



CLONING AND CHARACTERIZATION OF GENES ENCODING TWO DETOXYFING ENZYMES, GLUTATHIONE S-TRANSFERASE AND CARBOXYLESTERASE, FROM BURROWING NEMATODE (*RADOPHOLUS SIMILIS*)

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Abstract- Background: *Radopholus similis* (Cobb) Thorne is a migratory endoparasitic nematode infesting several tropical and sub-tropical plant species. Computational screening and conserved domain annotation of assembled EST sequences from the burrowing nematode (*R. similis*) revealed seven contigs similar to glutathione S-transferases (GSTs) and four contigs for carboxylesterases / cholinesterases (CESs). One of the contigs corresponding to each gene were cloned from *R. similis* cDNA (KM670018, KP027005) and characterized by phylogeny and structural motif comparison. Glutathione S-transferase is a critical antioxidant and detoxification enzyme and carboxylesterase is responsible for controlling the nerve impulse, detoxification and various developmental functions. Detoxifying ability of these proteins makes them as major targets of pesticides used for plant parasitic nematode (PPN) management.

Methodology: In the present work, both molecular biology and bioinformatics approaches have been used to study two potential target genes of *R. similis*. The study presents 3D-structural models for *Rs*-GST and *Rs*-CES proteins using conventional molecular modeling techniques and structural motifs have been characterized with motif elucidation and sequence analysis methods. Subsequently, consense based phylogenetic analysis approach was followed to define the evolutionary relationships for each target proteins.

Results: We report for the first time the presence and amplification of two novel target genes (GSTs and CESs) from *R. similis*. The structural motif characterization of the two genes with corresponding nematode genes indicated the functional diversity of the conserved motifs present in *Rs*-GST and *Rs*-CES. The search for protein signature motifs through InterProScan analysis confirmed the presence of thioredoxin-like fold (IPR012336), glutathione S-transferase, N-terminal (IPR004045), glutathione S-transferase, C-terminal (IPR010987) and glutathione S-transferase domain (PF00043) for *Rs*-GST and carboxylesterase, type B (IPR002018), alpha/beta hydrolase fold (IPR029058) and carboxylesterase domain (PF00135) for *Rs*-CES. The 3D protein models of each protein were developed through homology modeling and the active-site residues were predicted. Phylogenetic analysis revealed the evolutionary relationships of each target proteins.

Conclusions: Identifying and cloning of genes involved in nematode survival and determining their functions are vital to elucidate the parasitism and host invasion processes of PPNs. The comparative structural analysis and motif characterization of studied targets will probably offer a novel approach for controlling plant nematodes.

Keywords- Burrowing nematode, *Radopholus similis*, plant parasitic nematode, target genes, detoxifying enzymes, glutathione S-transferase, carboxylesterase, motif characterization..

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Introduction

The burrowing nematode *Radopholus similis* (Cobb, 1893) Thorne, 1949 is a migratory endoparasite infesting several tropical and sub-tropical plant species and causing massive plant tissue necrosis because of their active migration through root tissues and destructive feeding on plant cells [1-2]. *R. similis* is known to invade and feed in the cortex of roots of more than 365 plant species [3-6] and is distributed throughout most tropical and subtropical areas, where it severely harms bananas, citrus crops, peppers and many other economically important crops [1-2,7].

Today plant parasitic nematodes (PPNs) are managed with chemical pesticides which are a short-term solution because of the soil inhabitation of nematodes and the perennial nature of crops [8-9]. Presently, the research focus has transferred to molecular biology approaches [10-13] rather than traditional approaches [14-15] for management of nematodes. The esophageal gland secretion proteins [16] and pathogenic genes have been the focus of many studies [10,17-19]. Another target group is antioxidant enzymes that help to defend themselves against reactive oxygen species (ROS) which are produced by the incomplete reduction of oxygen during respiration in mitochondria and as side products of a variety of normal metabolic reactions or from the immune response of the host. Secretory

antioxidant proteins produced by PPNs play an important role in diminishing plant defense system and invading host plants [20]. Cloning corresponding genes and determining their functions are important for elucidating the host invasion processes of plant nematodes and will likely provide a new approach to control PPNs [21].

Glutathione S-transferases (GSTs) are widely distributed among all living cells and are a major cellular detoxification system via catalyzing toxin conjugation with reduced glutathione (GSH) or passively binding to various exogenous /endogenous toxic molecules. They comprise a large family of multifunctional dimeric enzymes that are encoded by multigene families involved in the metabolism of a broad variety of xenobiotics and reactive endogenous compounds. GSTs have been exploited as potential target for many chemotherapeutic agents [22-26]. Detoxifying enzyme, GSTs belongs to a protein family involved in critical antioxidant and detoxification of xenobiotics, protection from oxidative damage and intracellular transport of hormones, endogenous metabolites and exogenous chemicals [27-28]. Functions of GSTs have been investigated in helminth parasites, not least because they have been identified as potential vaccine candidates in digeneans [29].

Carboxylesterases (CESs) is a cytoplasmic enzyme which plays a critical role in

neutralizing xenobiotics and plays an important role in the metabolism of organophosphates [30-31]. Carboxylesterases detoxify organophosphate (OP) and carbamates (CB) pesticides and synthetic pyrethroid (SPs) by two main ways, hydrolysis of the ester bond and binding of the pesticide (OP) to the active site of CESs [32].

