# In-silico study to identify the pathogenic single nucleotide polymorphisms in the coding region of *CDKN2A* gene

Farzaneh Ghasemi<sup>1</sup>, Mehri Khatami<sup>1,\*</sup>, Mohammad Mehdi Heidari<sup>1</sup>, Reyhane Chamani<sup>1</sup>, Hadi Zare-Zardini<sup>2,3</sup>

- 1. Department of Biology, Faculty of Science, Yazd University, Yazd, Iran
- 2. Hematology and Oncology Research Center, Shahid Sadoughi University of Medical Sciences, Yazd, Iran
- 3. Department of Sciences, Farhangian University, Isfahan, Iran

\*Corresponding author: Dr Mehri Khatami, Department of Biology, Faculty of Science, Yazd University, Pajoohesh Street, Yazd, Iran. Email: m.khatami@yazd.ac.ir. ORCID ID: 0000-0002-5840-5399

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#### Abstract

**Background:** *CDKN2A*, encoding two important tumor suppressor proteins p16 and p14, is a tumor suppressor gene. Mutations in this gene and subsequently the defect in p16 and p14 proteins lead to the downregulation of RB1/p53 and cancer malignancy. To identify the structural and functional effects of mutations, various powerful bioinformatics tools are available. The aim of this study is the identification of high-risk non-synonymous single nucleotide variants in the *CDKN2A* gene via bioinformatics tools.

**Materials and Methods:** Among the identified polymorphisms in this gene, 353 missense variants are retrieved from the national center for biotechnology information/single nucleotide polymorphism database (NCBI/dbSNP). Then, the pathogenicity of missense variants are considered using different bioinformatics tools. The stability of these mutant proteins, conservation of amino acids and the secondary and tertiary structural changes are analyzed by bioinformatics tools. After the identification of high-risk mutations, the changes in the hydrophobicity of high-risk amino acid substitutions are considered.

**Results:** Deleterious single nucleotide polymorphisms (SNPs) were screened step by step using the bioinformatics tools. The results obtained from the set of bioinformatics tools identify high-risk mutations in *CDKN2A* gene.

Conclusion: 18 high-risk mutations including L16R/Q, G23D/R/S, L32P, N42K, G55D, G67D/R, P81R, H83R, G89D/S, A102E, G101R, G122R, and V126D were identified. According to the previous experimental studies, the association of L16R, G23D/R/S, L32P, G67R, H83R, G89D, G101R, and V126D amino acid substitutions with various cancers has been confirmed.

**Keywords:** Computational biology, *CDKN2A*, Gene, SNP, Tumor suppressor protein.

### Introduction

CDKN2A, localized on 9p21.3 and encoding different tumor suppressor proteins by alternate reading frame mechanism, is an important cell cycle regulator (1). The tumor suppressor p16 is a low molecular weight protein that contains 156 amino acids including four ankyrin (Ank) repeats (Ank1: codons 11-40; Ank2: codons 44-72; Ank3: codons 77-106; Ank4: codons 110-139) (2). The p16 binds to a cyclin-dependent kinase (CDK) 4/6 and inhibits the CDK4/6 and cell cycle at the G1 phase (3, 4). CDK4/6cyclin D and CDK2-cyclin E complexes inactivate RB1 by phosphorylation mechanism. So, E2F1 inhibited by RB1 is

released and leads to cell growth and division. Therefore, p16 activates RB1 by inhibition of CDK4/6 (5). The tumor suppressor p14 containing 132 amino acids activate p53 through binding to MDM2. MDM2 as an antagonist of p53 protein downregulates p53 through ubiquitination (6). Downregulation or inactivation of the mentioned tumor suppressor proteins has been considered in several cancers such as colon cancer (7), lung cancer (8), melanoma (9), pancreatic cancer (10), head and neck squamous cell carcinoma (HNSCC) (11), glioma (12), (13-15).and leukemia Different mechanisms such promoter hypermethylation (7), sequence deletion (16), and point mutation (4) lead to the decreased expression and dysfunction of p16 and p14.

The single nucleotide variants (SNVs) are responsible for about 90% of human variability (17). Also, due to the existence of SNVs, there exist alterations in protein structure and subsequently its function. The size, charge, and hydrophobicity value in amino acids are unique traits. The original wild type and mutant residues often differ in these properties (18). Therefore, the amino acid replacement may impair the function, structure, and stability of protein, and ultimately proteinprotein interaction. Eventually, loss of function and modified function (or deficiency) of tumor suppressor proteins lead to abnormal proliferation and overgrowth. But, some SNVs are benign polymorphism, and these variants have no effects on protein function (19). The experimental methods, as the most reliable approaches, are expensive and timeconsuming processes. Also, in some cases, the methods of mutagenesis and extraction of mutant protein are impossible in vitro and in vivo (20). So, bioinformatics tools can help researchers to predict the effects of mutations. But, there is no single tool for this purpose and to get reliable results, various tools can be used. There are powerful bioinformatics tools to evaluate the alterations of protein structure and function. Using these tools, the distance of atoms (20-22) and pathogenesis of mutations (23-28) as well as changes in protein structure and polar contacts (22) can be predicted. Also, the stability of mutant proteins can be investigated by the evaluation of the total energy of proteins The determination of protein structure in the presence of the mutation is the most important challenge in biology. This study included a comprehensive investigation to identify the pathogen p16 protein nsSNVs bioinformatics tools. Also, the high-risk missense variants were introduced via an in-silico study. To confirm the obtained

results, the changes in structure and hydrophobicity of mutant amino acids were compared with native residues. The conservation of the native residue was also considered. This computational approach can be used as a prelude to planning a targeted molecular method to prove the obtained results from the bioinformatics study.

This paper organizes as follows: In the next section, the used methods have been introduced to consider the deleterious mutations. The related results obtained from the bioinformatics tools were collected in section 3. Also, the correlation of the high-risk mutations with various cancers was gathered in section 4. Finally, the paper was ended with conclusions in section 5.

## **Materials and Methods Materials**

### Data collection

The identified nsSNVs in *CDKN2A* gene were retrieved from the national center for biotechnology information/single nucleotide polymorphism database (NCBI/dbSNP)

(https://www.ncbi.nlm.nih.gov/snp/). Also, the structure of p16 (PDB ID: 1DC2) was retrieved from the protein databank (PDB), (https://www.rcsb.org/).

### Amino acid substitution effects Sorting Intolerant From Tolerant (SIFT)

Using SIFT analyzer (https://sift.bii.a-star.edu.sg/), it is possible to predict the effects of amino acid substitution on protein function through considering the sequence homology and the physical properties of amino acids (29). The mutation is introduced as an affecting protein function if the score evaluated by SIFT is lower than 0.05. 'Seq Rep' is a fraction of sequences that contain one of the basic amino acids. The low fraction indicates insufficient information in this position. The presence of severely gapped or unalignable in this position can cause low confidence prediction.

#### **Polymorphism Phenotyping** v2(PolyPhen-2)

PolyPhen-2

(http://genetics.bwh.harvard.edu/pph2/), as a web-based tool, predicts the effect of an amino acid substitution on the structure and function of the protein, based on the multiple sequence alignment of the 3D protein structure. The score of positionspecific independent count (PSIC) from 0 to 1 can be calculated using this tool. PolyPhen-2 reports the result as a benign (with 0-0.15 score), possibly damaging (with 0.15-0.85 score), and probably damaging (with 0.85-1 score) (30). This tool provides two values of "sensitivity" and "specificity" for Confidence predictions.

### **Protein Variation Effect ANalyzer** (PROVEAN)

PROVEAN (http://provean.jcvi.org/index. php) web server is used to analyze protein variants via an alignment-based score approach. This online tool predicts the effect of missense variants and indel on the protein function (31). If the calculated score of the amino acid substitution is lower than -2.5, this mutation deleterious.

### Predicting Human Deleterious SNPs in the human genome (PHD-SNPg)

PHD-SNPg (https://snps.biofold.org/phdsnpg/) is a machine learning method that depends on sequence-based features. This tool considers the impact of SNVs in the coding and non-coding regions. The SNV is identified as a pathogenic or benign mutation. A probabilistic score is from 0 to 1. If the score is >0.5, the variants are predicted to be pathogenic mutations (32).

### SNPs&GO

SNPs&GO (http://snps.biofold.org/snpsand-go/) is a web server tool that predicts the effect of amino acid substitution as a disease-associated variation or neutral variation effects based on protein structure, and sequence (33). The evaluated score is from 0 to 1 that SNV with a score >0.5 is identified as a disease association variation.

### Protein Analysis through Evolutionary **Relationships (PANTHER)**

**PANTHER** The (http://www.pantherdb.org/) classification system is a genomic analysis system based on gene function, ontology, pathways, and statistical analysis tools (34).

### I-Mutant2.0

(http://folding.biofold.org/i-I-Mutant2.0 mutant/i-mutant2.0.html) is a support vector machine (SVM) that predicts protein stability changes due to the single point mutations based on protein structure sequence. Using empirical thermodynamic data. I-Mutant2.0 calculates the free energy changes of the protein, i.e., delta delta G (DDG) (35). Accordingly, protein stability is decreased or increased if DDG is lower or upper than 0, respectively.

