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## An Initial Event in Insect Innate Immune Response: Structural and Biological Studies of Interactions between β-1,3-glucan and the N-terminal Domain of β-1,3-glucan Recognition Protein

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## Abstract

In response to invading microorganisms, insect  $\beta$ -1,3-glucan recognition protein ( $\beta$ GRP), a soluble receptor in the hemolymph, binds to the surfaces of bacteria and fungi and activates serine protease cascades that promote destruction of pathogens by means of melanization or expression of antimicrobial peptides. Here we report on the NMR solution structure of the N-terminal domain of βGRP (N-βGRP) from Indian meal moth (*Plodia interpunctella*), which is sufficient to activate the prophenoloxidase (proPO) pathway resulting in melanin formation. NMR and isothermal calorimetric titrations of N- $\beta$ GRP with laminarihexaose, a glucose hexamer containing  $\beta$ -1,3 links, suggest a weak binding of the ligand. However, addition of laminarin, a glucose polysaccharide ( $\sim 6 \text{ kDa}$ ) containing  $\beta$ -1,3 and  $\beta$ -1,6 links that activates the proPO pathway, to N- $\beta$ GRP results in the loss of NMR cross-peaks from the backbone <sup>15</sup>N-<sup>1</sup>H groups of the protein. suggesting the formation of a large complex. Analytical ultra centrifugation (AUC) studies of formation of N-BGRP:laminarin complex show that ligand-binding induces self-association of the protein:carbohydrate complex into a macro structure, likely containing six protein and three laminarin molecules (~ 102 kDa). The macro complex is quite stable, as it does not undergo dissociation upon dilution to sub-micromolar concentrations. The structural model thus derived from the present studies for N-BGRP: laminarin complex in solution differs from the one in which a single N- $\beta$ GRP molecule has been proposed to bind to a triple helical form of laminarin on the basis of an X-ray crystallographic structure of N-βGRP:laminarihexaose complex [Kanagawa, M., Satoh, T., Ikeda, A., Adachi, Y., Ohno, N., and Yamaguchi, Y. (2011) J. Biol. Chem. 286, 29158-29165]. AUC studies and phenoloxidase activation measurements carried out with the designed mutants of N-βGRP indicate that electrostatic interactions involving Asp45, Arg54, and Asp68 between the ligand-bound protein molecules contribute in part to the stability of NβGRP:laminarin macro complex and that a decreased stability is accompanied by a reduced activation of the proPO pathway. Increased  $\beta$ -1,6 branching in laminarin also results in

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Atomic coordinates have been deposited into the Protein Data Bank (PDB) with accession number 2KHA.

SUPPORTING INFORMATION

The supporting information includes a table of statistics for a structural ensemble of 20 lowest- energy structures of Plodia interpunctella N-BGRP; MALDI mass spectrum of laminarin, figure of B-1,3-glucan-binding activities of N-BGRP and mutants as measured by curdlan pull-down assay and isothermal titration calorimetry, g(s\*) profiles of sedimentation velocity studies of N-β GRP in the presence of varying amounts of laminarihexaose, and sedimentation velocity profiles of N-BGRP:laminarin complex with increasing amounts of laminarin. This material is available free of charge via the Internet at http://pubs.acs.org.

destabilization of the macro complex. These novel findings suggest that ligand-induced selfassociation of  $\beta$ GRP: $\beta$ -1,3-glucan complex may form a platform on a microbial surface for recruitment of downstream proteases, as a means of amplification of the initial signal of pathogen recognition for the activation of the proPO pathway.

#### Keywords

pathogen recognition; innate immunity; β-1,3-glucan; βGRP

## INTRODUCTION

 $\beta$ GRPs, also called Gram-negative bacteria binding proteins (GNBPs), are a family of insect pathogen recognition receptors that bind to  $\beta$ -1,3-glucan, a structural component of fungal cell walls and bacterial surfaces (1–6). The protein-carbohydrate binding event triggers activation of serine protease cascades leading to the activation of the proPO pathway in the hemolymph and intracellular Toll signaling (7,8). The activated proPO pathway produces melanin that encapsulates pathogenic microorganisms, whereas the Toll pathway results in the expression of antimicrobial peptides/proteins.

βGRPs share a conserved primary structure comprising an amino-terminal carbohydratebinding domain (N-βGRP) and a carboxyl-terminal β-1,3-glucanase-like domain. N-βGRP binds to curdlan, a linear water-insoluble β-1,3-glucan polysaccharide, and to laminarin, a water-soluble β-1,3-glucan polysaccharide containing β-1,6 branches (1,9). *P. interpunctella* N-βGRP mixed with laminarin generates a significant synergistic activation of the proPO pathway (9). N-βGRP also induces aggregation of microorganisms such as *Saccharomyces cerevisiae* and *Escherichia coli*, albeit less effectively than does the full-length protein (9). Aggregation of pathogens *in vivo* may create a superior trigger for activating biochemical cascades of cellular immunity (9) or may provide a platform to assemble effector complexes (10). To gain insight into the molecular mechanism of such pathogen recognition events requires, as a first step, characterization of the structural basis and consequences of binding of β-1,3-glucan by N-βGRP.