OP and CB pesticides obtain toxicity from their ability to inhibit many metabolic and physiological enzymes like acetylcholinesterase (AChE), cytochrome P450, protein kinase C, CESs and GSTs, causing toxicity [33]. Both the enzymes (GSTs and CESs) are potentially involved in the pesticide detoxification [34-35]. Various studies reports that, these enzymes (CESs and GSTs) are commonly used as biomarkers in ecological risk assessment of pesticide contaminated environment [36-38]. Kale and Krishnamoorthy (1982) showed up to 80% inhibition of CESs by carbaryl in the earthworm [39]. Hans *et al.*, (1993) reported the induction of GSTs activity in worms by other pesticides, which showed a maximum induction of 250% by endosulphan [40]. Muthusamy *et al.*, (2011) studied the activity of GSTs and CESs in treated samples at lower and higher pesticide concentration and showed that an increased activity in lower dose for CESs and the activity were decreased in high dose [27]. These studies indicate that xenobiotics can elicit changes in the secretory proteins and this makes both the proteins as major targets for pesticides. Comparative sequences, structural and functional analysis of these enzymes are thus important.

Despite the devastating effects of *R. similis* on crop yields, this nematode is still poorly characterized at the molecular level. This paper reports the cloning and characterization of two genes encoding detoxifying enzymes glutathione S-transferase and carboxylesterase in *R. similis*. The 3-D structural and motif characterization of each protein have been done to understand the detoxification systems. We present structural model for each proteins using conventional molecular modeling techniques. The genetic relationships of each gene between corresponding genes from nematoda, vertebrata and arthropoda have also been examined through phylogenetic analysis.

Materials and methods

Radopholus similis culture and collection

The burrowing nematode, *R. similis* was isolated from black pepper roots and was maintained in carrot disk cultures at ICAR-Indian Institute of Spices Research, Kozhikode. Nematodes aseptically teased out of the carrot disks were rinsed using sterile demineralized water; actively moving nematodes from carrot disk cultures were taken after surface sterilization [41] and transferred to a sterile DEPC treated eppendorf tube.

RNA extraction and cDNA synthesis

Total RNA was extracted from mixed stages of *R. similis* (30,000 nos.) using Qiagen RNeasy Lipid Tissue Mini Kit (QIAGEN GmbH, Germany) according to the manufacturer's instructions. First-strand cDNA synthesise was carried out with 2µg RNA as template extracted using ThermoScientific RevertAid Reverse Transcriptase (Thermo Fisher Scientific Inc., MA USA) in the presence of an oligo (dT) primer (Thermo Fisher Scientific Inc., MA USA). The reaction mixture was taken in sterile, nuclease free tube on ice; the mixture contained 2 µg RNA as template, 0.5 µg oligo (dT)₁₈ primer, DEPC-treated water and incubated at 65°C for 5 min and chilled on ice and centrifuged. The mixture was then added to another sterile, nuclease free tube containing 5X reaction buffer, ThermoScientific RiboLock RNase inhibitor (Thermo Fisher Scientific Inc., MA USA), 10 mM dNTP mix and RevertAid reverse transcriptase. The mixture was then mixed gently and incubated for 60 minutes at 42°C and then terminated the reaction by heating at 70°C for 10 minutes. Additionally for the preparation of DNA-free RNA for amplification, removal of genomic DNA from isolated RNA preparation was done using ThermoScientific DNaseI, RNase-free according to manufacturer's protocol.

PCR amplification, cloning and sequencing of genes

Genes of *R. similis* coding for glutathione S-transferase and carboxylesterase were amplified from a mixed stage cDNA pool under standard PCR conditions using corresponding primers designed from assembled *R. similis* EST contigs. The sequences of the primer sets for amplification of *Rs*-GST were forward primer:

5'-CTCATCGCTACATTCATGGTG 3'; reverse primer: 5'- TGTGCTCAGAATTTCTTCATCAG 3'. For amplification of *Rs*-CES, the forward primer: 5'- ATGAGCAGATTCCTACTCGTTTCG 3'; reverse primer: 5'- CTGACTGCATCCGGCTATC 3'. PCR amplification reactions were performed in 25 µl reaction mixture containing 1X buffer, 1.5 µM MgCl₂, 2, 4 µM dNTPs, 0.5 µM forward and reverse primers, 1 U Taq DNA polymerase, and 1 µl cDNA. The PCR cycling parameters were 94°C initial denaturation for 5 minutes, followed by 35 cycles of amplification, denaturation at 94°C for 1 min; annealing at 60°C (*Rs*-GST) and 62°C (*Rs*-CES) for 1 min; and extension at 72°C for 1 min; and additional polymerization at 72°C for 10 min and hold at 4°C. Amplified fragments were purified from agarose gels using ThermoScientific GeneJET Gel Extraction Kit (Thermo Fisher Scientific Inc., MA USA) and cloned into pGEM-T vector (Promega, Madison USA) with standard protocol. Freshly prepared competent cells of *Escherichia coli* DH5α were transformed with recombinant plasmids and positive clones were selected for plasmid isolation. They were sequenced at Eurofins Genomics India Pvt. Ltd., Bangalore, India and the sequences were then assembled using CAP3 [42].

Sequence analysis and motif characterization

Putative protein sequences were obtained by translating the cDNA sequences using the EMBOSS program TRANSEQ [43]. Protein structure analysis was done with ExPasy-ProtParam tool [44] to analyze the various physical and chemical parameters of these protein sequences. The computed parameters were theoretical isoelectric point (PI), extinction coefficient, molecular weight, amino acid composition, atomic composition, estimated half-life, instability index, aliphatic index and grand average of hydropathicity (GRAVY). Signal peptides were predicted using SIGNALP 3.0 [45] and SLP-Local (Subcellular Location Predictor based on local features of amino acid sequence) was used to predict the subcellular location proteins [46]. Conserved domains were predicted by NCBI Conserved Domain Database and InterProScan [47]. Sequence alignment of identified *Rs*-GST and *Rs*-CES with corresponding known nematode genes was created with MUSCLE program [48]. Conserved motifs and their relative positions of both the proteins were predicted using MEME Version 4.9.1 [49].