### Structural consideration NetSurfP-2.0

NetSurfP-2.0

(http://www.cbs.dtu.dk/services/NetSurfP/ ), as a sequence-based tool, predicts the secondary structure, surface accessibility, structural disorder, and backbone dihedral angles (Phi and Psi angels) for each residue of the protein sequence. This tool predicts 2-class relative solvent accessibility (RSA) for an amino acid (buried or exposed) with a threshold of 25%. Also, absolute solvent accessibility (ASA) output is calculated by multiplying RSA and ASA<sup>max</sup> (36).

### PvMOL software

PyMOL written in Python is a molecular visualization system for the evaluation of structural biology. This software is used to create mutations in p16 and consider the polar contacts and hydrogen bond (Hbond) length in the native and mutant proteins.

#### Conservation **Surface** mapping (ConSurf)

ConSurf web server (https://consurf.tau.ac.il/) evaluates the conservation level of the amino acids, based on the evolutionary relations between the protein and its homologs (37).

ConSurf maps the 3D structure of the protein with a color scale extending from 1 to 9 that 1 (9) is related to a hypervariable (highly conserved) amino acid (38).

### **Project HOPE**

HOPE (https://www3.cmbi.umcn.nl/hope/) is a web service that analyzes the structural effects of the point mutation in a protein sequence by combining the available information obtained from a series of web services and databases (39).

### Analyzing hydrophobicity changes

The hydrophobic changes are analyzed using the web-based PEPTIDE (https://www.peptide2.com/N peptide hy drophobicity hydrophilicity.php) and ExPASy/ProtScale with Kyte & Doolittle acid amino scale (https://web.expasy.org/protscale/). Based on the chemical and physical properties of amino acids, ExPASy/ProtScale predicts the hydrophobic or hydrophilic scale of the protein structure parameters (40).

### Results

### **Data collection**

The 8405 SNPs (single nucleotide polymorphisms) have been identified in the *CDKN2A* gene but, there were 465 SNPs in the coding region of this gene. From 465 missense variants, the 353 nsSNVs have been identified in p16 transcript and these 353 nsSNVs were selected for bioinformatics analysis.

### Amino acid substitution effects

Using SIFT, out of 353 amino acid substitutions, the 143 mutations were predicted as an "affected protein function", and 210 amino acid substitutions were predicted as a "tolerated substitution". The SIFT 'Seq Rep' score for all nsSNPs except 2 positions was 1.00, but for 2 positions including 125, 126, this score was 0.94. These high Seq Rep scores indicated high confidence prediction. Then, all of the 143 acid damaging amino substitutions identified by SIFT were screened by PolyPhen-2. According to the results of PolyPhen-2, 119 the amino acid

substitutions were predicted "damaging" mutations. The "sensitivity" and "specificity" obtained from this tool for all of the predictions have been gathered in Table I. One can see that the evaluated sensitivity for the majority of predictions was very low, so these results had low confidence. But, this problem was solved using the combination of various bioinformatics tools. Using the PROVEAN analyzer, it was found that 19 amino acid substitutions had neutral effects, but 100 amino acid substitutes had deleterious effects. This tool used 166 related sequences classified in 30 clusters as the supporting sequence set for predictions. In the next step, the stability of 100 mutant proteins was studied via the I-Mutant2.0 tool. I-Mutant2.0 calculated the DDG value of 100 amino acid substitutions, also this tool predicted a decrease in the stability of 79 mutations (see Table I).

Also, the deleterious or neutral effects of the amino acid substitutions shown in Table II were considered by SNPs&GO, PANTHER, and PHD-SNPg. Accordingly, the 48 amino acid substitutions produced damaging effects on protein function.

### Structural consideration

The secondary structure, RSA, ASA, and class assignment of the 48 damaging amino acid replacements were studied via NetSurfP-2.0 web service. Also, the conservation of residues was considered by ConSurf. Out of the 48 amino acid substitutions. the amino 18 substitutions (L16R/Q, G23D/R/S, L32P, N42K, G55D, G67D/R, P81R, H83R, G89D/S, A102E, G101R, G122R, and V126D) displayed a huge increase in RSA and also these amino acid substitutions highly conserved. So. were mutations were identified as high-risk mutations. It is necessary to mention that the mutant protein stability was decreased when the solvent accessibility increased (41). As shown in Table III, the class assignment, RSA, ASA, and the secondary structure of the native and

mutant amino acids were considered. According to the NetSurfP-2.0 results, Asn at position 42 and Gly at positions 23, 55, 67, 89, 101, and 122 were located on the surface of the protein. But, Leu at positions 16 and 32, Pro at position 81, His at position 83, Ala at position 102, and Val at position 126 are buried in the core of the protein.

The structural alterations of the 18 amino acid substitutions were considered using the HOPE web service. According to the HOPE results, in the L16R/Q, G23D/R/S, N42K, G55D, G67D/R, P81R, H83R, G89D/S, G101R, A102E, G122R, and V126D amino acid substitutions, the mutant residues were bulkier or larger than the wild type residues. The helix structure might be unstable in the L16R due to the placement of two large amino acids next to each other (amino acids sequence at positions 15 and 16: WL to WR). About L32P, substituted Pro was localized in the  $\alpha$ -helix structure and disrupted  $\alpha$ -helix due to the missing H-bond.

In L16R, G23D/R, N42K, G55D, G67D/R, P81R, H83R, G89D, G101R, A102E, G122R, and V126D, the wild type residues were neutral, but the mutant residues were charged (either positive or negative). In L16R, P81R, H83R, A102E, and V126D, the mutant residues introduced a charge in buried residues leading to probable defects in protein folding.

Gly at positions 23, 55, 67, 89, 101, and 122 was wild type residue. Among the amino acids, Gly was the simplest and more flexible one that played an important role in the secondary structures specially β-turns. So, the mutation of this residue might lead to protein dysfunction or decreased protein stability (42). According to the ConSurf server, the wild type residues at positions 16, 32, 101, and 102 were completely conserved. So, the alteration of these positions was probably damaging to the protein function (37, 43).

### **Evaluation of 3D structure by PyMOL**

The investigated structure of p16 has been retrieved from the PDB databank with ID:

1DC2. In the study of mutations using PyMOL, no change in polar contacts was observed in L16Q, G55D, P81R, and G101R mutations. As shown in Figures 1, 2, and 3, the polar contacts of mutant residues the mutations in G23D/R/S, L32P, N42K, G67D/R, H83R, G89D/S, A102E, G122R, and V126D in comparison with wild type residues were changed. According to Figure 1, new Hbonds have been created in the Arg substituted Leu at position 16 (between R16 and L63, length=1.9 Å), Arg substituted His at position 83 (between R83 and T77 with length=2.5 Å, between R83 and D108 with length=0.9 Å), Ser substituted Gly at position 89 (between S89 and G122, length=2.9 Å), and Glu substituted Ala at position 102 (between E102 and L97, length=2.1 Å). H-bond lengths between G89 with A85 (H-bond length=2.2 Å) and G89 with A86 (H-bond length=1.8 Å) were changed with replacing Ser at position 89. From Figure 1c one can see that H-bond length between S89 and A85 (S89 and A86) was equal to 2.3 (1.7) Å. According to Figure 1d, there was a polar contact between A102 and L104 (length=2.1 Å), this bond has been destroyed by replacing Glu at position 102. Figure 2 shows that H-bonds length was changed in G23D/R/S, G67R, and G89D amino acid substitutions. There was a polar contact between G23 and A20 with a length of 2.0 Å that by substitution Gly to Asp/Arg/Ser this distance was reduced to 1.9 Å (Figure 2a). As shown in Figure 2b, in G67D, the H-bond length between G67 and L63 was 1.7 Å and between G67 and L64 was 2.0 Å. But, in the mutant forms (G67R), the H-bond length between R67 and L63 (between R67 and L64) was equal to 1.4 (2.6) Å. Gly as a wild type residue at position 89, created polar contacts between G89 and A85 with a length of 2.2 Å and between G89 and A86 with a length of 1.8 Å. Placement of Asp at position 89 leds to an increase in distance between D89 and A85 to 2.3 Å. Also, as shown in Figure 2c,

the distance between D89 and A86 was reduced to 1.7 Å.

Figure 3 shows that L32P, N42K, G67D, G122R, and V126D amino acid substitutions led to the loss of H-bonds. Also, in mutations L32P and G67D, Hbonds length was changed. Leu at position 32 as a wild type residue had polar contacts with G35 (length=2.6 Å) and V28 (length=1.5 Å). From Figure 3a one can see that by changing the Leu to Pro at position 32 the polar contact with G35 would be destroyed and the length of Hbond with V28 was noticeably changed (length=2.5 Å). According to Figure 3b, there was an H-bond between N42 and S43 with a length of 1.9 Å. While by the substitution of Asn to Lys at position 42, the polar contact with S43 has been destroyed. As shown in Figure 3c, Gly 67 formed an H-bond with Asn 39 (H-bond length=2.4 Å), an H-bond with L63 (Hbonds length=1.7 Å), and an H-bond with L64 (H-bond length=2.0 Å). The H-bond with L64 has been destroyed by changing the Gly to Asp at position 67. Also, the Hbond length between D67 and L63 was changed and was equal to 1.4 Å and a new

H-bond has been created between D67 and N39 with a length of 2.5 Å. There was a polar contact between G122 and A118 (H-bond length=2.6 Å) that this polar contact has been missed by replacing Arg at position 122 (see Figure 3d). Figure 1e shows that at position 126, Val had H-bonding with H123 (H-bond length=1.9 Å) in the wild type form, but this polar contact was destroyed by replacing ASP at position 126.

