Insight to the Mode of Action of *Allium sativum* Leaf Agglutinin (ASAL) Expressing in T3 Rice Lines on Brown Planthopper

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**ABSTRACT**

Brown planthopper, the sap sucking hemipteran pest, is one of the major contributors to the yield loss of rice through the world. To combat the situation researchers are interested identifying genes from plant origin having potentiality to develop hemipteran pest resistance. Interestingly, it was observed that rice plants expressing ASAL, a monocot mannose binding lectin, showed significant resistance to brown planthopper and green leafhopper. Additionally, antibiotic resistant marker gene free ASAL expressing rice lines were developed to overcome the biosafety issues. However, the basis behind the resistance against planthoppers is still not clearly understood. Ligand blot assay was performed with total BBMV protein from BPH and a ~56 kDa receptor protein was detected. LC MS/MS analysis revealed that the receptor protein is NADH quinone oxidoreductase (NQO), a key player in electron transport chain, insect defense response and male/female gametogenesis. Presumably interaction of ASAL with NQO may lead to toxicity and loss of fecundity among BPH feeding on ASAL expressing transgenic rice plants. These findings provide a stable scientific basis for considering these transgenic ASAL expressing rice plants as significant product for combating BPH attack associated yield loss of rice.

Keywords: *Allium sativum* Leaf Agglutinin (ASAL); Hemipteran Pest Management; Brown Planthopper; NADH Quinone Oxidoreductase (NQO); Fecundity; Transgenic Rice; Brush Boarder Membrane Vesicle (BBMV) Protein

1. Introduction

Rice (*Oryza sativa* L.) is the most important food crops in the world and about 90% of the world’s rice is grown and consumed in Asia [1]. Production of this crop is greatly hampered by several biotic and abiotic factors of which sap sucking hemipteran pests namely green leafhopper (*Nephotettix virescens*) and brown planthopper (*Nilaparvata lugens*) are highly devastating that cause huge amount of yield loss every year [2].

Between the two hemipterans, BPH is the most damaging pest of rice in several Asian countries as reported in the last 10 years which may be avoided if rice ecosystem is better managed [3]. Direct feeding practice of BPH drains out plant fluids and essential nutrients. During high infestation, plants become brown and finally die.

These hoppers not only damage the plants by sucking the plant sap but also act as vectors of plant viruses [4]. To develop BPH resistant lines, several genes such as Bph1 and Bph2 conferring resistance to the pest have been identified and incorporated in different popular rice varieties by conventional breeding method [5]. Unfortunately, the major genes resistant to BPH were not stable [6]. Therefore, introduction of exotic pest-resistant genes into popular rice cultivars through *Agrobacterium* mediated genetic transformation technique has been preferred.

Among different important insect resistant genes, Bt toxins are one of the most promising proteins but unfortunately Bt toxins remain ineffective against this group of sap-sucking pests [7]. On the other hand plant lectins, specially monocot mannose binding lectins like GNA, ASAL etc. showed significant insecticidal efficacy both in native as well as expressed conditions [8-12]. Interestingly, *Allium sativum* leaf agglutinin (ASAL) from garlic leaf expressed in transgenic rice exhibited significant anti-metabolic effect towards BPH and GLH [13].
However, scientists have great concern about the public acceptance of these plants due to unwanted incorporation of antibiotic resistance gene, *Hygromycin phosphotransferase (hpt II)*, as plant selection marker. Therefore attempts were made to achieve successful elimination of hpt II marker gene by using the cre/lox recombination system. Bacteriophage P1 specific loxP sites were introduced into the gene construct, flanking the antibiotic resistance gene. Independent ASAL-lox-hpt II-lox plants were generated. Cre recombinase gene was introduced separately in rice line. Both cre/lox were brought together by crossing between ASAL-lox T₀ and cre T₀ plants. The resultant ASAL expressing hybrid rice plants were obtained without antibiotic resistant marker gene sequences which showed significant degree of resistance against BPH and GLH. The T₃ progeny plants of one such line were particularly used in the present study. Among the several strategies used for eliminating antibiotic or herbicide resistant marker gene, cre/lox P site-specific recombination system has the key advantage where no commercial enzyme mixture is required for recombination. The 38 kDa cre recombinase specifically recognizes & induces precise recombination & DNA excision between two directly repeated asymmetric 34 bp loxP sites flanking the target DNA sequences to be eliminated [14-16].