Motif characterization of *Rs*-GST (KM670018) have been studied using alignment with related glutathione S-transferase protein sequences from other nematodes such as *Caenorhabditis elegans* (NP_506983), *Ancylostoma duodenale* (KIH55339), *Necator americanus* (ACX53261), *Haemonchus contortus* (CDJ94821), *Caenorhabditis breneri* (EGT52653), *Caenorhabditis remanei* (XP_003109030), *Caenorhabditis briggsae* (XP_002630606) and *Meloidogyne incognita* (ABN64198). Motif characterization of *Rs*-CES (KP027005) have been studied using alignment with related carboxylesterase protein sequences from other nematodes such as *Necator americanus* (ETN81995), *Caenorhabditis elegans* (CAA46899, NP_503411), *Caenorhabditis briggsae* (XP_002647363), *Caenorhabditis remanei* (XP_003110383), *Ascaris suum* (ERG83158, ERG83160), *Strongyloides ratti* (CEF63325).

Evolutionary analysis

A final alignment of 199 amino acids for glutathione S-transferases and 308 for carboxylesterases were used in the analyses. The evolutionary history was inferred using the Bayesian, minimum evolutionary and maximum likelihood analysis. Bayesian analysis was performed in MrBayes version 3.1 [50] with two searches run simultaneously for at least two million generations. Flat Dirichlet priors were used for the gamma shape parameter and the proportion of invariable sites. Three heated chains (temperature 0.2) and one cold chain were used in each search. The parameter was then fixed for a bootstrap analysis with 10,000 replicates. Maximum likelihood and minimum evolutionary analyses were performed using MEGA5 [51]. The analysis involved 25 protein sequences for *Rs*-GST and 23 for *Rs*-CES. Out-group was selected for both the genes as *C. elegans peroxidase gene with accession no. Q95003.1*.

The consensus tree was identified using consense package of Phylip 3.69 [52]. The majority-rule consensus of the bootstrap replicate trees was calculated using consense and seqboot package in the Phylip; which generates a majority rule consensus tree that retains the relationships found in majority of the trees.

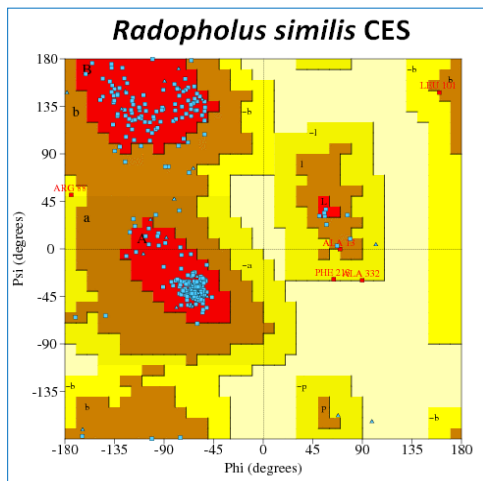


Fig-4b Ramachandran plot for modeled *Rs-CES* showing 90.5% residues in most favored region.

Motif characterization

The comparison of structural features for both the protein sequence revealed that the proteins retained most of the consensus sequence motifs that are crucial in their functions. Distinct motifs were identified to be conserved for both *Rs-GST* and *Rs-CES*.

Comparison of *Rs-GST* (KM670018) with related glutathione S-transferase protein sequences from other nematodes identified conserved motifs for nematode GSTs. Five distinct motifs were identified to be conserved for *Rs-GST* all along the

length, which are related to the function and specificity of GSTs. The conserved motifs of *Rs-GST* and their relative positions are displayed in [Fig-5a] and [Fig. 5b]. The InterProScan results showed that the N-terminal domain (G-site) lies between 1-71 bp, so the motif-1, motif-2, motif-3 and motif-4 constitute the G-site and C-terminal domain (H-site) lies between 73-201 bp containing only motif-5. Additionally, the conserved GSH binding site (G-site) (Tyr⁴, Arg¹⁶, Pro⁵², Asp⁴⁸, Ser⁵⁶), and substrate binding pocket (H-site) (Asp¹⁴⁹, Pro¹⁷²) residues in nematode GSTs were identified in the sequence through multiple sequence alignment (MSA) shown in [Fig-6].

The motifs that are conserved among nematode carboxylesterase are identified by aligning *Rs-CES* (KP027005) with related carboxylesterase protein sequences from other nematodes. Four distinct motifs were identified to be conserved for *Rs-CES* all along the length, which are related to the function and specificity of carboxylesterases. The three catalytic residues (catalytic triad): a serine, a glutamate or aspartate and a histidine were contained in the motif regions. The catalytic triad residues (Ser, Asp, His) were contained in Motif 1, motif 3 and motif 4. The conserved motifs of *Rs-CES* and their relative positions are displayed in [Fig-7a] and [Fig-7b]. The three catalytic triad residues of nematode carboxylesterase: a serine, a glutamate or aspartate and a histidine in the motif regions were displayed in multiple sequence alignment [Fig-8]. Mapping conserved motifs of *Rs-GST* and *Rs-CES* onto the 3D structure is displayed in [Fig-9a] and [Fig-9b]; mapping revealed that most of the motif residues were positioned near the catalytic centre of the proteins forming active-sites. Whereas less conserved amino acids occurred more often in the alpha helices of two proteins.



Fig-5a Schematic diagram of conserved amino acid motifs within the nematode GSTs along with *Rs-GST* as analyzed through MEME 4.0 software tool. The black solid line represents different glutathione S-transferase sequences and their length, while colored boxes represent conserved motifs along the length of each sequence.

