Prediction and Analysis of Breast Cancer Related Deleterious Non-Synonymous Single Nucleotide Polymorphisms in the PTEN Gene

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Abstract

One of the most common cancer types faced by the women around the world is breast cancer. Among the several low, moderate and high penetrance genes conferring susceptibility to breast cancer, PTEN is one which is known to be mutated in many tumor types. In this study, we predicted and analyzed the impact of three deleterious coding non-synonymous single nucleotide polymorphisms rs121909218 (G129E), rs121909229 (R130Q) and rs57374291 (D107N) in the PTEN gene on the phenotype of breast tumors using computational tools SIFT, Polyphen-2, PROVEAN, MUPro, POPMusic and the GETAREA server.

Keywords: Breast cancer - PTEN - nSNPs - genetic factors - computational tools

Asian Pac J Cancer Prev, 17 (4), 2199-2203

Introduction

Breast cancer is one of the common cancer types faced by the women in the modern life. In countries like United States, one in 8 women develops breast cancer in her lifetime and its incidence rates is slightly increasing among African American women (De Santis et al., 2014). Along with colorectal cancer, it ranks high in all countries. Its incidence in the United States and Europe is twice as high as it is in Asian countries, and its incidence rates have been increasing in all countries (Saika and Sobue, 2013). Tumor heterogeneity, lifestyle factors including obesity, breastfeeding, and alcohol consumption are some of the traditional risk factors associated with breast cancer (Kwan et al., 2009). Some of the genetic and hormonal factors constitute risk to breast cancer (Martin and Weber, 2000).

Phosphatase and tensin homolog deleted on chromosome ten (*PTEN*) is one of the frequent mutated gene found in many primary and metastatic malignancies including breast cancer (Kechagioglou et al., 2014). Human estrogen receptor-positive (ER+) breast cancer cell lines containing inducible *PTEN* short hairpin RNAs, result in the hyperactivation of the PI3K pathway and a concomitant change in gene expression similar to luminal B breast cancer types (Maggi and Weber, 2015). In breast cancer cells, reduced expression of *PTEN* is known to confer susceptibility to inhibitors of the PI3 kinase/Akt pathway (De Graffenried et al., 2004). Triple-negative breast cancers are aggressive forms, CIB1 plays a broad role in its cell survival, tumor growth and a low expression of *PTEN* is a key predictor of sensitivity to CIB1 depletion

(Black et al., 2015). So, treatment with trastuzumab to improve the disease-free and overall survival has been a standard approach for HER2-overexpressing breast cancer patients and *PTEN* status was suggested to be one of the indicators (Adamczyk et al., 2015).

Variations on the promoter region of PTEN are known to affect the progression of breast cancer and the survival of the patients (Heikkinen et al., 2011). Previously several single nucleotide polymorphisms such as rs1234212, rs11202586, rs1234221, rs1903860, rs1234220,rs1234219,rs1903858,rs2299939,rs1234224, rs1234223, rs1234213, rs2673832 with a *PTEN* haplotype associated with breast cancer risk were predicted (Haiman et al., 2006). Previous study showed that a high level of discordance in PTEN level, PIK3CA mutations and receptor status between primary tumors and metastases influenced the patient selection and response to PI3Ktargeted therapies (Gonzalez-Angulo et al., 2011). An in vivo study showed that non catalytic PTEN missense mutation predisposes the organ-selective cancer development (Caserta et al., 2015). In the present study, we aim to predict the breast cancer-associated nSNPs in PTEN and to further to analyze

Materials and Methods

SNP datasets used for the study

SNP datasets for *PTEN* were retrieved from the dbSNP database (http://www.ncbi.nlm.nih.gov/projects/SNP/, Build 138; access date: August 22, 2015) (Sherry et al., 2001) for our study.

