

Amplification, Cloning and *In silico* Prediction of Full Length Elicitin Gene from *Phytophthora capsici*, the Causal Agent of Foot Rot Disease of Black Pepper

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Abstract

Elicitins are a family of small proteins secreted by *Phytophthora*, which induce leaf necrosis in infected plants. Here, we report the cloning of Elicitin gene from *P. capsici*, an Oomycete plant pathogen which causes significant damage to a broad range of host plants. Elicitin sequence was amplified using primers designed from the known Elicitin genes of other *Phytophthora* organisms based on their conserved motifs. The PCR amplified product size of 256 bp length and the BLAST analysis of the sequenced product showed perfect match with alpha-elicitin sequences of *P. capsici*. Subsequently, attempt was made to characterize the complete gene of elicitin from genome sequence information of *P. capsici*, by querying the amplified product against the genome. Local BLAST search against full genome identified entire coding sequence. Further sequence analysis identified promoter sequence, transcription start site, a leader signal sequence and a core elicitin domain, with a conserved 6 Cysteine residues. In addition, the three dimensional structure of capsicein was modelled, and the binding affinity of sterol and capsicein was studied using molecular docking. The developed model predicted strong binding affinity for Tyr 47.

Keywords: Elicitin; *P. capsici*; Cloning; *In silico*

Introduction

The genus *Phytophthora* comprises over 100 phytopathogenic species, although 100-500 undiscovered species are estimated to exist [1]. *Phytophthora* are classified as oomycetes, a diverse group of fungus-like eukaryotes that share phylogenetic similarity with brown algae and diatoms [2,3].

A common feature of many different types of plant pathogens is the secretion of a variety of extracellular effectors or elicitor molecules into the plant apoplast [4]. Many of these proteins, called elicitors elicit plant defense responses, and, in particular, a form of programmed cell death called the hypersensitive response (HR). *Phytophthora* species ubiquitously secrete a unique class of highly conserved effector molecules named elicitins. Elicitins are widespread in *Phytophthora* species and closely related *Pythium* species [5]. Elicitins are low molecular weight proteins (10 kDa) secreted into liquid minimal medium [6]. Two classes of elicitins have been identified; alpha-elicitins, which are acidic and induce only necrosis, whereas beta-elicitins are basic and also induce distal necrosis [7]. Acidic elicitins (capsicein and parasiticein) are reported from *P. capsici* and *P. parasitica* respectively, while basic-elicitin (cryptogein and cinnamomin) are from *P. cryptogea* and *P. cinnamomi*, respectively. Five classes of elicitin have been defined from the primary structure. Class IA and Class IB encompass elicitin with only the elicitin domain and are 98-amino acid-long proteins. Class II contains HAE (hyperacidic elicitin), with a short hydrophilic C-terminal tail and 103-104 amino acid long ORFs. Class III contains elicitin sequence about 165-170 amino acid long that consist of 98-amino acid elicitin sequence and ~70 amino acid long C-terminal domain, which represents an O-glycosylated domain. Elicitins from *Pythium* spp. have been classified into a distinct *Pythium* spp. group [8]. The elicitin from *P. capsici* known as capsicein belong to class IA.

These various types of elicitor molecules induce biochemical changes as part of the resistance response. Electrolyte leakage, oxidative

burst, production of phytoalexin and PR proteins, and increased biosynthesis of ethylene have been described in leaf tissue treated with non-specific elicitors [9], and with specific elicitors. Treatment of suspension cultured tobacco cells with elicitins from *P. megasperma* leads to rapid protein phosphorylation, Ca²⁺ influx, extracellular and transient H₂O₂ production, alkylinization of the extracellular medium, acidification of the cytosol, lipid peroxidation, gene expression, disruption of microtubular cytoskeleton and cell wall modification [10,11].

Elicitins are also sterol carrier proteins which bind sterols and catalyze their transfer between membranes [12]. The initial complex of elicitin binding to sterols was confirmed by the crystal structures of cryptogein in complex with dehydroergosterol (DHE) [13]. Binding of elicitin to ligands appears to be essential for induction of a biological response in plants [14]. Interaction between the protein and ergosterol involves several residues, among which tyrosine residues are the most denoted [15].