Although insecticidal activity of transgenic marker gene free ASAL positive rice lines was well documented [17] against *N. lugens* and other leafhoppers, the mode of action of ASAL against leafhoppers are still unknown and needs to be explored before utilization of the transgenic lines in future IPM programme. Therefore, to achieve a sound scientific basis behind the on field application of transgenic plants, the actual mode of action of ASAL against leaf hoppers needs to be well documented. The aim of this study is to elucidate the mode of action of expressed ASAL in transgenic rice plants against *N. lugens* and establish the potentiality of transgenic antibiotic resistant marker gene free ASAL expressing rice plants for minimizing planthopper associated yield loss of the crop.

2. Materials and Methods

2.1. Materials Used

Transgenic rice seeds cv IR64 used for this study were developed by the present investigating group [17]. BPH were obtained from Regional Rice Research Station at Chinsurah, West Bengal for all the experiments.

2.2. Seed Germination

Rice seeds (T₃) were surface sterilized for 30 minutes in 1.5% (v/v) sodium hypochlorite solution, rinsed thoroughly (3 - 4 times) in sterile water and sown on half strength MS [18] germinating medium with 2% sucrose and 0.7% w/v agar [17]. 30 days old plants were used for molecular analysis.

2.3. PCR Screening and Segregation Analysis of Transgenic Line

Multiplex PCR was carried out with seven T₃ plants (randomly chosen) of T₂ parental line No. L03C04 (6) using ASAL gene specific primer pairs. Genomic DNA was isolated from young green leaves of T₃ progeny plants and control rice plants followed by CTAB extraction method [19]. Multiplex PCR was subjected using Qiagen Multiplex PCR Kit (GmbH, Hilden, Germany). Each 50 µl PCR reaction mixture contained 1x PCR master mix including Hot Star Taq DNA polymerase. Primers (Table 1) were used in 0.2 μM concentration. The reaction started with incubation for 15 minutes at 94°C followed by 35 cycles of [30 s at 94°C, 90 s at 58°C, 90 s at 72°C] and final extension with 10 minutes at 72°C in My Cycler (Bio Rad, Hercules, CA, USA). DNA of pBKkg ASAL and pBK16.2 plasmids was taken as positive control for ASAL and cre gene respectively. DNA of control rice plant was taken as negative control. The segregation patterns of ASAL and cre genes in the T₃ hybrid lines were observed by analysing the presence or absence of respective gene-specific amplicons resolved in 1.2% Agarose gel.

2.4. Western Blotting

Protein from leaves of 1 month old control and T₃ progeny plants were extracted in Tris-cl buffer pH-7. After quantification using Bradford reagent [20], western blotting was carried out with total protein extracted from each of the T₃ progeny plants probed with anti-ASAL antibody according to the protocol described by Dutta et al. 2005 [11]. About 50 μg of total plant protein re-

### Table 1. List of PCR primers.

<table>
<thead>
<tr>
<th>Primers</th>
<th>Sequence</th>
<th>Annealing temperature (°C)</th>
<th>Amplicon size (bp)</th>
</tr>
</thead>
<tbody>
<tr>
<td>ASAL forward</td>
<td>5’ATGGCCAGGAAACTACTGACGA 3’</td>
<td>58</td>
<td>362</td>
</tr>
<tr>
<td>ASAL reverse</td>
<td>5’CTATCCTCTGTAGTGACTCCAGTA 3’</td>
<td></td>
<td></td>
</tr>
<tr>
<td>cre forward</td>
<td>5’ATGTCGAATTTAATGACC 3’</td>
<td>58</td>
<td>1100</td>
</tr>
<tr>
<td>cre reverse</td>
<td>5’CTAATCCCATTCCTCCAGC 3’</td>
<td></td>
<td></td>
</tr>
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solved in 15% SDS-PAGE was electrophoretically transferred to Hybond-C membrane (GE Healthcare, UK) using Hoeffer (Hoefer Inc., Holliston, MA, USA) electroblot apparatus. The membrane was transiently stained with Ponceau S (Sigma-Aldrich, USA) for confirming the transfer of proteins onto it. Blocking of membrane was done with 5% non-fat milk (Merck, Germany) dissolved in 1x PBST at 37°C for 3 hours. After washing with 1x PBST, the membrane was then incubated with anti-ASAL polyclonal antibody (1:4000) for 2 hours at 37°C. Further probing of the primary antibody was done for 2 hours with anti-rabbit IgG-HRP conjugate (1:20,000) (Sigma-Aldrich, USA) and developed using Hybond ECL kit on Hybond ECL film (GE Healthcare, Germany). Purified native ASAL protein (0.5 µg) was used as positive control and total protein (50 µg) from untransformed plants was used as negative control.

