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In Silico Analysis of Rpp1 Locus-Derived NB-ARC-LRR Domains Reveals Insights into Rust Resistance in Soybean

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Authors' contributions

This work was carried out in collaboration among all authors. All authors read and approved the final manuscript.

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ABSTRACT

Soybean rust poses significant economic challenges to soybean producers, resulting in decreased seed size, weight, oil content, and overall crop yield. This study aimed to comprehensively characterize the rust resistance locus Rpp1 identified in soybean genotype EC 241780 and its

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associated genes within the soybean genome. Our investigation revealed distinctive features of three genes (Glyma18G281700, Glyma18G281600, and Glyma18G281500) located within the Rpp1 locus on chromosome 18. Phylogenetic and domain analyses confirmed the presence of the ULP1 domain in these genes, while motif analysis identified unique patterns exclusive to them. Notably, these genes possessed exclusive isoleucine and glutamic acid residues at positions 836 and 904, along with an unusual expansion of 11 glutamic acid residues at position 861. Expression-data showed significantly higher expression levels of Glyma18G281700 and Glyma18G281600 during leaf trifoliate stages I and II, suggesting their active involvement under adverse conditions. Protein structure analysis revealed that Glyma18G281500 exhibited distinct characteristics compared to the other two genes and displayed a more stable protein structure. This observation may indicate a distinct functional role for Glyma18G281500. In conclusion, our findings suggest that the genes within the Rpp1 locus may contribute to rust resistance, particularly against specific rust races, due to their unique characteristics associated with the ULP1 domain.

Keywords: Rpp1; rust resistance in soybean; ULP1 domain.

1. INTRODUCTION

Soybean (Glycine max) holds a prominent position as a globally significant crop, serving as a vital source of both protein and oil for human and animal consumption. However, soybean plants continually face formidable challenges from a spectrum of pathogens encompassing bacteria, fungi, and viruses, which pose substantial threats to crop yield and quality [1]. Among these adversaries, soybean rust, incited by the biotrophic fungus Phakopsora pachyrhizi, stands out as one of the most deleterious foliar diseases afflicting soybean [2]. Over the years, extensive screening efforts involving at least 200 germplasm accessions and breeding lines have led to the identification of seven resistance loci denoted as Rpp (signifying resistance to P. pachyrhizi). These loci include Rpp1 [3], Rpp2 [4], Rpp3 [5], Rpp4 (Hartwig, 1986), Rpp5 [6], Rpp6 [7], and Rpp7 [8]. Nevertheless, it is crucial to acknowledge that none of these resistance genes exhibit efficacy against all currently known soybean rust pathotypes [8,9,10].

In response to the pervasive onslaught of these pathogens, plants have evolved a sophisticated immune system centered around the recognition of pathogen-derived molecules through specific receptors. Notably, a prominent category of receptors that play a pivotal role in soybean's defense against diseases are the leucine-rich repeat-containing (LRR) domain proteins. These proteins are characterized by the presence of nucleotide-binding site (NBS) and leucine-rich repeat (LRR) domains, complemented bv variable amino- and carboxy-terminal domains. all of which collectively facilitate a range of critical protein-protein interactions and ligand binding events (McHale et al. 2006). Their significance in plant defense mechanisms becomes apparent as they serve as the vanguard, recognizing and responding to pathogen-associated molecular patterns (PAMPs) or effector molecules released by invading pathogens [11]. Notably, a majority of disease resistance genes (R genes) identified in plants thus far are comprised of nucleotidebinding site leucine-rich repeat (NBS-LRR) Furthermore, LRR proteins [11]. domaincontaining proteins have also been implicated in reinforcing soybean's defense against fungal pathogens, as evidenced by previous studies [12,13].