### Analyzing hydrophobicity changes

As shown in Table IV, the wild type residues at L16R/Q, G23D/R, L32P, G55D, G67D/R, G89D, G101R, A102E, G122R, and V126D amino substitutions were more hydrophobic than the mutant residues. So, because of the reduction in hydrophobicity values, the probability of hydrophobic interactions was reduced. So, these mutations can disrupt the structure of the protein. Changes in hydrophobicity calculated by ExPASy were noticeable for L16R/Q, G23R, L32P, G67R, G101R, A102E, G122R, and V126D, but can be ignored in G23S, N42K, and G89S mutations.

Table I. The summary of pathogenicity predictions of nsSNVs obtained via SIFT, PolyPhen-2, I-Mutant 2.0. and PROVEAN.

| Mutant 2.0, a | mu I KO / L              | 211 V.                     |                                  |                                       |                                 |                               |
|---------------|--------------------------|----------------------------|----------------------------------|---------------------------------------|---------------------------------|-------------------------------|
| SNP ID        | Amino acid change in p16 | SIFT<br>SCORE <sup>1</sup> | POLYPHEN-2<br>SCORE <sup>2</sup> | POLYPHEN-2<br>Sensitivity/Specificity | I-MUTANT2.0<br>DDG <sup>3</sup> | PROVEAN<br>SCORE <sup>4</sup> |
| rs864622263   | p.L16R                   | APF                        | Probably Damaging                | 0.00/1.00                             | Decrease                        | Deleterious                   |
|               |                          | 0.00                       | 1.000                            |                                       | -1.67                           | -4.818                        |
|               | p.L16P                   | APF                        | Probably Damaging                | 0.00/1.00                             | Decrease                        | Deleterious                   |
|               |                          | 0.00                       | 1.000                            |                                       | -1.04                           | -5.55                         |
|               | p.L16Q                   | APF                        | Probably Damaging                | 0.00/1.00                             | Decrease                        | Deleterious                   |
|               |                          | 0.00                       | 1.000                            |                                       | -2.07                           | -4.842                        |
| rs760065045   | p.A20P                   | APF                        | Probably Damaging                | 0.00/1.00                             | Decrease                        | Deleterious                   |
|               |                          | 0.02                       | 1.000                            |                                       | -0.16                           | -4.06                         |
| rs864622484   | p.A20G                   | APF                        | Probably Damaging                | 0.78/0.95                             | Decrease                        | Deleterious                   |
|               |                          | 0.03                       | 0.966                            |                                       | -1.20                           | -3.21                         |
| rs1329324238  | p.A21D                   | APF                        | Probably Damaging                | 0.00/1.00                             | Decrease                        | Deleterious                   |
|               |                          | 0.02                       | 1.000                            |                                       | -1.40                           | -4.90                         |
| rs1064794292  | p.G23D                   | APF                        | Probably Damaging                | 0.00/1.00                             | Decrease                        | Deleterious                   |
|               |                          | 0.04                       | 1.000                            |                                       | -2.41                           | -5.899                        |
| rs1131691186  | p.G23R                   | APF                        | Probably Damaging                | 0.00/1.00                             | Decrease                        | Deleterious                   |
|               |                          | 0.01                       | 1.000                            |                                       | -1.22                           | -6.72                         |
|               | p.G23S                   | APF                        | Probably Damaging                | 0.27/0.99                             | Decrease                        | Deleterious                   |
|               |                          | 0.03                       | 0.998                            |                                       | -1.40                           | -5.029                        |
|               | p.G23C                   | APF                        | Probably Damaging                | 0.00/1.00                             | Decrease                        | Deleterious                   |
|               |                          | 0.00                       | 1.000                            |                                       | -0.70                           | -7.535                        |
| rs748780473   | p.V25G                   | APF                        | Possibly Damaging                | 0.87/0.91                             | Decrease                        | Deleterious                   |