2.5. ELISA

100 µg of crude protein extracted from T3 transgenic leaves of each plant were coated in the wells of microtiter plate with coating buffer (15 mM Sodium carbonate, 35 mM sodium bicarbonate, 3 mM sodium azide). After charging the protein extract with primary and secondary antibody, substrate [O-phenylene diamin dia mine hydrochloride (OPD, BRL)] dissolved in citrate buffer (pH 9.6) was added and coloured reaction was developed. Optical density was measured at 415 nm in a Bio-Rad microtiter plate reader model no. 680 [11].

2.6. Hemagglutination Assay

The heamagglutination assay was done with the total protein extracts (100 µg/well) of five T3 ASAL expressing lines. Blood sample was collected from the rabbit by a syringe and immediately transferred to a glass tube pre filled with 0.9% saline as described by Banerjee et al. [21]. The erythrocytes were incubated with plant extracts in microtiter well and after 1 hour of incubation at 37°C, the hemagglutination activity was checked.

2.7. In planta Insect Bioassay

T3 plants expressing ASAL and control plants were subjected to insect bioassay with BPH as described by Sengupta et al. [17]. Five marker gene free plants (line no. SD2, SD4, SD5, SD7 and SD8) were tested with BPH nymph and compared the mortality of the insects incubated on control plants. Each set was infested with 20 BPH nymphs of first instar into an individual insect-proof cage and survivability of insects was monitored at every 2 days from the release of insects up to 16 days. Effect of ASAL on fecundity of BPH was assessed by measuring total nymph production by adult insects on the transgenic rice plants (line no. SD2, SD4, SD5, SD7 and SD8). Each experimental set was repeated five times and statistical analysis was done in order to compare the significance of differential mortality among all transgenic and controls plants.

2.8. Ligand Blot Assay

Total brush border membrane vesicle (BBMV) from *Nilaparvata lugens* was isolated according to Bandopadhyay et al. [22] using MgCl2 sequential precipitation method suited for luminal membranes. About 20 µg of total BBMV resolved in 12% SDS-PAGE was electrophoretically transferred to Hybond-C membrane (GE Healthcare, UK) using Hoeffer (Hoefer Inc., Holliston, MA, USA) electroblot apparatus. The membrane was temporarily stained with Ponceau S (Sigma-Aldrich, USA) to ensure the protein transfer from gel to Hybond-C membrane. Blocking of membrane was done with 5% non-fat milk (Merck, Germany) dissolved in 1x PBST at 37°C for 2 hours. The membrane was further subjected to incubation with 100 µg of total protein extracted from transgenic ASAL positive SD7 line (ASAL expression level ~5 ng·µg−1) for 2 hours following extensive washing with 1x PBST. The membrane was then incubated with anti-ASAL antibody (1:4000) for 2 hours at 37°C. Further probing of the secondary antibody was done with anti-rabbit IgG-HRP conjugate (1:20,000) (Sigma-Aldrich, USA) and film was developed using Hybond ECL kit (GE Healthcare, Germany) as described by Mondal et al. [23]. Ligand blot assay with total protein (100 µg) from non-transgenic rice plant served as negative control.

2.9. LC MS/MS Analysis of Insect Receptor

Receptor protein band detected via ligand blot assay when challenged with total protein from ASAL positive transgenic rice line were excised, trypsinned and processed for LC MS/MS analysis as described in details by Sarkar et al. [24]. The extracted peptides were analyzed by LC MS/MS on Orbitrap Velos. Individual MS/MS spectra were searched through Mascot search using UNIPROT_2011_08 database with preset parameters such as standard global modifications, minimum of 5 peptide match and 99% protein thresholds. The results were observed using Scaffold (version Scaffold_3.6.4, Proteome software Inc., Portland, OR).

3. Results

3.1. PCR Analysis

Seven T3 plants were taken from T2 parental line no.
Insight to the Mode of Action of *Allium sativum* Leaf Agglutinin (ASAL) Expressing in T3 Rice Lines on Brown Planthopper

L03C04 (6) (randomly chosen) for multiplex PCR analysis. Five plants were found to be positive for ASAL and two plants were found to be negative suggesting about the segregation of ASAL in T3 generation (Figure 1). Absence of *cre* gene was also observed in the ASAL positive lines.

### 3.2. Expression of ASAL in T3 Progenies

Western blot of total protein extract from 1 month old leaves of the five T3 lines were analysed which showed ASAL positive protein band at 12.5 kDa region (Figure 2(a)). ELISA result of five plants showed ASAL expression level in the range of 0.4% to 0.5% of the total soluble protein fraction (Figure 2(b)). Among these five lines, SD7 rice line showed highest expression level of ASAL (0.5%).