Three-dimensional structures have been reported for N- $\beta$ GRP from three insect species, Bombyx mori (11,12), Drosophila melanogaster (13) and P. interpunctella (12): all three N- $\beta$ GRPs adopt a common immunoglobulin-like fold with two sheets forming a  $\beta$ -sandwich (Figure 1A). However, different models have been proposed for the binding of  $\beta$ -1,3-glucan by N- $\beta$ GRP. In the model based on the solution structure of *B. mori* N- $\beta$ GRP, as determined by means of Nuclear Magnetic Resonance (NMR) spectroscopy, the  $\beta$ -1,3-glucan binding site is located on the concave sheet (strands A, B and E) (11), while in the models based on the X-ray crystal structure of ligand-free *D. melanogaster* GNBP3 N-domain (13) and that of *P. interpunctella* N- $\beta$ GRP complexed with laminarihexaose (12), the binding site is confined to the convex sheet (strands C, C', F, G and G'). Furthermore, these three models do not provide any clues for understanding the molecular basis of the synergistic activation of the proPO pathway by  $\beta$ -1,3-glucan and N- $\beta$ GRP.

In the present work, we describe the NMR solution structure determination of *P. interpunctella* N- $\beta$ GRP (N-terminal region of 118 residues) and mapping of the  $\beta$ -1,3-glucan binding site. We have characterized the binding of laminarihexaose, and that of laminarin, to N- $\beta$ GRP by NMR, isothermal titration calorimetry (ITC), analytical ultracentrifugation studies (AUC), site-directed mutagenesis, and prophenoloxidase activity measurements. Our results demonstrate, for the first time, ligand-induced self-association of N- $\beta$ GRP: $\beta$ -1,3glucan complex. This process has a significant contribution from electrostatic interactions

between N- $\beta$ GRP molecules in the complex. Reduction in the extent of self-association between N- $\beta$ GRP: $\beta$ -1,3-glucan complex molecules leads to a decrease in the rate of proPO activation by the macro complex. Thus it is suggested that such a protein:carbohydrate macro complex formation is a structural initiation signal for serine protease cascade activation of insect immune response and a likely prerequisite for aggregation of microorganisms.

## MATERIALS AND METHODS

#### Materials

Laminarin from *Laminaria digitata* was purchased from Sigma Aldrich (L9634, St. Louis, MO), and its  $\beta$ -1,3 to  $\beta$ -1,6 cross-link number ratio was 7 (14). Laminarihexaose and curdlan (a water-insoluble  $\beta$ -1,3-glucan) were from Megazyme (Wicklow, Ireland). A much more branched laminarin from *Eisenia bicyclis* with a  $\beta$ -1,3 to  $\beta$ -1,6 cross-link number ratio of 3 (15) was purchased from Tokyo Chemical Industry (Tokyo, Japan).

#### **Protein Expression and Purification**

The DNA sequence of *P. interpunctella* N-βGRP was cloned via BamHI/HindIII sites into Invitrogen pPROEX HTb plasmid (9). The resulting recombinant protein contains an Nterminal hexahistidine tag followed by a TEV protease cleavage site before the N-βGRP sequence. Expression and purification of His<sub>6</sub>-N-βGRP was performed as described previously (9) with a slight modification. His<sub>6</sub>-N-βGRP was expressed in *E. coli* BL21 cells by induction with isopropyl-β-D-thiogalactopyranoside and purified using a Ni<sup>2+</sup>affinity column. The hexahistidine tag was cleaved by incubating His<sub>6</sub>- N-βGRP with His<sub>6</sub>-TEV protease (GE Helthcare), and the resultant mixture was passed through a Ni<sup>2+</sup> affinity column to remove the cleaved hexahistidine tag, the TEV protease and uncleaved protein. Purity of N-BGRP was confirmed by sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE), and the amino acid sequence of recombinant N-BGRP was confirmed by mass spectrometry. The recombinant protein had five extra N-terminal and seven extra C-terminal residues from the vector used. Proteins uniformly labeled either with <sup>15</sup>N or with both <sup>13</sup>C and <sup>15</sup>N were produced by growing *E. coli* in M9 medium containing <sup>15</sup>N-ammonium chloride and either D-glucose or uniformly <sup>13</sup>C-labeled Dglucose (Cambridge Isotope Laboratories), respectively.

N-βGRP containing mutation D45A or D45K was prepared according to the instructions of QuikChange mutagenesis kit (Stratagene). The mutation was confirmed by DNA sequencing and mass spectrometry of the purified protein.