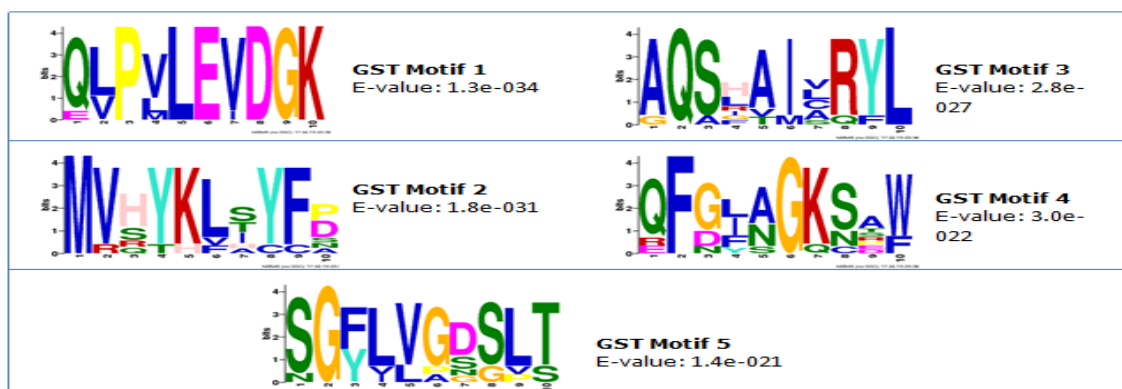


Fig-5b Sequence logo of five prominent conserved motifs in *Rs-GST* and other nematode GSTs.

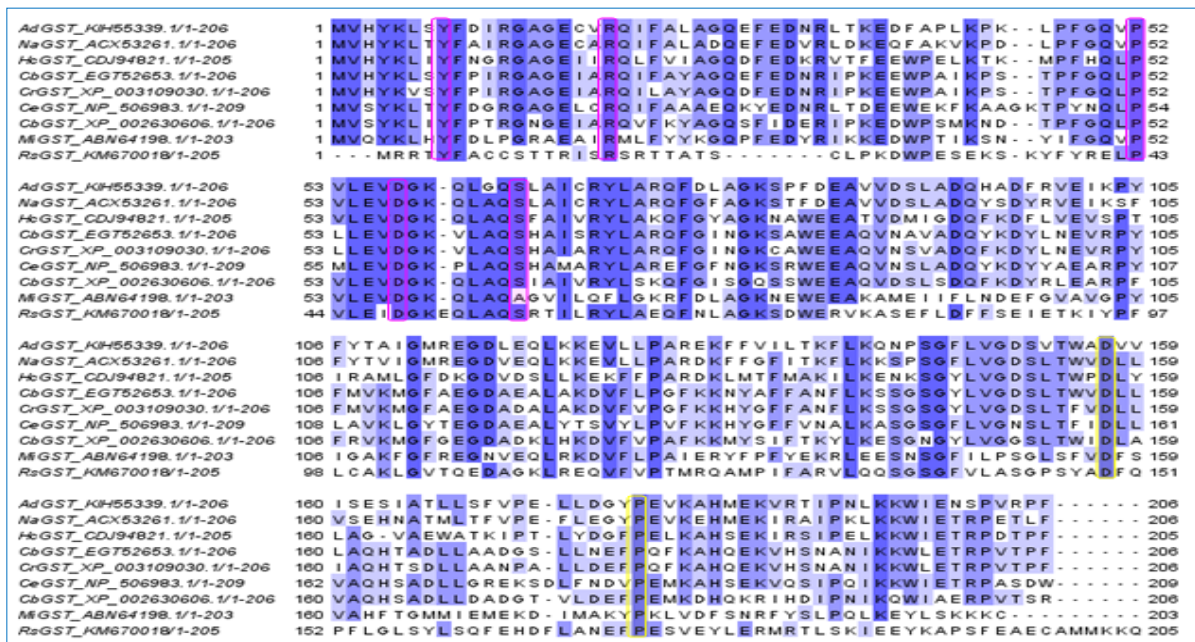


Fig-6 Protein alignment of *Radopholus similis* glutathione-s transferase (RsGST) with other nematodes is displayed according to percentage identity. GSH binding sites (G-sites) (Tyr⁴, Arg¹⁶, Pro⁵², Asp⁴⁸, Ser⁵⁶), are indicated in pink box and substrate binding pocket (H-sites) (Asp¹⁴⁹, Pro¹⁷²) are indicated in yellow box.



Fig-7a Schematic diagram of conserved amino acid motifs within the nematode carboxylesterase sequences along with Rs-CES as analyzed through MEME 4.0 software tool. The black solid line represents different carboxylesterase sequences and their length, while colored boxes represent conserved motifs along the length of each sequence.