Although, elicitin gene has been cloned and sequenced from *P. capsici* infecting *Nicotiana glutinosa* [16], there is no report on cloning of elicitin from *P. capsici* infecting black pepper. Despite the identification of elicitin gene from *P. capsici*, transcription start site (TSS) and promoters for the elicitin (capsicein) have not been clearly identified.

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In this paper, we report the amplification, cloning and complete gene information of elicitin gene from *P. capsici* infecting black pepper. In addition, we used the complete genome sequence information of *P. capsici* to identify the Transcription Start Site (TSS) and promoters. Further, the docking analysis revealed the binding mode of capsicein and ergosterol at atomic level. Data presented here provide the first *in silico* structural description of alpha elicitin-sterol interaction of *P. capsici* infecting black pepper.

Materials and Methods

Culturing *Phytophthora*

Ten mm discs were cut from 4-day old cultures of *Phytophthora capsici* (culture nos: 98-177, 03-10, 05-14, 06-17, 96-03, 96-09, 96-10, 98-185, 05-03, 97-11 and 99-144 grown in carrot agar, and transferred to 100 ml of Ribeiros medium amended with 100 µl of thiamine solution (1 mg/ml). This was allowed to grow for five days, and then the grown up mycelium was used for DNA extraction.

Nucleic acid extraction

DNA was isolated from the following isolates *viz.* 98-177, 03-10, 05-14, 06-17, 96-03, 96-09, 96-10, 98-185, 05-03, 97-11 and 99-144, using protocol [17]. The mycelium was first dried on a sterile filter paper and 100 mg of mycelium was crushed using glass powder in sterile microfuge tubes, using a micropestle in STE buffer (1 M Tris-cl pH 8, 5 M NaCl, 0.5 M EDTA and 10% SDS). Phenol:Chloroform:Isoamylalcohol (Sigma) extraction was done followed by isopropanol precipitation to get the DNA, and later the DNA was purified using RNase (5 mg/ml) (Fermentas). The DNA was finally quantified using biophotometer (Eppendorf).

Oligonucleotide primer designing

For primer designing, the elicitin nucleotide sequence from *P. infestans* was used initially to do a Blastn database search for homologous elicitin genes in other *Phytophthora* species. Multiple sequence alignment was performed using ClustalW to align the sequences, and to locate well conserved regions. Primers were designed based on the conserved regions and primer properties were studied using OligoCalc.

PCR amplification and cloning

Elicitin genes from *P. capsici* were amplified by polymerase chain reaction (PCR), using primer ELICPHYF6 and ELICPHYR6: AGCATCCTCTCGGACTCGT, CGAGAACCCGTTTCGCGTA. About 100 ng of DNA was taken for PCR amplification along with 2 µl of 10X taq buffer (Bangalore Genei), 1 µl of forward and reverse primers, 10 picomole (Sigma Genosys), 0.3 µl of Taq polymerase (Bangalore Genei) 1.5 units, 0.75 µl of 10 mM dNTP (Fermentas) in a total volume of 25 µl. The reaction conditions were 94°C for 2 minutes initial hold, and then, 94°C for 30 seconds, 55°C for one minute, 72°C for 1 minute for about 30 cycles, and a final extension at 72°C for about 10 minutes. The amplified fragment was cloned in a T/A cloning vector (Fermentas). Both the clones, as well as the PCR product from all the isolates, were sequenced using Sanger sequencing at SCIGENOM, Cochin, India. The sequencing results were analysed through BLAST programme.

Search for full length elicitin (capsicein) gene

A local BLAST search against the complete genome of *P. capsici* was made to search for the complete elicitin gene sequence using Bioedit [18]. The Indian initiative of sequencing the complete genome of *P. capsici* was taken up by Indian Institute of Spices Research (IISR),

funded by Indian Council of Agricultural Research (ICAR). For this, a native isolate of *Phytophthora* (Is. No. 98-93) infecting black pepper was completely sequenced using next generation sequencing platform, Illumina-Solexa GA II. The complete genome of *P. capsici* was taken as a reference, and the amplified elicitin sequence was queried against the complete genome using Bioedit. The positive match was annotated to find the portion coding for elicitin gene.

