

## ***In silico* analysis and 3-D structure prediction of non-synonymous single nucleotide polymorphisms (nsSNPs) in the human *MED12* gene associated with uterine leiomyomas**

### **ABSTRACT**

**Aims.** Uterine leiomyomas are one of the most common benign gynecologic tumors, but the exact causes are not completely understood. *MED12* mutation was discovered in approximately 71% of uterine leiomyomas. Our recent studies confirmed the high frequency of *MED12* exon2 deleterious mutations in uterine leiomyoma in senegalese patients. In this context, molecular dynamics simulations of wild-type and non synonymous variants were conducted to access their structural dynamic and stability characteristics.

**Methodology.** Uterine leiomyoma tissues were obtained from symptomatic women who underwent hysterectomy or myomectomy for medically indicated reasons. We collected 50 uterine leiomyomas and sequenced *MED12* exon2 region after DNA extraction and amplification. To understand the mutation's role in this gene, we utilized computational tools based on different algorithms, including missense tools to predict pathogenic variants, stability tools to analyze the impression of a mutated gene on the function of the protein, and molecular architecture research analysis to detect the gene interactions.

**Results.** A significant genetic alteration of the *MED12* gene has showed with a high frequency of mutations noted in particular codon 44 of exon 2. All these mutations being predicted as deleterious testify to their implication in the pathobiology of uterine fibroids. In addition, the noted alterations lead to instability of the *MED12* protein and thus a change in its biological function in uterine fibroids.

**Key words:** *MED12* ; leiomyomas ; *in silico* prediction ; 3D structure modeling

### **INTRODUCTION**

Uterine fibroids, more commonly known as myomas, are the most common benign tumours of female reproductive organs [1]. They are associated with significant morbidity and constitute a real public health problem. Several studies have also indicated the involvement of hormonal status in the development of uterine fibroids [2,3]. Although the molecular mechanisms involved in their etiology remain completely elucidated, several studies have focused on the *MED12* gene [4]. Component of the Mediator complex, a coactivator involved in the regulated transcription of nearly all RNA polymerase II-dependent genes. Mediator functions as a bridge to convey information from gene-specific regulatory proteins to the basal RNA polymerase II transcription machinery. Mediator is recruited to promoters by direct interactions with regulatory proteins and serves as a scaffold for the assembly of a functional preinitiation complex with RNA polymerase II and the general transcription factors. This subunit may specifically regulate transcription of targets of the Wnt signaling pathway and SHH signaling pathway [5,6].

One the major challenges in modern genetics is predicting the effect of the overwhelming number of variants being revealed through sequencing projects. This is particularly important in analyzing variants occurring in the human population that could be involved in the pathogenesis of disease. Since the structure of a protein is intimately linked to its stability, function and interactions, many *in silico* prediction methods employ knowledge of protein structure, either exclusively or in combination with sequence-based features, with the aim of providing high-quality predictions [7,8,9]. Knowledge of protein structure can be used to predict the phenotypic consequence of a missense variant. It has been noted [3] that in many studies the challenges and costs arise more from the analysis of the data than the actual sequencing. The aims were to provide an analysis describing structurally damaging changes introduced by human missense variants, which can suggest that a variant is likely to be disease causing, and to compare results obtained using experimental and 3D model structures.

## 1. METHODS

### 1.1. Samples

50 patients with uterine fibroids were recruited from the Grand Yoff General Hospital. After obtaining informed consent, each patient was given a biopsy of the tumour tissue and a blood sample (to serve as a control) on an EDTA tube.

### 1.2. DNA Extraction, *MED12* Gene Amplification and Sequencing

The total DNA of each sample was extracted using the Qiagen protocol (Qiagen Dneasy Blood and Tissues kit). After extraction, exon 2 of the *MED12* gene was amplified. Primers forward 5'GCCCTTTCACCTTGTTCTT3' and reverse 5'TGTCCCTATAAGTCTTCCCAACC3' were used for PCR reaction under the conditions previously described by Mäkinen et al., [4].

