



Molecular Cloning and Characterization of the Phenylalanine Aminomutase Gene From *Taxus baccata* L.

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ABSTRACT

Background: Taxol is one of the most important anti-cancer drugs, which is obtained from yew trees (*Taxus* sp.). The first step in side chain assembly of taxol is catalyzed by phenylalanine aminomutase, which converts α -phenylalanine to β -phenylalanine.

Objectives: In this study, for the first time, we report on the cloning, preliminary expression and characterization of a full-length gene and cDNA encoding phenylalanine aminomutase from *Taxus baccata* L.

Materials and Methods: Comparison of the full-length gene with other ones identified from the *Taxus* species showed high similarity, particularly with *Taxus x media*.

Results: The results showed that the expression level of this gene in *Taxus baccata* is very low and therefore this enzymatic step could be a rate limiting step in the taxol biosynthesis pathway. Successful amplification of the cDNA was only obtained from RNA samples isolated from methyl jasmonate elicited suspension cells of *Taxus baccata*. The cloned cDNA contained a 2064 bp open reading frame encoding a protein composed of 687 amino acids. Sequence comparison analysis revealed that the gene is very similar (98 - 99 %) with respect to the nucleotide and amino acid sequences in different *Taxus* species and also share the signature active site motif (175ASG177).

Conclusions: The predicted structure of TbPAM was analyzed using bioinformatic tools. The results indicated that the protein has similar overall folding to tyrosine aminomutase.

Keywords: Molecular Cloning; Phenylalanine Aminomutase; Taxol; *Taxus baccata* L.

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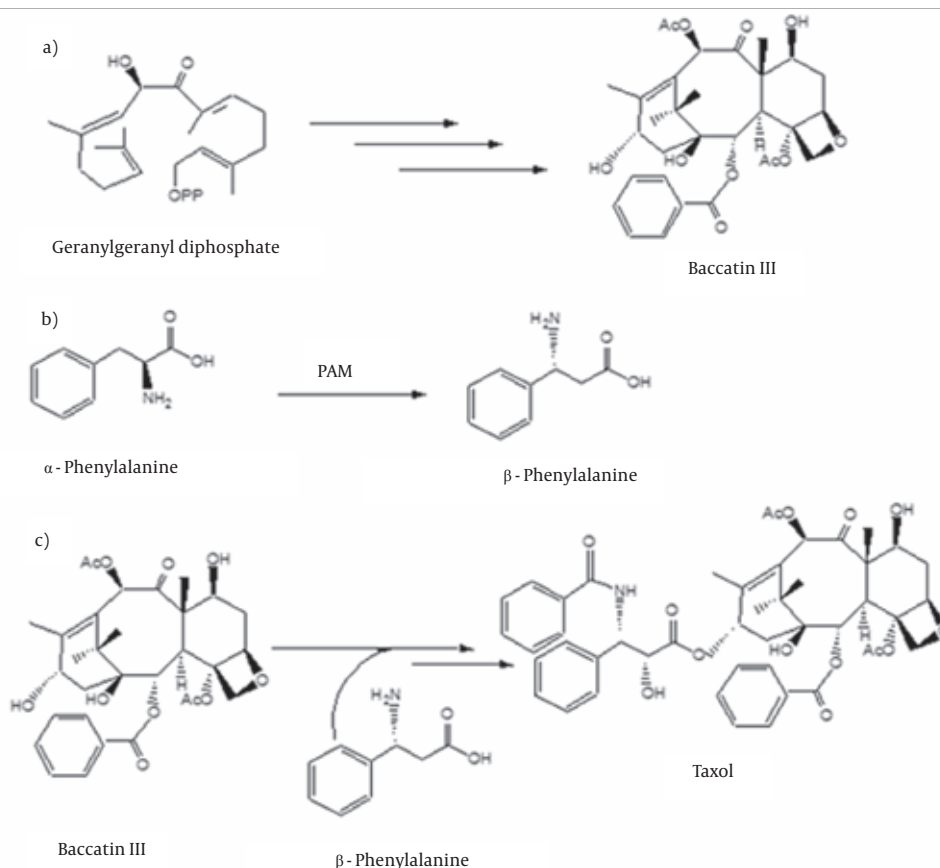
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1. Background