#### NMR Spectroscopy

A typical solution sample of N-βGRP used for three-dimensional NMR experiments was 1.7 – 2.0 mM uniformly <sup>13</sup>C/<sup>15</sup>N-labeled protein in 20 mM sodium phosphate buffer, pH 6.5. NMR data were collected at 25°C using a Bruker Avance 800 MHz spectrometer equipped with a cryogenic probe at Structural Biology Center, University of Kansas. The data sets were processed using NMRPipe (16) and analyzed using Sparky (17). A series of standard 2D- and 3D-NMR spectra was collected for backbone and side-chain resonance assignments and structural constraint measurements (18). These include 2D <sup>15</sup>N-HSQC and <sup>13</sup>C-HSQC, 3D CBCA(CO)NH, HNCACB, HN(CO)CA, HNCA, CCCONH, HCCCONH, HCCH-TOCSY, <sup>15</sup>N-edited TOCSY, <sup>15</sup>N-edited NOESY, and <sup>13</sup>C-edited NOESY. An additional 3D <sup>15</sup>N-edited NOESY-HSQC data set was gathered from the <sup>15</sup>N-labeled sample in order to identify HN-HN contacts. Sequence-specific backbone <sup>1</sup>H, <sup>15</sup>N, and <sup>13</sup>C resonance assignments were made from an analysis of the 3D NMR data sets, CBCA(CO)NH, HNCACB, HNCA. Side-chain resonances were assigned using the data

from 3D CCCONH, HCCCONH, HCCH-TOCSY, and <sup>15</sup>N-edited TOCSY. Unambiguous NOE constraints were manually assigned using 3D <sup>15</sup>N-edited and <sup>13</sup>C-edited NOESY spectra. Dihedral constraints for the backbone torsion angles ( $\phi$ ,  $\psi$ ) were obtained using the software TALOS (19). Hydrogen bond constraints were employed for  $\beta$ -stand regions based on the NOE patterns and the chemical shift index prediction of the secondary structure (20). Structures of N- $\beta$ GRP were calculated by a simulated annealing procedure using CNS (21). One hundred structures were calculated from which twenty lowest-energy structures were chosen for the final structural ensemble.

For NMR titration experiments, 2D <sup>1</sup>H-<sup>15</sup>N HSQC spectra were collected of 0.5–1.0 mM <sup>15</sup>N-labeled N- $\beta$ GRP in 20 mM sodium phosphate buffer (pH 6.5) at 25°C. Laminarihexaose or laminarin solutions (3 mM and 1mM, respectively) prepared in the same buffer and then added stepwise to the protein sample (0.5 mM). The chemical shift difference between ligand-free and ligand-saturated protein was calculated for each backbone NH group as [( $\Delta$ H2+( $\Delta$ N/5)2)/2]1/2.

#### Fluorescence Spectroscopy

Fluorescence emission spectra were recorded with a Cary Eclipse fluorescence spectrophotometer (Varian, Inc.) equipped with dual monochromators. The experiments were performed using a 1 cm path length 3.5 ml quartz cuvette. The samples were excited at 295 nm, and emissions were measured between 300 and 400 nm to monitor Trp fluorescence.

#### Analytical Ultracentrifugation

Sedimentation velocity experiments were conducted with an Optima XL-I ultracentrifuge (Beckman Coulter, Inc. Brea, CA) using an An-60 Ti rotor at 20°C with 50 mM Tris-HCl (pH 7.3) buffer containing 50 mM NaCl (22). Sedimentation was monitored by absorbance or interference optics using double-sector aluminum cells with a final loading of 400  $\mu$ l per sector. Sedimentation was performed at 49,000 rpm with scans made at 5 min intervals. Data were analyzed using DCDT+ software version 1.16 (www.jphilo.mailway.com). Sedimentation coefficients were calculated using  $g(s^*)$  and dc/dt fitting functions in DCDT+ software. Buffer density and viscosity were calculated by SEDNTERP version 1.08 (www.jphilo.mailway.com). Bovine serum albumin (BSA) was used as a standard to account for effects of changes in solution density and viscosity when high concentrations of laminarin or laminarihexaose were used. The partial specific volume of a protein was calculated from its amino acid composition using SEDNTERP (0.7306 ml/g for N- $\beta$ GRP at 20°C). The partial specific volume used for laminarin was 0.622 ml/g (23).

#### **Isothermal Titration Calorimetry**

ITC measurements were carried out using MCS-ITC system (MicroCal, Northampton, MA) at 30°C (24). Recombinant N- $\beta$ GRP and laminarin solutions were dialyzed overnight at 4°C against 50 mM Tris-Cl (pH 7.5) buffer containing 50 mM NaCl. Laminarihexaose was directly dissolved in the buffer. All solutions were degassed before use. A typical experiment consisted of 20 injections of 10  $\mu$ l laminarin or laminarihexaose solution (1.67 mM) into 1.38 ml protein solution. The heat of dilution of laminarin or laminarihexaose solution upon injection into the buffer solution was subtracted from the experimental titration data. Baseline corrections and integration of the calorimeter signals were performed using the software, Origin (MicroCal).