|   |  | 0.00   | 0616  |  | -3.29   | -3.84   |
|---|--|--|---|--|---|---|
| rs775176191   | p.V28G   | APF  | Probably Damaging   | 0.00/1.00  | Decrease  | Deleterious   |
| 1==1<=<200  | Daou   | 0.00   | 1.000   | 0.00/1.00  | -3.46   | -5.482  |
| rs1554656382  | p.R29W   | APF<br>0.00  | Probably Damaging 1.000   | 0.00/1.00  | Decrease<br>-0.68   | Deleterious -5.26   |
| rs878853650   | p.L32P   | APF  | Probably Damaging   | 0.00/1.00  | Decrease  | Deleterious   |
| 1307000000  | P.2321   | 0.00   | 1.000   | 0.0071.00  | -2.03   | -5.743  |
| rs745827714   | p.L32V   | APF  | Probably Damaging   | 0.41/0.98  | Decrease  | Deleterious   |
|   |  | 0.00   | 0.997   |  | -2.18   | -2.50   |
| rs746834149   | p.G35V   | APF  | Probably Damaging   | 0.00/1.00  | Decrease  | Deleterious   |
|   |  | 0.02   | 1.000   |  | -0.17   | -5.95   |
| rs200382984   | p.A36G   | APF  | Probably Damaging   | 0.80/0.94  | Decrease  | Deleterious   |
|   | •  | 0.01   | 0.937   |  | -1.30   | -2.76   |
| rs752731682   | p.N39H   | APF  | Probably Damaging   | 0.27/0.99  | Decrease  | Deleterious   |
|   | •  | 0.01   | 0.998   |  | -1.45   | -3.93   |
| rs1554656306  | p.N39S   | APF  | Possibly Damaging   | 0.83/0.93  | Decrease  | Deleterious   |
|   | 1  | 0.03   | 0.864   |  | -0.69   | -3.80   |
| rs864622439   | p.N39K   | APF  | Probably Damaging   | 0.68/0.97  | Decrease  | Deleterious   |
| 1500102210  | Piritoria  | 0.02   | 0.995   | 0.00,01,7  | -1.04   | -4.66   |
| rs1060501264  | p.N42S   | APF  | Probably Damaging   | 0.14/0.99  | Decrease  | Deleterious   |
| 131000301204  | p.11125  | 0.00   | 0.999   | 0.1 1/ 0.55  | -0.92   | -4.31   |
| rs1587339638  | p.N42K   | APF  | Probably Damaging   | 0.00/1.00  | Decrease  | Deleterious   |
| 181307337030  | p.11421X   | 0.00   | 1.000   | 0.00/1.00  | -1.03   | -5.17   |
| ne 1597330662   | p.N42D   | APF  | Probably Damaging   | 0.14/0.99  | Decrease  | Deleterious   |
| rs1587339662  | p.1442D  | 0.00   | 0.999   | 0.14/0.99  | -1.00   | -4.31   |
| 1220700460  | C450   |  |   | 0.00/1.00  |   |   |
| rs1328708469  | p.G45S   | APF  | Probably Damaging   | 0.00/1.00  | Decrease  | Deleterious   |
| 17/200401/  | DAGIN  | 0.03   | 1.000   | 0.00/1.00  | -0.16   | -4.66   |
| rs1563892916  | p.R46W   | APF  | Probably Damaging   | 0.00/1.00  | Decrease  | Deleterious   |
|   |  | 0.01   | 1.000   |  | -0.66   | -6.69   |
| rs763804037   | p.P48R   | APF  | Probably Damaging   | 0.00/1.00  | Decrease  | Deleterious   |
|   |  | 0.04   | 1.000   |  | -0.10   | -4.84   |
| rs199907548   | p.I49T   | APF  | Probably Damaging   | 0.00/1.00  | Decrease  | Deleterious   |
|   |  | 0.01   | 1.000   |  | -3.20   | -3.773  |
|   | p.I49S   | APF  | Probably Damaging   | 0.00/1.00  | Decrease  | Deleterious   |
|   |  | 0.00   | 1.000   |  | -2.94   | -4.666  |
| rs587778189   | p.Q50P   | APF  | Probably Damaging   | 0.00/1.00  | Decrease  | Deleterious   |
|   |  | 0.02   | 1.000   |  | -0.67   | -5.185  |
| rs561034503   | p.G55D   | APF  | Probably Damaging   | 0.00/1.00  | Decrease  | Deleterious   |
|   |  | 0.04   | 1.000   |  | -0.09   | -6.49   |
| rs104894099   | p.V59G   | APF  | Probably Damaging   | 0.27/0.99  | Decrease  | Deleterious   |
|   | •  | 0.00   | 0.000   |  | 4.72  | 5.000   |
|   |  |  | 0.998   |  | -4.72   | -5.983  |
|   | p.V59E   |  |   | 0.14/0.99  | Decrease  |   |
|   | p.V59E   | APF  | Probably Damaging   | 0.14/0.99  | Decrease  | Deleterious   |
| rs36204594  |  | APF<br>0.00  | Probably Damaging<br>0.999  |  | Decrease<br>-2.28   | Deleterious<br>-5.242   |
| rs36204594  | p.V59E<br>p.A60E   | APF<br>0.00<br>APF   | Probably Damaging<br>0.999<br>Probably Damaging   | 0.14/0.99  | Decrease<br>-2.28<br>Decrease   | Deleterious<br>-5.242<br>Deleterious  |
|   | p.A60E   | APF<br>0.00<br>APF<br>0.01   | Probably Damaging<br>0.999<br>Probably Damaging<br>1.000  | 0.00/1.00  | Decrease<br>-2.28<br>Decrease<br>-1.24  | Deleterious<br>-5.242<br>Deleterious<br>-4.617  |
|   |  | APF<br>0.00<br>APF<br>0.01<br>APF  | Probably Damaging<br>0.999<br>Probably Damaging<br>1.000<br>Probably Damaging   |  | Decrease -2.28 Decrease -1.24 Decrease  | Deleterious<br>-5.242<br>Deleterious<br>-4.617<br>Deleterious   |
|   | p.A60E<br>p.A60P   | APF<br>0.00<br>APF<br>0.01<br>APF<br>0.02  | Probably Damaging<br>0.999<br>Probably Damaging<br>1.000<br>Probably Damaging<br>1.000  | 0.00/1.00<br>0.00/1.00   | Decrease -2.28 Decrease -1.24 Decrease -0.39  | Deleterious<br>-5.242<br>Deleterious<br>-4.617<br>Deleterious<br>-4.68  |
|   | p.A60E   | APF<br>0.00<br>APF<br>0.01<br>APF<br>0.02<br>APF   | Probably Damaging<br>0.999<br>Probably Damaging<br>1.000<br>Probably Damaging<br>1.000<br>Probably Damaging   | 0.00/1.00  | Decrease -2.28 Decrease -1.24 Decrease -0.39 Decrease   | Deleterious -5.242 Deleterious -4.617 Deleterious -4.68 Deleterious   |
| rs769382085   | p.A60E<br>p.A60P<br>p.A60T                                     | APF<br>0.00<br>APF<br>0.01<br>APF<br>0.02<br>APF<br>0.04   | Probably Damaging<br>0.999<br>Probably Damaging<br>1.000<br>Probably Damaging<br>1.000<br>Probably Damaging<br>1.000  | 0.00/1.00<br>0.00/1.00<br>0.00/1.00  | Decrease -2.28 Decrease -1.24 Decrease -0.39 Decrease -1.38   | Deleterious -5.242 Deleterious -4.617 Deleterious -4.68 Deleterious -3.73   |
| rs769382085   | p.A60E<br>p.A60P   | APF<br>0.00<br>APF<br>0.01<br>APF<br>0.02<br>APF<br>0.04<br>APF  | Probably Damaging<br>0.999 Probably Damaging<br>1.000 Probably Damaging<br>1.000 Probably Damaging<br>1.000 Probably Damaging<br>1.000 Probably Damaging  | 0.00/1.00<br>0.00/1.00   | Decrease -2.28 Decrease -1.24 Decrease -0.39 Decrease -1.38 Decrease  | Deleterious -5.242 Deleterious -4.617 Deleterious -4.68 Deleterious -3.73 Deleterious   |
| rs769382085   | p.A60E<br>p.A60P<br>p.A60T<br>p.G67R                           | APF<br>0.00<br>APF<br>0.01<br>APF<br>0.02<br>APF<br>0.04<br>APF  | Probably Damaging 0.999 Probably Damaging 1.000   | 0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.00/1.00   | Decrease -2.28 Decrease -1.24 Decrease -0.39 Decrease -1.38 Decrease -0.96  | Deleterious -5.242 Deleterious -4.617 Deleterious -4.68 Deleterious -3.73 Deleterious -7.56   |
| rs769382085   | p.A60E<br>p.A60P<br>p.A60T                                     | APF<br>0.00<br>APF<br>0.01<br>APF<br>0.02<br>APF<br>0.04<br>APF<br>0.00<br>APF   | Probably Damaging 0.999 Probably Damaging 1.000 Probably Damaging   | 0.00/1.00<br>0.00/1.00<br>0.00/1.00  | Decrease -2.28 Decrease -1.24 Decrease -0.39 Decrease -1.38 Decrease -0.96 Decrease   | Deleterious -5.242 Deleterious -4.617 Deleterious -4.68 Deleterious -3.73 Deleterious -7.56 Deleterious   |
| rs769382085<br>rs758389471  | p.A60E p.A60P p.A60T p.G67R p.G67S                             | APF<br>0.00<br>APF<br>0.01<br>APF<br>0.02<br>APF<br>0.04<br>APF<br>0.00<br>APF   | Probably Damaging 0.999 Probably Damaging 1.000   | 0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.00/1.00  | Decrease -2.28 Decrease -1.24 Decrease -0.39 Decrease -1.38 Decrease -0.96 Decrease -1.35   | Deleterious -5.242 Deleterious -4.617 Deleterious -4.68 Deleterious -3.73 Deleterious -7.56 Deleterious -5.70   |
| rs769382085<br>rs758389471  | p.A60E<br>p.A60P<br>p.A60T<br>p.G67R                           | APF<br>0.00<br>APF<br>0.01<br>APF<br>0.02<br>APF<br>0.04<br>APF<br>0.00<br>APF<br>0.00<br>APF  | Probably Damaging 0.999 Probably Damaging 1.000 Probably Damaging   | 0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.00/1.00   | Decrease -2.28 Decrease -1.24 Decrease -0.39 Decrease -1.38 Decrease -0.96 Decrease -1.35 Decrease  | Deleterious -5.242 Deleterious -4.617 Deleterious -4.68 Deleterious -3.73 Deleterious -7.56 Deleterious -5.70 Deleterious   |
| rs769382085<br>rs758389471<br>rs863224605                                 | p.A60E p.A60P p.A60T p.G67R p.G67S p.G67D                      | APF<br>0.00<br>APF<br>0.01<br>APF<br>0.02<br>APF<br>0.04<br>APF<br>0.00<br>APF<br>0.00<br>APF  | Probably Damaging 0.999 Probably Damaging 1.000   | 0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.00/1.00                           | Decrease -2.28  Decrease -1.24  Decrease -0.39  Decrease -1.38  Decrease -0.96  Decrease -1.35  Decrease -2.26  | Deleterious -5.242 Deleterious -4.617 Deleterious -4.68 Deleterious -3.73 Deleterious -7.56 Deleterious -5.70 Deleterious -6.70   |
| rs769382085<br>rs758389471<br>rs863224605                                 | p.A60E p.A60P p.A60T p.G67R p.G67S                             | APF<br>0.00<br>APF<br>0.01<br>APF<br>0.02<br>APF<br>0.04<br>APF<br>0.00<br>APF<br>0.00<br>APF<br>0.00<br>APF                               | Probably Damaging 0.999 Probably Damaging 1.000 Probably Damaging   | 0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.00/1.00  | Decrease -2.28 Decrease -1.24 Decrease -0.39 Decrease -1.38 Decrease -0.96 Decrease -1.35 Decrease -2.26 Decrease                                     | Deleterious -5.242 Deleterious -4.617 Deleterious -4.68 Deleterious -3.73 Deleterious -7.56 Deleterious -5.70 Deleterious -6.70 Deleterious   |
| rs769382085<br>rs758389471<br>rs863224605<br>rs1060501260                 | p.A60E p.A60P p.A60T p.G67R p.G67S p.G67D p.A68G               | APF<br>0.00<br>APF<br>0.01<br>APF<br>0.02<br>APF<br>0.04<br>APF<br>0.00<br>APF<br>0.00<br>APF<br>0.00<br>APF                               | Probably Damaging 0.999 Probably Damaging 1.000 Probably Damaging 0.998                         | 0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.27/0.99              | Decrease -2.28 Decrease -1.24 Decrease -0.39 Decrease -1.38 Decrease -0.96 Decrease -1.35 Decrease -2.26 Decrease -0.82                               | Deleterious -5.242 Deleterious -4.617 Deleterious -4.68 Deleterious -3.73 Deleterious -7.56 Deleterious -5.70 Deleterious -6.70 Deleterious -3.842                                      |
| rs36204594 rs769382085 rs758389471 rs863224605 rs1060501260 rs559848002   | p.A60E p.A60P p.A60T p.G67R p.G67S p.G67D                      | APF<br>0.00<br>APF<br>0.01<br>APF<br>0.02<br>APF<br>0.04<br>APF<br>0.00<br>APF<br>0.00<br>APF<br>0.00<br>APF                               | Probably Damaging 0.999 Probably Damaging 1.000 Probably Damaging 1.098 Probably Damaging                               | 0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.00/1.00                           | Decrease -2.28 Decrease -1.24 Decrease -0.39 Decrease -1.38 Decrease -0.96 Decrease -1.35 Decrease -2.26 Decrease -0.82 Decrease                      | Deleterious -5.242 Deleterious -4.617 Deleterious -4.68 Deleterious -3.73 Deleterious -7.56 Deleterious -5.70 Deleterious -6.70 Deleterious -3.842 Deleterious                          |
| rs769382085<br>rs758389471<br>rs863224605<br>rs1060501260                 | p.A60E p.A60P p.A60T p.G67R p.G67S p.G67D p.A68G p.N71T        | APF<br>0.00<br>APF<br>0.01<br>APF<br>0.02<br>APF<br>0.04<br>APF<br>0.00<br>APF<br>0.00<br>APF<br>0.00<br>APF<br>0.00<br>APF                | Probably Damaging 0.999 Probably Damaging 1.000 Probably Damaging 0.998 Probably Damaging 0.998                         | 0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.27/0.99              | Decrease -2.28 Decrease -1.24 Decrease -0.39 Decrease -1.38 Decrease -0.96 Decrease -1.35 Decrease -2.26 Decrease -0.82 Decrease -0.54                | Deleterious -5.242 Deleterious -4.617 Deleterious -4.68 Deleterious -3.73 Deleterious -7.56 Deleterious -5.70 Deleterious -6.70 Deleterious -3.842 Deleterious -5.500                   |
| rs769382085<br>rs758389471<br>rs863224605<br>rs1060501260                 | p.A60E p.A60P p.A60T p.G67R p.G67S p.G67D p.A68G               | APF<br>0.00<br>APF<br>0.01<br>APF<br>0.02<br>APF<br>0.04<br>APF<br>0.00<br>APF<br>0.00<br>APF<br>0.00<br>APF<br>0.00<br>APF<br>0.00<br>APF | Probably Damaging 0.999 Probably Damaging 1.000 Probably Damaging 0.998 Probably Damaging 0.998 Probably Damaging/0.993 Probably Damaging/      | 0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.27/0.99              | Decrease -2.28 Decrease -1.24 Decrease -0.39 Decrease -1.38 Decrease -0.96 Decrease -1.35 Decrease -2.26 Decrease -0.82 Decrease -0.54 Decrease       | Deleterious -5.242 Deleterious -4.617 Deleterious -4.68 Deleterious -3.73 Deleterious -7.56 Deleterious -5.70 Deleterious -6.70 Deleterious -3.842 Deleterious -5.500 Deleterious       |
| rs769382085 rs758389471 rs863224605 rs1060501260 rs559848002              | p.A60E p.A60P p.A60T p.G67R p.G67S p.G67D p.A68G p.N71T p.T79P | APF<br>0.00<br>APF<br>0.01<br>APF<br>0.02<br>APF<br>0.04<br>APF<br>0.00<br>APF<br>0.00<br>APF<br>0.00<br>APF<br>0.00<br>APF                | Probably Damaging 0.999 Probably Damaging 1.000 Probably Damaging 0.998 Probably Damaging 0.998                         | 0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.27/0.99              | Decrease -2.28 Decrease -1.24 Decrease -0.39 Decrease -1.38 Decrease -0.96 Decrease -1.35 Decrease -2.26 Decrease -0.82 Decrease -0.54                | Deleterious -5.242 Deleterious -4.617 Deleterious -4.68 Deleterious -3.73 Deleterious -7.56 Deleterious -5.70 Deleterious -6.70 Deleterious -3.842 Deleterious -5.500                   |
| rs769382085 rs758389471 rs863224605 rs1060501260 rs559848002              | p.A60E p.A60P p.A60T p.G67R p.G67S p.G67D p.A68G p.N71T        | APF<br>0.00<br>APF<br>0.01<br>APF<br>0.02<br>APF<br>0.04<br>APF<br>0.00<br>APF<br>0.00<br>APF<br>0.00<br>APF<br>0.00<br>APF<br>0.00<br>APF | Probably Damaging 0.999 Probably Damaging 1.000 Probably Damaging 0.998 Probably Damaging 0.998 Probably Damaging/0.993 Probably Damaging/      | 0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.27/0.99              | Decrease -2.28 Decrease -1.24 Decrease -0.39 Decrease -1.38 Decrease -0.96 Decrease -1.35 Decrease -2.26 Decrease -0.82 Decrease -0.54 Decrease       | Deleterious -5.242 Deleterious -4.617 Deleterious -4.68 Deleterious -3.73 Deleterious -7.56 Deleterious -5.70 Deleterious -6.70 Deleterious -3.842 Deleterious -5.500 Deleterious       |
| rs769382085 rs758389471 rs863224605 rs1060501260 rs559848002 rs1554654113 | p.A60E p.A60P p.A60T p.G67R p.G67S p.G67D p.A68G p.N71T p.T79P | APF<br>0.00<br>APF<br>0.01<br>APF<br>0.02<br>APF<br>0.04<br>APF<br>0.00<br>APF<br>0.00<br>APF<br>0.00<br>APF<br>0.00<br>APF<br>0.00<br>APF | Probably Damaging 0.999 Probably Damaging 1.000 Probably Damaging 0.998 Probably Damaging 0.998 Probably Damaging 0.993 Probably Damaging 0.993 | 0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.00/1.00<br>0.27/0.99<br>0.70/0.97 | Decrease -2.28 Decrease -1.24 Decrease -0.39 Decrease -1.38 Decrease -0.96 Decrease -1.35 Decrease -2.26 Decrease -0.82 Decrease -0.54 Decrease -0.58 | Deleterious -5.242 Deleterious -4.617 Deleterious -4.68 Deleterious -3.73 Deleterious -7.56 Deleterious -5.70 Deleterious -6.70 Deleterious -3.842 Deleterious -5.500 Deleterious -3.81 |