Taxus species are known to produce a wide range of natural diterpenoids known as taxoids (taxanes), with approximately 350 identified forms. One of the best-known taxoids, taxol, has been well proven to be a potent anticancer drug with powerful effects against a range of cancers. The production of taxol from the original plant source is very limited due to the low abundance and slow growth of *Taxus* trees and the low concentration of taxol in the trees (1). On the other hand, demand for the drug has increased considerably due to the expansion of clinical trials and treatments (2). Therefore, it seems that the only way to increase the yields of taxol and other clinically important taxoids by genotypes, is to study and manipulate the biochemical and enzymatic reactions in the taxoid pathway (3). This is a precondition for the eventual over expression of the slow steps and/or silencing of the subsidiary steps of the pathway (3). Generally, the taxol molecule consists of two components (Figure 1): the taxane ring moiety (10-deacetyl baccatin III) derived from the terpenoid pathway and C-13 side chain derived from the phenylpropanoid pathway (4). Since, the taxane ring moiety is much more readily available

than taxol itself (4), the low content of taxol may be related to its side chain assembly. The taxol biosynthesis pathway contains about 20 distinct enzymatic steps (2). Tremendous progress has been made in the cloning and functional expression of the genes involved in the pathway. The first step in the side chain assembly of taxol is catalyzed by phenylalanine aminomutase (PAM) which converts α -phenylalanine to β -phenylalanine. The activity of the enzyme was demonstrated in the soluble fraction of crude cell-free extracts of *T. brevifolia* (4). Feeding studies showed that β -phenylalanine is incorporated into both phenylisoserine and benzoate moieties of the side chain (4). Thus, PAM may limit the rate of taxol production at the commercial scale and could be an important target for genetic engineering in yew and its derived cell cultures to increase the taxol productivity. On the other hand, characterization of the enzyme is essential to understand the nature of taxol side chain biosynthesis (5). PAM gene from *Taxus cuspidata* and *Taxus chinensis* has been cloned and functionally expressed in *Escherichia coli* (4, 5). Until now, there is no report on the cloning of PAM gene from *T. baccata*. This species contains comparatively lower content of taxol and higher content of baccatin III in its needles than the other *Taxus* species (6).

Figure 1. Taxol Biosynthesis Pathway



a) biosynthesis of taxane ring moiety from the terpenoid pathway, b) conversion of α -phenylalanine to β -phenylalanine by phenylalanine aminomutase (PAM), c) attachment of C-13 side chain to the Baccatin III and biosynthesis of Taxol.

2. Objectives

In this paper, we described the cloning and characterization of the PAM gene from *T. baccata* (TbPAM) as an initial step to investigate the physiological role of the gene in this species.

3. Materials and Methods

3.1. Materials

General molecular biology reagents were obtained from Fermentas (Germany), Merck (Germany) or Sigma (St. Louis, MO), unless otherwise indicated. Enzymes for PCR and RT-PCR were purchased from Roche (Roche Applied Science, Germany). Cloning of the full-length gene and cDNA were carried out using pTZ57R/T vector (InsTAclone™ PCR Cloning Kit) from Fermentas (Germany). Plant growth regulators, nutrients and other plant tissue culture materials were purchased from Duchefa (Biochemie B.V., Netherlands).

3.1.1. Plant Material, Media and Culture Conditions

Young leaves and stems were collected from adult *Taxus baccata* trees at the flower garden of Isfahan in Isfahan, Iran. Callus was obtained from the leaf explants in B5 medium containing 2 mg.L⁻¹ NAA and 0.2 mg.L⁻¹ kinetin in dark condition as previously described (7). In order to prepare cell suspension culture of *Taxus*, three grams of two-month old callus were added to 50 mL of the same liquid B5 medium in a 250 mL flask. The flask was incubated for 21 days in darkness at 25°C, shaking at 110 rpm. The culture was elicited at day 14 by methyl jasmonate with a final concentration of 100 µM. The elicited cells were separated from the remaining medium at day 21 by centrifugation at 5000 rpm for 10 min and used for RNA isolation procedure.

3.2. DNA and RNA Isolation

Total RNA was extracted from the elicited cell suspension using a special method for *Taxus* species (3). Briefly, the cells were ground to a fine powder in liquid nitrogen and added to 10 mL of preheated extraction buffer containing a high concentration of β-mercaptoethanol (4%) and PVP (4%). The homogenate was then extracted with chloroform: isoamylalcohol (24:1 v/v), 3 times. The upper phase was carefully mixed with a 0.5 volume of 3 M sodium acetate (pH 4.8), incubated at -20 °C for 30 min and centrifuged. The supernatant was mixed gently with 1/4 volume of 10 M LiCl and placed at 4°C overnight. The RNA was pelleted by centrifugation, dissolved in 100 µL DEPC-treated water, and then treated with RNase-free DNase I (Fermentas). The homogenate was extracted twice with chloroform: isoamylalcohol and the RNA was pelleted using ice-cold absolute ethanol, and finally dissolved in DEPC treated water. Genomic DNA was extracted using

a CTAB (hexadecyltrimethylammoniumbromide)-based method (3) and used for cloning of the full length PAM gene. The quality and concentration of RNA and DNA samples were examined by agarose gel electrophoresis and spectrophotometer (ND-1000, USA) analysis.