The objective of the present investigation was to conduct a comparative analysis of the Rpp1 locus with other NB-ARC-LRR domain-containing genes in soybean. This comprehensive analysis aimed to elucidate the variations and specificities associated with the Rpp1 locus, shedding light on its unique role within the intricate framework of soybean's defense mechanisms against the formidable soybean rust pathogen.

2. MATERIALS AND METHODS

2.1 Isolation and Comparison of NBS-LRR Containing Genes

In pursuit of identifying the nucleotide-binding site leucine-rich repeat (NBS-LRR) domain, a systematic search was conducted across a sovbean genome in UNIPROT database (https://www.uniprot.org/) (supplementary file 1). То gain insights into the evolutionary relationships among various leucine-rich repeat (LRR) domain-containing proteins, including those within the Rpp1 locus, a rigorous multiple sequence alignment was executed. This alignment was carried out using ClustalW, into the BIOEDIT software (version 7.2). The precise

conserved regions of amino acid sequences were compared among all LRR domaincontaining proteins.

2.2 Phylogenetic Analysis

To construct a phylogenetic tree, Neighbor Joining analysis was employed. This analysis was carried out using the Simple phylogeny tool (https://www.ebi.ac.uk/Tools/phylogeny/simple_p hylogeny/). Subsequently, the resultant phylogenetic tree was refined and visually enhanced using the Interactive Tree of Life (ITOL) platform (https://itol.embl.de/upload.cgi), providing a clear depiction of the relationships and distinctions among the analysed proteins.

2.3 Identification of Protein Domains

To ascertain the presence of both single and multiple domains within the protein sequences under investigation, a dedicated analysis was conducted using the batch-CD tool for leucinerich repeat (LRR) domain-containing proteins, including those within the Rpp1 locus, which is accessible through the National Center for Biotechnology Information (NCBI) website (https://www.ncbi.nlm.nih.gov/Structure/bwrpsb/b wrpsb.cgi). Further, TB-tool was employed for the visualization of domain structures present within the analyzed protein sequences. This visualization facilitated comprehensive а understanding of the arrangement and distribution of domains. contributing to the elucidation of potential functional roles.

2.4 Motif Analysis

To uncover conserved motifs within the protein sequences, a motif search was diligently carried out for leucine-rich repeat (LRR) domaincontaining proteins, including those within the Rpp1 locus. This search was performed using the MEME tool (http://meme.nbcr.net/meme/). By identifying conserved motifs, this analysis shed light on potential functional elements shared among the studied proteins.

2.5 Expression Data Analysis

Expression data pertinent to the various stages and tissues associated with the studied gene sequences were procured from the Legume Mine database via the Legume Information System (LIS). These data resources provided valuable insights into the expression patterns of the analyzed genes across different developmental stages and tissues. To effectively visualize the expression patterns and relationships among the studied genes, the acquired expression data were transformed into a heatmap format. This transformation was accomplished using the Clustvis online tool (https://biit.cs.ut.ee/clustvis/), which enabled the creation of a graphical representation facilitating the interpretation of gene expression profiles.

2.6 Protein Structure Analysis

The Phyre2 automatic fold recognition server was utilized to predict the structure and/or function of three genes (Glyma18G281700, Glyma18G281600, and Glyma18G281500) with modeling confidence of 100.0%. These а predictions were generated using the single highest scoring template (http://www.sbg.bio.ic.ac.uk/phyre2/html/page.cgi ?id=index). Furthermore, additional structurebased analyses were conducted using Phyre2, including relative conservation analysis based on Jensen-Shannon divergence, evaluation of clashes between atoms in specific residues, Ramachandran analvsis. and mutational sensitivity assessment using the SuSPect tool to predict potential missense mutations in the protein sequences.

3. RESULTS AND DISCUSSION

Asian soybean rust (ASR) is a highly destructive ailment impacting soybean production on a global scale [14]. Genes that confer resistance against Phakopsora pachyrhizi (Rpps) have been harnessed to safeguard soybeans from this disease. Seven loci resistances to P. pachyrhizi (Rpp) have been identified on genetic linkage maps of soybeans [15,16,8]. Due to the Ρ. pachyrhizi variation in considerable races/isolates, relying on a single Rpp locus/gene proves insufficient to offer lasting and comprehensive resistance against ASR disease.