### 3.3. Agglutination Assay

In the haemagglutination assay, ASAL positive T3 lines showed agglutination of rabbit erythrocytes by forming a carpet of erythrocytes cells in the wells (Figure 3: Panel A, well 1 - 5) whereas the control line showed a tight button of rabbit erythrocytes cells at the centre of the well (Figure 3: Panel B, well 1 - 5) indicating the negative agglutination reaction. This result confirmed the retention of lectin activity of the expressed ASAL from transgenic rice lines.

### 3.4. Insect Bioassay

*In planta* insect bioassay was carried out on 1-month-old five ASAL-expressing T3 progeny rice plants. All five transgenic plants inoculated with BPH, showed significant reduction in survivability and fecundity of the pest compared to that of the insects incubated on untransformed control plants. The number of living insect declined to 5.3 ± 0.33 (27%) to 6 ± 1.0 (30%) whereas in control plant the number of living insect was 13.66 ± 0.3 (68.3%) (Figure 4(a)). The mean number of survival of BPH on transgenic plants [t(0.01) 18.37] was significantly (p < 0.01) different than the control plants (unpaired t test). After 16 days of the experiment SD7 line represents maximum mortality of BPH. Fecundity of BPH on ASAL expressing plants was also examined by counting the total nymphs produced by the adult insects on transformed and untransformed control rice plants. After 30 days of assay period, the mean number of BPH nymphs on ASAL positive T3 plants was found to be 50 ± 2.8 (42.4%) to 40.67 ± 1.2 (35%) where as in control plants the number increased up to 118 ± 4.1 (100%) (Figure 4(b)) suggesting the lower reproductive rate of BPH fed on transgenic rice plants compared to the control rice plants.

### 3.5. Ligand Blots Assay and LC MS/MS Analysis

Approximately 56 kDa receptor protein from the brush border membrane vesicle (BBMV) of *N. lugens* was lightened up after ligand blot assay carried out with total protein from transgenic ASAL positive SD7 rice lines (Figure 5). Identified receptor protein band was subjected to LC MS/MS analysis. Sequence information of peptide fragments obtained through MS/MS were individually searched against Uniprot database and receptor protein identified as NADH-quinone oxidoreductase from *N. lugens* [Accession no: Q5W1K6_NILLU] with a sequence coverage of 45% (Table 2, Supply Figure 1).

### 4. Discussion

Lectin has been defined as carbohydrate-binding protein or glycoprotein of non-immune origin which agglutinates or precipitates glycoconjugates linked to the erythrocytes [25]. Lectins are generally multivalent in nature and the specificity of which is largely determined by monosaccharide termini. Several plants possess lectin particularly in the seeds and tubers as storage protein. Some of which also serve as defence proteins when infested by enemies. A broad spectrum of plant lectins has been tested on several insect species more specifically on hemipterans. Usually the first experiments to check the effects of a particular lectin on an insect are conducted with an artificial diet. A major drawback of this approach is the composition of the artificial diet, which often differs considerably from the natural insect diet. Furthermore, for rearing of many insects no artificial diet had been worked out yet. However for controlling hemipteran pests, efforts have been made to target lectin ex-
Insight to the Mode of Action of *Allium sativum* Leaf Agglutinin (ASAL) Expressing in T3 Rice Lines on Brown Planthopper

4.1. Molecular Analysis of Transgenic T3 Lines

Analysing T3 marker free transgenic rice lines, it was found that 71% of the transgenic rice plants showed presence of *ASAL* gene and absence of *cre* gene in this generation suggesting about segregation of ASAL among T3 progenies which is close to attain homozygosity. Significant level of ASAL expression was observed in T3 rice lines revealed by ELISA and western blot assay.

4.2. Insecticidal Activity of T3 ASAL Positive Rice Lines

*In planta* insect bioassay experiments demonstrated significant insecticidal efficacy of different transgenic ASAL positive rice lines against BPH. Furthermore, fecundity rate was also observed to be reduced among the insects (BPH) fed on transgenic rice plants which seem to be comparable with the earlier reports from the present investigating group [13,17].

4.3. Basis behind the Observed Toxicity of Expressed ASAL against *N. lugens*

The insecticidal efficacy of ASAL in SD2, SD4, SD5, SD7 and SD8 T3 rice lines had already been documented by *in planta* insect bioassay experiment. To understand the basis of this toxic effect, ligand blot assay had been done and ~56 kDa putative receptor protein band from *N. lugens* BBMV fraction was detected. LC MS/MS analysis has shown the receptor protein band (56 kDa) was of NADH quinone oxidoreductase enzyme of *N. lugens* [27].