3.3. PCR and RT-PCR

The PCR was done in a total volume of 25 µL containing 2 mM MgCl₂, 10 pmol of each primer of PAM (Forward primer: 5'-TTTGAATTCATGGGGTTTGCCGTGGAATC-3'; reverse primer: 5'-TTTGGTACCCTAGACGCCGTGGCGCA-3'), 200 µM of each dNTPs, 1 unit Taq DNA polymerase (Roche Applied Science, Germany), and 100 ng of genomic DNA in 1x PCR buffer. The PCR amplifications were carried out using a MyCycle™ Thermal cycler (Bio-Rad, USA). Each PCR was initiated by a preheating step for 3 min at 94°C followed by 30 cycles consisting of a denaturation step (30 sec at 94°C), primer-annealing step (30 sec at 61°C), and elongation step (3 min at 72°C). The final elongation step was extended for 8 min. The first-strand cDNA synthesis was done using AMV reverse transcriptase with Oligo-(dT)₁₇ primer and approximately 2 µg of total RNA according to the manufacturer's instructions (Roche Applied Science, Germany). The second strand amplification was carried out using high fidelity PCR system (Roche Applied Science, Germany) and the PCR conditions similar to that explained before. The PCR products were then analyzed by gel electrophoresis.

3.4. Cloning of Full-Length cDNA and Genomic Gene

The PCR product of appropriate size was gel purified and extracted carefully using a DNA Extraction Kit (Fermentas). The purified fragment was then cloned into the pTZ57R/T vector according to the manufacturer's instructions (InsTAclone™ PCR Cloning Kit, Fermentas). The vectors were introduced into *E. coli* XL1-Blue cells by electroporation with a Gene Pulser Xcell™ apparatus (Bio-Rad, USA) and the cells were cultured overnight at 37°C in Luria Bertani (LB) medium containing 50 µg.mL⁻¹ ampicillin. The positive clones were selected based on lacZ gene marker and finally sent for sequencing to the Macrogen Company (Seoul, Korea).

3.5. Preliminary expression and Analysis of Recombinant Protein

The recombinant plasmid containing PAM cDNA was co-transformed into *E. coli* BL21 (iDE3) cells (Merck Biosciences, Schwalbach, Germany) and cultivated at 30°C overnight in 5 mL LB medium containing 100 µg.mL⁻¹ of ampicillin. The overnight culture was then inoculated into 500 mL of the same medium and cultivated at 30°C. The culture was grown at 30°C to A₆₀₀ = 0.5 – 0.6, and then induced by addition of 0.5 mM isopropyl β-D-thiogalactopyranoside (IPTG) to express the PAM gene. The cells were grown for another 4 h at 30°C and finally the presence of the respec-

tive protein was detected using SDS-PAGE (12% Laemmli gels). Protein concentrations were determined on the basis of the calculated extinction coefficient at 280 nm: 51185 M⁻¹ cm⁻¹.

3.6. Bioinformatics Analysis

BLAST was used to investigate for homology in the GenBank database and the Gene Runner program was used to deduce the amino acid sequence of the gene. Multiple sequence alignments were performed using the DNASTAR Lasergene core suit software. Analysis and characterization of the protein were performed using protparam belonging to the Swiss Institute of Bioinformatics [http://www.expasy.ch/tools/protparam.html/]. Secondary structure of TbPAM was analyzed by the Swiss-pdb viewer tool (version 4.01). Tertiary structure of TbPAM was predicted by using phyre webserver (Version 0.2) and Swiss Model 8.05 [http://www.expasy.org/swissmodel/].