The NBS-LRR protein, encoded by the NBS-LRR gene in plants, plays a critical role in enabling plants to counteract pathogen invasions [17]. There have been numerous extensive reviews since the first NBS-LRR-encoding genes were cloned from plants [18]. This gene is notably abundant across various plant species such as Arabidopsis thaliana and soybean, encompassing 149 and 319 NBS-LRR genes family, respectively [19]. The NBS-LRR protein is also referred to as NB-ARC-LRR (nucleotide binding adaptors composed of Apaf-1, R protein, and CED-4) protein due to the presence of nucleotide binding and ATPase domains (NB-ARC), which function as signal transduction

domains within a multifaceted protein superfamily [19].

The study undertaken current was to comprehensively characterize the Rpp1 locus in sovbeans. with the primary objective of elucidating the domain specificity underlying rust resistance. In a prior investigation [20], the Rpp1 locus was identified in soybean accession EC 241780 harbouring three distinct NB-ARC-LRR Glvma18G281700. containing genes: Glyma18G281600. and Glyma18G281500. Subsequently, a detailed analysis of these three was conducted, coupled with genes а comparative examination against other LRR domain-containing genes within the soybean aenome.

To elucidate the genetic complexities within this genomic region, we commenced a multiple sequence alignment encompassing 21 distinct genes encoding leucine-rich repeat (LRR) domains. This systematic analysis unveiled significant disparities in conserved segments when juxtaposed with genes localized within the Rpp1 genomic region, as illustrated in Fig. 1. Notably, within this gene set, exclusive to the Rpp1 locus were the occurrences of isoleucine and glutamic acid residues at positions 836 and 904, respectively. Furthermore, a distinctive feature emerged, with an expansion of 11 glutamic acid residues observed at position 861, setting it apart from all other genes under examination (Fig. 1). This distinct pattern suggests its singularity and implies a potential involvement defensive mechanisms. in Phylogenetic analysis was executed to investigate the evolutionary relationships between the genes associated with rust resistance (Rpp1) and other LRR-containing genes in soybean. Our findings unveiled the presence of four distinct clades, with the Rpp1 genes (Glyma18G281700, Glyma18G281600, and Glyma18G281500) forming a distinct clade of their own, highlighting their unique functional roles among the diverse LRR genes (Fig. 2). In addition to phylogenetic insights, we conducted domain analysis to probe the roles of various domains in soybean rust resistance. Our examination unveiled several domains present in the 21 genes, including the NB-ARC superfamily, RX-CC like, LRR superfamily, Rx N, PLN03210 superfamily, ULP1 superfamily, Retrotrans_gag superfamily, and PLN00113 superfamily, as depicted in Fig. 3. Of particular interest, the ULP1 domain was identified within the Rpp1 locus genes, suggesting its potential association

with the defense against rust disease. Notably, Pedley et al. [3] corroborated our findings by reporting that each candidate gene within the Rpp1 locus encodes an N-terminal ubiquitin-like protease 1 (ULP1) domain. Their research further demonstrated that co-silencing of these Rpp1 candidates resulted in the abrogation of the immune response in the Rpp1-resistant soybean accession PI 200492, underscoring the pivotal role of Rpp1 as a ULP1-NBS-LRR protein in orchestrating the immune response. Motif analysis was also performed, revealing distinct patterns among the three genes within the Rpp1 locus, as depicted in Fig. 4. There was absence of several motif was observed in Rpp1 locus containing genes.