Sequencing reactions were performed in a MJ Research PTC-225 Peltier thermocycler with ABIPRISM BigDye TM Terminator Cycle kits. Each sample was sequenced using forward primer.

### 1.3. Detection of Mutations

To determine the presence of any mutation and its position relative to the *MED12* gene, the raw sequencing data were submitted to the Mutation Surveyor software version 5.0.1 (www.softgenetics.com), which compares the submitted chromatograms with the reference sequence of *MED12* gene (NT\_011669\_70337906). Non-synonymous deleterious mutations were analysed as previously described by Kenem et al., [10].

### 1.4. Prediction of deleterious nsSNP

The functional impact of variants may lead to a wide range of molecular changes, even within a single protein, including disrupted stability and structure, disrupted macromolecular binding, ablation of post-translational modification (PTM) sites, among others. To help prioritize potentially deleterious variants, groups have developed *in silico* prediction programs. Many of these methods rely primarily or exclusively on sequence-based features, such as amino-acid evolutionary conservation.

- **ClinVar** (<http://www.ncbi.nlm.nih.gov/clinvar/>) provides a freely available archive of reports of relationships among medically important variants and phenotypes [11]. ClinVar accessions submissions reporting human variation, interpretations of the relationship of that variation to

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human health and the evidence supporting each interpretation. For this analysis the reference NP\_005111.2 was used to find clinical significance of nsSNP.

- **ProtParam** computes various physico-chemical properties that can be deduced from a protein sequence [12]. The protein can either be specified as a Swiss-Prot/TrEMBL accession number or ID, or in form of a raw sequence. Using this tool it is possible to compute an instability index. A protein whose instability index is smaller than 40 is predicted as stable, a value above 40 predicts that the protein may be unstable.
- **Missense 3D** can be used to predict the phenotypic consequence of a missense variant [13]. The Mutant and Wild-type structures were analyzed to identify whether the structural consequence of the substitution is expected to be damaging in terms of the stability of the folded protein. Based on well-established principles of protein conformation and previous studies on the structural consequences of disease-associated substitutions, 17 structural features were considered. The aims were to provide an analysis describing structurally damaging changes introduced by human missense variants, which can suggest that a variant is likely to be disease causing, and to compare results obtained using experimental and 3D model structures.
- **MutPred2** is a machine learning-based method, and software package that integrates genetic and molecular data to reason probabilistically about the pathogenicity of amino acid substitutions [14]. This is achieved by providing (1) a general pathogenicity prediction, and (2) a ranked list of specific molecular alterations potentially affecting the phenotype. The loss and gain of structural and functional properties are modelled via posterior probabilities. Through assessment of these probabilities, MutPred can predict the molecular cause of disease-associated substitution. Analysis was done with wild-type protein sequence in FASTA format ID NP\_005111.2 and the substitution sites were identified. The probability of the mutation being deleterious is reported. The final prediction score is the average of the scores from all networks and ranges between 0 and 1; higher scores reflect a higher probability of pathogenicity. Any molecular mechanisms that are likely to be disrupted due to the mutation are reported, with corresponding P value. Functional analysis includes the prediction of DNA-binding site, catalytic domains, calmodulin-binding targets, and post-translational modification sites [15-17].

### 1.5. Prediction effects of nsSNP on protein stability

To see whether the nsSNP of exon 2 may affect the stability of MED12 protein, each mutant was compared to wild type using *in silico* prediction softwares. The Uniprot ID (Q93074 or MED12\_HUMAN) and AlphaFold structure prediction for PDB file (AF-Q93074-F1) were used for analysis.