4. Results

4.1. DNA and RNA Isolation

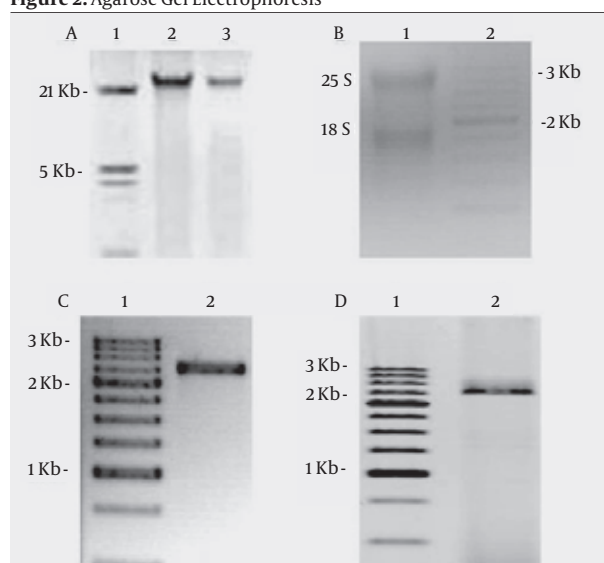
Isolating high-quality samples of nucleic acid is a critical step in many molecular biology experiments especially in cDNA synthesis of low expressed genes or rare mRNAs. Degradation and contamination of the RNA samples during the isolation process may cause the failure of the following experiments. Different tissues of *Taxus* contain high levels of secondary metabolites especially phenolic compounds that are rapidly oxidized and vigorously bond nucleic acids during isolation and finally result in brown and insoluble nucleic acid pellets. The presence of such powerful oxidizing agents in the samples may result in decreased purity and productivity of the final extracted nucleic acids. Agarose gel electrophoresis of nucleic acids also showed good quality of DNA and RNA samples with no considerable polysaccharide, polyphenol and protein contaminations (Figure 2). The results indicate that high quality DNA and RNA samples with appropriate yields can be obtained using the protocol.

4.2. Cloning and Sequence Analysis of the Full-Length Gene

For identification and amplification of the full length of the PAM gene from *Taxus baccata*, forward and reverse primers were designed based on highly conserved 5' and 3' terminal sequences of ORF regions of the PAM genes in other *Taxus* species. Digestion sites of *EcoRI* (under lined sequence) and *KpnI* (bold sequence) restriction enzymes were also added to the ends of primers for cloning the gene in the vector (pASK-IBA43plus) for subsequent experiments. In order to improve the digestion efficiency, four thymine nucleotides were also added to the 5'-ends of primers. As a result of these nonspecific nucleotides

in primers, the annealing temperature was reduced considerably. PCR reaction using the primers and *Taxus baccata* genomic DNA (as template) resulted in the specific amplification of a single fragment of about 2250 bp (Figure 2) that was cloned into the vector and sequenced subsequently. Investigating the sequence of the cloned fragment in Gene Bank showed high similarity to the sequences of PAM genes identified in other *Taxus* species. Moreover, specific amplification of the fragment in the PCR reaction confirmed the assumption that the 3'- and 5'-ends of PAM gene from *Taxus baccata* are similar to those reported previously from other *Taxus* species. The cloned full-length of PAM gene from *Taxus baccata* was 2233 bp in length (designated as TbPAM, Genbank accession number: GU214709) and contained an exon within the gene from nucleotide 1092 to 1260. Searches against Gen Bank using the BLAST algorithm revealed that the similarity between TbPAM and the gene obtained from *Taxus x media* is more than that from *Taxus canadensis* and *Taxus chinensis* with respect to the sequence, length and exon site. Since *Taxus x media* originated from the hybridization of *T. baccata* and *T. cuspidate* (8), such a result was predictable.

Figure 2. Agarose Gel Electrophoresis



A, *Taxus baccata* Genomic DNA (1: Marker and 2, 3: Isolated DNA); B, RNA isolated from *Taxus baccata* (1: 1 µg total RNA and 2: Marker); C, PCR product (1: Marker and 2: Full length of Phenylalanine aminomutase gene (2233 bp)); D, RT-PCR product (1: Marker and 2: Phenylalanine aminomutase cDNA (2064 bp)).

4.3. Cloning and Sequence Analysis of the TbPAM cDNA

Isolating RNA samples from the native tissues and also non-elicited cell suspensions of *Taxus baccata* to use in RT-PCR reactions for synthesis of PAMcDNA, didn't result in any amplified product and all of the experiments failed. This may be related to the low expression level of the gene in *Taxus baccata* L. Moreover, the lower productivity of taxol