Within LLR proteins, several highly conserved and meticulously ordered motifs are recognized, including the P-loop, kinase-2, and Gly-Leu-Pro-Leu motifs [17]. The LRRs (leucine-rich repeats) constitute structurally adaptable domains specializing in protein-protein interactions, with capacity to evolve diverse bindina the specificities. These LRRs are subject to diversifying selection [21,22,18], particularly at the level of the projected solvent-exposed residues. Intriguingly, in this region, the NBS-LRR proteins not only lack uniformity but exhibit significantly greater diversity than anticipated through random genetic drift. This suggests the presence of selective pressures that promote the development of novel pathogen-specificities, catering to the recognition of distinct pathogen Avr proteins. The evolution of this gene family is also influenced by processes such as clustering of NBS-LRR genes through tandem duplications and ectopic duplications followed by local rearrangements and gene conversion [19].

Furthermore, we delved into the functional aspects of these genes by analyzing their expression data across various tissues and developmental stages. The expression data of soybean variety Williams 82 was used for the genes studied, analysis. Among the 21 Glyma18G281700 and Glyma18G281600 exhibited relatively higher expression levels during leaf trifoliate Stage-I and stage II, implying their active involvement during stress conditions (Fig. 5). This study also aligns with recent research by Si et al. [13], which delved into the expression patterns of genes encoding LRR-RLKs in oomycetes. Their findings indicate a dual pattern of gene expression, with active transcription during vegetative growth stages and

host infection. This suggests the pivotal roles played by LRR-RLKs in various facets of the oomycete life cycle, encompassing growth, development, and interactions with host organisms. This aspect of our research contributes valuable insights into the functional significance of LRR-RLKs in oomycetes across different phases of their life cycle and their potential implications in host-pathogen interactions.



Fig. 1. Depicting conserved region in different NB-ARC- LRR containing genes in Soybean Multiple sequence alignment was executed by using ClustalW into the BIOEDIT software (version 7.2) for 21 different LRR containing genes including Rpp1 locus underlined genes. Conserved regions of amino acid sequences were highlighted among all LRR domain-containing proteins. Rpp1 locus genes (Glyma18G281700, Glyma18G281600, and Glyma18G281500) contained exclusive isoleucine and glutamic acid residues at positions 836 and 904, along with an unusual expansion of 11 glutamic acid residues at position 861.

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Fig. 2. Phylogenetic tree of NB-ARC- LRR containing domain protein family in Glycine Max Phylogenetic tree was constructed for 21 different LRR containing genes including Rpp1 locus underlined genes by using Simple phylogeny, a tool provided by EMBL-EBI

(https://www.ebi.ac.uk/Tools/phylogeny/simple_phylogeny) and a life tree was built using 1000 bootstrap tests on iTOL (Interactive tree of life) (https://itol.embl.de/). Phylogenetic tree showed four distinct clades, with the Rpp1 genes (Glyma18G281700, Glyma18G281600, and Glyma18G281500) forming a distinct clade of their own.





Domain analysis was conducted using the batch-CD tool in NCBI

(https://www.ncbi.nlm.nih.gov/Structure/bwrpsb/bwrpsb.cgi) for 21 different LRR containing genes including Rpp1 locus underlined genes. Further, TB-tool was employed for the visualization of domain structures. Each domain is represented in the different colored box as per legends. A distinct domain ULP1 was observed within the Rpp1 locus genes.





Motif analysis was done for 21 different LRR containing genes including Rpp1 locus underlined genes by using MEME tool (http://meme.nbcr.net/meme/). Different motif was shown in different color as per legends for each genes. All three Rpp1 genes (Glyma18G281700 and Glyma18G281600) exhibits distinct pattern of motif than other tested genes.



Fig. 5. Expression patterns of NB-ARC- LRR containing genes in Soybean

Expression data of soybean variety Williams 82 were collected from Legume Mine database via the Legume Information System (LIS) of various stages and tissues for 21 different LLR containing genes including Rpp1 locus. Data were presented in the form of heatmap. Two Rpp1 genes (Glyma18G281700 and Glyma18G281600) exhibited relatively higher expression levels at leaf trifoliate Stage-I and stage II, suggesting their active involvement during stress conditions.



