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Gloverins of the silkworm *Bombyx mori*: Structural and binding properties and activities

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Abstract

Gloverins are basic, glycine-rich and heat-stable antibacterial proteins (~14-kDa) in lepidopteran insects with activity against *Escherichia coli*, Gram-positive bacteria, fungi and a virus.

Hyalophora gloveri gloverin adopts a random coil structure in aqueous solution but has α -helical structure in membrane-like environment, and it may interact with the lipid A moiety of lipopolysaccharide (LPS). *Manduca sexta* gloverin binds to the O-specific antigen and outer core carbohydrate of LPS. In the silkworm *Bombyx mori*, there are four gloverins with slightly acidic to neutral isoelectric points. In this study, we investigate structural and binding properties and activities of *B. mori* gloverins (*BmGlv*s), as well as correlations between structure, binding property and activity. Recombinant *BmGlv*1–4 were expressed in bacteria and purified. Circular dichroism (CD) spectra showed that all four *BmGlv*s mainly adopted random coil structure (>50%) in aqueous solution in regardless of pH, but contained α -helical structure in the presence of 1,1,1,3,3,3-hexafluoro-2-propanol (HFIP), smooth and rough mutants (Ra, Rc and Re) of LPS and lipid A. Plate ELISA assay showed that *BmGlv*s at pH 5.0 bound to rough mutants of LPS and lipid A but not to smooth LPS. Antibacterial activity assay showed that positively charged *BmGlv*s (at pH 5.0) were active against *E. coli* mutant strains containing rough LPS but inactive against *E. coli* with smooth LPS. Our results suggest that binding to rough LPS is the prerequisite for the activity of *BmGlv*s against *E. coli*.

Keywords

Gloverin; Lipopolysaccharide; Circular dichroism; Antibacterial; Random coil; α -helix; *Bombyx mori*

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Appendix A. Supplementary data

Supplementary data related to this article can be found at <http://dx.doi.org/10.1016/j.ibmb.2013.03.013>.

1. Introduction

Insects have the largest numbers and species on earth, and they combat a variety of pathogens mainly relying on sophisticated innate immune system. Insect defense system consists of three major parts: structural barriers, cellular and humoral immune responses (Lemaitre and Hoffmann, 2007). Structural barriers, the first protective lines, refer to cuticle, midgut epithelium and trachea. Cellular immune responses, including phagocytosis, nodulation and encapsulation, are mediated by several types of hemocytes (Lavine and Strand, 2002; Sideri et al., 2007). Humoral immune responses include melanization of hemolymph and secretion of antimicrobial peptides (AMPs) (Hoffmann, 1995). AMPs, the major and best known immune effectors induced by infection, are synthesized by fat body, hemocytes and other tissues, and regulated by the Toll and immune deficiency (IMD) pathways (Bulet et al., 1999; Lemaitre and Hoffmann, 2007).

At least 150 insect AMPs have been purified or identified. Most insect AMPs are small and cationic, and they show activities against bacteria and/or fungi (Hoffmann, 1995; Bulet and Stocklin, 2005). Based on the sequences, structures and activities, insect AMPs can be classified into four families, the α -helical peptides (e.g., cecropin, moricin and sarcotoxin), cysteine-rich peptides (e.g., insect defensin, drosomycin and heliomicin), proline-rich peptides (e.g., apidaecin, drosocin and lebecin), and glycine-rich peptides (e.g., attacin and gloverin) (Laszlo-Otvos, 2000; Bulet and Stocklin, 2005). Glycine-rich peptides with molecular masses of more than 10 kDa have become a large family of insect AMPs, including attacin, sarcotoxin II, gloverin, hymenoptaecin, coleopteracin, hemiptericin and tenecin 4 (Hultmark et al., 1983; Ando et al., 1987; Axen et al., 1997; Casteels et al., 1993; Bulet et al., 1991; Cociancich et al., 1994; Chae et al., 2012).

So far, small cationic AMPs (~4 kDa) have been the focus of structural and activity study. Cysteine-rich peptides, which are cyclic by formation of disulfide bonds, can form stable structures in aqueous solution. However, linear AMPs do not form stable structures in aqueous solution but can change to stable structures in membrane mimic environment (Nguyen et al., 2008; Haney and Vogel, 2009). Cysteine-rich AMPs, such as insect defensin A, drosomycin, termicin and heliomicin, contain "cysteine stabilized $\alpha\beta$ motif" (CS $\alpha\beta$) structure with antiparallel β -sheet connected to a single α -helix by two disulfide bridges (Cornet et al., 1995; Landon et al., 1997; Silva et al., 2003; Lamberty et al., 2001). But nearly all the structures of α -helical linear AMPs are obtained by using micelle suspensions or in the presence of organic solvents. For example, *Hyalophora cecropia* cecropin A exists as random coil in aqueous solution but forms an amphipathic helical structure in 1,1,1,3,3,3-hexafluoro-2-propanol (HFIP)/water solution (Holak et al., 1988). The α -helical structure of moricin is also obtained in methanol and in solution containing 2,2,2-trifluoroethanol (TFE) or sodium dodecylsulphate (SDS) (Dai et al., 2008).

Since the first purification and characterization of gloverin from *Hyalophora gloveri* pupal hemolymph (Axen et al., 1997), a gloverin has been isolated from *Helicoverpa armigera* (Mackintosh et al., 1998), and two gloverins have been detected in hemolymph of septic injured *Diatraea saccharialis* larvae (Silva et al., 2010). Gloverin genes have also been identified in *Antheraea mylitta* (Gandhe-Archana et al., 2006), *Galleria mellonella* (Seitz et al., 2003; Brown et al., 2009), *Manduca sexta* (Abdel-Latif and Hilker, 2008; Xu et al., 2012; Zhu et al., 2003), *Plutella xylostella* (Eum et al., 2007; Etebari et al., 2011), *Spodoptera exigua* (Hwang and Kim, 2011), and *Trichoplusia ni* (Lundstrom et al., 2002). In the silkworm *Bombyx mori*, four gloverin genes (*BmgIv1-4*) have been identified, and *BmgIv2-4* genes are derived from duplication of *BmgIv1* (Cheng et al., 2006; Kaneko et al., 2007; Kawaoka et al., 2008; Mrinal and Nagaraju, 2008). Among the gloverins with known activities, *H. gloveri* gloverin is active against *Escherichia coli* D21f2 and D21 mutant

strains with rough LPS (Axen et al., 1997), *H. armigera* gloverin is active against *E. coli* strains with smooth LPS and D22 strain that is defective in lipid A (Mackintosh et al., 1998), *T. ni* gloverin 1 and 2 have activity against *E. coli* D21f2 and D22 strains and a virus (Lundstrom et al., 2002; Moreno-Habel et al., 2012). However, *S. exigua* gloverin is active against a Gram-positive bacterium (*Flavobacterium* sp.) but inactive against *E. coli* strain with smooth LPS (Hwang and Kim, 2011), *M. sexta* gloverin shows activity against a Gram-positive *Bacillus cereus* and two fungi (*Saccharomyces cerevisiae* and *Cryptococcus neoformans*) but inactive against *E. coli* strain with smooth LPS (Xu et al., 2012).

The majority of gloverins is basic or highly basic (pI ~8.3 for *H. gloveri* gloverin, *T. ni gloverin* 1 and *S. exigua* gloverin, pI > 9.0 for most other gloverins) and heat-stable with high content (>18%) of glycine residues (Xu et al., 2012). *H. gloveri* gloverin (*HgGlv*, pI ~8.3) can inhibit the growth of *E. coli* by inhibiting synthesis of bacterial outer membrane proteins and increasing permeability of the membrane (Axen et al., 1997). Basic gloverins may interact with lipopolysaccharide (LPS) via charge–charge interaction with negatively charged lipid A (Axen et al., 1997). But direct binding of gloverin to microbial components including LPS has only been reported for *M. sexta* gloverin (*MsGlv*, pI ~9.3) (Xu et al., 2012). Recombinant *MsGlv* can bind to the O-specific antigen and outer core carbohydrate moieties of LPS, Gram-positive lipoteichoic acid (LTA) and peptidoglycan (PG), and laminarin, but does not bind to lipid A (Xu et al., 2012). Known gloverins with acidic or neutral pI include *Heliothis virescens* gloverin (pI ~7.2) (Genbank accession number: ACR78446), *A. mylitta* gloverin 2 (pI ~6.8) (Genbank accession number: ABG72700), and four *B. mori* gloverins (*BmGlv*s) (pI ~ 5.5, 7.0, 6.3 and 7.0 for *BmGlv*1–4, respectively) (Kawaoka et al., 2008). Recombinant *BmGlv*s show activity against *E. coli* strains with smooth LPS (Kawaoka et al., 2008; Mrinal and Nagaraju, 2008). However, it is not clear whether *BmGlv*s can interact with LPS and whether they also adopt random coil structures in aqueous solution and undergo conformational transitions in the hydrophobic environment. In this study, we investigate structural transitions of *BmGlv*s in the hydrophobic environment (organic solvent, detergent micelles and LPS), binding properties of *BmGlv*s to LPS and other microbial cell wall components, and antibacterial activities of *BmGlv*s against *E. coli* strains with smooth and rough mutant forms of LPS.