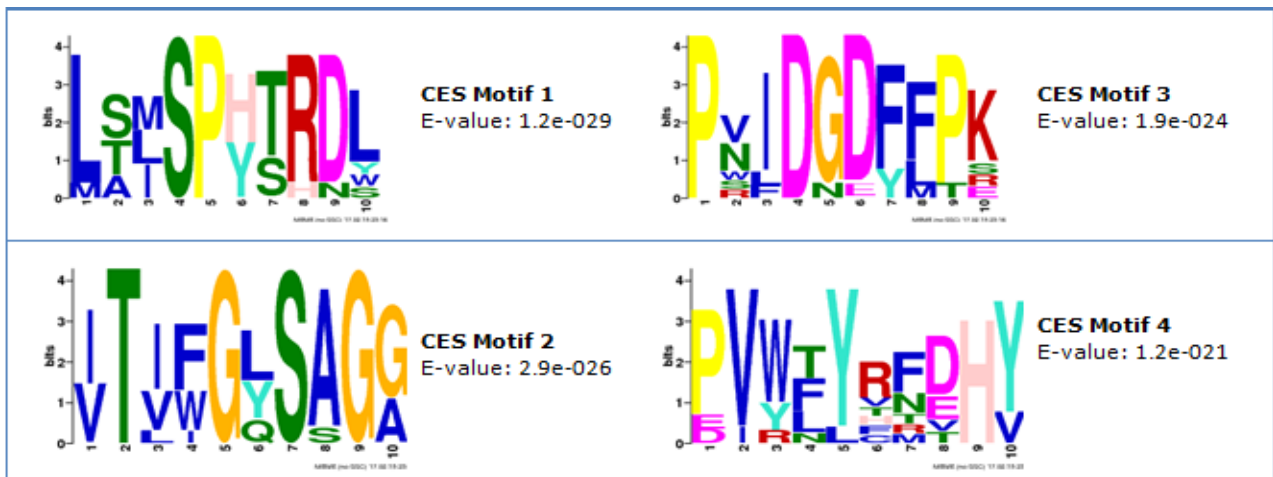


Fig-7b Sequence logo of four prominent conserved motifs in Rs-CES and other nematode CESs.

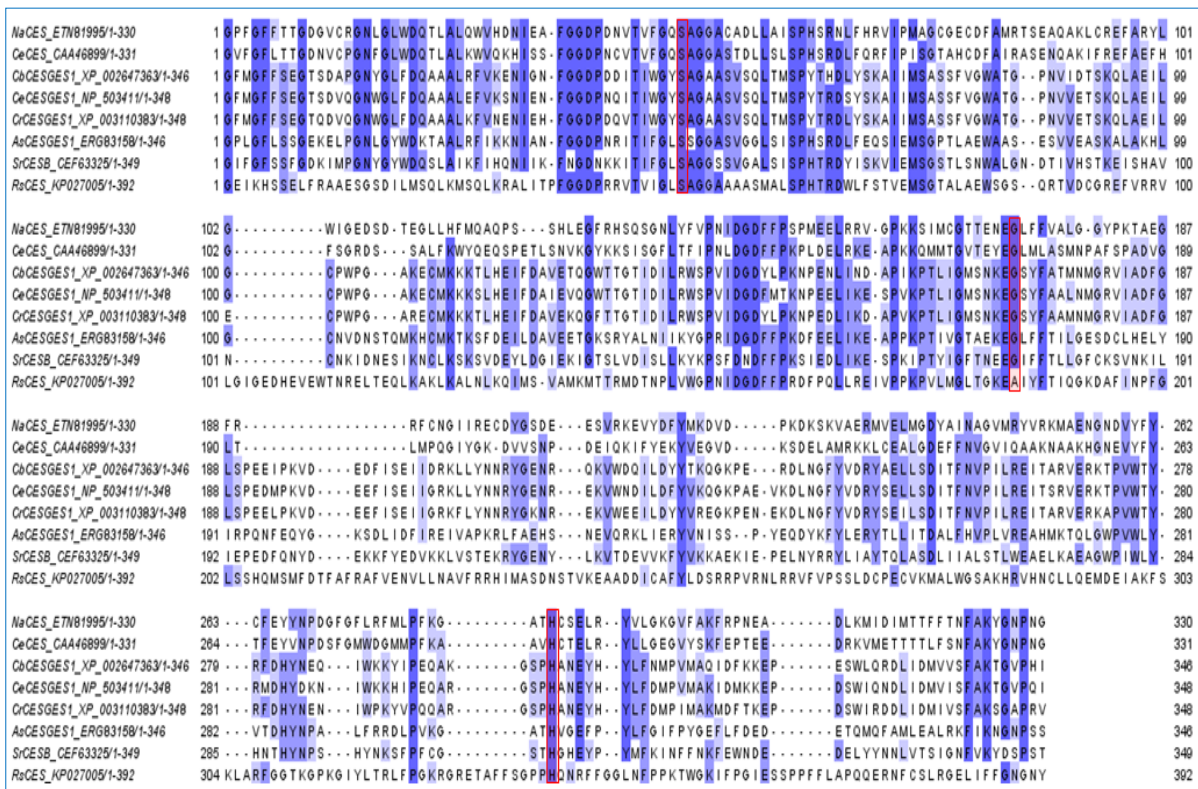


Fig-8 Protein alignment of *Radopholus similis* carboxylesterase (RsCES) with other nematodes is displayed according to percentage identity. The three catalytic residues (catalytic triad): a serine, a glutamate or aspartate and histidine residues were indicated in red color.

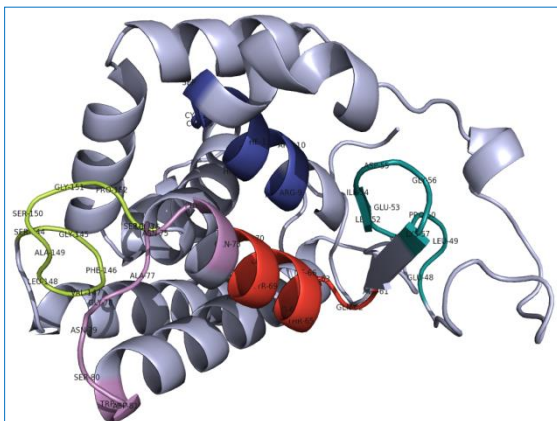


Fig-9a Three-dimensional model of Rs-GST, the conserved motifs are indicated in cyans (Motif-1), blue (Motif-2), red (Motif-3), magentas (Motif-4) and Yellow (Motif-5) and non conserved in light blue.