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#### **Curdlan Pull-down Assay**

The procedure described by Fabrick *et al.* (9) was employed. Briefly, for each assay, 20  $\mu$ g of purified protein ( $\beta$ GRP-N wide type, D45A, or D45K, respectively) was incubated with 0.5 mg of curdlan for 10 min. The protein-curdlan mixture was centrifuged at 10,000 × g for 5 min, and the supernatant corresponding to the unbound fraction was saved. The bound protein was eluted from curdlan by heating at 95°C for 5 min in SDS sample buffer. Equal volumes of purified, unbound and bound proteins were analyzed by SDS-PAGE with Coomassie Blue staining.

#### Activation of the ProPO Pathway

A method as described by Ma and Kanost (2) and Fabrick *et al.* (4) was used, incorporating the modified protocol recently developed by Laughton and Siva-Jothy (25) to follow proPO activation in the absence or presence of laminarin. Briefly, 10  $\mu$ l of recombinant proteins (0.4 mg/ml) was incubated with 5 [l of buffer or laminarin (10 mg/ml) and mixed with 5  $\mu$ l of *Manduca sexta* plasma. The volume of each sample well was brought up to 130  $\mu$ l with sodium phosphate buffer (pH 6.8). After incubation for 15 min at room temperature, 20  $\mu$ l of 30 mM dopamine hydrochloride was added and phenoloxidase activity was determined by measuring absorbance at 490 nm. The phenoloxidase activity was performed using Prism 5 (GraphPad Software).

## **RESULTS AND DISCUSSION**

#### Homogeneity of N-βGRP

The homogeneity of extensively purified N- $\beta$ GRP was characterized by AUC. The sedimentation analysis of N- $\beta$ GRP exhibited a homogeneous species with an  $S_{20, w}$  value of 1.90 S corresponding to a weight average molecular weight of 14,500 by *dc/dt* analysis--a value in good agreement with the calculated mass of 14,601 Da. N- $\beta$ GRP sedimentation pattern remained the same for a 1.7 mM solution (data not shown), thus indicating that N- $\beta$ GRP existed as a monomer in solution at concentrations used for NMR experiments.

#### N-βGRP Solution Structure and Mapping of β-1,3–Glucan Binding-Site

The monomer structure of N- $\beta$ GRP was determined using solution NMR methods (Figure 1B). The chemical shifts of backbone and side-chain <sup>1</sup>H, <sup>15</sup>N, and <sup>13</sup>C nuclei of backbone were assigned by the application of standard multi-dimensional heteronuclear NMR methods (26). Upfield C $\gamma$  chemical shift and downfield C $\beta$  chemical shift of Pro19, together with the strong NOE correlation between its H $\alpha$  and H $\alpha$  of preceding Tyr18, indicate that the peptide bond between Tyr18 and Pro19 adopts a *cis* conformation, consistent with the reported crystal structures (12,13), but in contrast to the *trans* conformation reported for the corresponding Pro in the solution structure of *B. mori* N- $\beta$ GRP (11). The heteronuclear steady-state {<sup>15</sup>N}-<sup>1</sup>H NOE values are sensitive to the dynamics of the peptide NH bonds on a sub-nanosecond time scale (27). For residues Val8-Thr110, the average value of heteronuclear NOE is 0.84. The decreased NOE values observed for the N- and C- terminal regions reflect their relatively increased flexibility (Figure 1C).

Structural statistics for the twenty lowest-energy structures are summarized in Supplementary Table 1. We have identified the eight  $\beta$ -strands according to the standard immunoglobulin nomenclature (28): Three  $\beta$ -strands, A (Lys13-Ile17), B (Gly21-Pro27) and E (Arg63-Asp68), comprise the concave sheet;  $\beta$ -strands C (Ser32-Leu40), C' (His51-Ile56), F (Lys78-Ile86), G (Gly91-Gln94) and G' (Gly97-Thr100) comprise the convex sheet. Strands G and G' are split by a kink (Asp95-Gln96). Two long loops, C-C' and E-F, have relatively higher flexibility than the strands as indicated by their slightly lower heteronuclear NOE values. However, the large number of inter-residue <sup>1</sup>H-<sup>1</sup>H NOE distance constraints observed for residues in these two loops indicates that the loops are ordered and fairly rigid (Figure 1B), and hence are unlikely to undergo a large conformational change suggested by Mishima *et al.* (13). Furthermore, we measured the intensity of fluorescence emitted by tryptophans, among which Trp81 is located on the strands F and shielded by loop C-C', before and after addition of laminarin to the protein sample, and found no change, thus ruling out any conformational changes for the C-C' and E-F loops brought about by laminarin-binding. Interestingly, the crystal structure of *P. interpunctella* N- $\beta$ GRP complexed with laminarihexaose (12) also shows no conformational alteration for these loops.