2. Materials and methods

2.1. Microorganisms and microbial components

E. coli DH5 α (TIANGEN, China) and *E. coli* RosettaTM (DE3) (Transgen, China) strains were used to clone and express recombinant *BmGlv*s. *E. coli* (ATCC 25922), *Pichia pastoris*, *Serratia marcescens* and *Bacillus thuringiensis* were from American Type Culture Collection (ATCC). *Staphylococcus aureus* and *B. cereus* were kindly provided by Professor Brian Geisbrecht, *S. cerevisiae* (BY4741) and *C. neoformans* (alpha) were provided by Professor Alexander Idnurm, *Bacillus subtilis* was provided by Professor Michael O'Connor, School of Biological Sciences at University of Missouri – Kansas City. *E. coli* D21, D21e7, D21f1 and D21f2 strains with rough mutants of LPS were purchased from *E. coli* Genetic Resources at Yale CGSC, The coli Genetic stock center (USA).

Smooth LPS from *Salmonella enterica*, *S. marcescens*, *E. coli* 055:B5, *E. coli* 026:B6 and *E. coli* 0111:B4, rough mutants of LPS from *E. coli* EH100 (Ra mutant), *E. coli* J5 (Rc mutant), *E. coli* F583 (Rd mutant) and *S. enterica* serotype minnesota Re 595 (Re mutant), as well as lipid A monophosphoryl from *E. coli* F583 (Rd mutant), laminarin, mannan, and zymosan were from Sigma–Aldrich (MO, USA) and used for binding assay. TLRgrade LPS and PG from *E. coli* K12 (LPS-K12 and PG-K12), TLRgrade peptidoglycan (PG) and lipoteichoic acid (LTA) from *B. subtilis* (LTA-BS and PG-BS) and *S. aureus* (LTA-SA and PG-SA) were from Invivogen (CA, USA) and also used for binding assay. LPS from *E. coli*

serotype 055:B5 (smooth LPS), *E. coli* serotype EH100 (Ra), *E. coli* serotype J5 (Rc) and *E. coli* serotype R515 (Re), and monophosphoryl lipid A from *E. coli* serotype R515 were purchased from Alexis (Alexis, Switzerland) and used for circular dichroism (CD) experiments.

2.2. Construction of expression vectors for recombinant *B. mori* gloverins

Total RNAs were isolated from the fat body of day-3 fifth instar *B. mori* larvae collected at 24 h after injection of *E. coli* (ATCC 25922) (1×10^5 cells per larva) using TRIzol Reagent (Invitrogen), and the first strand cDNA was synthesized using M-MLV Reverse Transcriptase (TOYOBO, Japan). RT-PCR was performed to obtain cDNA sequences encoding *B. mori* mature gloverins using the following primer pairs: 5'-CAT GCC ATG GAT ATT CAC GAC TTT GTC AC-3' and 5'-CGC CTC GAG CCA CTC GTG AGT AAT CTG-3' (for mature *BmGlv1*, residues 44–178), 5'-CAT GCC ATG GAC GTC ACT TGG GAC AAA CAA-3' and 5'-CAG CTC GAG CCA ATC ATG GCG GAT CTC TG-3' (for mature *BmGlv2*, residues 43–173), 5'-CGA TCC ATG GAC GTC ACG TGG GAC ACG-3' and 5'-CCG CTC GAG CCA CTC ATG CCG GAT CTC-3' (for mature *BmGlv3*, residues 43–173), 5'-CAT GCC ATG GAC GTC ACC TGG GAC AAA CAA G-3' and 5'-CCG CTC GAG CCA ATC ATG GCG GAA CTC T-3' (for mature *BmGlv4*, residues 41–171). PCR products were purified using the EZNA cycle-pure kit (Omega, USA) and digested with *Nco*I and *Xho*I enzymes. After purification by agarose gel electrophoresis, these DNA fragments were ligated into the *Nco*I/*Xho*I-digested expression vector pET-21d (+) (Novagen), and the ligated products were then transformed into competent *E. coli* DH5 α cells. The recombinant expression vectors containing target genes from positive bacterial colonies were extracted and confirmed by restriction enzyme digestion and DNA sequencing.

2.3. Expression and purification of recombinant *B. mori* gloverins and production of polyclonal antibody against *BmGlv2*

To express soluble recombinant *BmGlv*s, recombinant plasmids were transformed into *E. coli* Rosetta™ (DE3) cells. Positive bacterial colonies were inoculated into LB medium containing ampicillin (100 μ g/ml), chloramphenicol (34 μ g/ml) and incubated overnight at 37 °C. The overnight cultures were diluted 1:100 into fresh LB medium containing ampicillin (100 μ g/ml), chloramphenicol (34 μ g/ml) and incubated at 37 °C to OD₆₀₀ = 0.6–0.8, then isopropyl-D-thiogalactoside (IPTG) (1 mM final concentration) was added to induce protein expression. After incubation for another 4 h at 37 °C, bacterial cells were harvested by centrifugation at 8,000g for 10 min at 4 °C.

The bacterial pellets were re-suspended in 10 mM phosphate, 200 mM NaCl, pH 8.0 (10 ml/g) and sonicated for 3 min. After centrifugation at 16,000g for 30 min at 4 °C, the supernatant was collected and applied to Ni²⁺-NTA column (Amersham Bio-sciences). The column was washed sequentially with 10, 25 and 50 mM imidazole in 10 mM phosphate, 200 mM NaCl, pH 8.0. Recombinant proteins were eluted with 500 mM imidazole in 10 mM phosphate, 200 mM NaCl, pH 8.0, and then loaded to a Superdex 75 column (16/60 mm) pre-equilibrated with 10 mM phosphate, 200 mM NaCl, pH 8.0. Fractions containing recombinant *BmGlv*s were pooled, desalted, and concentrated to 1 mg/ml in 10 mM phosphate (pH 5.0 or 8.0) or 10 mM phosphate, 100 mM NaCl (pH 5.0 or 8.0). Protein concentrations were determined by absorbance at 280 nm with the theoretical molar extinction coefficients (<http://www.expasy.org>).

To produce polyclonal antibody against *BmGlv2* in a rabbit, purified recombinant *BmGlv2* (600 μ g) was applied to a preparative SDS-PAGE, and the gel slice containing recombinant

BmGlv2 was cut out and used as an antigen to inject a rabbit for polyclonal antibody production (Cocalico Biologicals, Inc., Reamstown, PA, USA).

2.4. SDS-PAGE and western blot analyses

Day 3 fifth instar *B. mori* Dazao larvae were injected with saline, *E. coli* (ATCC 25922) (1×10^5 cells/larva), *S. aureus* (ATCC 27217) (1×10^5 cells/larva), or yeast (*Pichia pastoris*) (1×10^5 cells/larva), or without treatment (naïve) and hemolymph was collected at 24 h post-injection. Hemolymph from at least four larvae of each group was combined, hemocytes were removed by centrifugation and cell-free hemolymph samples were used for Western blot analysis. Recombinant *BmGlv1–4* purified from *E. coli* (1 μ g each for SDS-PAGE analysis, 0.1 μ g each for Western blot) and cell-free hemolymph samples (2 μ l each) were separated on 15% SDS-PAGE and proteins were stained with Coomassie Brilliant Blue or transferred to nitrocellulose membranes (162–0097, Bio-Rad). The membrane was blocked with 5% BSA in Tris-buffered saline (100 mM Tris-HCl, pH 7.6, 150 mM NaCl) containing 0.1% Tween-20 (TBS-T) at room temperature for at least 3 h and then incubated overnight with rabbit anti-*BmGlv2* antibody (1:2000) at 4 °C in TBS-T containing 5% BSA with gentle rocking. Then, the membrane was washed four times with TBS-T and incubated with goat anti-rabbit IgG conjugated to alkaline phosphatase (1:10,000) in TBS-T containing 5% BSA for 2 h at room temperature. After washing four times with TBS-T (10 min each time), the signal was developed by using alkaline phosphatase (AP) conjugate color development Kit (170–6432, Bio-Rad).

2.5. CD spectroscopy

CD experiments were performed on a Jasco-810 spectropolarimeter (Jasco, Tokyo, Japan) at 25 °C using a quartz cell with a path length of 0.1 cm. Spectra were recorded over a wavelength of 190–260 nm. Each spectrum was obtained after subtracting the signal from protein-free solution. Proteins were dissolved to 0.15 mg/ml in 10 mM phosphate, pH 5.0 or 8.0, in the presence or absence of 40% HFIP, 10 mM SDS, 10 mM dodecylphosphocholine (DPC), smooth LPS, Ra-LPS, Rc-LPS, Re-LPS, or lipid A (w/w = 1/1), or in the presence of increasing concentrations of HFIP (10–40%), SDS (0.5–100 mM) or DPC (1–20 mM). Percentages of secondary structures were estimated using the Jasco protein secondary structure estimation program by the method of Yang et al. (1986).