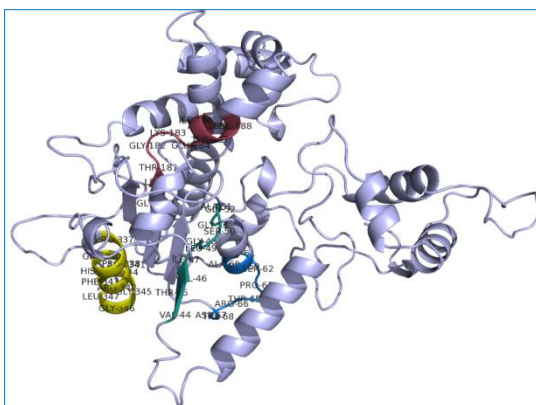


Fig-9b Three-dimensional model of Rs-CES, the conserved motifs are indicated in cyans (Motif-1), blue (Motif-2), red (Motif-3), and Yellow (Motif-4) and non conserved in light blue.

Evolutionary analysis

Multiple sequence alignment and motif characterization with other nematodes revealed that the *Rs-GST* and *Rs-CES* retained most of the consensus sequence motifs that are crucial for their function. The phylogenetic results indicated that the GSTs of plant parasites are forming in to a single clade, separated from other nematodes [Fig-10], indicating sequence similarity; *Rs-GST* showed higher similarity to plant-parasitic nematode, *Meloidogyne incognita*, both are grouped together in the phylogenetic tree. Phylogenetic analysis of *Rs-CES* [Fig-11] revealed that it is related to CESs of other nematodes such as *Caenorhabditis elegans*, *C. remanei*, *C. briggsae*, *Strongyloides ratti* and *Ascaris suum*, indicating their sequence similarity. Since carboxylesterase (CES) is sequenced for the first time in *R. similis* and among plant parasitic nematodes, carboxylesterase sequences of other plant parasitic nematodes were not available for the study. *Rs-CES* are closely related to gut esterase 1 (GES) protein of *C. elegans*, *C. remanei*, *C. briggsae*, *S. ratti* and *A. suum*.

Discussion

The migratory plant-parasitic nematode, *R. similis* is a serious problem in many crops and our options to control this nematode are very meager. There is an urgent need to develop novel measures to control them by identifying new molecules and bioagents that act against specific target genes of the nematode. In the present study, an attempt was made to clone and characterize the two detoxification genes viz. glutathione S-transferase and carboxylesterase from *R. similis*. Both the genes had signal peptide for secretion and catalytic domains for functional specificity.

The *Rs-GST* had a predicted molecular mass of 23.702 kDa and was well within the range of 21–29 kDa reported for other GSTs [66] and agreed with the previous findings that the average molecular mass of sigma GSTs in parasites is 22 kDa [67]. Further analyses revealed that *Rs-GST* possess all the specific structural features of representatives of GSTs in the Sigma class: a coding domain for the GST_N_Sigma_like (PSSM: cd03039) and another for the GST_C_Sigma_like (PSSM: cd03192). GSTs that share greater than 40% sequence identity are generally included in the same class, and those that possess less than 20–30% sequence identity are assigned to separate classes [68-70]. The signal peptide