Sequence analysis of capsicein

An upstream gene search was done to look for the promoters and transcription start site, using Neural Network Promoter Prediction tool [19]. The complete CDS of elicitin sequence was subjected to protein domain analysis, using Interproscan [20], and disulfide bonds were found out using Disulfind tool [21]. The probable signal sequence was predicted using SignalP [22]. To further consider the relationship between capsicein and elicitins from different *Phytophthora* species, a phylogenetic analysis of 26 elicitins and elicitin like sequences were carried out by the neighbour joining method using MEGA5 [23].

Homology modelling of capsicein

Certain Class IA and Class IB elicitins bind sterols and lipids, and act as sterol carrier proteins. Thus, in order to study the sterol binding activity, an attempt was made to model the three dimensional structure of capsicein. Homology modelling of capsicein was performed using Modeller 9v8. Beta-cryptogein chain A (PDB id: 1BXM) was used as template to generate the three-dimensional structure of capsicein. Elicitins also behave like sterol carrier proteins. Hence, to predict whether *P. capsici* elicitin can bind to ergosterol, the docking program Molegro Virtual Docker (MVD) [24] was used. The capsicein protein was prepared by MVD and ergosterol was cleaned and optimized using Argus Lab. The 3-D structure of docked complex and the binding mode was analyzed using MVD.

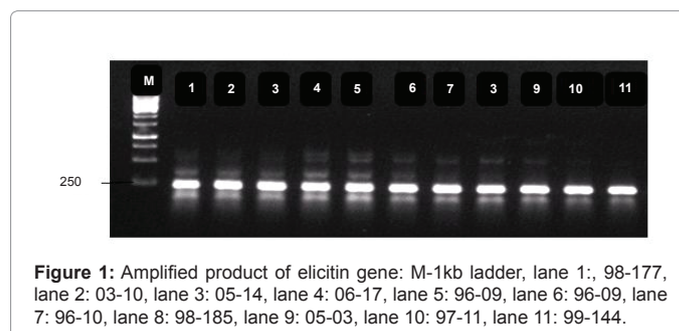
Results

Cloning of elicitin (capsicein) sequence

The PCR amplified product size was about ~256 bp (Figure 1). BLAST analysis of the sequenced product showed perfect match with alpha-elicitin sequences from *P. capsici* present in the NCBI database. The sequence was deposited in NCBI and the accession number is Accession No: JF495405. The blast analysis of the sequence (Accession no: JF495405) revealed a perfect match with alpha elicitin capsicein mRNA from *P. capsici* showing 96% homology.

Determination and analysis of full length elicitin (capsicein) gene

Local BLAST search of amplified product against complete genome of *P. capsici* showed a perfect match to Scaffold 38. Further annotation



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1  TTCITTTQQTAA YVALVLSLS DSSFNQCATD SGYSMLTATA LPTTAQYKLM CASTACNTMI
61  SKIVSLNPPD CELTVPTSLG VLNVYSYANG FSATCASL
    
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Figure 2: *Phytophthora capsici* elicitin complete CDS.

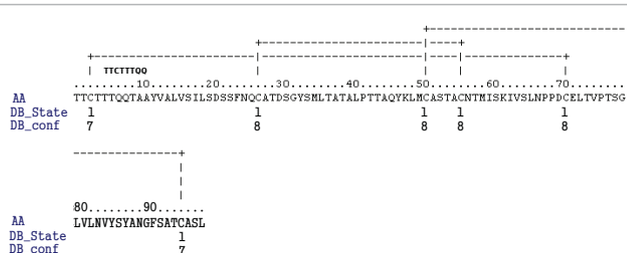


Figure 3: Cysteine disulfide bonding state and connectivity in *P. capsici* elicitin.