|              |         | 0.00        | 1.000                      |           | -0.97             | -7.51                 |
|--------------|---------|-------------|----------------------------|-----------|-------------------|-----------------------|
| rs1057519881 | p.H83R  | APF<br>0.00 | Probably Damaging 1.000    | 0.00/1.00 | Decrease<br>-0.47 | Deleterious<br>-7.494 |
| rs121913385  | p.H83D  | APF<br>0.00 | Probably Damaging<br>0.999 | 0.14/0.99 | Decrease<br>-1.62 | Deleterious<br>-8.475 |
| rs1064796336 | p.A85F  | APF<br>0.00 | Probably Damaging<br>1.000 | 0.00/1.00 | Decrease<br>-0.14 | Deleterious<br>-5.350 |
| rs878853646  | p.A85S  | APF<br>0.00 | Probably Damaging<br>1.000 | 0.00/1.00 | Decrease<br>-0.28 | Deleterious<br>-2.74  |
|              | p.A85T  | APF<br>0.00 | Probably Damaging<br>1.000 | 0.00/1.00 | Decrease<br>-0.87 | Deleterious<br>-3.53  |
| rs1190283873 | p.A86T  | APF<br>0.01 | Probably Damaging<br>1.000 | 0.00/1.00 | Decrease<br>-0.67 | Deleterious<br>-3.71  |
| rs749714198  | p.R87W  | APF<br>0.01 | Possibly Damaging<br>0.675 | 0.86/0.92 | Decrease<br>-0.48 | Deleterious<br>-7.439 |
| rs137854597  | p.G89S  | APF<br>0.00 | Probably Damaging<br>1.000 | 0.00/1.00 | Decrease<br>-1.04 | Deleterious<br>-5.600 |
| rs137854599  | p.G89A  | APF<br>0.00 | Probably Damaging<br>1.000 | 0.00/1.00 | Decrease<br>-0.84 | Deleterious<br>-5.650 |
|              | p.G89D  | APF<br>0.00 | Probably Damaging<br>1.000 | 0.00/1.00 | Decrease<br>-1.34 | Deleterious<br>-6.592 |
| rs1563889362 | p.L91Q  | APF<br>0.00 | Probably Damaging<br>0.999 | 0.14/0.99 | Decrease<br>-2.65 | Deleterious<br>-5.39  |
| rs34886500   | p.R99W  | APF<br>0.01 | Probably Damaging<br>1.000 | 0.00/1.00 | Decrease<br>-0.36 | Deleterious<br>-4.76  |
| rs104894094  | p.G101W | APF<br>0.00 | Probably Damaging<br>1.000 | 0.00/1.00 | Decrease<br>-0.78 | Deleterious<br>-6.134 |
|              | p.G101R | APF<br>0.00 | Probably Damaging<br>1.000 | 0.00/1.00 | Decrease<br>-0.64 | Deleterious<br>-5.890 |
| rs35741010   | p.A102T | APF<br>0.01 | Probably Damaging<br>0.995 | 0.68/0.97 | Decrease<br>-0.53 | Deleterious           |
| rs137854598  | p.A102E | APF<br>0.00 | Probably Damaging<br>1.000 | 0.00/1.00 | Decrease<br>-0.38 | Deleterious<br>-4.47  |
| rs767642535  | p.R103W | APF<br>0.02 | Probably Damaging 1.000    | 0.00/1.00 | Decrease<br>-0.21 | Deleterious -5.10     |
| rs1554654028 | p.V106E | APF<br>0.04 | Probably Damaging 1.000    | 0.00/1.00 | Decrease<br>-1.93 | Deleterious -3.55     |
| rs1339792331 | p.D108V | APF<br>0.00 | Probably Damaging 1.000    | 0.00/1.00 | Decrease<br>-0.39 | Deleterious<br>-8.24  |
| rs121913381  | p.D108Y | APF<br>0.01 | Probably Damaging 1.000    | 0.00/1.00 | Decrease<br>-1.46 | Deleterious<br>-8.201 |
| rs778971134  | p.G111R | APF<br>0.00 | Probably Damaging 1.000    | 0.00/1.00 | Decrease<br>-1.05 | Deleterious<br>-5.43  |
|              | p.G111S | APF<br>0.00 | Probably Damaging 1.000    | 0.00/1.00 | Decrease<br>-1.20 | Deleterious -3.67     |
| rs876660436  | p.R112C | APF<br>0.05 | Probably Damaging 1.000    | 0.00/1.00 | Decrease<br>-0.59 | Deleterious<br>-4.05  |
| rs104894104  | p.P114S | APF<br>0.00 | Probably Damaging 1.000    | 0.00/1.00 | Decrease<br>-1.39 | Deleterious<br>-7.479 |
|              | p.P114T | APF<br>0.00 | Probably Damaging 1.000    | 0.00/1.00 | Decrease<br>-1.86 | Deleterious<br>-7.495 |
| rs121913386  | p.P114L | APF<br>0.00 | Probably Damaging 1.000    | 0.00/1.00 | Decrease<br>-0.66 | Deleterious<br>-9.361 |
|              | p.P114H | APF<br>0.00 | Probably Damaging<br>1.000 | 0.00/1.00 | Decrease<br>-1.83 | Deleterious<br>-8.407 |
| rs750655995  | p.V115G | APF<br>0.01 | Probably Damaging<br>1.000 | 0.00/1.00 | Decrease<br>-3.55 | Deleterious<br>-6.16  |
|              | p.V115E | APF<br>0.01 | Probably Damaging<br>0.999 | 0.14/0.99 | Decrease<br>-2.20 | Deleterious<br>-5.32  |
| rs1060501270 | p.A118G | APF<br>0.05 | Probably Damaging 0.996    | 0.55/0.98 | Decrease<br>-1.03 | Deleterious<br>-3.73  |
| rs1554653960 | p.A118T | APF<br>0.02 | Probably Damaging<br>1.000 | 0.00/1.00 | Decrease<br>-1.07 | Deleterious<br>-3.71  |
| rs113798404  | p.G122R | APF         | Probably Damaging          | 0.00/1.00 | Decrease          | Deleterious           |

