase activities. APEX1 is also involved in numerous critical cellular responses including tumor occurrence and development, oxidative stress, cell cycle regulation and apoptosis (Zhang and Wang, 2010). It is the only known repair enzyme with self-regulation system (Dyrkheeva et al., 2007). Objective of this study was mutational screening of APEX1 gene and its association with other clinicohistopathological parameters in breast cancer patients.

In this study all exons of APEX1 gene were screened for genomic variation in 530 breast cancer cases and 395 control individuals. A total of fourteen mutations were identified in patients and in some control individuals. This includes seven mutations in 5’UTR region, one in 3’UTR region, two intronic and four missense mutations. Out of these observed mutations 11 were novel and three already reported mutations (rs41561214, rs17112002 and Ser129Arg) by Mahjabeen et al., 2013. Most of the mutations in 5’UTR, 3’UTR and intronic region were significantly higher in breast cancer patients compared to healthy control individuals. Promoter region and 5’ UTR is important for the regulation of translation of transcript (Lodish and Harvey, 2004). Polymorphisms in promoter region can influence interactions between transcriptional factors and their ability—recognize DNA sequences in a promoter region and thus affect gene expression (Luo et al., 2014) so mutations in these regions might be very crucial for cellular functioning. Avery (2007) suggested that variants outside the DNA repair domain such as 5’UTR or promoter regions of APEX1 are informative and may be markers for the disease-causing single nucleotide polymorphisms (SNPs). Various SNPs have been reported in 5’UTR, 3’UTR and promoter regions of APEX1 gene and found-influence the susceptibility and progression of lung cancer (Lo et al., 2009; Lu et al., 2009), bladder cancer (Wang et al., 2010), glioblastoma (Zhou et al., 2011), osteosarcoma (Zhang et al., 2010) and prostate cancer (Kang et al., 2013;}

### Table 4. Association of APEX1 Mutations with Family History and Menopausal Age in Breast Cancer Patients in Present Study

| Mutations          | Patients (%) | OR (95%CI)   | p-Value | Patients (%) | OR (95%CI)   | p-Value |
|--------------------|--------------|--------------|---------|--------------|--------------|---------|
| g.20923559G>A 5'UTR | 9 (9.9)      | 4.10 (1.44-11.76) | 0.008   | 4 (4.4)      | 1.47 (0.14-14.49) | 1.00    |
| g.20923481G>A 5'UTR | 6 (6.6)      | 2.54 (0.43-14.99) | 0.39    | 16 (10.32)   | 4.01 (1.21-13.33) | 0.02    |
| g.20923529C>T 5'UTR | 2 (2.2)      | 1.47 (0.14-14.49) | 1.00    | 06 (5.2)     | 3.22 (0.72-14.46) | 0.24    |
| g.20923416C>T 5'UTR (rs41561214) | 4 (4.4) | 2.54 (0.43-14.99) | 0.39    | 10 (6.45)    | 4.01 (1.21-13.33) | 0.02    |
| g.20925669A>T 3'UTR (rs17112002) | 12 (12.8) | 4.26 (1.41-12.88) | 0.009   | 08 (5.16)    | 3.68 (0.98-13.86) | 0.08    |
| g.20924881G>A Exon 4 p.Glu101Lys | 12 (12.8) | 4.26 (1.41-12.88) | 0.009   | 08 (5.16)    | 3.68 (0.98-13.86) | 0.08    |
| g.20924941G>C Exon 4 p.Ala121Pro | 0 (0.0)      | 1.47 (0.14-14.49) | 1.00    | 02 (1.3)     | 1.47 (0.14-14.49) | 1.00    |
| g.20924948C>G Exon 4 p.Ser123Trp | 04 (2.6)     | 2.54 (0.43-14.99) | 0.4     | 0 (0.0)      | 1.47 (0.14-14.49) | 1.00    |
| g.20924967T>G Exon 4 p.Ser129Arg | 09 (9.6)     | 6.80 (0.86-53.94) | 0.05    | 06 (5.2)     | 4.51 (0.54-57.62) | 0.01    |

Discussion

Almpaired base-excision repair (BER) function can give rise to the accumulation of DNA damage and initiation of cancer. Epidemiologic studies have linked single nucleotide polymorphisms (SNPs) in BER pathway genes—increased human cancer risk including breast cancer which have been evaluated in different populations in past by focusing on three key genes: X-ray repair cross-complementing protein 1 (XRCC1), 8-oxoguanine DNA glycosylase (OGG1) and apurinic/apyrimidinic endonuclease 1 (APEX1) (Hu et al., 2001; Zhang et al., 2006; Lo et al., 2009; Lu et al., 2009; Li et al., 2011; Karahalil et al., 2012; Mahjabeen et al., 2013; Kim et al., 2013; Popanda et al., 2013). APEX1 is a multifunctional protein playing an important role in DNA repair process through AP-endonucleolytic 3’ phosphodiesterase, 3’-5’ exonuclease and 3’ phosphatase activities. APEX1 is also involved in numerous critical cellular responses including tumor occurrence and development, oxidative stress, cell cycle regulation and apoptosis (Zhang and Wang, 2010). It is the only known repair enzyme with self-regulation system (Dyrkheeva et al., 2007). Objective of this study was mutational screening of APEX1 gene and its association with other clinicohistopathological parameters in breast cancer patients.