in *Taxus baccata* compared to the other taxus species may also be related to the low expression of the PAM gene in this species and perhaps PAM would be a rate limiting enzyme in taxol biosynthesis pathway. However, more studies on the taxol biosynthesis pathway and the functions of the enzyme in the pathway could be useful in this context. Several studies have previously emphasized on the elicitor effects of methyl jasmonate on taxol synthesis from culture suspension of *Taxus* cells (9). Wang et al. (9) presumed that methyl jasmonate may activate the biosynthesis pathway from baccatin III to taxol and inhibit biosynthesis from baccatin III to cephalomannine. Based on this assumption, it seemed that methyl jasmonate is a suitable choice to induce the expression of PAM gene and consequently to increase the probability of gene amplification. Therefore, we isolated RNA samples from *T. baccata* cell suspensions, which were induced to result maximal taxol synthesis with 100 μ M methyl jasmonate at day 14 of culture. RT-PCR reactions by using these RNA samples resulted in amplification of a single fragment with length of about 2050 bp (Figure 2). Sequencing result of the fragment (Figure 3) coincided with the sequences from ORF region of PAMcDNAs identified

previously from other *Taxus* species (AY582743, AY724735). The cloned cDNA of TbPAM (Genbank accession number: GU214708) was 2064 bp in length. Nucleotide BLAST of the sequence from the NCBI website indicated that the gene shows only very minor sequence differences (almost as silent mutation) with the other known sequences of *Taxus* PAM (99% similarity with *Taxus chinensis* and *Taxus x mediana* and 98% with *Taxus canadensis*). Since all *Taxus* species are known to be very closely related and are considered to represent geographic variants of the basic species, *T. baccata*, such a result had been anticipated. The protein contained 687 amino acids (Genbank accession number: ADA57703) with a molecular weight of about 75.25 kDa and an isoelectric point of 5.7. As also shown by Figure 4, amino acid sequence of TbPAM had high similarity (98%) with the other known *Taxus* Phenylalanine aminomutases (AAU01184, AAU01183, and AAU01185). The minor differences are likely attributable to the species differences or to allelic variations. The secondary structure of TbPAM was analyzed by Swiss-pdb viewer tool (version 4.01) and the results showed that the putative TbPAM peptide contained 51% alpha helix, 7% beta sheet, and 42% random coil.

E S K S R V R D I L G D I N I F N E V R K I I

GTCGCGGCGCTGGCCCGGAGGCATGACGTGAAGGTTGCGTTGGAGGCGGAGCAATGCAGAGCCCGTGT
V A A L A R R H D V K V A L E A E Q C R A R V

GAAGACGGCGCCGACATATACGGCGTCACCACGGGCTTCGGCGCGTGCCTCGAGCCGGAGGACCAACCA
E D G A D I Y G V T T G F G A C S S R R T N Q

CTGCTCGCGGGGGTGTCTTACTAAAGGATGCGCTTCCTCCGTCGACGAGCTCCCCGCGACCGCCACCCG
L L A G V F T K G C A S S V D E L P A T A T R

TATGGATGTTCCGGCATCCGGTGGGAGGTCATGGAAGCGCTGGAAGGCTTCTCAACAGCAATGTCTC
Y G C S G I R W E V M E A L E K L L N S N V S

TCGGGAGACCTCATCCCGCTCGCCTACATTGCAGGGCTCCTGATCGGGAAGCCTAGCGTAATCGCTCG
S G D L I P L A Y I A G L L I G K P S V I A R

GCGTTGAGCAGGGTGGGGCTGCGGCCATTCAAGCTCCAGGCCAAAGAAGGGCTGGCGCTCGTCAACGG
A L S R V G L R P F K L Q A K E G L A L V N G

ATGTACGACGCCAATGTTCTGTTGCTGCTCGTGAAACGCTTTGCGGAATGTTCTGCGAGGTGATCTT
M Y D A N V L L L L V E T L C G M F C E V I F

AAAGTGAAGCGCCACCCGGGCCAGATCGAATCGCGGAGCTGCTCGAGTGGCTGCTGCGGTGCGAGCCC
K V K P H P G Q I E S A E L L E W L L R S S P

GATAAGCTGAAGAAACCGAAACAGGATCGCTATGCTCTGAGGTGCGAGCCCGCAGTGGTTGGCTCCTCT
D K L K K P K Q D R Y A L R S S P Q W L A P L

GAGACGGAGGTCAATTCCGCCAATGATAACCCCATCATTTGACCACGCCAATGACAGGGCTCTCCATGG
E T E V N S A N D N P I I D H A N D R A L H G

ATGGACTACGTGCGCATCGCAGTCGCCGGGCTGGGGAACTCTTGTTCGCTCAGTTCACGGAGCTGAT
M D Y V R I A V A G L G K L L F A Q F T E L M

CTCTCCCTGGGGCCGGACCTGAGCGTGGACTACGGCCTCAAGGGGCTCGACATCGCCATGGCCGCTTA
L S L G P D L S V D Y G L K G L D I A M A A Y

ACCACACACGTTTACAGCGCGGAACAGCACAACCAGGACATCAACTCTCTGGCGCTCATCTCCGCCCG
T T H V H S A E O H N O D I N S I A I T S A R

Figure 3. The Full-Length cDNA Sequence and Deduced Amino Acid Sequence of *Taxus baccata* Phenyl Alanine Aminomutase