NADH quinone oxidoreductase (NQO) is the first enzyme of electron transport respiratory chain. It contributes to ATP synthesis. Studies revealed that some secondary metabolites like rotenone and piericidin A from microbial and plant sources are important components that act on the respiratory chain by inhibiting NQO [28]. It was also documented by earlier investigators that NQO gene expression was enhanced in BPH, feeding on B5 (BPH resistance) rice lines, indicating that there might be some compounds having a similar structure and function as rotenone or piericidin A generated in B5 plants, which might inhibit the respiratory chain enzymes in BPH to...
Insight to the Mode of Action of *Allium sativum* Leaf Agglutinin (ASAL) Expressing in T₃ Rice Lines on Brown Planthopper

**Figure 4.** *In planta* insect bioassay. (a) The graph shows the percentage of survival of BPH on untransformed control plant and five T₃ progeny plants SD2, SD4, SD5, SD7, SD8; (b) Bar diagram showing fecundity pattern of BPH fed on untransformed (control) and ASAL expressing rice plants (SD2, SD4, SD5, SD7, SD8).

**Figure 5.** SDS-PAGE (12%) profile and subsequent ligand blot assay of total BBMV protein from *N. lugens*. Lane M: Standard protein molecular weight marker. Lane 1: SDS PAGE profile of total BBMV protein fraction of *N. lugens* stained by Coomassic brilliant blue; Lane 2: Ligand blot analysis of total BBMV protein of *N. lugens* challenged with total protein from SD7 transgenic rice lines showing putative ~56 KDa receptor protein band of ASAL. Lane 3: Ligand blot analysis of total BBMV protein of *N. lugens* challenged with Control non-transgenic rice line (negative control).

Certain degree. This result emphasises that NQO may play important role in the defence response of BPH against resistant rice lines [27]. So it is quite obvious that binding of ASAL to NQO may upset insect defence mechanism and costs its survival.

It has also been reported in *Drosophila melanogaster* that the mitochondrial acyl carrier protein (mtACP), a subunit of NADH ubiquinone oxidoreductase is involved in *de novo* fatty acid synthesis in the mitochondrion. AP-element-induced loss-of-function mutation in the mtacp1 gene resulted in lethality, indicating that the gene is essential for viability of the insect [29]. Additional studies showed that mtacp1 is expressed at higher levels during late embryogenesis, in the pupa and adult mosquitoes, indicating its requirement for both male and female gametogenesis and possibly for tracheal development in *Drosophila*. Correlating the earlier observations with our present findings, it is assumed that ASAL binding to NQO of BPH can affect normal development and gametogenesis, which results into loss of fecundity of insects feeding on presently, mentioned ASAL rice lines.

**5. Conclusion**

Antibiotic resistant marker gene free ASAL expressing rice lines are found to be highly resistant to BPH. ASAL, having binding affinity to NQO (a respiratory chain enzyme) of BPH, exerts its toxicity resulting in high mortality and loss of fecundity. Above findings provide adequate information about the potentiality of ASAL expressing advanced progenies of rice plants to minimize the crop loss due to sucking insect attack.

**6. Acknowledgements**

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Table 2. Receptor identification through LC MS/MS.

<table>
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<tr>
<th>Target insect</th>
<th>Uniport accession no</th>
<th>Receptor</th>
<th>Organism</th>
<th>Obs KDa/Exp KDa</th>
<th>Amino acid match</th>
<th>Sequence coverage</th>
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<tbody>
<tr>
<td>Brown Planthopper (N. lugens)</td>
<td>Q5W1K6_NILLU</td>
<td>NADH-quinone oxidoreductase</td>
<td>N. lugens</td>
<td>56/51.2 KDa</td>
<td>209/462</td>
<td>45%</td>
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Swarnava Das and Mr. Sudipta Basu are also duly acknowledged.

REFERENCES


Insight to the Mode of Action of *Allium sativum* Leaf Agglutinin (ASAL) Expressing in T3 Rice Lines on Brown Planthopper


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Supplementary Figure 1

Q6W1K6_NILLU (100 %), 5120.0 Da

(Q6W1K6) NADH-quinone oxidoreductase OS=Niliaparvata lugens PE=2 SV=1
14 unique peptides, 17 unique spectra, 37 total spectra, 209462 amino acids (45 % coverage)