2.6. Binding of *B. mori* gloverins to LPS

To test binding of *BmGlv*s to LPS and other microbial components, plate ELISA assays were performed. Briefly, wells of a flat bottom 96-well plate (Polysorp, Nunc) were coated with different forms of LPS, LTA, PG, laminarin, mannan, zymosan, or lipid A (2 μ g/well) as described previously (Yu and Kanost, 2000; Yu et al., 2005). The plates were placed overnight at room temperature until the water evaporated completely, heated to 60 °C for 30 min, and then blocked with 1 mg/ml BSA in Tris buffer (TB) (50 mM Tris-HCl, 50 mM NaCl, pH 8.0) for 2 h at 37 °C. Then, plates were rinsed four times with TB, purified *BmGlv1*, *BmGlv2*, *BmGlv3*, *BmGlv4*, or CP36 (a recombinant cuticle protein from *M. sexta* as a control protein) was diluted to 1 μ g/ml in 10 mM phosphate, 100 mM NaCl, pH 5.0 or 8.0, containing 0.1 mg/ml BSA and added to the coated plates (50 μ l/well). Binding was allowed to occur for 3 h at room temperature, and the plates were rinsed four times with TB. Then monoclonal anti-polyhistidine antibody (Sigma-Aldrich, USA) (1:2000 in TB containing 0.1 mg/ml BSA) was added (100 μ l/well) and incubated overnight at 4 °C. The plates were rinsed four times with TB, and alkaline phosphatase-conjugated goat anti-mouse IgG (Sigma-Aldrich, USA) (1:3000 in TB containing 0.1 mg/ml BSA) was added (100 μ l/well) and incubated for 2 h at 37 °C. The plates were rinsed, p-nitro-phenyl phosphate (1 mg/ml in 10 mM diethanolamine, 0.5 mM MgCl₂) was added (50 μ l/well), and absorbance at 405 nm of each well was determined every minute for 30 min period using a microtiter

plate reader (Bio-Tek Instrument, Inc.). Specific binding of BmGlv3 to each microbial component was obtained by subtracting the total binding of the control CP36 protein from the total binding of each BmGlv. These experiments were repeated at least three times. Figures were made with the GraphPad Prism software (GraphPad, CA, USA) with one representative set of data. Significance of difference was determined by an unpaired t-test or by one way ANOVA followed by a Tukey's multiple comparison test using the same software (GraphPad, CA, USA).

2.7. Antimicrobial activity assays

Antimicrobial activity of purified BmGlv3 was tested against two Gram-negative and four Gram-positive bacteria (*E. coli* DH5a, *S. marcescens*, *S. aureus*, *B. subtilis*, *B. cereus*, and *B. thuringiensis*), two fungal strains (*S. cerevisiae* (BY4741) and *C. neoformans* (alpha)), and four isogenic *E. coli* K-12 strains with different rough mutants of LPS: *E. coli* D21 (with Ra-LPS), *E. coli* D21e7 (with Rc-LPS), *E. coli* D21f1 (with Rd-LPS), and *E. coli* D21f2 (with Re-LPS). The activities were determined using bacterial clearance assays as described by Mrinal and Nagaraju (2008) with slight modifications. Briefly, overnight bacterial or fungal cultures were subcultured in LB or YPD medium (1% yeast extract, 2% peptone and 2% dextrose) until mid-log phase. The bacterial and fungal cultures were centrifuged at 1000 g for 10 min at 4 °C and washed once with 10 mM phosphate, 100 mM NaCl, pH 5.0 or 8.0. The bacterial and fungal cells were adjusted to OD₆₀₀ = 0.4 and 0.2, respectively, in 10 mM phosphate, 100 mM NaCl, pH 5.0 or 8.0. Then the prepared cell cultures (85 µl each) were mixed with purified BmGlv3 (15 µl of 1 mg/ml in 10 mM phosphate, 100 mM NaCl, pH 5.0 or 8.0) (final concentrations of BmGlv3 were 150 µg/ml or ~ 10 µM) or buffer alone (10 mM phosphate, 100 mM NaCl, pH 5.0 or 8.0) as controls in 96-well plates. Bacterial cells were incubated at 37 °C with 220 rpm shaking and fungal cells were cultured at 30 °C with 220 rpm shaking. OD₆₀₀ was measured every hour by Powerwave XS plate reader (BioTek, VT, US). The activity assay was also performed with increasing concentrations of BmGlv3 (0.4, 2 and 10 µM) against *E. coli* D21 and D21f2 strains. In addition, antibacterial activity of BmGlv3 against *E. coli* D21 and D21f2 strains was verified by bacterial viability assay. At 2 h after incubation with BmGlv3 (10 µM each), aliquots of *E. coli* D21 and *E. coli* D21f2 cells were serially diluted with 10 mM phosphate, 100 mM NaCl, pH 5.0 and plated on LB-agar plates containing streptomycin sulfate (50 µg/ml) and ampicillin (10 µg/ml). The plates were incubated at 37 °C overnight and the numbers of viable bacterial cells were recorded. Microbial growth curves were generated using the Graphpad Prism version 4.0 for Windows (GraphPad Software, CA, USA).

3. Results

3.1. Expression and purification of recombinant *B. mori* gloverins

Based on the amino acid sequences, all reported gloverins have three forms: prepro-, pro-, and mature gloverins. The theoretical molecular masses and isoelectric points (pI) of mature BmGlv1–4 are 14.4 kDa and 5.49, 14.1 kDa and 7.03, 14.1 kDa and 6.32, and 14.1 kDa and 6.94, respectively (Kawaoka et al., 2008). In order to obtain gloverins for structural and functional studies, mature BmGlv1 (residues 44–178), BmGlv2 (residues 43–173), BmGlv3 (residues 43–173), and BmGlv4 (residues 41–171) with a C-terminal six-histidine tag were expressed as soluble proteins in *E. coli* Rosetta™ (DE3) cells. The recombinant proteins were purified by Ni-NTA affinity column followed by a Superdex 75 column using phosphate buffer (pH 5.0 or 8.0) (Fig. S1A), and all four BmGlv3 were recognized by rabbit polyclonal antibody against recombinant BmGlv2 (Fig. S1B). The calculated masses and pI of recombinant BmGlv1–4 are 15.6 kDa and 6.02, 15.3 kDa and 6.75, 15.3 kDa and 6.48, and 15.3 kDa and 6.7, respectively. Western blot analysis of cell-free hemolymph samples from *B. mori* larvae using anti-BmGlv2 antibody showed that

*BmGlv*s were induced by *E. coli* but not by injury (saline-injection), *S. aureus* or yeast (Fig. S1C). The estimated total concentration of *BmGlv*s in *E. coli*-induced hemolymph was about 0.1–0.12 mg/ml (7–9 μ M) (Fig. S1C).

3.2. CD spectroscopy and secondary structural prediction of *BmGlv*s in aqueous solution

CD spectroscopy is an important tool to determine secondary structures of proteins in solutions. We performed CD experiments for *BmGlv*1–4 in phosphate buffer from pH 3–8, and all four *BmGlv*s adopted random coil conformation in aqueous solution in regardless of pH values (Fig. S2), indicating that pH alone cannot induce conformational transitions. Fig. 1 shows CD spectra of *BmGlv*1–4 in phosphate buffer at pH 5.0 and 8.0, in which *BmGlv*1–4 are positively and negatively charged, respectively, based on their theoretical pI. The following CD experiments, ELISA binding assays and antibacterial activity assays were all performed at pH 5.0 and 8.0. CD spectra of *BmGlv*1–4 at both pH 5.0 and 8.0 displayed negative ellipticity around 200 nm (Fig. 1), which is typical for random coil conformation (Tiffany and Krimm, 1972; Yang et al., 1986), and all four *BmGlv*s contained over 50% random coil as well as certain contents of β -strands (25–40%) and turns (9–19%), but did not contain α -helix conformation (Table 1).

3.3. CD spectroscopy and secondary structural prediction of *BmGlv*s in membrane-like environment

Most linear antimicrobial peptides adopt random coil structures in aqueous solution, but change to more defined structures when encountering bacterial membrane. Organic solvents such as trifluoroethanol (TFE) and HFIP, and detergent micelles like SDS and DPC, are commonly used to mimic a membrane environment (Haney and Vogel, 2009). *HgGlv* undergoes conformational transition to α -helix in hydrophobic environment (HFIP solution) (Axen et al., 1997). To test whether *BmGlv*s also undergo conformational transitions in a membrane-like environment, CD experiments were performed for *BmGlv*1–4 in the presence of HFIP, SDS and DPC.

We first collected CD spectra of *BmGlv*s at pH 5.0 and 8.0 in the presence of increasing concentrations of HFIP (10–40%), SDS (0.5–100 mM) or DPC (1–20 mM). HFIP at 10% already caused conformational transitions of *BmGlv*s, but DPC and SDS even at high concentrations did not cause conformational transitions of *BmGlv*s (Fig. S3) with the exception of *BmGlv*4, which underwent conformational transitions at pH 5.0 even with 1 mM DPC (Fig. S3F). Fig. 2 shows the CD spectra of *BmGlv*s in 40% HFIP, 10 mM SDS and 10 mM DPC. CD spectra of *BmGlv*1–3 in phosphate buffer at both pH 5.0 and 8.0 containing 40% HFIP showed two minima at 208 and 222 nm (Fig. 2A–C, E–G) and CD spectra of *BmGlv*4 in the presence of 40% HFIP also showed a major negative peak around 222 nm (Fig. 2D and H), which are characteristics of α -helix (Holzwarth and Doty, 1965; Yang et al., 1986), indicating that *BmGlv*1–4 adopted α -helix conformation in the presence of HFIP in a pH-independent manner. The predicted secondary structures showed that *BmGlv*1–4 contained 57.6, 57.6, 32.7 and 59.5% α -helix at pH 5.0 (Table 2), and 44.9, 25.6, 26.5 and 46.6% α -helix at pH 8.0 (Table 3), respectively. These results suggest that more *BmGlv*s converted to α -helical structure at pH 5.0 when proteins were positively charged than at pH 8.0 when proteins were negatively charged in the presence of HFIP.