- **DynaMut2** is a web server that combines Normal Mode Analysis (NMA) methods to capture protein motion and graph-based signatures to represent the wild-type environment to investigate the effects of single and multiple point mutations on protein stability and dynamics [18,19]. DynaMut2 was able to accurately predict the effects of missense mutations on protein stability, achieving Pearson's correlation of up to 0.72 (RMSE: 1.02 kcal/mol) on a single point

and 0.64 (RMSE: 1.80 kcal/mol) on multiple-point missense mutations across 10-fold cross-validation and independent blind tests. For single-point mutations, DynaMut2 achieved comparable performance with other methods when predicting variations in Gibbs Free Energy ( $\Delta\Delta G$ ) and in melting temperature ( $\Delta T_m$ ) [20]. Value binding free energy change ( $\Delta\Delta G$ ) for mutation were calculated in kcal/mol. Calculation of correlation matrix is frequently utilized to illustrate dynamical information of proteins in two dimension. To observe the correlation in the dynamics, correlation matrices for each of the nsSNP were calculated through DynaMut web server.

- **NetsurfP3**, after the analysis of protein modifications, the structural analysis of protein was predicted using the tool NetSurf-3.0 (<http://www.cbs.dtu.dk/services/NetSurfP/>) [21]. This tool works on a neural network algorithm and predicts the secondary structure of amino acids in a sequence, as well as its structural disorder and backbone dihedral angles for each residue in the sequence. This server predicts the surface accessibility with Relative Surface Area (RSA), Accessible Surface Area (ASA, secondary structure, disorder, and phi / psi dihedral angles of amino acids in an amino acid sequence. Amino acid sequence of MED12\_Human was retrieved with its genbank accession no NP\_005111.2 and a length of 2177 amino acids. Each nsSNP was inserted manually to assess its impact on protein function.

## 2. RESULTS

### 2.1 MED12 exon 2 mutations

Out of 50 patients with uterine fibroids, 88% have mutations in exon2 of the *MED12* gene. All single-position mutations affecting exon 2 induce an amino acid change. The frequency of non-synonymous mutations identified are listed in Table 1.

Interpretations of variants in ClinVar show that all nsSNP except F45V, K60M and N61Y are associated with uterine leiomyomas (Table 2).

Table 1. *MED12* mutation frequencies

Mutation position	Amino acid change	Number of patients	Frequency
c.107T>G	p.L36R	1	2%
c.128A>C	p.Q43P	1	2%
c.130G>A	p.G44S	12	24%
c.130G>T	p.G44C	4	8%
c.130G>C	p.G44R	3	6%
c.131G>A	p.G44D	10	20%
c.131G>T	p.G44V	3	6%
c.131G>C	p.G44A	1	2%
c.133T>G	p.F45P	1	2%
c.179A>T	p.K60M	1	2%
c.181A>T	p.N61Y	1	2%

Table 2. Clinical significance of nsSNP

Protein	Molecular consequence	Clinical significance
p.L36R	Missense	Associated with leiomyomas (SCV000109663.1)
p.Q43P	Missense	Associated with leiomyomas (SCV000109676.1)
p.G44S	Missense	Others (SCV000599938.1)
p.G44C	Missense	Associated with leiomyomas (SCV000109681.1)
p.G44R	Missense	Associated with leiomyomas (SCV000109680.1)

p.G44D	Missense	WILMS Tumor (SCV000599933.1)
p.G44V	Missense	Associated with leiomyomas (SCV000109684.1)
p.G44A	Missense	Associated with leiomyomas (SCV000109683.1)
p.F45P	Missense	Uncertain significance
p.K60M	Missense	Uncertain significance
p.N61Y	Missense	Uncertain significance

## 1.2. Effect of nsSNP on protein stability

For each nsSNP, ProtParam results identifies the protein with more number of negatively charged amino acids than positive (Table 3). The physico-chemical annotation of the protein revealed the protein as unstable (instability index > 40).

Table 3. Physico-chemical properties of nsSNP using ProtParam

Protein	Negatively charged residues	Positively charged residues	Instability index
p.L36R	228	218	57.72
p.Q43P	228	217	57.58
p.G44S	228	217	57.93
p.G44C	228	217	57.63
p.G44R	228	218	57.72
p.G44D	229	217	57.78
p.G44V	228	217	57.69
p.G44A	228	217	57.72
p.F45P	228	217	57.72
p.K60M	228	216	57.72
p.N61Y	228	217	57.72

Analysis using Missense 3D show that no structural damage was detected for nsSNP L36P, F45V, K60M and N61Y. It was identified that all residues in codon 44 were identified to fall in disallowed regions (Table 4).