|              |         | 0.03         | 1.000                   |           | -0.61             | -6.347                |
|--------------|---------|--------------|-------------------------|-----------|-------------------|-----------------------|
| rs146179135  | p.D125Y | APF<br>0.04* | Possibly Damaging 0.743 | 0.85/0.92 | Decrease<br>-0.31 | Deleterious<br>-4.222 |
| rs104894098  | p.V126D | APF<br>0.00* | Probably Damaging 1.000 | 0.00/1.00 | Decrease -2.62    | Deleterious<br>-6.223 |
| rs1350305259 | p.V126L | APF<br>0.03* | Probably Damaging 0.898 | 0.82/0.94 | Decrease -1.10    | Deleterious<br>-2.57  |
| rs1563888826 | p.R128W | APF<br>0.01  | Probably Damaging 0.992 | 0.70/0.97 | Decrease<br>-0.42 | Deleterious<br>-3.24  |

\*There is low confidence in this prediction. ¹Score<0.05= Affected Protein Function (APF), ²Score 0.15-0.85= possibly damaging and score: 0.85-1= probably damaging, ³DDG<0= Decrease stability, ⁴Score<-2.5= deleterious.

Table II. Disease probability by PHD-SNP<sup>g</sup>, PANTHER, and SNPs&GO.

| SNP ID       | Amino acid change in | PHD-SNPg/SCORE*  | PANTHER           | SNPs&GO/SCORE** |
|--------------|----------------------|------------------|-------------------|-----------------|
|              | p16                  |                  |                   |                 |
| rs864622263  | p.L16R               | Pathogenic/0.979 | Probably Damaging | Disease/0.893   |
|              | p.L16Q               | Pathogenic/0.969 | Probably Damaging | Disease/0.858   |
|              | p.L16P               | Pathogenic/0.971 | Probably Damaging | Disease/0.679   |
| rs760065045  | p.A20P               | Pathogenic/0.972 | Possibly Damaging | Disease/0.531   |
| rs1329324238 | p.A21D               | Pathogenic/0.955 | Possibly Damaging | Disease/0.645   |
| rs1064794292 | p.G23D               | Pathogenic/0.946 | Probably Damaging | Disease/0.903   |
| rs1131691186 | p.G23S               | Pathogenic/0.932 | Probably Damaging | Disease/0.813   |
|              | p.G23C               | Pathogenic/0.973 | Probably Damaging | Disease/0.864   |
|              | p.G23R               | Pathogenic/0.963 | Probably Damaging | Disease/0.662   |
| rs775176191  | p.V28G               | Pathogenic/0.940 | Possibly Damaging | Disease/0.826   |
| rs878853650  | p.L32P               | Pathogenic/0.973 | Probably Damaging | Disease/0.861   |
| rs1587339638 | p.N42K               | Pathogenic/0.767 | Possibly Damaging | Disease/0.571   |
| rs1328708469 | p.G45S               | Pathogenic/0.916 | Probably Damaging | Disease/0.544   |
| rs199907548  | p.I49S               | Pathogenic/0.796 | Possibly Damaging | Disease/0.801   |
| rs587778189  | p.Q50P               | Pathogenic/0.974 | Possibly Damaging | Disease/0.903   |
| rs561034503  | p.G55D               | Pathogenic/0.963 | Probably Damaging | Disease/0.657   |
| rs36204594   | p.A60E               | Pathogenic/0.967 | Possibly Damaging | Disease/0.747   |
| rs758389471  | p.G67R               | Pathogenic/0.971 | Probably Damaging | Disease/0.643   |
|              | p.G67S               | Pathogenic/0.973 | Probably Damaging | Disease/0.594   |
| rs863224605  | p.G67D               | Pathogenic/0.963 | Probably Damaging | Disease/0.696   |
| rs559848002  | p.N71T               | Pathogenic/0.994 | Possibly Damaging | Disease/0.688   |
| rs11552823   | p.P81R               | Pathogenic/0.984 | Probably Damaging | Disease/0.521   |
| rs34968276   | p.H83Q               | Pathogenic/0.980 | Probably Damaging | Disease/0.753   |
| rs1057519881 | p.H83R               | Pathogenic/0.947 | Probably Damaging | Disease/0.905   |
| rs121913385  | p.H83D               | Pathogenic/0.974 | Probably Damaging | Disease/0.895   |
| rs1064796336 | p.A85F               | Pathogenic/1.000 | Probably Damaging | Disease/0.832   |
| rs878853646  | p.A85S               | Pathogenic/0.986 | Probably Damaging | Disease/0.649   |
|              | p.A85T               | Pathogenic/0.982 | Probably Damaging | Disease/0.617   |
| rs1190283873 | p.A86T               | Pathogenic/0.969 | Probably Damaging | Disease/0.597   |
| rs749714198  | p.R87W               | Pathogenic/0.905 | Probably Damaging | Disease/0.893   |
| rs137854597  | p.G89S               | Pathogenic/0.973 | Probably Damaging | Disease/0.886   |
| rs137854599  | p.G89A               | Pathogenic/0.979 | Probably Damaging | Disease/0.872   |
|              | p.G89D               | Pathogenic/0.981 | Probably Damaging | Disease/0.933   |
| rs1563889362 | p.L91Q               | Pathogenic/0.832 | Possibly Damaging | Disease/0.774   |
| rs104894094  | p.G101W              | Pathogenic/0.937 | Probably Damaging | Disease/0.873   |
|              | p.G101R              | Pathogenic/0.806 | Probably Damaging | Disease/0.810   |
| rs35741010   | p.A102T              | Pathogenic/0.598 | Probably Damaging | Disease/0.553   |

| rs137854598  | p.A102E | Pathogenic/0.705 | Probably Damaging | Disease/0.762 |
|--------------|---------|------------------|-------------------|---------------|
| rs767642535  | p.R103W | Pathogenic/0.612 | Possibly Damaging | Disease/0.567 |
| rs1339792331 | p.D108V | Pathogenic/0.981 | Possibly Damaging | Disease/0.806 |
| rs121913381  | p.D108Y | Pathogenic/0.986 | Possibly Damaging | Disease/0.902 |
| rs104894104  | p.P114S | Pathogenic/0.987 | Probably Damaging | Disease/0.819 |
|              | p.P114T | Pathogenic/0.990 | Probably Damaging | Disease/0.788 |
| rs121913386  | p.P114L | Pathogenic/0.993 | Probably Damaging | Disease/0.751 |
|              | p.P114H | Pathogenic/0.995 | Probably Damaging | Disease/0.840 |
| rs1554653960 | p.A118T | Pathogenic/0.972 | Probably Damaging | Disease/0.526 |
| rs113798404  | p.G122R | Pathogenic/0.941 | Possibly Damaging | Disease/0.822 |
| rs104894098  | p.V126D | Pathogenic/0.970 | Possibly Damaging | Disease/0.932 |

<sup>\*,\*\*</sup>Amino acid substitution with the score>0.5 is identified as a pathogenic or disease association variation.