Effect of  $\beta$ -1,3-glucan binding on N- $\beta$ GRP structure was characterized by titrating laminarihexaose, a  $\beta$ -1,3-linked glucose hexamer, into a solution of N- $\beta$ GRP and monitoring the NMR cross-peaks of the protein backbone <sup>15</sup>N-<sup>1</sup>H groups (Figure 2A & B). The perturbed residues are mainly localized on the convex sheet (Figure 2C), consistent with the hexasaccharide-binding site identified in the crystal structure of N- $\beta$ GRP:laminarihexaose complex (12). The crystal structure shows that this region interacts with three laminarihexaose molecules, which led the authors to propose that N- $\beta$ GRP binds to a triple helical structure of laminarin (12). However, to date no structural data are available for laminarin which is a branched polysaccharide. The X-ray fiber diffraction data collected for the linear  $\beta$ -1,3 polysaccharide, curdlan, have been interpreted to be indicative of a single-helical or a triple- helical structure, depending upon the hydration state of the polysaccharide (29, 30). In the presence of water, the diffraction data have been shown to be consistent only with a single-helix structure (29).

ITC experiments did not detect any heat exchange for the titration of laminarihexaose (up to 1.6 mM) into N- $\beta$ GRP (78  $\mu$ M; data not shown). This result is consistent with weak laminarihexaose - N- $\beta$ GRP interactions, as suggested by the NMR data (Figure 2A & B).

#### Interactions between N-βGRP and Laminarin

Laminarin is a water-soluble oligosaccharide containing both  $\beta$ -1,3 and  $\beta$ -1,6 glycosidic bonds, and the number of  $\beta$ -1,6 glycosidic bonds varies depending upon the source. *L. digitata* laminarin used in this work has a  $\beta$ -1,3/ $\beta$ -1,6 glycosidic bond ratio of 7 (14). It shows a broad mass distribution (up to 7,505 Da) with major mass spectral peaks in the 3932 – 4580 Da region as obtained by MALDI-TOF mass spectrometry (Supplementary Figure 1), consistent with the work of Barral *et al.* (31). Sedimentation velocity analysis of laminarin gave an  $S_{20, w}$  value of 1.02 S, which yielded an apparent molecular weight of 4.5 kDa by *dc/dt* analysis. At a high concentration (5mg/ml), laminarin had a similar S value, which suggests that laminarin does not associate/aggregate by itself. Young *et al.* determined an average molecular weight of 7,700 Da for laminarin from light-scattering experiments (32). For the purpose of calculating molarities of laminarin solutions used in our work, we adopted a value of 6 kDa for the molecular weight, as provided by the supplier (Sigma Aldrich).

When laminarin was titrated into N- $\beta$ GRP, nearly all the cross-peaks in the <sup>1</sup>H-<sup>15</sup>N HSQC spectrum broadened to a noise level (Figure 3). The NMR sample still remained clear with no precipitation. These observations indicate that in the presence of laminarin, N- $\beta$ GRP forms a higher order, water-soluble structure. The results also suggest that the binding-site identified for laminarihexaose could interact with the surface structure of a carbohydrate polymer. The few cross-peaks that still remained after addition of laminarin to N- $\beta$ GRP arise from flexible side-chain NH groups and peptide NH groups of some terminal residues.

Binding of laminarin to N- $\beta$ GRP was characterized using ITC (Supplementary Figure 2). Because of the heterogeneity in laminarin as well as self-association of the resulting protein:carbohydrate complex, a likely cooperative process, the ITC data, although very similar to those of earlier studies (12,13), were not fit to any simple binding equilibrium. Qualitatively, the ITC data provided evidence for the binding of laminarin by N- $\beta$ GRP, an exothermic process.

Interaction between N-BGRP and laminarin was characterized by sedimentation velocity analysis using absorption optics, which detects the distribution of the protein (Figure 4): As the concentration of laminarin added to a fixed concentration of N-βGRP (26.2 μM) increased from 8.3 to 24.9  $\mu$ M, the concentration of free N- $\beta$ GRP at 1.9 S decreased and a broad peak appeared between 4 and 9 S. With further increase in laminarin concentration (83.3  $\mu$ M), nearly all of N- $\beta$ GRP bound to laminarin and a major peak appeared at 5.5 S. A 20-fold excess of laminarin (1.87 mM) caused a slight decrease in S value (5.1 S) with a shoulder at ~8 S. A  $g(s^*)$  fitting analysis yielded an average molecular weight of ~95 kDa for the 5.5 S species, a value that is remarkably greater than would be expected for a complex that contains one molecule of N- $\beta$ GRP and three molecules of laminarin. Laminarin contains at least 30 glucose units, and thus each laminarin molecule could bind multiple N-BGRP molecules. However, the sedimentation profile around 5.5 S region changed only in a limited way even after addition of 20-fold excess of laminarin (1.87 mM). This suggests that formation of the 5.5 S species involves not only protein- carbohydrate interactions, but also protein-protein interactions. Because N-BGRP or laminarin alone does not oligomerize in solution, these results indicate that binding of laminarin causes selfassociation of N-BGRP: laminarin complex. Based on the sedimentation coefficient rule (33), which equates the ratio of average molecular weights of two proteins to the ratio of their sedimentation coefficients raised to the power of 3/2, the shoulder at ~8 S may be attributed to a higher order complex that is about twice the size of the 5.5 S macro complex. Similar association profiles were observed for the titration of laminarin into M. sexta N- $\beta$ GRP (data not shown).