Table 4. Structural damage detected using Missense 3D

Protein	Structural damage detected
p.L36R	No structural damage detected
p.Q43P	Clash (the mutant structure has a MolProbity clash score higher than 30 compared to the wild-type)
p.G44S	Disallowed (the substitution triggers disallowed : the mutant residue is in outlier region while the wild-type residue is in the favoured region)
p.G44C	Disallowed
p.G44R	Disallowed
p.G44D	Disallowed
p.G44V	Disallowed
p.G44A	Disallowed
p.F45P	No structural damage detected
p.K60M	No structural damage detected
p.N61Y	No structural damage detected

The loss and gain of structural and functional properties are modelled via posterior probabilities. Through assessment of these probabilities, MutPred predict the molecular cause of disease-associated substitution. Analysis show that all nsSNP except G44D, K60M and N61Y are associated with molecular mechanism such as gain of intrinsic disorder, altered disorder interface, gain or loss of helix, gain or loss of loop and post-translational modifications such as gain or loss of methylation, loss of ubiquitylation (Table 5).

Table 5. Pathogenicity prediction and molecular mechanism association of nsSNP using MutPred2

Substitution	MutPred2 score	Affected Prosite and ELM motifs	Molecular mechanism with $p \leq 0.05$
p.L36R	0.812	ELME 00146/00147	- Gain of intrinsic disorder ( $p=0.02$ ) - Altered disorder interface ( $p=0.04$ ) - Gain of $\beta$ -factor ( $p=0.03$ ) - Loss of ubiquitylation at K132 ( $p=0.03$ ) - Loss of methylation at K32
p.Q43P	0.550	None	- Loss of helix ( $p=0.04$ ) - Gain of loop ( $p=0.04$ ) - Gain of methylation at K42 ( $p=0.04$ )
p.G44S	0.700	None	- Altered disordered interface ( $p=0.04$ )
p.G44C	0.859	None	- Loss of intrinsic disorder ( $p=0.03$ ) - Altered disordered interface ( $p=0.01$ ) - Gain of helix ( $p=0.05$ ) - Loss of loop ( $p=0.02$ ) - Gain of disulfide linkage at G44 ( $p=0.04$ )
p.G44R	0.787	None	- Gain of helix ( $p=0.02$ ) - Altered disordered interface ( $p=0.05$ )
p.G44D	0.833	None	
p.G44V	0.847	ELME 000146	- Loss of intrinsic disorder ( $p=0.04$ ) - Loss of loop ( $p=0.03$ )
p.G44A	0.736	ELME 000146	- Altered disordered interface ( $p=0.03$ ) - Gain of helix ( $p=0.03$ ) - Loss of loop ( $p=0.05$ )
p.F45V	0.619	None	- Altered disordered interface ( $p=6,7^{e-3}$ ) - Gain of methylation at K42 ( $p=0.04$ )
p.K60M	0.339	None	
p.N61Y	0.397	None	

For all nsSNP analysed, only mutations in codon 44 are considered distabilising using DynaMut2 (Table 6). In comparison with the reference sequence, all the mutations show a conformational change in the 3D structure of the MED12 protein; in other words, there is a deformation of the protein and to a large extent a modification of the biological activity of the MED12 protein in uterine fibroids (Figure 1).