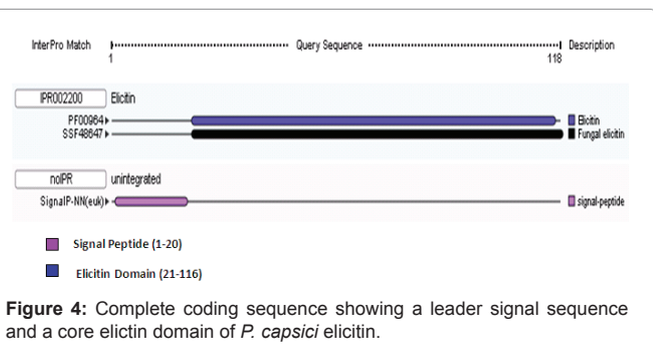


Figure 4: Complete coding sequence showing a leader signal sequence and a core elicitin domain of *P. capsici* elicitin.

of the matched region identified the portion coding for elicitin gene. Translation of the coding region gene identified 118 amino acid length elicitin gene, without any introns (Figure 2).

BLAST search of the protein sequence showed a perfect match to alpha elicitin of *P. capsici*. The predicted isoelectric point was 4.68 and the N-terminal portion has Valine at position 13. Disulfid tool predicted the cysteines disulfide bonding state and connectivity as follows: Cys3–Cys71, Cys27–Cys56 and Cys51–Cys95, which is characteristic of elicitin (Figure 3) [14,25]. InterproScan identified a typical elicitin domain of length 95 amino acid length from T²¹ to A¹¹⁶, which lacked protein kinase C-dependent phosphorylation sites found in the carboxy-terminal region of basic elicitins (Figure 4) [26]. Like other elicitins, capsicein contains 10 Leu, 6 Cys and 3 Met, and lacks Trp, His and Arg. Capsicein comprise only a signal peptide and an elicitin domain.

TSS analysis and promoter prediction

Alignment of relative TSS of various elicitin genes indicates a conserved amino acid pattern between the TSS of alpha and beta and HAE elicitins (Figure 5) [26]. Consensus TSS pattern of alpha elicitin [GCCATTGTGCA] was searched in the upstream portion of elicitin gene, and the transcription start site of capsicein was located 50 base pairs upstream of first start codon.

Further, sequence analysis was performed upstream the TSS to locate the promoter. However, sequences related to TATA boxes were found 900-930 bp upstream of TSS (Figure 6).

β-CRY	CTCATTCTGCAATTTGC <i>ccagtcac</i>
β-CIN	CTCATTCTGCAATTTgGC <i>tccgctcac</i>
α-CRY	GCCATTGTGCAATTTGC <i>tctgtcac</i>
α-CIN	GGCATTGTGCAATTTGC <i>tctgtcgc</i>
α-PAR	GCCATTGTGCAATTTGC <i>tctcatcc</i>
HAE-CRY1	CTCATTCTCAATTTcC <i>cttgccaa</i>
HAE-CIN1	CGCACTCCACAATTTGa <i>aatttgcc</i>
HAE-CRY2	CTCACTCCACAATTTGC <i>cttgccaa</i>
HAE-CIN2	TTCATTCTGCAATTTGC <i>tttgccga</i>

Figure 5: Conserved amino acid pattern of TSS of elicitins. TSS is underlined.