In the presence of 10 mM SDS, CD spectra of all four *BmGlv*s at both pH 5.0 and pH 8.0 did not display negative peaks typical for α -helix conformation (Fig. 2), and the predicted secondary structures showed that *BmGlv*1–4 at both pH 5.0 and 8.0 contained mainly random coils (38–45%) and β -strands (43–56%) with low contents of α -helix (2–18%) but no turns (Tables S1 and S2). In the presence of 10 mM DPC, CD spectra of *BmGlv*1–3 at both pH 5.0 and 8.0 and CD spectrum of *BmGlv*4 at pH 8.0 were similar to CD spectra of

BmGlv1–4 with SDS (Fig. 2A–C, E–H), but the spectrum of *BmGlv4* at pH 5.0 showed a distinct minimum around 220 nm (Fig. 2D). Predicted secondary structures showed that *BmGlv1–3* at pH 5.0 and 8.0 and *BmGlv4* at pH 8.0 in the presence of 10 mM DPC mainly contained random coils (38–44%) and β -strands (34–57%) with low contents of α -helix (2–16%) and turns (0–13%) (Tables S1 and S2). But *BmGlv4* at pH 5.0 (with positive net charge) contained 47% α -helical conformation and 17.5% random coil, indicating that most random coil in *BmGlv4* was converted to α -helix conformation in the presence of DPC similar to that in the presence of HFIP (Table S1).

3.4. Binding of BmGlvs to LPS

LPS is a major component of Gram-negative bacteria cell walls, and it is composed of three moieties: the O-specific antigen, the core (outer and inner core) carbohydrate, and the lipid A (Raetz, 1990; Yu and Kanost, 2002). HgGlv may interact with the lipid A moiety of LPS to exert its activity (Axen et al., 1997), and *MsGlv* can bind to the O-specific antigen and outer core carbohydrate moieties of LPS as well as Gram-positive LTA and PG, but does not bind to lipid A (Xu et al., 2012). To test binding of *BmGlvs* to different forms of LPS and other microbial components, plate ELISA assays were performed using purified recombinant *BmGlvs* in phosphate buffer at pH 5.0 and 8.0. Almost no or very little *BmGlv1–4* bound to *E. coli* LPS-K12 and PG-K12, *S. aureus* LTA-SA and PG-SA, *B. subtilis* LTA-BS and PG-BS, laminarin, mannan, or zymosan at both pH 5.0 and 8.0 (Fig. 3A – D), and no or very little *BmGlv1–4* bound to smooth LPS from several *E. coli* strains and Gram-negative bacteria (Fig. S4). Also, almost no or very little *BmGlv1–4* bound to Ra-, Rc-, Rd- and Re-LPS and lipid A at pH 8.0; however, all four *BmGlvs* bound to rough mutants of LPS and lipid A at pH 5.0, with more proteins bound to Rd-LPS and Re-LPS than to Ra-LPS, Rc-LPS and lipid A (Fig. 3E – H). Together, these results suggest that positively charged (at pH 5.0) *BmGlv1–4* may bind to the inner core carbohydrate of LPS and lipid A.

3.5. CD spectroscopy and secondary structural prediction of BmGlvs in the presence of LPS

To determine whether interaction between *BmGlvs* and LPS can induce conformational transitions to α -helix, CD experiments were performed for *BmGlv1–4* in the presence of smooth LPS, rough mutants (Ra, Rc and Re) of LPS and lipid A. In the presence of Ra-LPS at both pH 5.0 and 8.0, *BmGlv1–4* all underwent conformational transitions to α -helix (Fig. 4) with predicted secondary structures of mainly α -helix (55–67%) and turns (33–45%), but no random coil or β -strand (except *BmGlv1* at pH 8.0) (Tables 2 and 3). *BmGlv1* at pH 8.0 in the presence of Ra-LPS contained high contents of β -strands (52%) and random coil (33%) but low content of α -helix (14%) (Table 3).

In the presence of smooth LPS, Rc- and Re-LPS, the CD spectra of *BmGlv1–4* at pH 5.0 were very similar to those with Ra-LPS (Fig. 5), although *BmGlv1–4* bound to rough mutants of LPS but not smooth LPS (Fig. 3). Lipid A also changed the CD spectra of *BmGlv1–4* comparing to those in aqueous solutions (Fig. 5). The predicted secondary structures showed that in the presence of smooth LPS, rough LPS (Ra-, Rc- and Re-LPS) and lipid A at pH 5.0, *BmGlv1–4* contained predominantly α -helix (50–72%) and turns (28–50%), with no β -strand or random coil (except for *BmGlv2* and *BmGlv4* with smooth LPS) (Table 4), suggesting that LPS and lipid A can induce conformational transitions from random coil to α -helix in *BmGlv1–4*. The concentrations of LPS and lipid A used in the CD experiment were 0.15 mg/ml (~15 μ M for smooth LPS), which were higher than the critical micelle concentration (CMC) of LPS (CMC for smooth LPS from *E. coli* 0111:B4 is 1.3–1.6 μ M) (Yu et al., 2006). Together, these results suggest that hydrophobic property of LPS and lipid A is required for conformational transitions from random coil to α -helix in

BmGlv1–4. We also tried the CD experiments with LTA and PG, but could not obtain CD spectra due to poor solubility of LTA and PG and precipitation problem after mixing LTA and PG with *BmGlv*s proteins.

3.6. Activity of *BmGlv*s against *E. coli* with rough LPS

Gloverins from different lepidopteran species show activities against *E. coli*, Gram-positive bacteria, fungi or a virus (Axen et al., 1997; Mackintosh et al., 1998; Lundstrom et al., 2002; Hwang and Kim, 2011; Xu et al., 2012; Moreno-Habel et al., 2012). It is not clear whether the activity of gloverin against different microbes is related to pH (net charge), conformational transition to α -helix, and/or binding to microbial surface. We showed above that pH alone cannot cause conformational transitions to α -helix in *BmGlv*s (Fig. 1 and Fig. S2) but pH is important for binding of *BmGlv*s to rough LPS (Fig. 3), and hydrophobic environment (HFIP, smooth and rough LPS, or lipid A) is required for conformational transitions to α -helix in *BmGlv*s (Figs. 2, 4 and 5, Tables 2–4, S1 and S2). To determine correlations between the activity of *BmGlv*s and structural conformation/binding ability, antimicrobial activity of *BmGlv1–4* was tested against several Gram-negative and Gram-positive bacteria as well as fungi. *BmGlv1–4* were inactive against Gram-negative *E. coli* DH5 α and *S. marcescens* with smooth LPS at pH 5.0 and 8.0, or the fungi *S. cerevisiae* (at pH 5.0 and 8.0) and *C. neoformans* (at pH 8.0), but were active against *C. neoformans* at pH 5.0 (Fig. S5). *BmGlv1–4* were also inactive against Gram-positive *S. aureus*, *B. subtilis*, *B. cereus*, and *B. thuringiensis* at both pH 5.0 and 8.0 (Fig. S6). These results were consistent with the binding properties of *BmGlv1–4* as they did not bind to smooth LPS, LTA, PG, laminarin, mannan and zymosan (Fig. 3A – D and S4).

We then tested the activity of *BmGlv*s against *E. coli* mutant strains containing rough LPS since *BmGlv1–4* bound to rough LPS at pH 5.0 (Fig. 3E – H). *BmGlv1–4* (~10 μ M each) at pH 5.0 were active against *E. coli* D21 (with Ra-LPS), *E. coli* D21e7 (with Rc-LPS), *E. coli* D21f1 (with Rd-LPS) and *E. coli* D21f2 (with Re-LPS) mutant strains, but were inactive against these *E. coli* mutant strains at pH 8.0 (Fig. 6). The concentration of *BmGlv*s used in the bacteria clearance assay was 10 μ M for each protein, which was within the physiological concentration of total *BmGlv*s in *E. coli*-induced hemolymph (7–9 μ M) (Fig. S1C). Using lower concentrations of recombinant *BmGlv*s at pH 5.0, the activity of *BmGlv*s against *E. coli* D21 and *E. coli* D21f2 was observed at 2 μ M for *BmGlv1*, 2 and 4 and even at 0.4 μ M for *BmGlv1* and 2 (Fig. 7), indicating that the activity of *BmGlv*s against *E. coli* mutant strains is dose-dependent. We also confirmed the activity of *BmGlv*s using bacterial viability assay. Significantly lower numbers of viable *E. coli* D21 and *E. coli* D21f2 cells were observed when the bacterial cells were treated with *BmGlv*s (~10 μ M) at pH 5.0 for 2 h (Fig. 8). The number of viable *E. coli* D21f2 cells was decreased but not significantly after treating with *BmGlv3* compared to the control. In addition, the activity of *BmGlv3* was significantly lower than that of *BmGlv1*, 2 or 4 (Fig. 8). Since *BmGlv1–4* bound to rough mutants of LPS at pH 5.0 (Fig. 3E – H), together these results suggest that binding of *BmGlv*s to *E. coli* surface (via rough LPS) is required for the activity of *BmGlv*s against *E. coli*.