In this study all exons of APEX1 gene were screened for genomic variation in 530 breast cancer cases and 395 control individuals. A total of fourteen mutations were identified in patients and in some control individuals. This includes seven mutations in 5’UTR region, one in 3’UTR region, two intronic and four missense mutations. Out of these observed mutations 11 were novel and three already reported mutations (rs41561214, rs17112002 and Ser129Arg) by Mahjabeen et al., 2013. Most of the mutations in 5’UTR, 3’UTR and intronic region were significantly higher in breast cancer patients compared to healthy control individuals. Promoter region and 5’ UTR is important for the regulation of translation of transcript (Lodish and Harvey, 2004). Polymorphisms in promoter region can influence interactions between transcriptional factors and their ability—recognize DNA sequences in a promoter region and thus affect gene expression (Luo et al., 2014) so mutations in these regions might be very crucial for cellular functioning. Avery (2007) suggested that variants outside the DNA repair domain such as 5’UTR or promoter regions of APEX1 are informative and may be markers for the disease-causing single nucleotide polymorphisms (SNPs). Various SNPs have been reported in 5’UTR, 3’UTR and promoter regions of APEX1 gene and found—influence the susceptibility and progression of lung cancer (Lo et al., 2009; Lu et al., 2009), bladder cancer (Wang et al., 2010), glioblastoma (Zhou et al., 2011), breast cancer (Kang et al., 2013;
In addition to these mutations we observed four missense mutations (Glu101Lys, Ala121Pro, Ser123Trp, Ser129Arg) in exon 4 which cover both the redox domain and DNA-repairing domain of APEX1. Glu101Lys at residue 101 is in the redox domain while, Ala121Pro, Ser123Trp, Ser129Arg are found in DNA-repairing domain of APEX1. APEX1 protein has a spatial configuration and mainly constituted by two overlapping functional domains at N and C terminals. The Nterminal domain is considered be responsible for the redox function and modify the DNA binding capacity of multiple transcriptional factors and is involved in regulating gene expression (Wong et al., 2003; Dyrkheeva et al., 2007; Zhang and Wang, 2010). C-terminal domain performs DNA repair functions and protects cells from the cytotoxicity caused by the continuously accumulating exogenous and endogenous AP site mutation (Zhang and Wang, 2010). Structural prediction of Ser129Arg variant in APEX1 by Mahjabeen et al. (2013), has shown no major change in protein structure compared with wild-type protein but APEX1-compromised cells are more susceptible to oxidative stress mainly due to reduced redox and 3’ phosphodiesterase activity impacting cell survival, pushing it towards apoptosis (Fishe1 et al., 2007).

Presence of identified missense mutations in both Redox and Repair domains signify that how repair process of DNA might be affected due to inhibition of Redox and repair activity of APEX protein. Missense mutations observed in this study were also analysed via Alamut biosoftware to check the conservation level of mutated nucleotides and encoded amino acids. It was predicted that observed missense mutations especially of highly conserved nucleotides and amino acids have shown some deleterious, potentially disease causing effects resulting physicochemical alterations in structure of amino acids. In-silico predictions about mutations using PolyPhen-2 (Adzhubei et al., 2010), SIFT (Kumar et al., 2009) and Mutation Taster (Schwarz et al., 2010) softwares have previously been considered an important tool in exploration of possible effects of mutations.

APEX1 mutations observed in this study were significantly correlated with different clinicopathological parameters. Mutation frequency was found higher in patients with invasive ductal carcinoma, tumors grade II, family history of cancer, early menopause and negative ER, PR, HER-2/neu status which have been previously reported-contribute in breast cancer development in Pakistani populations (Naeem et al., 2008; Azizun-Nisa et al., 2008; Sharif et al., 2010; Ahmed et al., 2011) and worldwide (Dumitrescu et al., 2005; Ahmed et al., 2011; Popanda et al., 2013). Woman’s breast cancer risk is two or more times greater if she has a family history of cancer (Ahmed et al., 2011). Estrogen and progestrone receptors (ER, PR) and HER-2/neu greatly influence the management of the malignancy (Rampaul et al., 2001) and generally has been found-be lower in Asian countries than those in the Africa, western world, USA and Australia (Ahmed et al., 2007; Hedayati-Moghadam 2008; Ahmed et al., 2011). Relation of menopausal age with breast cancer has been reported inconsistently (Schultz et al., 2005; Shakeel et al., 2013) where we cannot ignore the effects of adjuvant treatment of breast cancer on their menopausal age (Rosenberg and Partridge, 2013).

In conclusion, our study demonstrates a significant association of germ line APEX1 mutations with breast cancer in Pakistani population. The data suggests that a combination of 5’UTR mutations and missense mutations, or any other mutation linked-it in the same gene or gene in vicinity, could conceivably play a role in the process of breast carcinogenesis in Pakistani population. Moreover in line with previous findings, inhibition of DNA repair or redox or both activities of APEX1 due to mutations or any inhibitor could potentially sensitize the tumour cells to therapeutic agents, making APEX1 an attractive molecular target in the treatment of cancer. These molecular and epidemiological findings provide evidence that APEX1, a DNA repairing gene could prove to be a good candidate of better diagnosis, treatment and prevention of breast cancer but more detailed studies are needed for a clearer picture of APEX1 gene and protein in pathogenesis of breast cancer.

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