Prediction of Deleterious SNPs

SIFT and Polyphen-2 database servers to screen out the deleterious coding nSNPs from other SNPs of *PTEN*. 'Sorting Tolerant From Intolerant' (SIFT) (http://sift.jcvi.org/) uses a sequence homology based approach for predicting the amino acid substitution in a protein affecting the protein function (Kumar et al., 2009). It assigns 0-0.05 score for intolerant or deleterious amino

acid substitutions and 0.05–1 scores for tolerant or neutral amino acid substitutions (Ng and Henikoff, 2003; Ng and Henikoff, 2006). PolyPhen-2 (http://genetics.bwh.harvard. edu/pph2/) on the other hand predicts the functional significance of variation using Naïve Bayes classifier. We used WHESS.db a quick access for precomputed set in PolyPhen-2 predictions was used for our analysis (Adzhubei et al., 2010). We submitted our query in the

Table 1. Functionally significant SNPs Predicted Using the SIFT Server

		Nucleotide change	Protein ID	Tolerance Index		
dbSNPID	Amino acid change			Using orthologues in the Protein alignment	Using homologues in the Protein alignment	
rs121909218	G129E	A/G	NP_000305	0	0	
rs121909221	S170R	A/T	NP_000305	0	0	
rs121909222	H123R	A/G	NP_000305	0	0	
rs121909223	C124R	C/G/T	NP_000305	0	0	
rs121909225	M35R	G/T	NP_000305	0	0	
rs121909226	L70P	C/T	NP_000305	0	0	
rs121909229	R130Q	A/C/G	NP_000305	0	0	
rs121909230	L112P	C/T	NP_000305	0	0	
rs121909233	D19N	A/G	NP_000305	0.18	0.04	
rs121909235	R234Q	A/G	NP_000305	0.31	0.05	
rs121909236	H61D	C/G	NP_000305	0	0	
rs121909237	A121G	C/G	NP_000305	0.08	0	
rs121909238	H93R	A/G	NP_000305	0	0	
rs121909239	D252G	A/G	NP_000305	0.01	0	
rs121909241	G132V	A/G/T	NP_000305	0	0	
rs57374291	D107N	A/G	NP_000305	0.05	0	
rs121913293	R173C	C/T	NP_000305	0	0	
rs121913294	R173H	A/C/G	NP_000305	0	0	

Table 2. Functionally Significant SNPs Predicted Using the Polyphen-2 Server

Nucleotide change	Amino acid change	Protein ID	HDivPred	HDiv Prob	HVarPred	HVarProb
G/A	G129E	P60484	Probably damaging	1	Probably damaging	1
T / A	S170R	P60484	Probably damaging	1	Probably damaging	0.999
A/G	H123R	P60484	Probably damaging	1	Probably damaging	0.998
T / C	C124R	P60484	Probably damaging	1	Probably damaging	0.999
T/G	M35R	P60484	Probably damaging	0.996	Probably damaging	0.974
T / C	L70P	P60484	Probably damaging	1	Probably damaging	1
G/A	R130Q	P60484	Probably damaging	1	Probably damaging	0.998
T / C	L112P	P60484	Probably damaging	1	Probably damaging	1
G/A	D19N	P60484	Probably damaging	0.988	Probably damaging	0.815
G/A	R234Q	P60484	Probably damaging	0.98	possibly damaging	0.617
C/G	H61D	P60484	Probably damaging	1	Probably damaging	0.998
C/G	A121G	P60484	probably damaging	0.999	probably damaging	0.968
A/G	H93R	P60484	probably damaging	1	probably damaging	0.998
A/G	D252G	P60484	probably damaging	0.989	possibly damaging	0.862
G/T	G132V	P60484	Probably damaging	1	Probably damaging	1
G/A	D107N	P60484	probably damaging	1	probably damaging	0.999
C/T	R173C	P60484	probably damaging	1	probably damaging	0.966
G/A	R173H	P60484	probably damaging	1	probably damaging	0.966