Figure 4. Multiple Sequence Alignment of the *Taxus baccata* Phenylalanine Aminomutase With the Other Genes From PAL/HAL Family

80

157 GDTDTGVTTGFGATSHRRTS--QGVELQKE LIRFINAGVFGS-CEGNV--LPEATTRAAMLVRTNTLPQGpal g

155 GDTDTGVTTGFGATSHRRTN--QGAELQKE LIRFINAGVFGK-CPENV--LSEDTTRAAMLVRTNTLLQGpal pm

133 GADIVGVTTFGAGCSRRTN--Q LSEIQES LIRCLLAGVFTKGCASSVDLPATATRSAMLRLNSFTYGpam ch

115 GADIVGVTTFGAGCSRRTN--Q LSEIQES LIRCLLAGVFTKGCASSVDLPATVTRSAMLRLNSFTYGpam xm

61 NIPYGVTTGFGEMITYMQVDKSKEVELQTNLVRSHSAGVGPL-----F--AEDEARAIVAARLNTLAKGTAM

76 GADIVGVTTFGAGCSRRTN--QLSELQES LIRCLLAGVFTKGCASSVDLPATATRSAMLRLNSFTYGpam tb

175-177

222 YSGIRWALLETIEKLLNAGITPKLPLRGTITASGDLVPLSYIAGLLTGRPN SKVRTRDGTMSGLEALKQpal g

220 YSGIRWDILETVEKLLNAGITPKLPLRGTITASGDLVPLSYIAGLLTGRPN SRVRSRDIEMSGAEALKKpal pm

201 CSGIRWEVMEALEKLLNSNVSPKVPLRGSVSASGDLIPLAYIAGLLIGKPSVIARIGDDVEVPAPEALSRpam ch

183 CSGIRWEVMEALEKLLNSNVSPKVPLRGSVSASGDLIPLAYIAGLLIGKPSVIARIGDDVEVPAPEALSRpam xm

123 HSAVRPIILERLAQYLNEGITPAIPEIGSLGX--DLAPLSHVASTLIGEGYVLR--DGRPVETAQVLAETAM

144 CSGIRWEVMEALEKLLNSNVSPKVPLRGSVSASGDLIPLAYIAGLLIGKPSVIARIGDDVEVPAPEALSRpam tb

214231

292 VGLEKPFELQPKKEGLAIVNGTSVGAALASIVCFDANVLA VLS EYMSAMFCEVMNGKPE-FTDPLTHRLKHpal g

290 VGLEKPFELQPKKEGLAIVNGTSVGAALASIVCFDANVLA LLS EVISAMFCEVMNGKPE-FTDPLTHRLKHpal pm

271 VGLR-PFKLQAKEGLALVNGTSFATAVASTVMYDANVLL L LVE TL CGMFCEVIFGREE-FAHPLIHVKVPpam ch

253 VGLR-PFKLQAKEGLALVNGTSFATALASTVMYDANVLL L LVE TL CGMFCEVIFGREE-FAHPLIHVKVPpam xm

188 RGIE-PLELRFKEGLALINGTSGMTGLSLV VGR ALEQAQAEIVTAL LIEAVRGSTSPFLAEGHDIARP TAM

214 VGLR-PFKLQAKEGLALVNGTSFATALASTVMYDANVLL L LVE TL CGMFCEVIFGREE-FAHPLIHVKVPpam tb

321-325

361 HPGQMEAAAIM EYVLDGS----SYMKAQAKLQELNPIQKPKQ-----QRYALRTSPQWLG PQVEVIRApal g

359 HPGQMEAAAIM EYVLDGS----SYMKAAKIHENMNLQKPKQ-----QRYALRTSPQWLG PQVEVIRSpal pm

339 HPGQIESAE LLEWL LRS----PFQEL SREYYSIDKLKPKQ-----QRYALRSSQWLA PLVQTIRDPam ch

321 HPGQIESAE LLEWL LRS----PFQEL SREYYSIDKLKPKQ-----QRYALRSSQWLA PLVQTIRDPam xm

257 HEGQIDTAANMRA LMRGSGLTVEHAD LRRELQKDKEAGKDVQRSEIVLQKAYSRLAIPQVVGAVRDTLYHTAM

282 HPGQIESAE LLEWL LRS----PFQEL SREYYSIDKLKPKQ-----QRYALRSSQWLA PLVQTIRDPam tb

371

420 ATHMIEREINSNDNPVIDVSRDKALHGGNFGGTPIGVSM DNLR LRLSIAAIGKLMFAQFSELVNDY YNGGLpal g

418 ATHMIEREINSNDNPVIDVARDKALHGGNFGGTPIGVSM DNLR LRLSIAAIGKLMFAQFSELVNDY YNGGLpal pm

358 ATT TVETEVSANDNPIIDHANDRALHGANFGQSSAVGFYMDYVRIA VAGLGKLLFAQFTELMIEYYS NGLpam ch

380 ATT TVETEVSANDNPIIDHANDRALHGANFGQSSAVGFYMDYVRIA VAGLGKLLFAQFTELMIEYYS NGLpam xm

327 ARHKRIELNSANDNPLFFEGKE-IFHGANFHGQPIAFAMDFVTIALTQLGVLAERQINRVLNRHLSYGLTAM

341 ATT TVETEVSANDNPIIDHANDRALHGANEFGQSSAVGFYMDYVRIA VAGLGKLLFAQFTELMIEYYS NGLpam tb

427

490 PSNLSGGPNPS LDYGLKGpal g

488 PSNLSGGPNPS LDYGLKGpal pm