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Promoter Sequence
1  CTGATTATCTATATGTCGTTGGCAATCCGCTGGTGGCCAGGTAAGTTTTCTTTTCAAA
61  ATGAAGCGCGCAAAACAACATAGAAACCAAGAGCAGTCAAAAATCTTCTAGGTGCACCTTGA
121 TGACAAAGTGGCATTGGCAACCAACAGATAAAGCCGCTGCACCAACCGCTTGAAGCAACGG
181 GTCCCTCAAGCAGAATCGAGTGGCGGTGATTTCGTGTAACAACAGTGAAGCTAGAGTGGCA
241 TGATCTCTGTCACTGGCAGAGCTTCCAGGCTGGAATCCTGCATATCCTCAATCGAACT
301 CGTTACGAAGCAGTGGCACACAGTGGAGCTCTGTGACCATGTCACTCATGGAGAAACTG
361 GCTTGGACAGGCGTGTGCTCCAGCAACGTCAGCTGCCAGCCATCCTGGTCACTGCGCAT
421 CCTCGTTAGTGAATGTATGAACTCCTCTAALAACAAAACTACTACATGTACRGTAC
481 AATGTAAGAGCTGTGCACTCTTAAGAGCACTTCACTGCTTCTCGAAGTCTTTCGGCT
541 GTCAATGCATCTGCTGCTGGCATTCAACAGCGCTGAGAATGAATAGAGCCCTATTGACGA
601 ACCCGAAGATACTCCTGATGGTGGCACTTCGATGGATCGGACCCCTCTCTGGTAAAT
661 GTGCTTCTCTGCTGCAGAACAGGTCGCTTCCTCTCTCTGCTCCTCGGAGGATACAGG
721 CGTTACAGCTTTGGTGTCTTGTGACTGTGCAACTCTCCGTAAGTCCGCGAAGCGGAG
781 CGGGTACATGTACGATGGTGCATGCTTGGCGTCCGATTCGGACTCCGAGGCGGCAAGG
841 CCGCCGATGCCACTGCCACAGCAGTGTCTCCATGGTCTACTCAAGATTCCTGACTCAG
901 GACTGCTGTCTCTAGCTaat tgtgcaATTGGCCCGCTCACCAGACCTTCCACTCC
961 CACCCACCCCTCCCAATGAACTTCCGTGCTGTGTGCGGCTACCGTCCCGCCCTC
Start codon      Signal Sequence
M N F R A L F A A T V A A A
1021 GTGGCTCCACTTCCGCGCACACAGTGCAGGACCCAGCAGACCGCGCGTACGTCGCGC
[V G S T S A ] T T C T T T Q Q T A A Y V A
1081 CTGTGGAGCATCCTCTCGGACTCGTTCGTTCAACAGTGGCGCACGACTCTGGCTACG
L V S I L S D S S F N Q C A T D S G Y S
1141 ATGCTGAGCGCTACGGCGCTGCCACGACGGCGCATCAAGCTCATGTGGCGTCCGAG
M L T A T A L P T T A Q Y K L M C A S T
1201 GCGTCAACAGGATGATCTCCAGATGTGTGCTGAAACCCCTGACTGGGAGCTGAGG
A C N T M I S K I V S L N P P D C E L T
1261 GTGCCACGAGCGGCTGTGCTCAACGTTACTCTGTAAGGACCGGTTTCTCGGCGCAG
V P T S G L V L N V Y S Y A N G F S A T
1321 TGCGCGTGGCTGTA
C A S L *
Stop codon
    