Table 6. Protein stability prediction with DynaMut2

Wild-type	Position	Mutant	Chain	Free energy change ( $\Delta\Delta G$ )	Prediction
L	36	R	A	0.76 Kcal/mol	Stabilising
Q	43	9	A	0.29 Kcal/mol	Stabilising
G	44	S	A	-0.18 Kcal/mol	Distabilising
G	44	C	A	-0.77 Kcal/mol	Distabilising
G	44	R	A	-0.25 Kcal/mol	Distabilising
G	44	D	A	-0.23 Kcal/mol	Distabilising
G	44	V	A	-1.25 Kcal/mol	Distabilising
G	44	A	A	-0.16 Kcal/mol	Distabilising
F	45	V	A	-0.05 Kcal/mol	Distabilising
K	60	M	A	0.21 Kcal/mol	Stabilising
N	61	Y	A	0.29 Kcal/mol	Stabilising

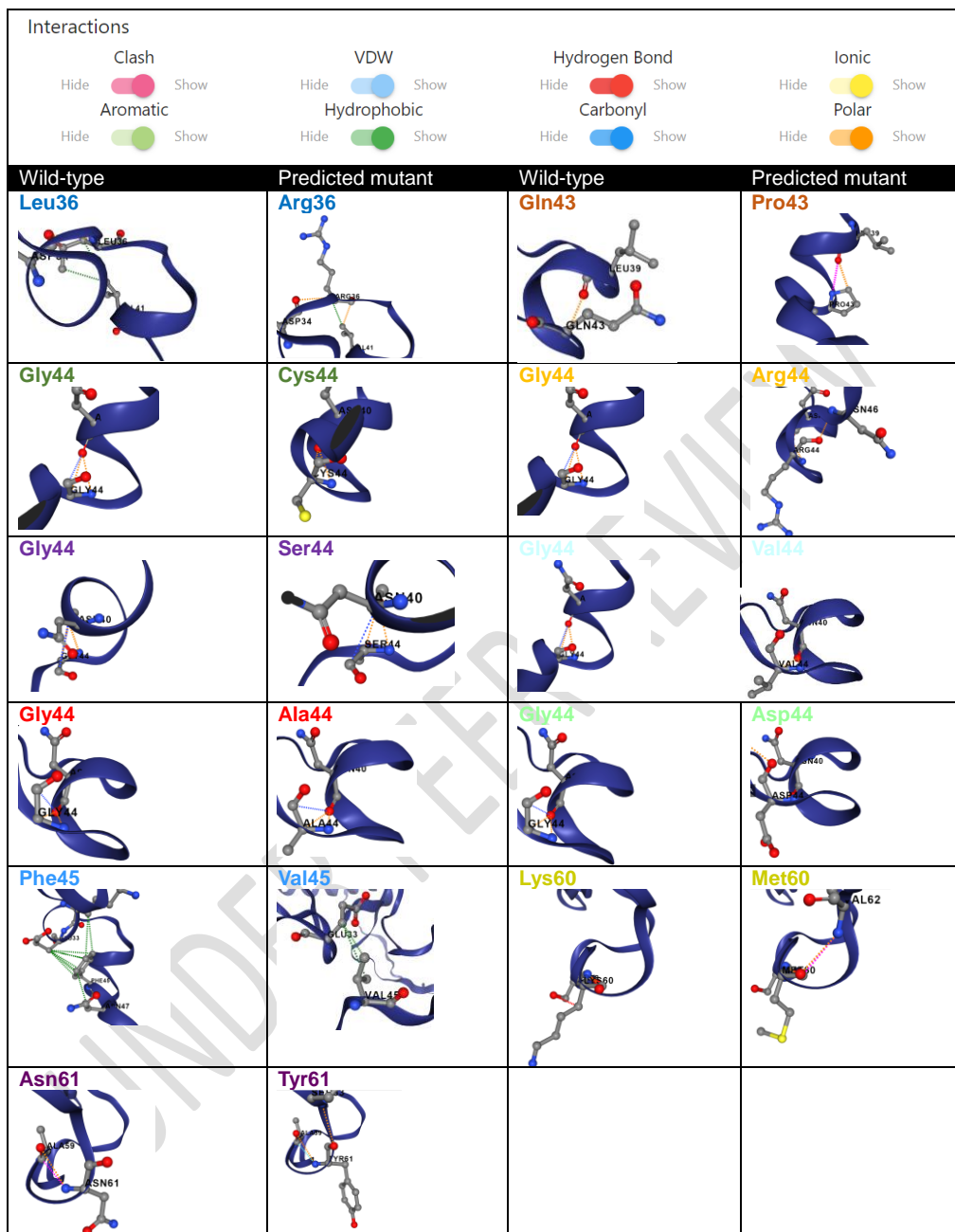


Figure 1. Impact of nsSNP of exon 2 MED12 on 3D-structure of the protein

Normal Mode Analysis (NMA) has been successfully applied to the study of the effects of mutations on protein dynamics (Figure 2).