4. Discussion

Gloverin belongs to the glycine-rich AMP family and has been identified only in lepidopteran insects so far. Gloverins from most lepidopteran species are basic (pI ~8.3) or highly basic (pI > 9.0) (Xu et al., 2012), but *H. virescens* gloverin, *A. mylitta* gloverin 2, and four *B. mori* gloverins are slightly acidic to neutral (pI ~ 5.5–7.2). Even basic gloverins from different lepidopteran species show activities against different microbes, including *E. coli*, Gram-positive bacteria, fungi, and a virus (Axen et al., 1997; Mackintosh et al., 1998; Lundstrom et al., 2002; Hwang and Kim, 2011; Xu et al., 2012; Moreno-Habel et al., 2012).

B. mori gloverins are also active against *E. coli* (Kawaoka et al., 2008; Mrinal and Nagaraju, 2008). Thus, it is not clear whether the activity of gloverin is related to its pI or not. *HgGlv* (pI ~8.3) is active against *E. coli*, and it adopts random coil conformation in solution but has α -helical conformation in the presence of HFIP (Axen et al., 1997). *MsGlv* (pI ~9.3) can bind to LPS but is inactive against *E. coli* (Xu et al., 2012). Therefore, it is also not clear whether the activity of gloverin is related to its binding ability to microbial surface and/or conformational transition to α -helix.

Most linear AMPs are unfolded in aqueous solutions and they can change to defined structures only after interacting with bacterial membrane (Nguyen et al., 2008). Organic solvents such as TFE and HFIP, and detergent micelles like SDS and DPC, are commonly used to mimic bacterial membrane. SDS is used to mimic bacterial membrane as it contains a negative head group, while DPC is used to mimic eukaryotic membrane that does not contain net negative charge (Haney and Vogel, 2009). We showed by CD experiments that *BmGlv1-4* in aqueous solution adopted mainly random coil (>50%) conformation with certain contents of β -strands and turns but no α -helix in regardless of pH values (pH 3–8) (Fig. 1 and Fig. S2, Table 1), suggesting that pH alone cannot cause conformational transitions to α -helix in *BmGlv1-4*. *BmGlv*s adopted α -helical conformations in the presence of organic solvent HFIP but not detergent micelles SDS or DPC (Fig. 2), a result consistent with α -helical conformations of *HgGlv* and *H. cecropia* cecropin in solution containing HFIP (Steiner, 1982; Holak et al., 1988; Axen et al., 1997). In addition, conformations of *BmGlv*s in the presence of HFIP differ from those in the presence of SDS and DPC, while moricin adopts similar conformations in the presence of TFE and SDS (Dai et al., 2008).

Interestingly, in the presence of smooth LPS, rough LPS (Ra-, Rc- and Re-LPS) and lipid A, *BmGlv1-4* at both pH 5.0 and 8.0 (except *BmGlv1* with Ra-LPS at pH 8.0) underwent conformational transitions from random coil to α -helix (Figs. 4 and 5) with predominantly α -helix (50–72%) and turns (28–50%), but almost no random coil or β -strand (Tables 2–4). The exception is *BmGlv1*, which contained mainly β -strands (52%) and random coil (33%) with only 14% α -helix in the presence of Ra-LPS at pH 8.0 (Fig. 4E and Table 3). Thus, in the hydrophobic membrane-like environment (smooth LPS, rough LPS and lipid A, as well as HFIP), *BmGlv*s adopt mainly α -helix structure. *BmGlv4* may differ from *BmGlv1-3*, since in the presence of HFIP and DPC, *BmGlv4* at pH 5.0 (with positive net charge) contained low contents of random coil, but *BmGlv1-3* at pH 5.0 contained high contents of random coil (Table S1).

The mode of action of basic AMPs may involve electrostatic interaction between the positively charged AMPs and the negatively charged head groups of bacterial membranes. For example, attacin (~20 kDa), another typical AMP of the glycine-rich family, acts on the outer membrane of *E. coli* by electrostatic binding to lipid A and interaction with the acyl chain of lipid A and phospholipids in the outer membrane (Carlsson et al., 1998). Our ELISA assays showed that *BmGlv*s did not bind to smooth LPS, LTA, PG, laminarin, mannan and zymosan at both pH 5.0 and 8.0 (Fig. 3A – D and S4), but bound to rough mutants of LPS and lipid A at pH 5.0 but not pH 8.0, with more proteins bound to Rd-LPS and Re-LPS than to Ra-LPS, Rc-LPS and lipid A (Fig. 3E – H). These results suggest that positively net charge in *BmGlv*s may contribute to binding of *BmGlv*s to rough LPS and lipid A, and *BmGlv*s may bind to the inner core carbohydrate of LPS. CD experiments showed that in the presence of smooth LPS, Ra-, Rc- and Re-LPS and lipid A, random coils in *BmGlv*s were converted to α -helix (Figs. 4 and 5), while *BmGlv*s did not bind to smooth LPS (Fig. 3A – D and S4). These results suggest that binding to LPS/lipid A is not required but membrane-like environment is necessary for conformational transitions of *BmGlv*s from random coil to α -helix.

Activity assays showed that BmGlv3 were inactive against *E. coli* DH5 α and *S. marcescens* with smooth LPS, and inactive against two fungal strains and several Gram-positive bacteria, but were active against *C. neoformans* at pH 5.0 (Figs. S5 and S6). These results are consistent with the binding properties of BmGlv3 as they did not bind to smooth LPS, LTA, PG, laminarin, mannan or zymosan (Fig. 3A – D and S4). However, BmGlv3 at pH 5.0 were active against *E. coli* mutant strains with rough LPS (Ra-, Rc-, Rd- and Re-LPS) (Fig. 6A – D), a result consistent with HgGlv (Axen et al., 1997), but inactive at pH 8.0 against these *E. coli* mutant strains (Fig. 6E – H). The activity of recombinant BmGlv3 against *E. coli* mutant strains was dose-dependent (Fig. 7) and within physiological concentrations in hemolymph, as the high concentration of recombinant BmGlv3 used in the assay was ~10 μ M, which was closed to the estimated concentration (7–9 μ M) of total native BmGlv3 in the *E. coli*-induced hemolymph of *B. mori* larvae (Fig. S1C). Compared the four BmGlv3, BmGlv1, 2 and 4 had similar high activity against *E. coli*, while BmGlv3 had lower activity than BmGlv1, 2 and 3 (Fig. 8).

BmGlv3 bound to rough mutants of LPS at pH 5.0 but not at pH 8.0 (Fig. 3E – H), but rough LPS caused conformational transition of BmGlv3 to α -helix at both pH 5.0 and 8.0 (Figs. 4 and 5). These results suggest that binding of BmGlv3 to rough LPS on *E. coli* surface is required for the activity of BmGlv3. BmGlv3 at acidic pH were active against *E. coli*, suggesting that they may have higher activity against bacteria in the midgut. We believe that conformational transition from random coil to α -helix in BmGlv3 is the key for pore formation on bacterial membrane, but binding of BmGlv3 to bacterial surface via rough LPS is the prerequisite for the activity of BmGlv3. Thus, whether a gloverin is active against Gram-negative bacteria, Gram-positive bacteria, fungi or viruses may depend on its binding to microbial surface and conformational transition to α -helix. We also tried CD experiments for BmGlv3 in the presence of LTA, PG, laminarin, mannan or zymosan to test interactions between BmGlv3 and these microbial components and determine whether conformational transitions to α -helix occur, but failed to obtain CD spectra due to poor solubility of these microbial components. In order to better understand the mode of action of gloverins and the mechanisms of conformational transitions to α -helix, it is necessary to determine three-dimensional structure of gloverin, which will be one of our future goals.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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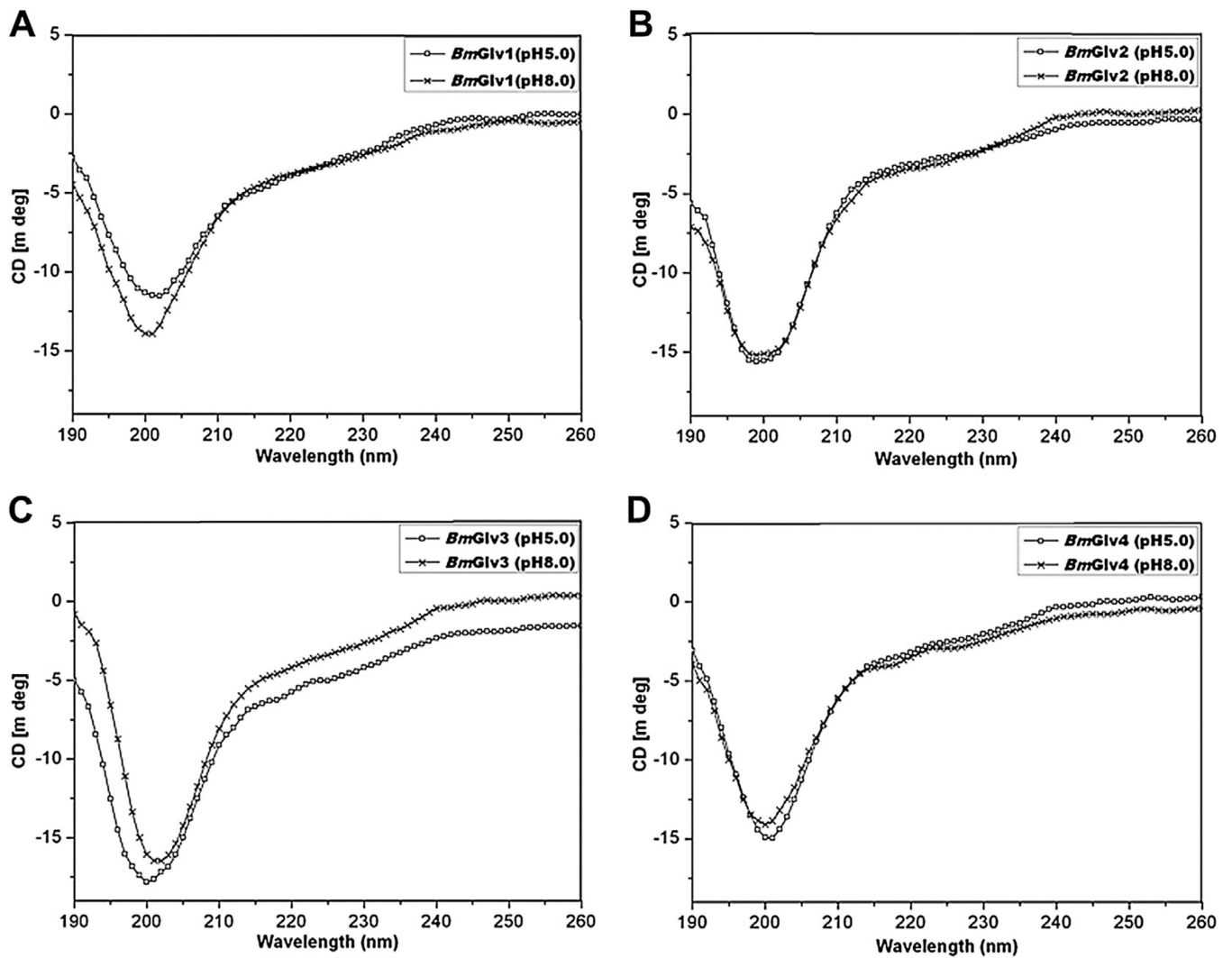