```

Figure 6: Complete gene characterization of capsicein. Transcription Start site is underlined. The transcription initiation consensus sequence is indicated in lowercase.

Phylogenetic analysis

Phylogenetic tree of the elicitin from *P. capsici* and 25 other elicitin and elicitin like sequences from other oomycete species is shown in Figure 7. Phylogenetic analysis of elicitins shows a clear classification into five groups based on their domain characterization.

Homology modelling and docking studies

Certain Class IA and Class IB elicitins bind sterols and lipids, and act as sterol carrier proteins [27]. Thus, in order to study the sterol binding activity, an attempt was made to model the three dimensional structure of capsicein. The modelled three dimensional structure of elicitins composed of two antiparallel sheet and five helices (Figure 8a). The overall quality of the structure analysed by Ramachandran plot, confirms 97.7% residues in the allowed regions (Figure 8b). The main elements of secondary structure of capsicein are a two-stranded,

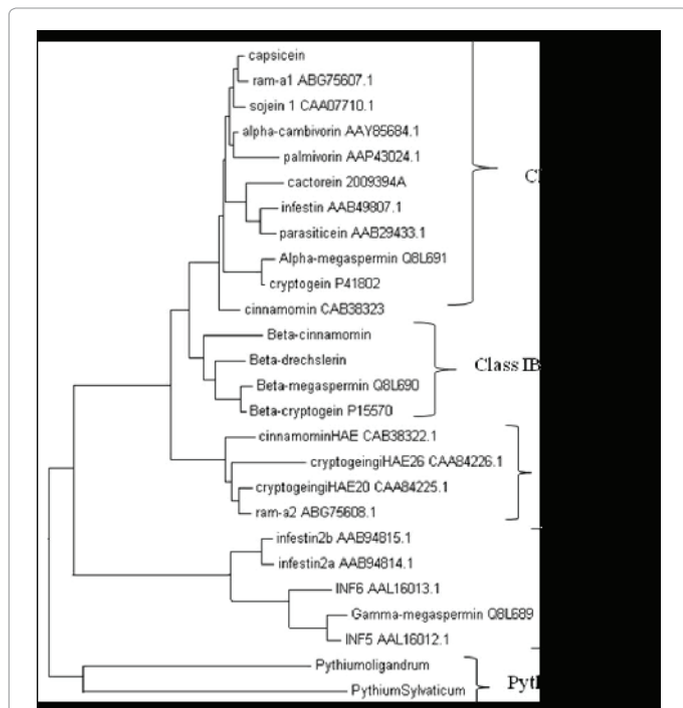


Figure 7: Phylogenetic analysis of the elicitin family proteins from *Phytophthora* spp. and *Pythium*.

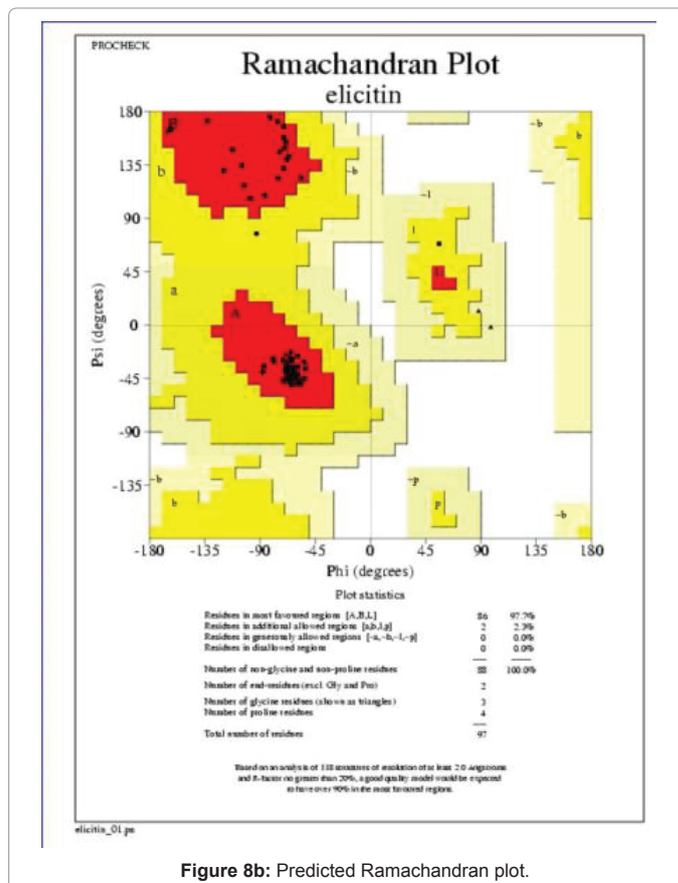


Figure 8b: Predicted Ramachandran plot.

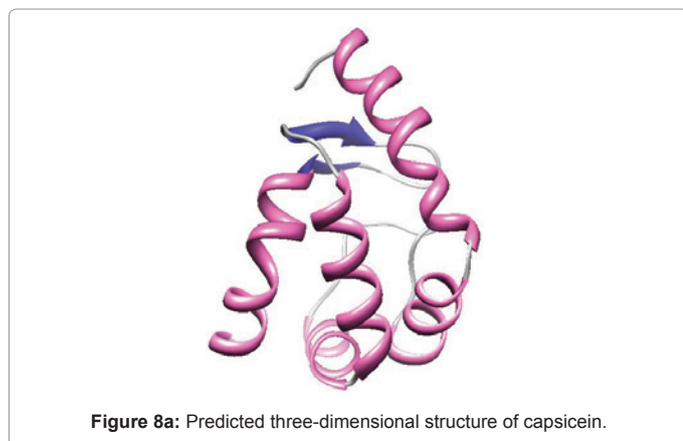


Figure 8a: Predicted three-dimensional structure of capsicin.

antiparallel sheet, consisting of residues Leu 73-Thr 74 and Val 81-Leu 82. Five helices consisting of residues Thr5-Ser20, Ser22-Ser31, Thr44-Ala52, Thr54-Leu 66 and Val84-Ser97 from N to the C terminus.

Ergosterol binds to the groove located inside the cavity of capsicin. Figure 9 shows the hydrogen bond between ergosterol and Tyr 47, correctly identified by docking.

Ergosterol bound to the cavity interacts mainly with the residues making a groove: Tyr 47, Tyr 12, Leu 41, Ile 63, and Met 35. Among which Tyr 47 binds with the least MolDock score of - 118.514 (Table 1), and hence, has better binding affinity for elicitin, and is very important.

Discussion

Despite the importance of *Phytophthora* species as devastating plant pathogens, the molecular mechanism that determines the outcome of interactions between *Phytophthora* and plants is to be fully elucidated.

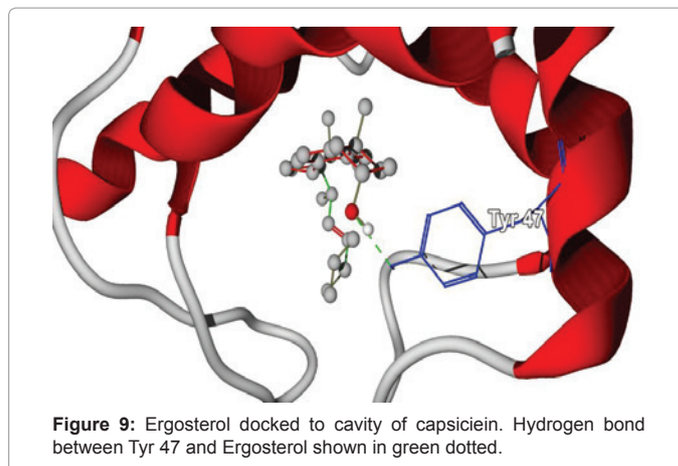


Figure 9: Ergosterol docked to cavity of capsicin. Hydrogen bond between Tyr 47 and Ergosterol shown in green dotted.

Many plant pathogens secrete a variety of elicitor molecules into plant apoplast [4]. Most of the *Phytophthora* species and some *Pythium* species ubiquitously secrete a unique class of highly conserved effector molecules named elicitors, which elicit plant defense responses, and in particular, a form of programmed cell death called hypersensitive response (HR) [5]. Here, we report the cloning and *in silico* prediction of full length elicitin gene from *P. capsici*, the causal agent of foot rot disease of black pepper.