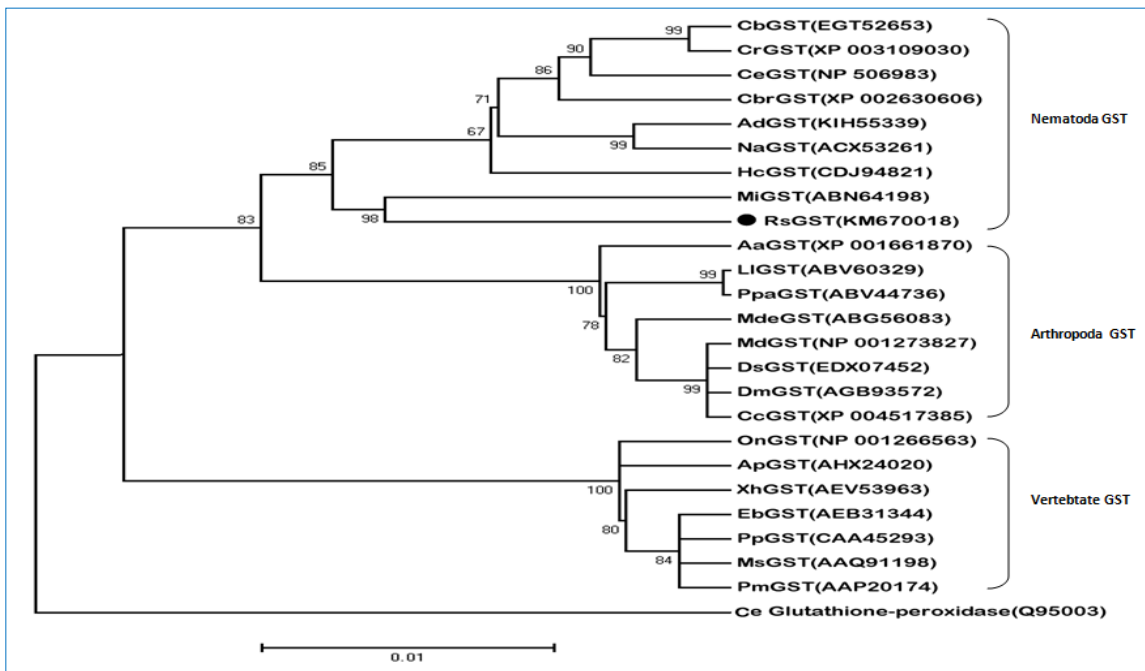


Fig-10 Consensus phylogenetic tree of amino-acid sequence for inferring relationships among GSTs, obtained by Bayesian analysis, Maximum Likelihood and Maximum evolutionary methods. The GenBank accession numbers of each gene are indicated in parentheses. Nodes with better than 60% bootstrap values are indicated with the number of frequency. The Rs-GST is indicated by black circle. Ce- *Caenorhabditis elegans*, Ll- *Lutzomyia longipalpis*, Ppa- *Phlebotomus papatasi*, Aa- *Aedes aegypti*, Dm- *Drosophila melanogaster*, Ds- *Drosophila simulans*, Md- *Musca domestica*, Cc- *Ceratitidis capitata*, Mde- *Mayetiola destructor*, Ad- *Ancylostoma duodenale*, Na- *Necator americanus*, Hc- *Haemonchus contortus*, Cb- *Caenorhabditis brenneri*, Cr- *Caenorhabditis remanei*, Cbr- *Caenorhabditis briggsae*, Mi- *Meloidogyne incognita*, Rs- *Radopholus similis*, Ms- *Micropterus salmoides*, Pm- *Pagrus major*, Eb- *Epinephelus bruneus*, Pp- *Pleuronectes platessa*, Ap- *Acanthochromis polyacanthus*, On- *Oreochromis niloticus*, Xh- *Xiphophorus hellerii*.

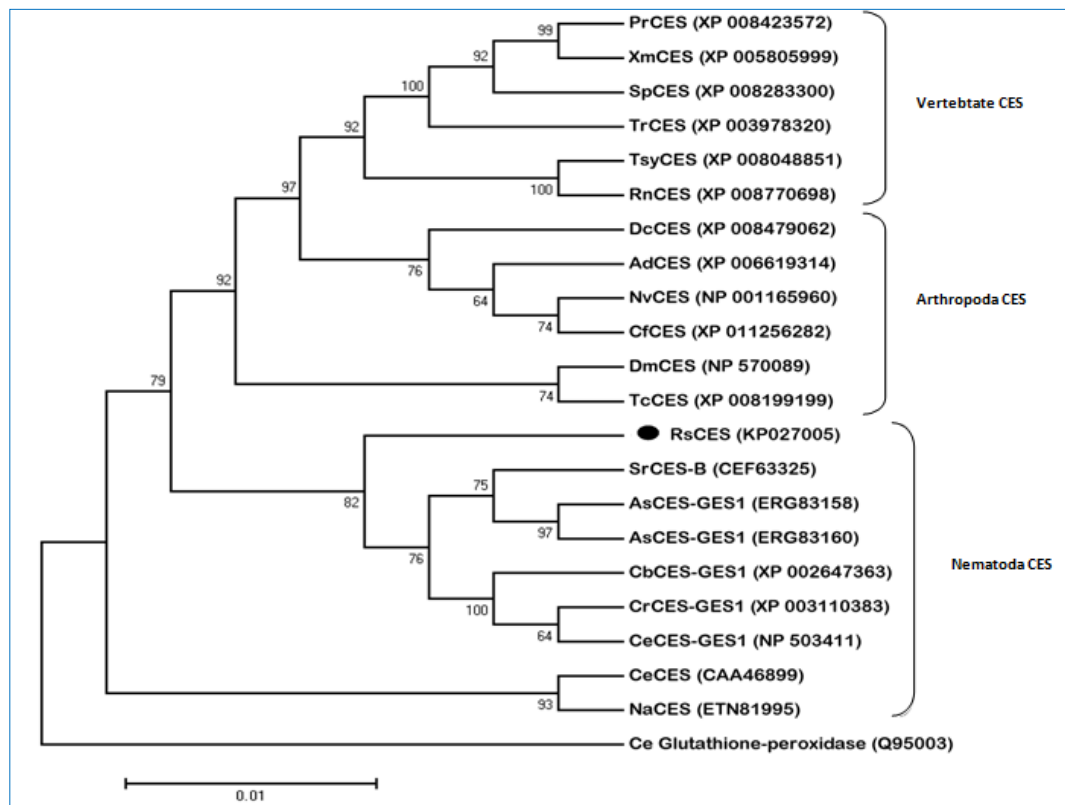


Fig-11 Consensus phylogenetic tree of the amino-acid sequence for inferring relationships among carboxylesterase sequences, obtained by Bayesian analysis, Maximum Likelihood and Maximum evolutionary methods. The GenBank accession numbers of each gene are indicated in parentheses. Nodes with better than 60% bootstrap values are indicated with the number of frequency. The Rs-CES is indicated by black circle. Na- *Necator americanus*, Rs- *Radopholus similis*, Ce- *Caenorhabditis elegans*, Cr- *Caenorhabditis remanei*, Cb- *Caenorhabditis briggsae*, Sr- *Strongyloides ratti*, As- *Ascaris suum*, Pr- *Poecilia reticulata*, Xm- *Xiphophorus maculatus*, Sp- *Stegastes partitus*, Tsy- *Tarsius syrichta*, Rn- *Rattus norvegicus*, Tr- *Takifugu rubripes*, Nv- *Nasonia vitripennis*, Tc- *Tribolium castaneum*, Ad- *Apis dorsata*, Dc- *Diaphorina citri*, Cf- *Camponotus floridanus*, Dm- *Drosophila melanogaster*.

analysis has indicated *Rs*-GST to be a secretory protein. Therefore, the results suggest that *Rs*-GST characterized here should be a secretory glutathione S-transferase and is affiliated with the Sigma class.