468 PGNLS LGPDL SVGYGLKGpam ch

450 PGNLS LGPDL SVGYGLKGpam xm

396 PE FLVSG-DPGLHSGFAGTAM

411 PGNLS LGPDL SVGYGLKGpam tb

Pal g, *Ginkgo biloba* Phenylalanine Ammonialyase (ABU49842); pal pm, *Pinus massoniana* Phenylalanine Ammonialyase (ACS28225.2); pam ch, *Taxus chinensis* phenylalanine aminomutase (AAU01183); pam xm, *Taxus x media* phenylalanine aminomutase (AAU01184); TAM, Tyrosine Aminomutase (ABY66005); pam tb, *Taxus baccata* phenylalanine aminomutase (ADA57703). Proposed active site (175-177) and binding sites (80, 214, 231, 322, 325, 371, 427 the numbering on the boxes based on pam tb sequence) showed in the boxes.

4.4. Predicted Structure and Active Site of TbPAM

Similarity searches using the BLAST algorithm indicated that the amino acid sequence of TbPAM is most similar (up to 98%) to phenylalanine ammonia-lyase (PAL). Among the other aminomutases, Tyrosine Aminomutase (TAM) showed most similarity (70%) to TbPAM with respect to the amino acid sequence. Based on PAL-derived TAL activity in monocots and fungi (10) and also high structure similarity between phenylalanine and tyrosine, it seems that the binding mechanism and mode of action could be similar in PAM and TAM. Based on this assumption and the scientific literature on the crystal structure of TAM (11), probable protein structure of TbPAM was predicted by using the phyre2 server from the Centre for Bioinformatics Imperial College, London (Figure 5). The predicted structure of TbPAM adopts a predominantly α -helical fold that is linked together by β -sheets. Such a structure could be essential for flexibility and plasticity of the enzyme and seems to be a functional requirement for substrate binding and catalysis. Data indicated that TbPAM has the same overall protein fold as tyrosine aminomutase (Sg-TAM) that was reported by Montavon and coworkers (11). Moreover, superposition of the theoretical structure of

TbPAM with TAM and PAL indicated significant homology where TbPAM contains a 4-methylidene-imidazole-5-one (MIO) cofactor. The MIO moiety formed in the active site of the enzyme by spontaneous (autocatalytic) cyclization and dehydration of an internal 175Ala-Ser-Gly177 tripeptide motif was similar to TAM and histidine/phenylalanine ammonia lyase (PAL/HAL) family (Figure 4). These similarities suggest a common mechanism in the vicinal interchange catalyzed by PAM and TAM, and also in the elimination of ammonia catalyzed by the related lyases. Furthermore, the amino acids involved in active site of SgTAM exactly conserved in TbPAM.

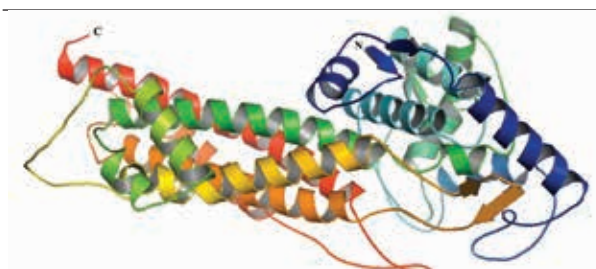


Figure 5. Predicted Structure of Phenylalanine Aminomutase from *Taxus baccata* L.