A 256 base pair PCR amplified product was cloned, sequenced, and BLAST analysis of the sequence showed perfect match with alpha-elicitin sequences from *P. capsici*. This sequence was compared to the

S. No.	Residue	MolDock Score
1	Tyr 47	-118.514
2	Met 35	-114.682
3	Ile 63	-109.994
4	Leu 41	-107.507
5	Tyr 12	-102.917

Table 1: MolDock Score of residues of capsicein interacting with ergosterol.

genome sequence information data of *P. capsici* by setting up a local BLAST search, which indeed showed perfect match with Scaffold 38 of the genome sequence data. Annotation was done further and matching region identified as the portion coding for elicitin gene. Translation of the coding region identified a 118 amino acid length elicitin gene, which lacked any introns in it. However, splicing probably occurs in some genes of *P. infestans* and *P. parasitica* [28,29]. Intron-exon boundary analyses indicated that splicing in *Phytophthora* is similar to other higher organisms [30].

BLAST search of the protein sequence showed a perfect match to alpha elicitin of *P. capsici*. The predicted isoelectric point of 4.68 and N-terminal sequence with Val at position 13 are consistent with the class acidic α elicitins [31]. Amino acid residue 13 is correlated with necrotic activity of elicitins. The beta elicitins contain a basic lysine residue, which is more necrotic, whereas alpha elicitins contain a hydrophobic valine residue at this position [32]. Though alpha and beta elicitins differ in their levels of necrotic activity, they induce similar levels of resistance [33]. The likely 20 amino acid length signal sequence was predicted to be M¹ to A²⁰. The signal peptide is highly conserved 18-20 amino acids sequence. Elicitins are synthesized as pre-proteins, which undergo post translational modifications through the removal of signal peptides [5,8]. Disulfid tool predicted the cysteine disulfide bonding state and connectivity characteristic to elicitin. The even number of cysteine residues can induce defence response in plants, and disulfide bridges formed by the cysteine residues are essential for HR induction and avirulence function [34,4]. The disulfide bridges might enhance stability in the plant apoplast, known to be rich in degradative proteases [35]. InterProScan identified a typical elicitin domain of length 95 amino acid length, which lacked protein kinase C-dependent phosphorylation sites found in the carboxy-terminal region of basic elicitins [26]. Like other elicitins, capsicein contains 10 Leu, 6 Cys and 3 Met, and lacks Trp, His and Arg. All elicitins share a 98 amino acid elicitin domain, with a conserved six cysteine residues. Capsicein comprise mainly a signal peptide and an elicitin domain. This class I elicitins are typically abundant, soluble, secreted proteins in mycelia cultures, and have the ability to transfer sterols [12], or bind phospholipids [14].