AUC experiments were similarly performed to monitor the effect of laminarihexaose on N-  $\beta$ GRP (Supplementary Figure 3): When 54  $\mu$ M N- $\beta$ GRP was mixed with 1.0 mM laminarihexaose, the  $g(s^*)$  vs  $s^*$  plot of N- $\beta$ GRP displayed no significant change, consistent with a weak-binding of the ligand. At a higher concentration of the mixture, 234  $\mu$ M N-  $\beta$ GRP and 14.1 mM laminarihexaose, a faster sedimenting species appeared in the 3–5  $s^*$ region, and with 1.4 mM N- $\beta$ GRP and 26 mM laminarihexaose, the sedimentation profile shifted further to a higher  $s^*$  region. The  $s^*$  value range (3 – 5 S) of the faster sedimenting species falls close to that  $s^*$  value of BSA (66.5 kDa, 4.3 S). These results suggest that high concentrations of laminarihexaose can mildly induce self-association of N-  $\beta$ GRP:laminarihexaose complex. It is of interest to note that *D. melanogaster* GNBP3 Nterminal domain does not bind to laminaritetraose (6), laminariheptaose or laminarihexaose (13). In contrast, several glucan-binding modules belonging to families other than GBM39, to which N- $\beta$ GRP belongs, possess high binding affinities for hexasaccharides (34).

#### Self-association of N-βGRP:Laminarin Complex

The stability of the macro-assembly of N- $\beta$ GRP:laminarin complex was assessed by dilution analysis: AUC experiments (Figure 5) with three different concentrations of N- $\beta$ GRP at a constant N- $\beta$ GRP:laminarin ratio of 1:22.9 reveal that the *s*\* profile of N- $\beta$ GRP:laminarin complex was unchanged at 5 S even after 50-fold dilution (from 34.0  $\mu$ M to 0.68  $\mu$ M of the protein). This result indicates that the macro complex does not undergo dissociation at submicromolar concentrations due to strong protein-protein and protein-carbohydrate interactions. Sedimentation velocity experiments were also performed with the more branched *E. bicyclis* laminarin (Supplementary Figure 4): While 83.3  $\mu$ M *L. digitata* 

laminarin with a  $\beta$ -1,3/ $\beta$ -1,6 ratio of 7 was sufficient for nearly complete conversion of N- $\beta$ GRP (26  $\mu$ M) into the 5.5 S protein:carbohydrate macro complex, a much higher concentration (0.8 mM) of *E. bicyclis* laminarin with a  $\beta$ -1,3/ $\beta$ -1,6 ratio of 3 was required for the macro complex formation. We infer that increased  $\beta$ -1,6 branching reduces the carbohydrate-binding strength of N- $\beta$ GRP and thus the stability of the protein:carbohydrate macro complex.

To determine the stoichiometry of N- $\beta$ GRP:laminarin complex, we performed sedimentation velocity analysis using interference optics. The sedimentation of both N- $\beta$ GRP and laminarin could be monitored, as these water-soluble molecules affect the refractive index of the solution. Fringe displacements caused by sedimentation provide a measure of weight concentration of sedimenting species (35). Sedimentation boundaries were compared between 75  $\mu$ M N- $\beta$ GRP and a mixture of 75  $\mu$ M N- $\beta$ GRP and 125  $\mu$ M laminarin (Figure 6). Fringe displacement of N- $\beta$ GRP alone was 2.5 arbitrary units, and that of the N- $\beta$ GRP:laminarin complex (the faster species than either N- $\beta$ GRP or laminarin alone) was 2.9. By using refractive indices of protein [(dn/dc) × 10<sup>3</sup> = 0.186 ml/mg] and dextran [(dn/dc) × 10<sup>3</sup> = 0.151 ml/mg], the weight ratio of N- $\beta$  GRP:laminarin in the complex was calculated to be approximately 1:0.20, which equals 2:0.92 as a molar ratio. This is consistent with each laminarin (L) molecule binding two N- $\beta$ GRP (P) molecules to form a P<sub>2</sub>L complex. Thus, the 5.5 S species is likely to be a trimer of P<sub>2</sub>L (~ 102 kDa; Figure 7A).

#### Biological Implication of Self-association of N-ßGRP:Laminarin Complex

Earlier work (9) has established that N- $\beta$ GRP and laminarin produce a pronounced synergistic activation of the proPO pathway. Formation of a macro assembly of the protein:carbohydrate complex in solution, as characterized in the current work, likely allows for ready recruitment of circulating  $\beta$ GRP molecules in the hemolymph to initiate protease cascades that function in defense against invading pathogens. The nature of protein-protein interactions present in the complex may be gleaned from a pseudoquadruplex observed in the crystal structure of N- $\beta$ GRP:laminarihexaose complex (12): In the unit cell N- $\beta$ GRP molecules pack in a side-by-side fashion due to strong electrostatic attractions such that the tip of the loop connecting strands C and C' of one molecule fits into a cleft formed between strands C' and E of another molecule. This structural arrangement is stabilized predominantly by the hydrogen bonds and the salt bridge formed between Leu47 and Asp45 of one protein molecule and Asp55, Arg54, and Asp68 of another (Figure 7 B & C).