Table III. The secondary structure analysis of high-risk mutations using NetsurfP-2.0 and the

conservation study by ConSurf.

| SNP ID       | Amino acid position and mutation | Conservation score* | Amino acid change | RSA <sup>1</sup> (%) | ASA <sup>2</sup><br>(Å) | Class<br>assignment | Secondary<br>structure |
|--------------|----------------------------------|---------------------|-------------------|----------------------|-------------------------|---------------------|------------------------|
| rs864622263  | L16R/Q                           | 9                   | L                 | 4                    | 8                       | Buried              | α Helix                |
|              |                                  |                     | R                 | 22                   | 51                      | Buried              | α Helix                |
|              |                                  |                     | Q                 | 18                   | 33                      | Buried              | α Helix                |
| rs1131691186 | G23R/S                           | 8                   | G                 | 30                   | 24                      | Exposed             | Turn                   |
|              |                                  |                     | R                 | 48                   | 110                     | Exposed             | Turn                   |
|              |                                  |                     | S                 | 37                   | 43                      | Exposed             | Turn                   |
| rs1064794292 | G23D                             | 8                   | G                 | 30                   | 24                      | Exposed             | Turn                   |
|              |                                  |                     | D                 | 51                   | 74                      | Exposed             | Turn                   |
| rs878853650  | L32P                             | 9                   | L                 | 12                   | 21                      | Buried              | α Helix                |
|              |                                  |                     | P                 | 25                   | 36                      | Buried              | α Helix                |
| rs1587339638 | N42K                             | 7                   | N                 | 15                   | 22                      | Exposed             | Coil                   |
|              |                                  |                     | K                 | 24                   | 50                      | Exposed             | Coil                   |
| rs561034503  | G55D                             | 8                   | G                 | 28                   | 22                      | Exposed             | Turn                   |
|              |                                  |                     | D                 | 48                   | 69                      | Exposed             | Turn                   |
| rs863224605  | G67D                             | 8                   | G                 | 69                   | 54                      | Exposed             | Turn                   |
|              |                                  |                     | D                 | 85                   | 122                     | Exposed             | Turn                   |
| rs758389471  | G67R                             | 8                   | R                 | 77                   | 177                     | Exposed             | Turn                   |
| rs11552823   | P81R                             | 7                   | P                 | 2                    | 3                       | Buried              | α Helix                |
|              |                                  |                     | R                 | 12                   | 27                      | Buried              | α Helix                |
| rs1057519881 | H83R                             | 8                   | Н                 | 4                    | 8                       | Buried              | α Helix                |
|              |                                  |                     | R                 | 11                   | 25                      | Buried              | α Helix                |
| rs137854599  | G89D/S                           | 8                   | G                 | 29                   | 23                      | Exposed             | Turn                   |
|              |                                  |                     | D                 | 51                   | 74                      | Exposed             | Turn                   |
|              |                                  |                     | S                 | 38                   | 45                      | Exposed             | Turn                   |
| rs104894094  | G101R                            | 9                   | G                 | 65                   | 51                      | Exposed             | Turn                   |
|              |                                  |                     | R                 | 76                   | 175                     | Exposed             | Turn                   |
| rs137854598  | A102E                            | 9                   | A                 | 6                    | 6                       | Buried              | Coil                   |
|              |                                  |                     | Е                 | 25                   | 44                      | Buried              | Coil                   |
| rs113798404  | G122R                            | 7                   | G                 | 56                   | 44                      | Exposed             | Turn                   |
|              |                                  |                     | R                 | 75                   | 172                     | Exposed             | Turn                   |
| rs104894098  | V126D                            | 7                   | V                 | 7                    | 11                      | Buried              | α Helix                |
|              |                                  |                     | D                 | 22                   | 32                      | Buried              | α Helix                |

\*The score is from 1 to 9. Amino acid substitution with the score=1 is identified as a hypervariable residue and with the score=9 is identified as a highly conserved residue. ¹RSA: Related solvent accessibility (the threshold for exposed or buried residue is 25%). ²ASA: Absolute solvent accessibility.

Table IV. Evaluation of the hydrophobicity changes via the PEPTIDE 2.0 tool and ExPASy resource portal.

| SNP ID       | Amino<br>acid<br>change | PEPTIDE-2<br>prediction/Hydrophobicity<br>index*(%) | Change of the nature of the amino acid | Hydrophobicity change in<br>substituted position by<br>ExPASy resource portal |
|--------------|-------------------------|---|--|---|
| rs864622263  | p.L16R                  | Hydrophobic: 49.36/Basic:14.74                      | Hydrophobic to<br>Hydrophilic          | -0.923  |
|              | p.L16Q                  | Hydrophobic: 49.36/Neutral: 22.44                   | Hydrophobic to<br>Neutral              | -0.811  |
| rs1064794292 | p.G23D                  | Acidic: 14.74/Neutral: 21.15                        | Neutral to<br>Hydrophilic              | -0.344  |
| rs1131691186 | p.G23S                  | Not change  | Neutral to Neutral                     | -0.044  |
|              | p.G23R                  | Basic: 14.74/Neutral: 21.15                         | Neutral to<br>Hydrophilic              | -0.455  |
| rs878853650  | p.L32P                  | Not change  | Hydrophobic to<br>Hydrophobic          | -0.600  |
| rs1587339638 | p.N42K                  | Basic: 14.74/Neutral: 21.15                         | Hydrophilic to<br>Hydrophilic          | -0.045  |
| rs561034503  | p.G55D                  | Acidic: 14.74/Neutral: 21.15                        | Neutral to<br>Hydrophilic              | -0.344  |
| rs863224605  | p.G67D                  | Acidic: 14.74/Neutral: 21.15                        | Neutral to<br>Hydrophilic              | -0.344  |
| rs758389471  | p.G67R                  | Basic: 14.74/Neutral: 21.15                         | Neutral to<br>Hydrophilic              | -0.455  |
| rs11552823   | p.P81R                  | Hydrophobic: 49.36/Basic:14.74                      | Hydrophobic to<br>Hydrophilic          | -0.322  |
| rs1057519881 | p.H83R                  | Not change  | Hydrophilic to<br>Hydrophilic          | -0.145  |
| rs137854597  | p.G89S                  | Not change  | Neutral to Neutral                     | -0.044  |
| rs137854599  | p.G89D                  | Acidic: 14.74/Neutral: 21.15                        | Neutral to<br>Hydrophilic              | -0.344  |
| rs104894094  | p.G101R                 | Acidic: 14.74/Neutral: 21.15                        | Neutral to<br>Hydrophilic              | -0.456  |
| rs137854598  | p.A102E                 | Hydrophobic: 49.36/Acidic: 14.74                    | Hydrophobic to<br>Hydrophilic          | -0.589  |
| rs113798404  | p.G122R                 | Basic: 14.74/Neutral: 21.15                         | Neutral to<br>Hydrophilic              | -0.455  |
| rs104894098  | p.V126D                 | Hydrophobic: 49.36/Acidic: 14.74                    | Hydrophobic to<br>Hydrophilic          | -0.856  |

<sup>\*</sup>Hydrophobicity index for wild type: Hydrophobicity: 50%, Acidic: 14.1%, Basic: 14.1%, Neutral: 21.79%.

Table V. Experimental studies on the correlation of high-risk mutations with various cancers.

| Amino acid change in p16 | Type of cancer   |
|--------------------------|--|
| p.L16R                   | Hereditary cutaneous melanoma (46)   |
| p.G23D                   | Multiple primary melanoma (47-49), Familial pancreatic cancer (50), Melanoma (51)  |
| p.G23R                   | Melanoma prone family (47), Multiple primary melanoma (48)                         |
| p.G23S                   | Familial melanoma (52)   |
| p.L32P                   | Primary familial melanoma (53), Pancreatic cancer (47), Melanoma (51), Familial    |
|                          | melanoma (54)  |
| p.G67R                   | Familial melanoma (55)   |
| p.H83R                   | Pancreatic cancer (56)   |
| p.G89D                   | Melanoma (57), Melanoma, HNSCC, and pancreatic cancer (58), Familial melanoma (54) |
| p.G101R                  | Cutaneous melanoma (59), Melanoma prone family (59), Pancreatic cancer (60)        |
| p.V126D                  | Melanoma (61), Pancreatic cancer (60)  |
| L16Q, N42K, G55D, G67D,  | There is no study about the association of these mutations with diseases.          |
| P81R, G89S, A102E, G122R |  |

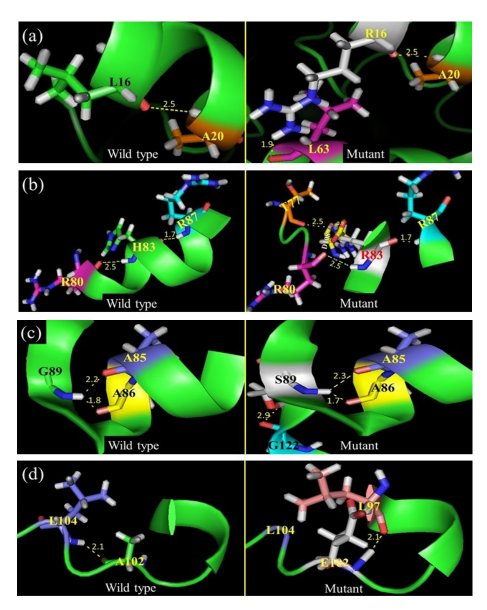


Figure 1. The created H-bonds in mutant variants (right column) while these bonds do not exist in the wild types (left column) (PDB ID=1DC2). (a): The created H-bond between R16 (mutant) and L63 in the L16R (H-bond length=1.9 Å). (b): The created H-bonds between R83 (mutant) and T77 (H-bond length=2.5 Å), between R83 and D108 (H-bond length=0.9 Å) in the H83R. (c): The created H-bond between S89 (mutant) and G122 in the G89S (H-bond length=2.9 Å). In the wild type, the H-bond length between G89 and A85 is 2.2 Å. By substituting Ser in the mutant form, the length of polar contact between S89 and A85 is equal to 2.3 Å. Also, in the wild type, the H-bond length between G89 and A86 is 1.8 Å, but between S89 and A86 is equal to 1.7 Å. (d): There was a polar contact between A102 and L104 with a length of 2.1 Å, but this bond has been destroyed by substituting of Glu and a new H-bond has been created between E102 and L97 with a length of 2.1 Å.