Fig. 1.

CD spectra of *BmGlv*s in aqueous solutions at pH 5.0 and 8.0. Purified *BmGlv1-4* were diluted to 0.15 mg/ml in 10 mM phosphate buffer (pH 5.0 or 8.0) and CD spectra were recorded on a Jasco-810 spectropolarimeter at 25 °C using a quartz cell with a path length of 0.1 cm. Each CD spectrum was obtained after subtracting the signal from protein-free solution.

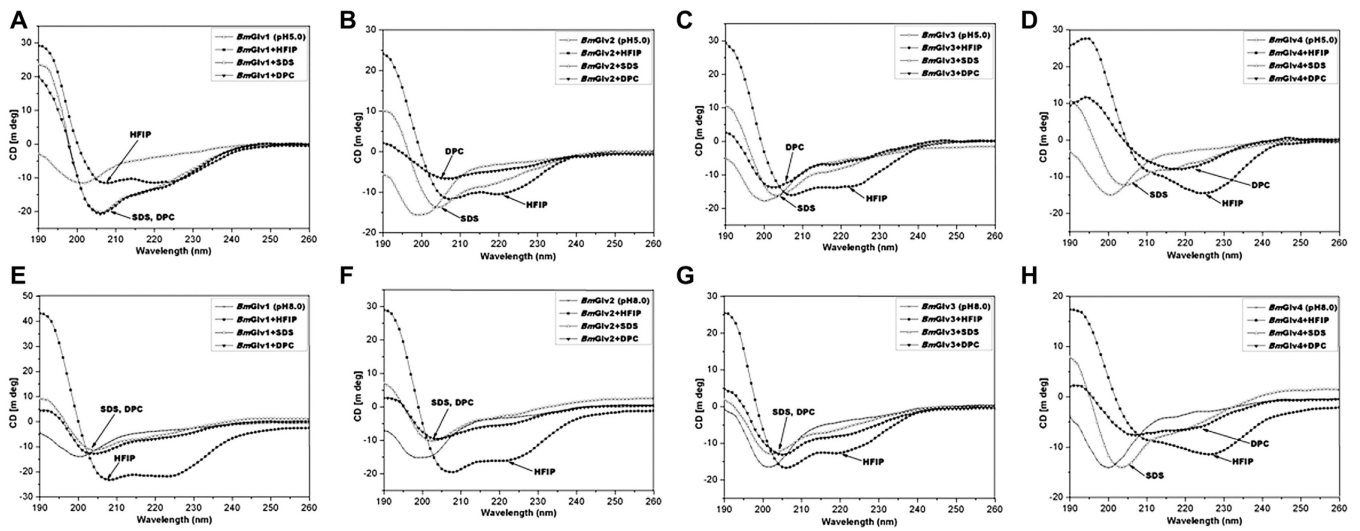


Fig. 2.

CD spectra of *BmGlv*s in the presence of HFIP, SDS and DPC at pH 5.0 and 8.0. Purified *BmGlv*1–4 were diluted to 0.15 mg/ml in 10 mM phosphate buffer (pH 5.0 or 8.0) in the presence or absence of 40% HFIP, 10 mM SDS or 10 mM DPC and CD spectra were recorded on a Jasco-810 spectropolarimeter at 25 °C. Each CD spectrum was obtained after subtracting the signal from protein-free solution.

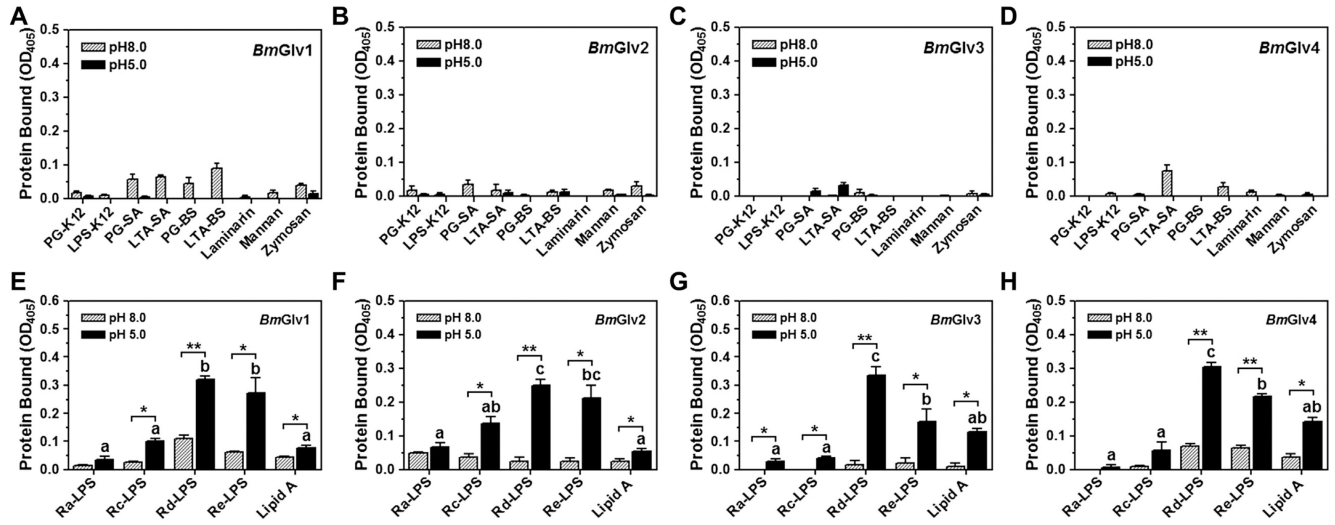


Fig. 3. Binding of *BmGlv*s to microbial cell wall components. Wells of 96-well plates were coated with different microbial components (2 µg/well) and blocked with BSA. Purified *BmGlv*s and the control CP36 (a recombinant cuticle protein from *M. sexta*) were diluted to 1 µg/ml in 10 mM phosphate, 100 mM NaCl, pH 5.0 or 8.0, and the diluted proteins were added to the coated plates. Binding of proteins to microbial components was determined by plate ELISA assays as described in the materials and methods. The figure showed specific binding of *BmGlv*s to each microbial component after subtracting the total binding of the control CP36 protein from the total binding of *BmGlv*s. Each bar represents the mean of at least three individual measurements ± SEM. Comparing binding of each *BmGlv* to rough LPS and lipid A at pH 5.0 (panels E–H), identical letters are not significant difference ($p > 0.05$), while different letters indicate significant difference (< 0.05) determined by one way ANOVA followed by a Tukey’s multiple comparison test. Comparing binding of *BmGlv* to each rough LPS or lipid A between pH 5.0 and 8.0 (panels E–H), ‘*’ ($p < 0.05$) and ‘**’ ($p < 0.01$) indicate significant differences determined by an unpaired *t*-test