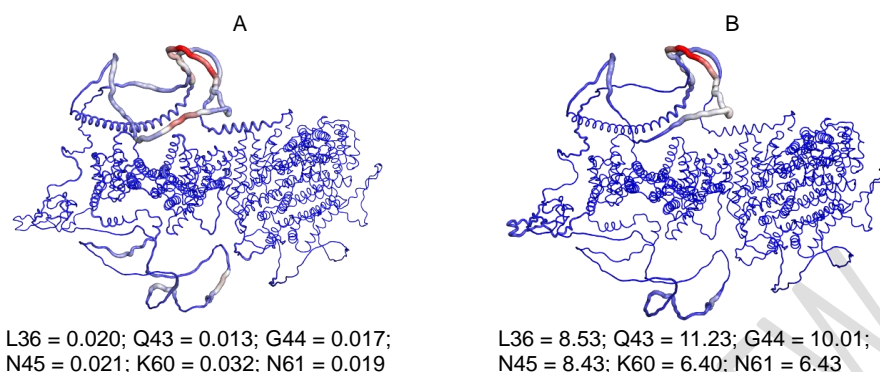


Figure 2. Structural deformation (A) and anatomic fluctuation (B) using Dynamut. The magnitude of the deformation/fluctuation is represented by thin to thick tube colored **blue** (low), **white** (moderate) and **red** (high).

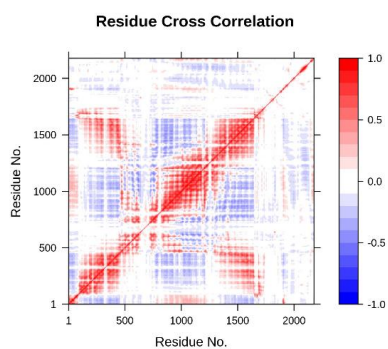


Figure 3. Dynamical Cross-Correlation Map (DCCM) All modes were used to calculate the residue cross-correlation. Correlation map revealing correlated (**red**) and anti-correlated (**blue**) regions in the protein structure.

NetSurfP tool revealed that surface accessibility of these nsSNPs in secondary structure remains affected due to amino acid substitutions (Table 7). Point mutations are in random coil and exposed regions (Figure 4). Below the secondary structure prediction disorder shows the probability of disorder related to that residue and codon 44 is most affected.

Table 7. Structural analysis of protein prediction using the tool NetSurf-3.0

Protein	RSA	ASA	SS	P. disorder
p.L36R	39%	90Å	Coil	68%
p.Q43P	55%	24Å	Coil	67%
p.G44S	23%	27Å	Coil	74%

p.G44C	17%	24Å	Coil	62%
p.G44R	42%	95Å	Coil	83%
p.G44D	40%	58Å	Coil	82%
p.G44V	28%	44Å	Coil	84%
p.G44A	17%	18Å	Coil	68%
p.F45V	32%	49Å	Coil	73%
p.K60M	33%	67Å	Coil	32%
p.N61Y	55%	111Å	Coil	17%

Relative Surface Area (RSA), Accessible Surface Area (ASA), Secondary structure (SS), protein disorder (P disorder)

**Relative Surface Accessibility:** ▲ Red is exposed and blue is buried, thresholded at 25%.  
**Secondary Structure:** 🌀 Helix, 📍 Strand, — Coil.  
**Disorder:** ● Thickness of line equals probability of disordered residue.