5. Discussion

Study and manipulation of the enzymes involved in metabolic pathways are considered as one of the most efficient methods to improve productivity and also to alter pathways for production of novel compounds. Study of substrate specificity and enzymatic chemistry is an essential step towards this aim. Aminomutases that are classified as isomerases, catalyze the vicinal interchange of an amino group and hydrogen either via radical-dependent homolytic or ion-dependent heterolytic mechanisms. *Taxus baccata* PAM (TbPAM) catalyzes the conversion of α -phenylalanine to β -phenylalanine; the first committed step in the biosynthesis of N-benzoyl phenylisoserinoyl side-chain of the anticancer drug, taxol. PAM revealed a multifunctional activity which catalyzes the isomerization of 2S- α -phenylalanine to 3R- β -phenylalanine with retention of configuration at C3, reversely the conversion of both R- and S- β -phenylalanine to the same 2S- α -phenylalanine, and also the α,β -elimination of ammonia from the α -isomer to form trans-cinnamic acid (4). Recent mechanistic studies on the aminomutases (PAM and TAM) revealed that 4-hydroxycinnamic acid (in the case of TAM) and cinnamic acid (in the case of PAM) are intermediates in the aminomutase reaction that can be released from the active site of the enzyme. These observations indicate that TAM and PAM might exhibit ammonia lyase activity and suggest the possibility of using the second half of the aminomutase reaction to synthesize β -amino acids from cinnamates (12). Comparison of the amino acid sequence of TbPAM with PAL and TAM indicate that the enzyme also contain a signature Ala-Ser-Gly motif which rearranges auto-catalytically to the MIO moiety in the active site of the enzyme. Unlike the aminomutases that require external cofactors, PAM activity depends on a MIO-derived cofactor. Based on crystallographic experiments on the MIO-dependent tyrosine aminomutase, Christanson et al. (13) suggested that MIO reacts with the amino group of the substrate during the initial deamination step of the aminomutase reaction. It seems that the α -amine of the substrate (α -phenylalanine) adds into the electrophilic moiety (MIO) via a conjugate addition that facilitates deprotonation of the β -hydrogen. In this condition, the carbon-nitrogen bond is broken and the MIO-NH₂ bond and α,β -unsaturated carboxylic acid are formed. Then the MIO-bound amine adds on at the β -position while these intermediates retain in the active site of the enzyme. Finally, the product (β -phenylalanine) is ultimately released and the MIO group regenerated. Walker et al. (4) revealed that potassium cyanide and sodium borohydride both eliminate the function of the MIO in PAM similar to ammonia lyases. The result suggests that the MIO moiety is essential for the activity of PAM similar to TAM and PAL. Comparison of PAM from *T. baccata* to those previously acquired from other *Taxus* species revealed that these enzymes are very identical (99 %) at the amino acid level and they share, as expected,

the signature active site motif 175Ala-Ser-Gly177. This may be related to the essential role of the enzyme in the taxol biosynthesis pathway. Similarity searches using the amino acid sequence of TbPAM indicated that the enzyme is most similar (up to 98%) to phenylalanine ammonia-lyase (PAL) that catalyzes the first committed step in the phenylpropanoid biosynthetic pathway. PAM and PAL share several enzymatic properties specially the lack of cofactor requirements and monomeric molecular weight (14) that suggest a common mechanism in the action of these enzymes. The higher *K_m* for PAM (1100 μ M) than PAL (29-260 μ M) indicates that if the phenylpropanoid and the taxoid pathways were in direct competition for substrate then the phenylpropanoid pathway would be favored (14). More importantly, the high *K_m* for PAM suggests that PAM may be a rate-limiting enzyme in the biosynthesis of taxol (5). Based on the points mentioned above and the relative abundance of some precursors such as baccatin III compared to taxol in the *Taxus* tissues, it seems that PAM catalyzes a rate limiting enzymatic step in taxol biosynthesis pathway and is considered as an important target for genetic engineering in yew. Over expressing the enzyme or genetic engineering to increase the affinity of the enzyme to the substrate in *Taxus*, may improve the biosynthesis of taxol. On the other hand, competition between PAM and PAL for α -phenylalanine may be an important factor that decreases the efficiency of PAM.

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Authors' Contribution

Abolghasem Abbasi Kajani: experimental procedures, preparing the manuscript, Mohammad Reza Mofid: bioinformatics analysis, the manuscript preparation, Khalil Alami Saeid: experimental procedures.

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