To test the hypothesis that these electrostatic interactions play a role in the formation of protein:carbohydrate macro complex, we selected Asp45 for mutagenesis because it makes a hydrogen bond with Asp68 and a salt bridge with Arg54 of another protein molecule through its side-chain carboxyl group (Figure 7C, inset). The sedimentation profiles of NβGRP and mutants, D45A and D45K, reveal that the complex formed with D45K shifted to 4 S compared to the 5 S complex formed with the wild type protein. On the other hand, the protein:laminarin macro complex formed by the D45A mutant does not show any change relative to the wild type complex. These results indicate that in the D45K mutant, electrostatic repulsion between positively charged Lys45 and Arg54 on one hand, and absence of a hydrogen bond between Lys45 and Asp68 on the other, perturb the protein:carbohydrate macro complex formation. Clearly, other protein-protein and proteincarbohydrate interactions contribute significantly to the stability of the macro protein:carbohydrate complex, as the D45A mutant shows no alteration in its ability to interact with laminarin and make the protein:carbohydrate macro complex, as compared to the wild type protein. It is of interest to note that while Arg54 is conserved in the threedimensional structures of N-BGRPs from P. interpunctella, B. mori, and D. melanogaster,

Asp45 in *P. interpunctella* N- $\beta$ GRP is replaced by a Glu in the other two species (Figure 1A).

 $\beta$ -1,3-glucan-binding activities of D45A and D45K mutants were compared with that of the wild-type protein by curdlan pull-down assay and ITC with laminarin (Supplementary Figure 2). Neither of these mutations has any effect on carbohydrate-binding activity. Thus, it is concluded that the electrostatic attractions involving Asp45 (Figure 7C), as deduced from the crystal structure of N- $\beta$ GRP:laminarihexaose (12), contribute to the self-association of N- $\beta$ GRP:laminarin complex.

Functional significance of the protein:carbohydrate macro complex formation was characterized by measuring proPO activation in insect blood plasma by the wild-type and mutant N- $\beta$ GRPs in the absence and presence of laminarin (Figure 8B & C). The D45K mutation decreased proPO activation both in the absence of, and, to a smaller extent, in the presence of, laminarin, while the D45A mutation showed no change, consistent with the AUC data (Figure 8A).

Formation of a macro structure of  $\beta$ GRP: $\beta$ -1,3-glucan complex, a likely cooperative process, might present a repeating pattern and thus provide an effective platform to initiate innate immune responses (36,37). Such an arrangement is reminiscent of peptidoglycan recognition by insect peptidoglycan recognition proteins (PGRPs) (38). It is also of interest to note that some lectins which have a low binding affinity for monosaccharides increase their affinities significantly for oligosaccharides by forming a cluster of lectinoligosaccharide complexes (39).

The present study thus presents a three-dimensional solution structure of *P. interpunctella* N- $\beta$ GRP that is highly similar to its crystal structure (12), provides supporting evidence for laminarihexaose-binding site, and presents data for a novel mechanism of proPO activation in insect immune response that involves self-association of the initial protein:carbohydrate complex, and finally demonstrates the biological relevance of this important step.

### CONCLUSION

Biophysical characterization of interactions between laminarin, a  $\beta$ -1,3-glucan, and Nterminal domains of insect  $\beta$ GRPs in solution has led to the novel finding that a stable macro complex results from self-association of the initially formed N- $\beta$ GRP:laminarin complex. Electrostatic interactions between bound protein molecules in the macro complex contribute to its stability and ability to influence the rate of activation of the prophenoloxidase pathway. An increase in  $\beta$ -1,6 branching of laminarin reduces the carbohydrate's  $\beta$ GRP-binding affinity. Macro protein:carbohydrate complexes appear to provide an efficient means for recruiting immune response proteins and thus amplifying the initial response.

#### Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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## ABBREVIATIONS

βGRP	$\beta$ -1,3-glucan recognition protein
GNBP	Gram-negative bacteria binding protein
NMR	nuclear magnetic resonance
ррт	parts per million
ITC	isothermal titration calorimetry
AUC	analytical ultracentrifugation
SDS-PAGE	sodium dodecyl sulfate polyacrylamide gel electrophoresis
pproPO	prophenoloxidase

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#### Figure 1.