NP\_005111.2\_mediator\_of\_RNA\_polymerase\_II\_transcription\_subunit\_12\_Homo\_sapiens

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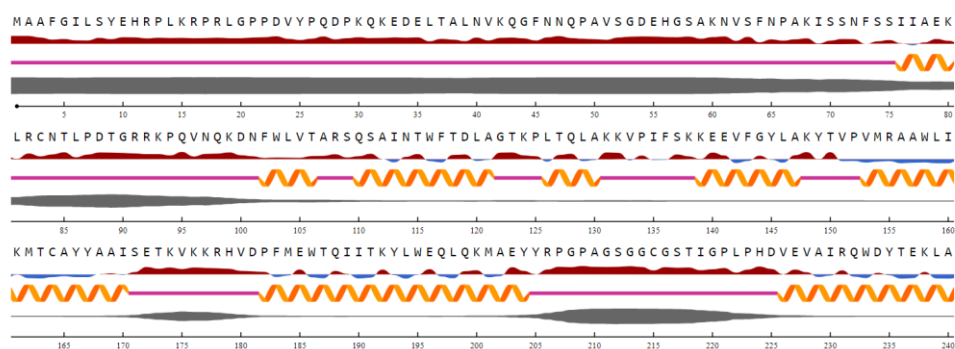


Figure 4. MED12 structural analysis through NetSurf-3.0. Surface Accessibility: The red upward elevation indicates the exposed residue, while the sky blue low elevation shows the buried residue in protein structure.

## DISCUSSION

*MED12*, a gene of 45 exons, located on the X chromosome (Xq13), expressed in all Eukaryotes, forms with Cyclin C, MED13 and CDK8, a subunit of this multiprotein complex. In this study, the *MED12* gene has been investigated in Senegalese women with uterine fibroids.

Analysis of the chromatograms revealed a presence of mutations of the *MED12* gene only in tumour tissues with a frequency of 74% (37/50). This confirms the hypothesis that the *MED12* gene is involved in the occurrence of uterine fibroids. Mäkinen et al. [4] first described the link between *MED12* mutations and fibroids. In 2011, and according to him, mutations in the *MED12* gene represent the largest genetic defect in uterine fibroids. In addition, mutations of the *MED12* gene in tumours other than uterine fibroids are rare. Only 0.3 to 0.5% of colorectal cancers have mutations in the *MED12*

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gene, stating that they are only passenger's mutations [22,23]. 5% of prostate cancers have different MED12 mutations [24,25].

All mutations detected are nsSNP and many of them are associated with uterine leiomyomas by Clinvar. The discovery of pathogenic variants, i.e., variants capable of causing disease, generally relies on a combination of family and population-based sequencing efforts. To assist genetic studies, particularly in characterizing rare variants and dissecting complex disease, machine learning methods have recently been developed to identify the signatures of pathogenicity and to predict the impact of variants of unknown significance. In this context, changes in the biological function of MED12 in uterine fibroids are also highlighted in protein function prediction analysis. Indeed, the mutations of exon 2 seem to induce gains and/or loss of function of the MED12 protein. These modifications constitute a proof of the biological modifications of the MED12 protein in women with uterine fibroids and therefore their implication in the occurrence and/or progression of these tumour cells.

All the mutations affecting exon 2 appear as deleterious mutations, in particular those affecting codon 44. In other words, all the mutations affecting exon 2 cause an aberrant function of the MED12 protein. A study by Bourbon et al. [26], involving 39 different species, showed that codon 44 is the most conserved codon of the *MED12* gene, which states that this codon plays an important role in the biological process of the protein. It has been evidenced that conserved regions are biologically very important, so variations in these regions may lead to potential functional changes. The missense mutations observed on this codon 44 can render the translated protein non-functional, indicating the specific importance of this amino acid for the MED12 function with respect to leiomyoma and suggesting that these mutations could represent alleles gain or loss of function. In Eukaryotes, the Mediator Complex consists of at least 30 proteins [26], structurally divided into four modules, which are the head, the middle, the tail and the kinase modules. The head and middle modules interact directly with RNA polymerase II while the tail module associates with several cofactors to facilitate transcription. The Kinase module interacts with the Mediator Complex to suppress transcription [27]. Indeed, the Mediator Complex exists in 2 forms. The L. mediator form contains 4 modules of the kinase subunit (MED12, MED13, Cyclin C, CDK8 or CDK19) and acts as a receptor. The S. mediator form (without the CDK8 module) stimulates basal transcription. The MED12 domain plays a vital role in connecting Cyclin C-CDK8 to the core of the complex, which activates CDK8 kinase. Moreover, according to the work in reference [27], the binding domain of Cyclin C resides at the level of the N-terminal region encoded by exons 1 and 2 of the *MED12* gene and codon 44 would play a role in this membership. This further confirms the transcriptional activation of MED12 aberrant function in uterine fibroids and that codon 44 is essential for this process.