Rpp1 locus

Fig. 6. Evaluation of different Protein structure based parameters for Rpp1 containing genes
Protein sequence were compared with Constraint-based Multiple Alignment Tool (COBALT)
(https://www.ncbi.nlm.nih.gov/tools/cobalt/cobalt.cgi?CMD=Web) (A). The Phyre2 automatic fold recognition
server was utilized to predict the structure and/or function of three Rpp1 containing genes (Glyma18G281700, Glyma18G281600, and Glyma18G281500) (http://www.sbg.bio.ic.ac.uk/phyre2/html/page.cgi?id=index) (B).
Furthermore, structure-based analyses were conducted that includes relative conservation analysis (C), clashes
analysis between atoms in specific residues (D), Ramachandran analysis (E), and mutational sensitivity
assessment (F) using the SuSPect tool to predict potential missense mutations in the protein sequences.
Glyma18G281500, one of the three genes within the Rpp1 locus, differs significantly from the other two genes
and also exhibits a more stable protein structure as it showed less number of clashes

Oomycetes exhibit a diverse array of LRR-RLKs (Leucine-Rich Repeat Receptor-Like Kinases), with each species demonstrating a unique complement of these receptor proteins. What's

intriguing is that these LRR-RLKs in oomycetes share a common structural pattern with their counterparts in extensively researched plant LRR-RLKs [23]. This structural similarity suggests potential functional parallels between oomycetes and plants in terms of signal transduction and cellular communication mechanisms [24,25,26]. Further investigation into these shared characteristics could provide valuable insights into the roles these receptors play in both oomycete biology and plantpathogen interactions [27-30]. In our study, various parameters related to protein structure also analyzed to comprehensively were understand the functional diversity among three Rpp1-containing genes. namelv Glyma18G281700. Glyma18G281600, and Glyma18G281500 (Fig. 6). Among these three genes, Glyma18G281500 exhibited notable differences in terms of sequence compression (Fig. 6A) and three-dimensional protein folding Additionally, Glyma18G281500 (Fia. 6B). displayed a relatively lower number of clashes between atoms in different residues compared to the other two genes. However, non-significant differences were observed among all three genes in terms of relative conservation analysis. Ramachandran analysis, and mutational sensitivity [31-34]. This comprehensive protein structure analysis suggests that Glyma18G281500 differs significantly from that of the other two genes and more stable in terms of atomic interactions. these differences do not necessarily translate into significant variations in terms of overall protein conservation, geometric conformations, or sensitivity to mutations across the three genes [35-38].

4. CONCLUSION

Our study integrates phylogenetic analysis, domain analysis, motif analysis, and expression data to elucidate the distinct behavior of genes within the Rpp1 locus, particularly due to the presence of the ULP1 domain. Additionally, among three Rpp1 genes Glyma18G281500 differs significantly from other two genes and also found with more stable protein structure, may suggest its distinct role over two other genes. While our findings suggest a potential contribution of these genes to rust resistance, it is crucial to acknowledge the possibility of racespecific responses. Further investigations in this realm hold promise for advancing our understanding of rust resistance mechanisms in soybeans.

DISCLAIMER (ARTIFICIAL INTELLIGENCE)

Author(s) hereby declares that NO generative Al technologies such as Large Language Models

(ChatGPT, COPILOT, etc) and text-to-image generators have been used during writing or editing of manuscripts.

SUPPLEMENTARY MARTIALS

Supplementary martials file 1 available in this link:https://journaljabb.com/index.php/JABB/librar yFiles/downloadPublic/21

COMPETING INTERESTS

The authors of this manuscript declare that they have no financial or non-financial conflicts of interest that could influence their research or the interpretation of their results.

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