Elicitin genes, like other genes of *Phytophthora*, share a consensus transcription start site (TSS) [36,37]. TSS pattern of alpha elicitin [GCCATTGTGCA] was searched in the upstream portion of elicitin gene, and the transcription start site of capsicein was located 50 base pairs upstream of first start codon. Further, sequence analysis was performed upstream the TSS to locate the promoter. Sequences related to TATA boxes were found 900-930 bp upstream of TSS. The hypothesis put forward here, concerning the promoter is that the promoters of capsicein are found 900-930 bp upstream of TSS.

Phylogenetic analysis of elicitins from different *Phytophthora* spp. shows a clear classification into five groups based on their domain characterization. The proteins belonging to Class IA and Class IB appeared on separate cluster in the tree, confirming the division of elicitins as acidic and basic [8]. The highly acidic elicitins, having a short hydrophilic C-terminal tail, appeared on significantly distinct

branch as class II. The INF2A, INF2B, INF6, INF5 and γ -megaspermin clustered as Class III. Class III elicitins consists of elicitin domain and O-glycosylated domain. Elicitins like proteins belonging to *Pythium* groups formed a separate cluster in the phylogenetic tree [8].

Elicitins act as sterol carrier proteins [27]. Homology modelling and docking studies showed how sterol binds to the groove located inside the cavity of capsicein. In this work, a possible link between the elicitor activity of capsicein and its efficiency to load sterols, with the use of molecular docking were performed. Interaction between the protein and ergosterol involves several residues, among which, tyrosine residues are the most significant [15]. In latest reports [38], it indicates that Tyrosine-47 residue is involved in the sole hydrogen bond between the protein core and the sterol. Moreover, the sterol seems to contact its aromatic ring. Replacement of tyrosine residues significantly affected both the interaction rate and the equilibrium concentration of the elicitin-sterol complex. Mutation of tyrosine to phenylalanine increases the steric hindrance, and results in less stable complex. *Phytophthora* and *Pythium* do not synthesize sterols, required for their reproduction [39]. Elicitin acts as a shuttle, carrying sterols from the host. Moreover, elicitin-sterol complex formation is required to trigger defense mechanism [38], in plant. The information gained from this study, while limited, provides useful possibilities for further research on alpha elicitins.

Acknowledgments

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