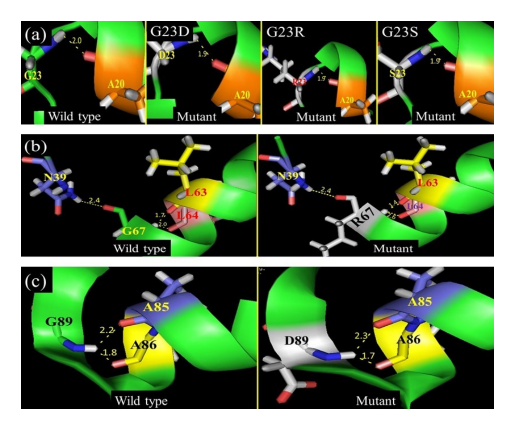


Figure 2. The change in the length of polar contact in G23D/R/S and G89D (PDB ID=1DC2) in the mutant form in comparison with the wild type protein. (a): In G23 (wild type), the H-bond length between G23 and A20 is 2.0 Å, but in the mutant forms, the H-bond length between D/R/S23 and A20 is 1.9 Å. (b): In G67 (wild type), the H-bond length between G67 and L63 is 1.7 Å and the H-bond length between G67 and L64 is 2.0 Å. But, in the mutant forms, the H-bond length between R67 and L63 is 1.4 Å and the H-bond length between R67 and L64 is 2.6 Å. (c): In G89 (wild type), the H-bond length between G89 and A85 is 2.2 Å, but the H-bond length between D89 and A85 is 2.3 Å. Also, in the wild type, the H-bond length between G89 and A86 in the mutant form is 1.7 Å.

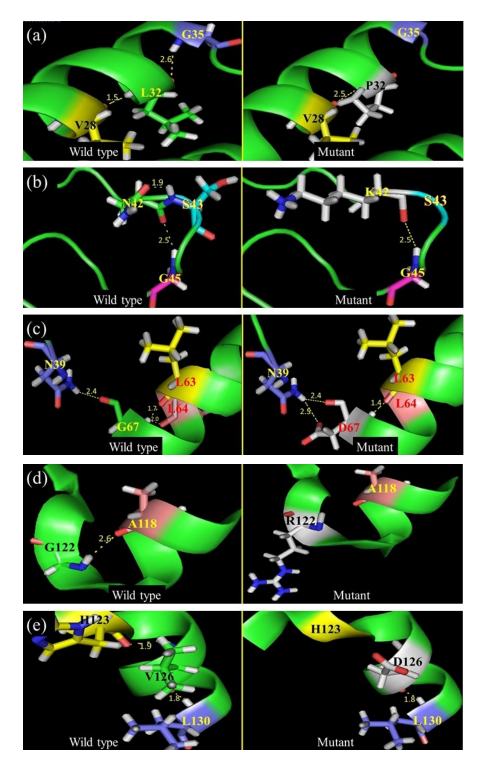


Figure 3. The loss of polar contact in the mutant form (right column) in comparison with the wild type protein (left column) (PDB ID=1DC2). (a): The H-bond between P32 and G35 has been missed while this bond exists between L32 and G35 in the wild type form with a length of 2.6 Å. Also, the H-bond length between L32 and V28 is 1.5 Å, but the H-bond length between P32 and V28 in the mutant form is 2.5 Å. (b): At position 42, Asn has H-bonding with S43 (H-bond length=1.9 Å) in the wild type form, but this polar contact is destroyed by replacing Lys at position 42. (c): Gly at position 67 has been H-bonding with N39 (H-bond length=2.4 Å), L63 (H-bond length=1.7 Å) and L64 (H-bond length=2.0 Å). The H-bond with L64 has been missed by replacing Asp. Also, the new H-bond has

been created between D67 and N39 with a length of 2.5 Å, while there is not this bond in wild type. Also, the distance of H-bond with L63 is changed to 1.4 Å by replacing Asp at position 67. (d): There is a polar contact between G122 and A118 (H-bond length=2.6 Å) that this polar contact has been missed by replacing Arg at position 122. (e): At position 126, Val has H-bonding with H123 (H-bond length=1.9 Å) in the wild type form, but this polar contact is destroyed by replacing ASP at position 126.

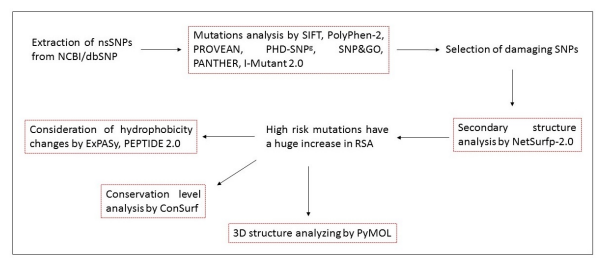


Figure 4. The flowchart of used methods for high-risk mutations identification.

### **Discussion**

Computational be approaches can extremely useful to plan the targeted molecular methods because identifying and studying the SNPs are quite expensive and time-consuming. Also, molecular procedures such as mutagenesis or protein extraction are occasionally impossible. Ou et al. showed that the high-risk SNPs can identified using the combined bioinformatics tools (with 94% sensitivity and 80% specificity) (27). Rajasekaran et al. considered 118 SNPs (in coding and untranslated region) in the CDKN2A gene in malignant melanoma via bioinformatics tools (44). In their study, the pathogenicity of nsSNVs was considered using only PolyPhen. They concluded that the missense variant with dbSNP rs11552822 (D84Y) could be the most deleterious SNP that leads to malignant melanoma (44). This study included a comprehensive investigation identified pathogen nsSNVs in p16 protein

using multiple bioinformatics tools with different approaches regardless of the specific disease. Also, conservation, hydrophobicity changes, and structural alterations of these high-risk mutations were considered. The used tools in this study were SIFT, PolyPhen-2, SNPs&GO, PROVEAN, I-Mutant2.0, PANTHER, PHD-SNPg, ConSurf. NetSurfP-2.0, PyMOL, PEPTIDE 2.0, ExPASy, and HOPE. The flowchart of used bioinformatics tools in this study has been shown in Figure 4. In the present consideration, according to the results of the bioinformatics tools, out of 353 amino acid substitutions, the 18 amino acid substitutions (L16R/Q, G23D/R/S, L32P, N42K, G55D, G67D/R, P81R, H83R, G89D/S, A102E, G101R, G122R, and V126D) have been identified as the highrisk mutations. It is worth mentioning that a constraint on the formation of the  $\alpha$ -helix is the presence of Pro residue, which has the least proclivity to form  $\alpha$ -helixes. In

Pro, the nitrogen atom is a part of a rigid ring, and rotation about the N—C α bond is not possible. Thus, a Pro residue introduces a destabilizing kink in the αhelix. In addition, the nitrogen atom of a Pro residue in a peptide linkage has no substituent hydrogen to participate in Hbonds with other residues. For these reasons, Pro is only rarely found in the αhelix (45).So, L32P amino substitution can have severe effects on the disruption of the protein structure. About V126D amino acid substitution, because of placement of two negatively amino acids next to each other in the V126D (amino acid sequence at positions 125 and 126: DV to DD) and placement of two positively amino acids next to each other in the P81R (amino acid sequence at positions 80 and 81: RP to RR), the helix structures might be unstable.

Correlation of the mutations of the CDKN2A with various cancers such as colon cancer (7), lung cancer (8), melanoma (9), pancreatic cancer (10), leukemia (13-15), glioma (12), HNSCC (11) have been determined. As shown in Table V, the association of L16R, G23D/R/S, L32P, G67R, H83R, G89D, G101R, and V126D amino acid substitutions with melanoma, pancreatic cancer, and HNSCC has been confirmed in previous studies (the related references have been gathered in Table V). It should be mentioned that the UTR regions in regulating protein expression are very important. So, it is suggested that mutations in these regions should be considered with appropriate tools.

### **Conclusions**

In this study, the nsSNVs of the CDKN2A have identified gene been from NCBI/dbSNP databank. the Also. pathogenicity of these nsSNVs was considered using powerful bioinformatics tools. The high-risk mutations were screened step by step via SIFT, PolyPhen-2, PROVEAN, I-Mutant2.0, PHD-SNPg, SNPs&GO, and PANTHER. Then, the secondary structure, amino acid conservation, and feature of amino acids including hydrophobicity, size, and polar contacts of 18 amino acid substitutions in protein were investigated via NetSurfP-2.0, ConSurf, HOPE, ExPASy, PEPTIDE 2.0, and PyMOL. Out of 353 missense variants, the 18 amino acid replacements including L16R/Q, G23D/R/S, L32P, N42K, G55D, G67D/R, P81R, H83R, G89D/S, A102E, G101R, G122R, and V126D were determined as the high-risk mutations. According to the previous studies, there is an association between ten replacements amino acid G23D/R/S, L32P, G67R, H83R, G89D, G101R, and V126D) and some diseases including melanoma, pancreatic cancer, and HNSCC.

### **Conflict of interests**

The authors declare no conflict of interest.

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