Study the dynamic nature and the role of flexibility/rigidity and accessible conformational landscapes in proteins is essential for understanding their function, as well as to evaluate how changes in a protein might impact its structure, function and interactions, giving rise to different phenotypes. Our analysis of structural impact of nsSNP show that mutations are located at the random coil. Some regions of the

protein chain do not form regular secondary structure and are not characterized by any regular hydrogen bonding pattern. These regions are known as random coils and are found in two locations in proteins: (i) Terminal arms - both at the N-terminus and the C-terminus of the protein; (ii) Loops - Loops are unstructured regions found between regular secondary structure elements [28]. Most loops are exposed to the solvent and are have polar or charged side-chains. In some cases loops have a functional role, but in many cases they do not. As a result, loop regions are often poorly conserved (i.e. more prone to change) during evolution.

Additionally, the human proteome is dynamic and changes in response to a stimuli, and post-translational modifications are commonly employed to regulate cellular activity. PTMs occur at distinct amino acid side chains or peptide linkages, and they are most often mediated by enzymatic activity [29,30]. Indeed, it is estimated that 5% of the proteome comprises enzymes that perform more than 200 types of post-translational modifications. These enzymes include kinases, phosphatases, transferases and ligases, which add or remove functional groups, proteins, lipids or sugars to or from amino acid side chains; and proteases, which cleave peptide bonds to remove specific sequences or regulatory subunits. Many proteins can also modify themselves using autocatalytic domains, such as autokinase and autoprotolytic domains. Significant gain or loss of methylation had been observed in L36R, Q43P, F45V variations, and gain of disulfide linkage at G44 in G44C. DNA methylation is a key regulator in transcription and altered effect of methylation behavior has been implicated in many diseases like cancer, atherosclerosis, aging etc. [31,32]. Post-translational modification can occur at any step in the "life cycle" of a protein. For example, many proteins are modified shortly after translation is completed to mediate proper protein folding or stability or to direct the nascent protein to distinct cellular compartments (e.g., nucleus, membrane). Other modifications occur after folding and localization are completed to activate or inactivate catalytic activity or to otherwise influence the biological activity of the protein. Proteins are also covalently linked to tags that target a protein for degradation. Besides single modifications, proteins are often modified through a combination of post-translational cleavage and the addition of functional groups through a step-wise mechanism of protein maturation or activation.

## CONCLUSION

Results obtained show a significant genetic alteration of the *MED12* gene with a high frequency of mutations noted in particular codon 44 of exon 2. All these mutations being predicted as deleterious testify to their implication in the pathobiology of uterine fibroids. In addition, the noted alterations lead to instability of the MED12 protein and thus a change in its biological function in uterine fibroids.

The results obtained thus open up avenues for understanding the molecular mechanisms involved in the occurrence and/or progression of uterine fibroids. They also provide a glimpse of treatment strategies because MED12 proves to be an indispensable biomarker in the progression of uterine fibroids.

## Ethical Statements

All procedures on human genetic material and data within this study were performed in accordance with the ethical principles of the local Ethics Committee of the Cheikh Anta Diop University. Informed consent was obtained from all individual participants prior to the inclusion in the study.

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