(A) Superimposition of ribbon structures of N- $\beta$ GRP from several insects: *P. interpunctella* purple, 2KHA (NMR, current work); *B. mori* - gray, 2RQE [NMR; (11)]; *D. melanogaster* green, 3IE4 [X-ray; (13)]; *B. mori* (laminarihexaose-bound) – orange, 3AQX [X-ray; (12)]; *P. interpunctella* - blue, 3AQY [X-ray; (12)]; *P. interpunctella* (laminarihexaose-bound) – red, 3AQZ [X-ray; (12)]; The NMR structure of *P. interpunctella* N- $\beta$ GRP reported here differs from that of the *B. mori* protein (11) in the configuration of a Pro, as indicated. (B) Ensemble of twenty lowest-energy structures of *P. interpunctella* N- $\beta$ GRP (2KHA). (C) A plot of backbone heteronuclear {<sup>15</sup>N}-<sup>1</sup>H NOE values *vs.* residue number for *P. interpunctella* N- $\beta$ GRP. Pro residues do not give rise to {<sup>15</sup>N}-<sup>1</sup>H NOEs. A couple of Cterminal residues had hetero NOE peak intensities barely above the noise level. Each  $\beta$ strand is represented by an arrow.



#### Figure 2.

Mapping of ligand-binding site on *P. interpunctella* N- $\beta$ GRP by NMR titration of <sup>1</sup>H-<sup>15</sup> laminarihexaose at 25°C, pH 6.5: (A) N HSQC spectra of N- $\beta$ GRP (0.5 mM) in the absence (*black*) or presence (*red*) of laminarihexaose (3 mM); (B) Chemical shift changes undergone by the backbone <sup>15</sup>N-<sup>1</sup>H groups; the weighted average of chemical shift changes of an <sup>15</sup>N-<sup>1</sup>H group was calculated by using the formula, [( $\Delta$ H2+( $\Delta$ N/5)2)/2]1/2, where  $\Delta$ H and  $\Delta$ N represent <sup>1</sup>H and <sup>15</sup>N chemical shift change, respectively; (C) Residues undergoing a chemical shift perturbation of > 0.02 ppm upon laminarihexaose titration are shown in red.





**Figure 3.** <sup>1</sup>H-<sup>15</sup>N HSQC spectra of N- $\beta$ GRP (0.5 mM) in the absence (*left*) or presence (*right*) of laminarin (1 mM) at 25°C, pH 6.5.



#### Figure 4.

Effect of addition of varying concentrations of laminarin on N- $\beta$ GRP, as monitored by sedimentation velocity experiments: Sedimentation profiles of N- $\beta$ GRP (26.2  $\mu$ M) in the absence (A) or presence of different levels of laminarin (B–H). Sedimentation at 49,000 rpm and 20 °C was monitored by measuring absorbance at 280 nm.



#### Figure 5.

Effect of dilution on N- $\beta$ GRP:laminarin complex as monitored by sedimentation velocity profiles: (A) 34  $\mu$ M N- $\beta$ GRP and 780  $\mu$ M laminarin at 280 nm; (B) 3.4  $\mu$ M N- $\beta$ GRP and 78  $\mu$ M laminarin at 220 nm; (C) 0.68  $\mu$ M N- $\beta$ GRP and 15.6  $\mu$ M laminarin at 205 nm. The concentration ratio of N- $\beta$ GRP:laminarin was maintained at 1:22.9. Sedimentation was carried out at 49,000 rpm and 20°C.



#### Figure 6.

Sedimentation velocity profiles for N- $\beta$ GRP, laminarin and their mixtures, as monitored at 49,000 rpm and 20°C by absorbance at 280 nm (A–C) and interference optics (D–F) at 5 min. intervals: 75  $\mu$ M N- $\beta$ GRP (A & D) and 125  $\mu$ M laminarin (B & E) were sedimented separately or together (C & F). Fringe displacements in D–F are given in arbitrary units.



## Figure 7.

A schematic representation of formation of N- $\beta$ GRP:  $\beta$ -1,3-glucan macro complex: (A) Laminarin binding to N- $\beta$ GRP and self-association of N- $\beta$ GRP:laminarin complex; (B) & (C) N- $\beta$ GRP packing and concomitant electrostatic interactions as can be observed in the crystal structure of N- $\beta$ GRP:laminarihexaose (12). Negative and positive potential surfaces are shown in red and blue, respectively.



#### Figure 8.

Electrostatic interactions play a role in the formation of N- $\beta$ GRP:  $\beta$ -1,3-glucan macro complex: (A) Sedimentation velocity profiles for N- $\beta$ GRP (wide type, black; D45A, blue; and D45K, red) in the presence of laminarin; Activation of the prophenoloxidase pathway by N- $\beta$ GRP and mutants without (B) or with laminarin (C). Samples of plasma (5  $\mu$ l) were mixed with protein alone or with protein and laminarin. After incubation at room temperature for 15 min, phenoloxidase activity was measured using dopamine hydrochloride as a substrate, as described under "Experimental Procedures". The bars represent the means  $\pm$  S.E. of data from three sets of measurements on a pooled plasma sample. Bars labeled with different letters (*a*, *b*, and *c*) are significantly different [analysis of variance (ANOVA), p < 0.05].