Table 3. Phenotype of the Predicted Deleterious SNPs

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dbSNPID	Amino acid change	Phenotype	References
rs121909218	G129E	Cowden disease, Breast cancer	(Liaw et al., 1997)
rs121909221	S170R	Bannayan- riley-ruvalcaba syndrome	(Marsh et al., 1997)
rs121909222	H123R	Cowden disease	(Nelen et al., 1997)
rs121909223	C124R	Cowden disease	(Nelen et al., 1997)
rs121909225	M35R	Cowden disease	(Olschwang et al., 1998)
rs121909226	L70P	Cowden disease	(Marsh et al., 1998)
rs121909229	R130Q	Cowden disease, Breast cancer	(Kurose et al., 1999; Baig et al., 2011)
rs121909230	L112P	Lhermitte-duclos disease	(Sutphen et al., 1999)
rs121909233	D19N	Malignant melanoma	(Celebi et al., 2000)
rs121909235	R234Q	Glioma	(Staal et al., 2002)
rs121909236	H61D	Macracephaly	(Reardon et al., 2001)
rs121909237	A121G	Squamous cell carcinoma	(Poetsch et al., 2002)
rs121909238	H93R	Autism	(Butler et al., 2005)
rs121909239	D252G	Autism	(Butler et al., 2005; Nagy et al., 2014)
rs121909241	G132V	Hamartoma tumor syndrome	(Tekin et al., 2006)
rs57374291	D107N	Breast cancer	(Baig et al., 2011)
rs121913293	R173C	Hereditary cancer- predisposing syndrome	Clinvar
rs121913294	R173H	Hereditary cancer- predisposing syndrome	Clinvar

Table 4. Impact of the SNPs Predicted Using the PROVEAN Server

dbSNPID	Amino acid change	PROVEAN score	Prediction (cutoff= -2.5)
rs121909218	G129E	-7.772	Deleterious
rs121909229	R130Q	-3.858	Deleterious
rs57374291	D107N	-4.675	Deleterious

form of dbSNP id for both SIFT and Polyphen-2.

Phenotype of predicted deleterious coding nSNPs
Search for phenotype information of the breast cancer

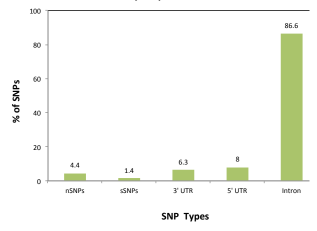


Figure 1. Distribution of *PTEN* **SNPs.** Coding nonsynonymous SNPs (nSNPs), coding synonymous SNPs (sSNPs), 3' UTR SNPs, 5' UTR SNPs and intronic SNPs

SNPs was performed using the databases SNPedia (Cariaso and Lennon, 2012), (Schaefer et al., 2012), Hapmap (International HapMap et al., 2010), Pubmed (Sood and Ghosh, 2006). The SNPs with breast cancer phenotype were further cross checked for deleterious nature using the PROVEAN software (Choi et al., 2012).

Modeling nSNPs locations in protein structure

Crystal structure of *PTEN* downloaded from the protein databank (PDB ID: 1d5r, chain A) (Lee et al., 1999) was used for modelling the nSNPs in the protein structure. All water molecules and the TLA ligand were removed from the crystal structure and the mutants (MTs) G129E, R130Q and D107N were created by replacing the wild-type (WT) protein residue with its polymorphic residue using PyMOL (PyMol, 2006). Mutants were optimized and energy minimized using Nomad-Ref server (Lindahl et al., 2006) with conjugate gradient method.

Analysis of the impact of mutant on the PTEN protein product

The effect of amino acid changes on the stability of *PTEN* protein was analyzed by using the MUpro (http://mupro.proteomics.ics.uci.edu/) (Cheng et al., 2006) web servers. Thermodynamic stability of the mutants was analyzed using POPMusic server (Dehouck et al., 2009). The solvent accessibility information of the WT and MTs was analyzed using GETAREA server (http://curie.utmb.edu/getarea.html) (Fraczkiewicz and Braun, 1998) which considers the residues which exceeds the ratio value 50% to be solvent and buried if it is less than 20% marked as "o" and "i" respectively.