All cytosolic GSTs have the same protein folding, which comprises two domains. The N-terminal domain (domain I) adopts *a/b* topology and provides the GSH-binding site (G-site). The C-terminal domain (domain II) is an all- α -helical structure and provides the structural element for recognition of a broad range of hydrophobic co-substrates (H-site). The H-site lies adjacent to the G-site and defines the substrate specificity of the enzyme [74-78]. The p-value of the different motifs in *Rs*-GST suggests that glutathione S-transferase motif-1 constituting G-site has diverse amino-acid residues among different nematode species, since it had a higher p-value (1.3e-034) when compared to the other motifs of *Rs*-GST.

The predicted molecular mass of *Rs*-CES, 59.449 kDa and isoelectric point of 4.37 were similar to those of intestinal carboxylesterase of *C. elegans* [71] and *A. suum* [72]. Also, in BLAST search, *Rs*-CES shared highest identity with gut esterase-1 of *Ascaris suum* (ERG83158) which is a parasitic nematode. Further analysis revealed that the *Rs*-CES possess conserved domain specific to carboxylesterase family (pfam00135) such as esterase_lipase (cd00312) and esterase_lipase superfamily (cl21494), which includes lipases, cholinesterases and carboxylesterases. All the three residues specific to catalytic apparatus were well conserved in catalytic triad of *Rs*-CES: a serine, a glutamate or aspartate and a histidine [73]. The presence of signal peptide for *Rs*-CES suggested that the characterized *Rs*-CES is a secretory protein and is affiliated with class of nematode esterase-1.

All the signature motifs of carboxylesterase such as alpha/beta hydrolase fold (IPR029058), carboxylesterase, type B serine active site/catalytic triad (Ser, Glu and His) (IPR002018), carboxylesterase domain (PF00135), esterase-lipase domain (cl21494) were present in characterized *Rs*-CES. The α/β hydrolase fold of carboxylesterase is responsible for the hydrolysis of carboxylesters and amides of various sizes. All carboxylesterase contains an active acylation, site a serine, a glutamate or aspartate and a histidine catalytic triad [79-80]. The p-value of the different motifs in *Rs*-CES suggests that carboxylesterase motif-4 (1.2e-021) constituting a part of catalytic triad is diverse than other motifs.

Rs-GST grouped along with nematode-specific GSTs in phylogenetic analyses and was distinct from those of the vertebrates and insects in the Sigma class GSTs [81-81]. It showed a close relationship with that of *Meloidogyne incognita*. Since both the genes are conserved across different species and genera, in spite of evolutionary pressure for diversification. Consensus based phylogenetic analysis revealed the genetic similarity/diversity of these enzymes with corresponding proteins of other nematodes, arthropoda and vertebrata. However, nematode GSTs are more conserved between the parasitic nematodes (*Ancylostoma duodenale*, *Necator americanus*, *Haemonchus contortus*, *Meloidogyne incognita* and *Radopholus similis*) and diverged from free-living nematodes indicating their possible role for parasitism [Fig-10]. Also phylogenetic analysis suggested that the evolution of nematode GSTs/*Rs*-GST from a common ancestor of Arthropoda GSTs were a relatively recent event. *Rs*-GST together with analyzed nematode GSTs belonged to a clade containing arthropoda GSTs, implying its functional similarity to arthropod GSTs. Whereas *Rs*-CES was closely related to gut esterase 1 (GES) protein of *C. elegans*, *C. remanei*, *C. briggsae*, *S. ratti* and *A. suum* forming a single clade, while normal carboxylesterase of *N. americanus* and *C. elegans* formed an independent clade suggesting their evolutionary distance [Fig-11]. *Rs*-CES was also closely related to CESs of parasitic nematodes. The application of RNAi to the identified detoxification genes in *R. similis* could open the door to unraveling the underlying biology of parasitism in plant parasitic nematodes. The Nematoda CES-GES genes, including *Rs*-CES shared a common ancestor with the arthropoda group including *Tribolium castaneum* and *Drosophila melanogaster*.

Significantly, phylogenetic analysis is in excellent agreement with the multiple sequence alignment result confirming the similarity. The multiple sequence alignment could reveal that, the GSH binding sites (G-sites) (Tyr4, Arg16, Pro52, Asp48, Ser56) and substrate binding pocket (H-sites) (Asp149, Pro172) were conserved in *Rs*-GST [Fig-6], as reported earlier [74-78]. In case of *Rs*-CES, the

three catalytic residues (catalytic triad): a serine, a glutamate or aspartate and a histidine residues were highly conserved [Fig-8] among CESs as reported by others [79-80]. The observed results of motif characterization and conservation confirm the phylogenetic relationship of these two genes determined by multiple sequence alignment thus, enhancing the confidence.

In summary, two detoxifying proteins, glutathione s-transferase and carboxylesterase were identified and characterized from *R. similis* that can form the basis towards pathogenicity of plant parasitic nematodes. The results of phylogenetic and motif characterization analysis suggested that the putative proteins encoded by sequenced *Rs*-GST and *Rs*-CES are involved in parasitism of nematode. The identified proteins can act as a valuable resource towards development of target-based nematicides. Along with the information on *C. elegans* glutathione S-transferase and carboxylesterase, the current findings on *Rs*-GST and *Rs*-CES from a burrowing nematode species should provide valuable insights into the evolutionary process and various physiological functions of both the detoxifying genes. The 3D-structures were constructed for both the proteins, as knowledge of the three-dimensional structure is essential for a better understanding of the functional mechanism. After comparative modeling and structural refinement, the general quality of structures was assessed and compared with known structures. We hope that our model will inspire new experimental effort in this area. Furthermore, a better understanding of the 3D-structure and conserved sites comparison of both the enzyme will be pertinent for developing organophosphate based synergists and alternative novel natural nematicides against *R. similis*.

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