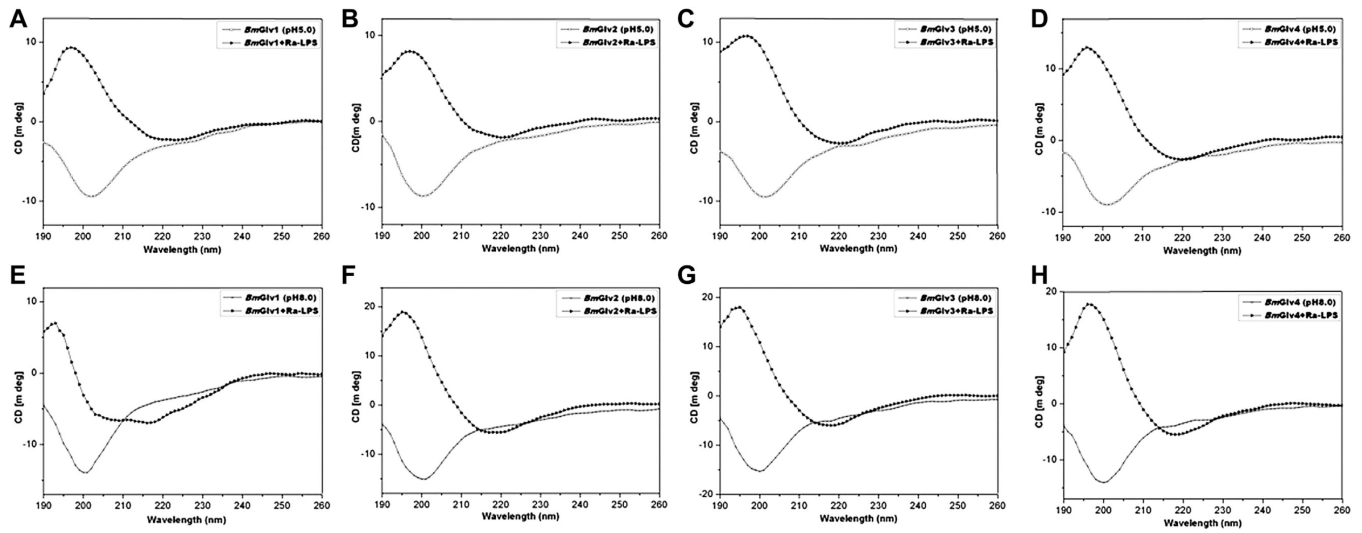


Fig. 4. CD spectra of *BmGlv*s in the presence of Ra-LPS at pH 5.0 and 8.0. Purified *BmGlv*1–4 were diluted to 0.15 mg/ml in 10 mM phosphate buffer (pH 5.0 or 8.0) in the presence or absence of Ra-LPS ($w/w= 1:1$) and CD spectra were recorded on a Jasco-810 spectropolarimeter at 25 °C. Each CD spectrum was obtained after subtracting the signal from protein-free solution.

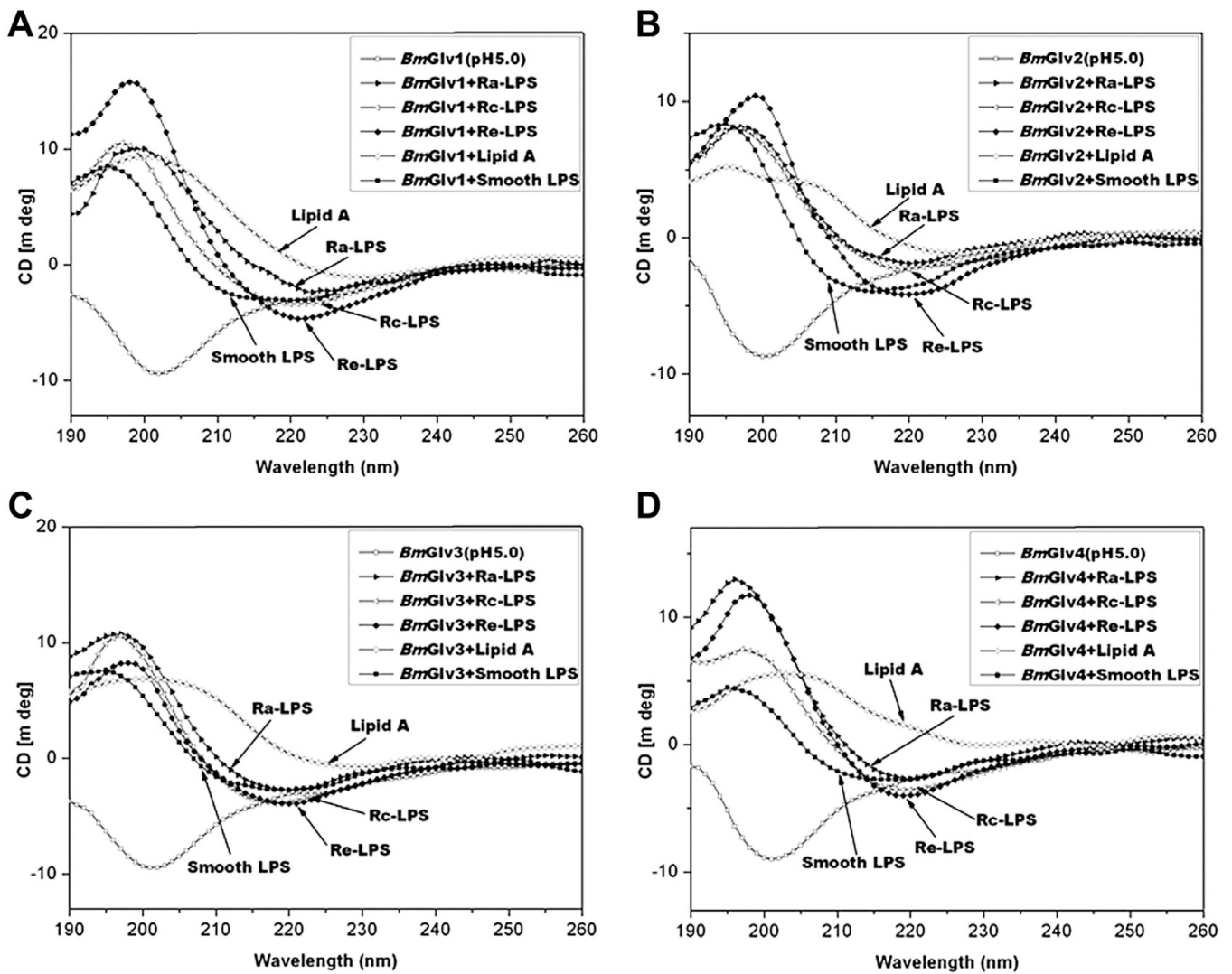


Fig. 5. CD spectra of *BmGlv*s in the presence of smooth LPS, Ra-, Rc- and Re-LPS and lipid A at pH 5.0. Purified *BmGlv*1–4 were diluted to 0.15 mg/ml in 10 mM phosphate buffer at pH 5.0 in the presence or absence of smooth LPS, Ra-, Rc- and Re-LPS or monophosphoryl lipid A ($w/w= 1:1$) and CD spectra were recorded on a Jasco-810 spectropolarimeter at 25 °C. Each CD spectrum was obtained after subtracting the signal from protein-free solution.

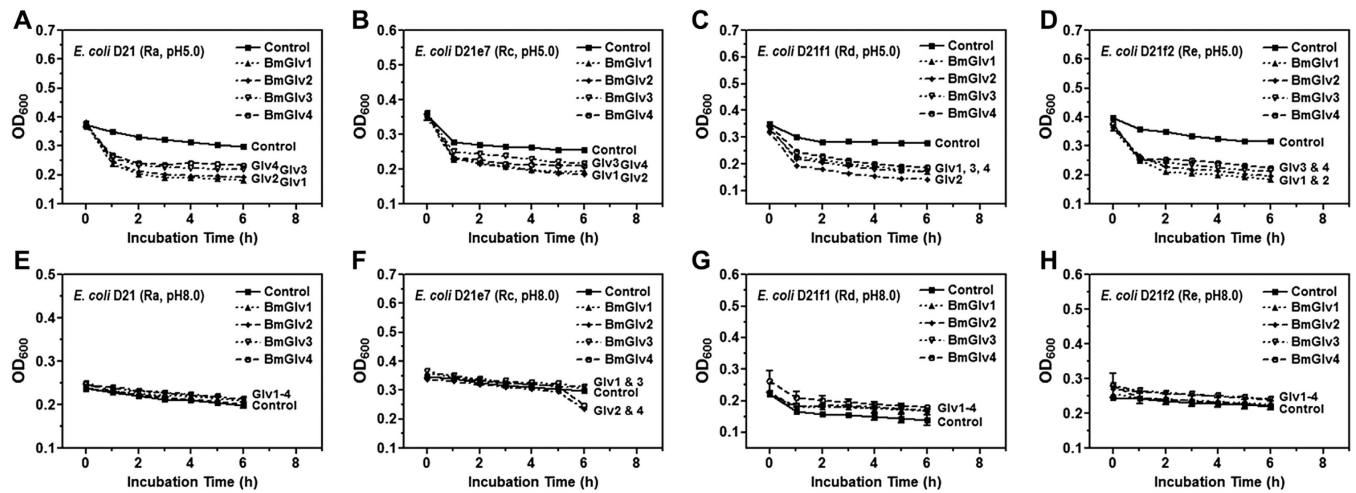


Fig. 6. The activity of *BmGlv*s against *E. coli* mutant strains containing rough LPS. Mid-log phase *E. coli* D21, D21e7, D21f1 and D21f2 cells with different rough mutant forms of LPS were diluted to $OD_{600} = 0.4$ in 10 mM phosphate, 100 mM NaCl (pH 5.0 or 8.0), and then incubated with purified *BmGlv*s (final concentration of 150 μ g/ml, ~ 10 μ M) or buffer (Control) in 96-well plates with 220 rpm shaking at 37 $^{\circ}$ C. OD_{600} was recorded every hour after incubation. The points represent the mean of four individual measurements \pm SEM.