Results and Discussion

nSNPs from dbSNP database

A search for total SNPs in *PTEN* against dbSNP database resulted in a total of 18242 SNPs, out of which 5597 were found to be Human (active) SNPs (i.e., Active Human RS and not including those that have been merged). Among the 5597 Human (active) SNPs, 247 were coding non-synonymous SNPs (nSNPs), 83 were coding synonymous, 358 SNPs occurred in the mRNA 3'

Table 5. Stability Changes of the Mutants Using Mupro Server

	Amino acid .	Stability Change			
dbSNPID	change	ΔG	Support Vector Machine	Neural Network	
rs121909218	G129E	Increase stability	0.35155369 (Increase stability)	0.5401244261305952 (Increase stability)	
rs121909229	R130Q	Increase stability	-0.81491724 (Decrease stability)	-0.998634501676394 (Decrease stability)	
rs57374291	D107N	Increase stability	-0.92320236 (Decrease stability)	-0.997148565146761 (Decrease stability)	

Table 6. Thermodynamic Stability of the Mutants Using PopMusic Server

dbSNPID	Amino acid change	solvent accessibility	Folding free energy (kcal/mol)
rs121909218	G129E	10.18	0.43
rs121909229	R130Q	2.9	1.06
rs57374291	D107N	21.05	0.69

Table 7. Total Area/Energy Tability of the Mutants Using GetArea Server

dbSNPID	Amino acid change	Total area/energy
rs121909218	G129E	15463.89
rs121909229	R130Q	15517.79
rs57374291	D107N	15491.66

UTR, 451 occurred in the mRNA 5' UTR and 4851 were occurred in intronic regions. It can be seen from the Fig. 1 that the vast majority of SNPs occur in the intronic region (86.6%) and more SNPs are nSNPs (4.4%) compared to synonymous SNPs (1.4%), SNPs occurring in the mRNA 3' UTR (6.3%) and 5' UTR (8%) regions. We selected coding nSNPs for our investigation.

Deleterious nSNPs in PTEN gene

Among the 247 coding nSNPs from dbSNP, 18 were found to be deleterious with a tolerance index score of less than or equal to 0.05 using SIFT server. Among these 18 deleterious SNPs, 13 had a highly deleterious tolerance index score of 0.00 using orthologues and homologues in the protein alignment and the remaining 5 deleterious nSNPs had a tolerance index score had a tolerance index score of 0.04, 0.05, 0.08, 0.01 and 0.05 using orthologues and homologues in the protein alignment respectively (Table 1). Among 18 nSNPs predicted to be deleterious using SIFT server, seven nSNPs showed a nucleotide change of A/G, three showed a change of C/T, two showed a change of C/G, one showed a change of A/T, one showed a change of G/T, two showed a change of A/C/G, one showed a change of C/G/T and one showed a change of A/G/T respectively. A/G and C/T changes occurred maximum number of times compared to the other nucleotide changes.

18 nSNPs that are predicted to be deleterious using SIFT server were submitted to Polyphen-2 to predict their respective functional significance of allele replacement. All the 18 nSNPs submitted to the Polyphen-2, were found

to be possibly damaging, or probably damaging by both HumDiv and HumVar predictions (Table 2).

Phenotype prediction of deleterious nSNPs

18 nSNPs that are predicted to be deleterious or probably damaging using SIFT and Polyphen-2 was subjected to phenotype prediction. Results showed that among the 18 nSNPs, three SNPs rs121909218 (G129E), rs121909229 (R130Q) and rs57374291 (D107N) showed a phenotype in breast tumors (Table 3). Results from PROVEAN server also showed that these SNPs as deleterious (Table 4) these SNPs were considered for further analysis.

Deleterious nSNPs impact on PTEN protein

To analyze the impact of deleterious nSNPs on the *PTEN* protein product, we have analyzed the stability of each mutant. Stability analysis of the mutants using the MUpro server showed that the mutants R130Q and D107N showed a decrease in the stability whereas the mutant G129E showed a increase in the stability (Table 5). Their respective change in the solvent accessibility and the free energy were provided in the Table 6 given below. Results from the Total area/energy of each mutant showed the mutants G129E and D107N major change compared to the wild type *PTEN* protein Total area/energy (15517.79) (Table 7).

In conclusion, the results from our study indicate that three mutations R130Q, D107N and G129E in *PTEN* are associated with the breast cancer phenotype. Results showed that these three mutants showed a change in stability. Overall, the present computational approach reported in this study allowed elucidation of the role of deleterious mutations in *PTEN* thereby providing useful information for the design of *PTEN* mutant-based therapeutic strategies against breast cancer.

Acknowledgements

The present study was supported under UGC Research Award 2014-16 funded by UGC, New Delhi (No. F. 301/2014(SAII)).

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