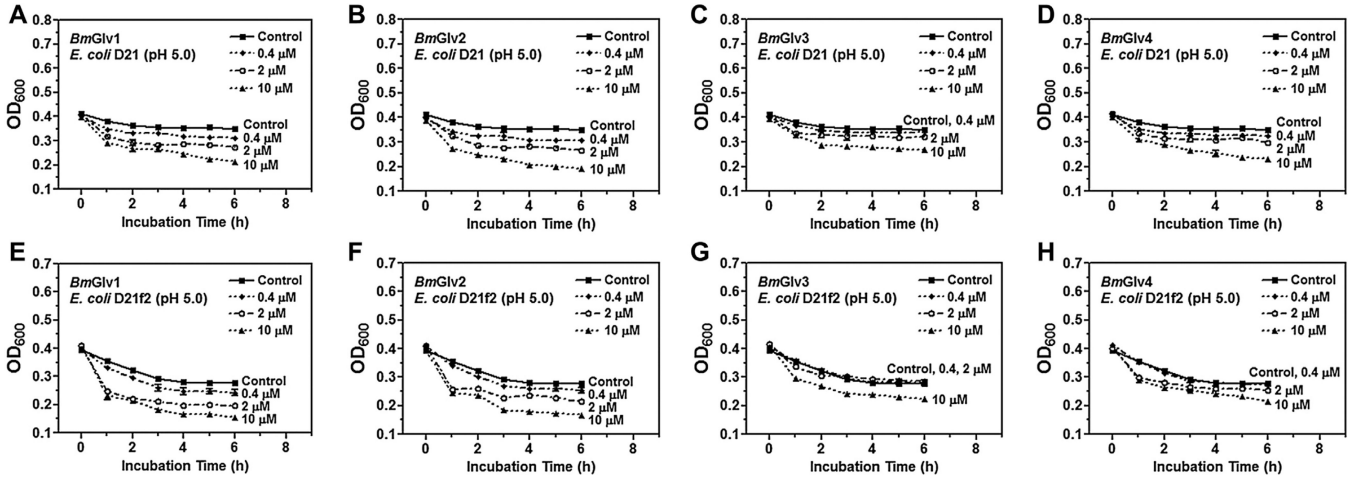
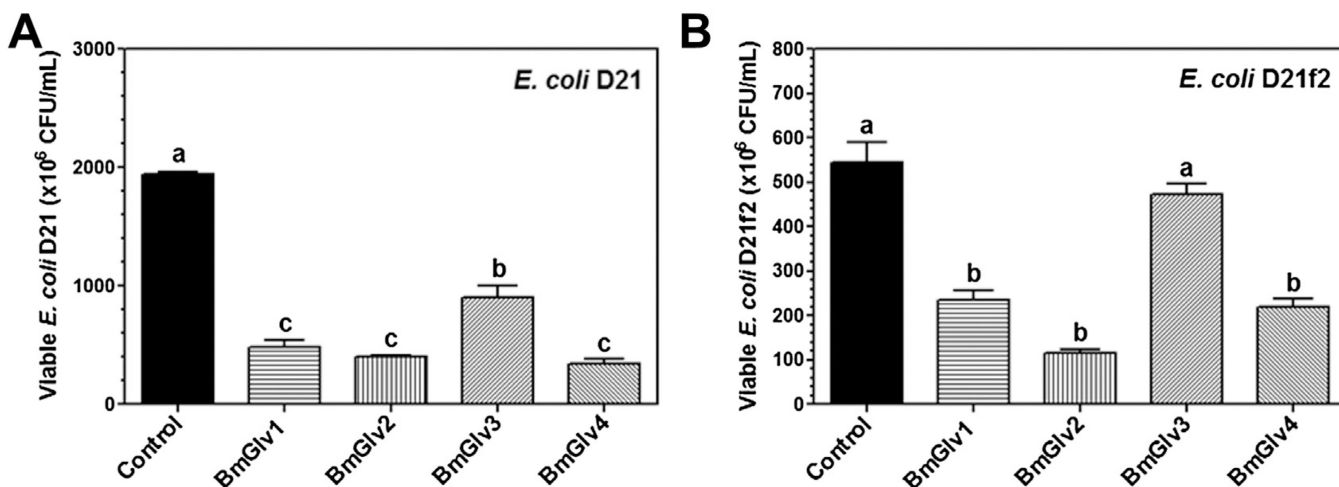


Fig. 7. Dose-dependent activity of *BmGlv*s against *E. coli* D21 and D21f2 mutant strains. Mid-log phase *E. coli* D21 and D21f2 cells were diluted to OD₆₀₀ = 0.4 in 10 mM phosphate, 100 mM NaCl, pH 5.0, and then incubated with increasing concentrations of purified *BmGlv*s (final concentration of 0.4, 2 and 10 μM) or buffer (Control) in 96-well plates with 220 rpm shaking at 37 °C. OD₆₀₀ was recorded every hour after incubation. The points represent the mean of four individual measurements ± SEM.

**Fig. 8.**

Viability of *E. coli* D21 and D21f2 cells after treatment with *BmGlv*s. Mid-log phase *E. coli* D21 and D21f2 cells were diluted to OD₆₀₀ = 0.4 in 10 mM phosphate, 100 mM NaCl, pH 5.0, and then incubated with purified recombinant *BmGlv*s (final concentration of ~10 mM) or buffer (Control) in 96-well plates with 220 rpm shaking at 37 °C for 2 h. The bacterial cells were serially diluted with 10 mM phosphate, 100 mM NaCl, pH 5.0, and aliquots of diluted bacterial cells were plated on LB-agar plates containing streptomycin sulfate (50 µg/ml) and ampicillin (10 µg/ml). The plates were incubated at 37 °C overnight and the numbers of viable bacterial cells were recorded. The bars represent the mean of three individual measurements ± SEM. Comparing the control group and *BmGlv*s treated groups, identical letters are not significant difference ($p > 0.05$), while different letters indicate significant difference ($p < 0.05$) determined by one way ANOVA followed by a Tukey's multiple comparison test

Table 1
 Predicted secondary structures of BmGlv1–4 in phosphate buffer at pH 5.0 and 8.0

	<i>BmGlv1</i>		<i>BmGlv2</i>		<i>BmGlv3</i>		<i>BmGlv4</i>	
	pH 5.0	pH 8.0	pH 5.0	pH 8.0	pH 5.0	pH 8.0	pH 5.0	pH 8.0
Helix (%)	0.0	0.0	0.0	0.0	0.0	0.0	0.0	0.0
Beta (%)	32.6	35.4	33.0	39.7	27.7	25.4	30.1	25.0
Turn (%)	12.6	12.0	12.4	9.1	17.6	19.2	15.8	17.7
Random (%)	54.8	52.6	54.6	51.2	54.7	55.4	54.1	57.3

Table 2
 Predicted secondary structures of *BmGlv1-4* in the presence of HFIP and Ra-LPS at pH 5.0

	BmGlv1		BmGlv2		BmGlv3		BmGlv4					
	pH 5.0	+HFIP	+Ra-LPS	+HFIP	+Ra-LPS	+HFIP	+Ra-LPS	+HFIP	+Ra-LPS			
Helix (%)	0.0	57.6	54.8	0.0	57.6	58.4	0.0	32.7	67.0	0.0	59.5	62.1
Beta (%)	32.6	0.0	0.0	33.0	0.0	0.0	27.7	25.9	0.0	30.1	0.0	0.0
Turn (%)	12.6	0.0	45.2	12.4	11.0	41.6	17.6	0.0	33.0	15.8	24.5	37.9
Random (%)	54.8	42.4	0.0	54.6	31.4	0.0	54.7	41.4	0.0	54.1	16.0	0.0

Table 3
 Predicted secondary structures of *BmGlv1-4* in the presence of HFIP and Ra-LPS at pH 8.0

	<i>BmGlv1</i>		<i>BmGlv2</i>		<i>BmGlv3</i>		<i>BmGlv4</i>					
	pH 8.0	+HFIP	+Ra-LPS	+HFIP	+Ra-LPS	+HFIP	+Ra-LPS	+HFIP	+Ra-LPS			
Helix (%)	0.0	44.9	14.0	0.0	25.6	63.0	0.0	26.5	65.4	0.0	46.6	63.4
Beta (%)	35.4	2.1	52.1	39.7	34.7	0.0	25.4	30.7	0.0	25.0	0.0	0.0
Turn (%)	12.0	10.9	0.7	9.1	0.9	37.0	19.2	0.0	34.6	17.7	19.5	36.6
Random (%)	52.6	42.0	33.2	51.2	38.8	0.0	55.4	42.8	0.0	57.3	34.0	0.0

