NMR Structure of Pardaxin, a Pore-forming Antimicrobial Peptide, in Lipopolysaccharide Micelles

MECHANISM OF OUTER MEMBRANE PERMEABILIZATION

Lipopolysaccharide (LPS), the major constituent of the outer membrane of Gram-negative bacteria, is an important element against permeability of bacterial agents, including antimicrobial peptides. However, structural determinants of antimicrobial peptides for LPS recognition are not clearly understood. Pardaxins (Pa1, Pa2, Pa3, and Pa4) are a group of pore-forming bactericidal peptides found in the mucus glands of sole fishes. Despite having a low net positive charge, pardaxins contain a broad spectrum of antibacterial activities. To elucidate the structural basis of LPS interactions of pardaxins, herein, we report the first three-dimensional structure of Pa4 bound to LPS micelles. The binding kinetics of Pa4 with LPS is estimated using [15N-Leu-19] relaxation dispersion NMR experiments. LPS/Pa4 interactions are further characterized by a number of biophysical methods, including isothermal titration calorimetry, 31P NMR, saturation transfer difference NMR, dynamic light scattering, and IR spectroscopy. In the LPS-Pa4 complex, Pa4 adopts a unique helix-turn-helix conformation resembling a "horseshoe." Interestingly, the LPS-bound structure of Pa4 shows striking differences with the structures determined in lipid micelles or organic solvents. Saturation transfer difference NMR identifies residues of Pa4 that are intimately associated with LPS micelles. Collectively, our results provide mechanistic insights into the outer membrane permeabilization by pardaxin.

Antimicrobial peptides (AMPs) are ubiquitous arsenals of all living organisms for combating against invading pathogens.

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The on-line version of this article (available at http://www.jbc.org) contains supplemental Figs. S1 and S2 and Table S1.

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3 The abbreviations used are: AMP, antimicrobial peptide; tr-NOE, transferred nuclear Overhauser effect; NOESY, nuclear Overhauser effect spectroscopy; TOCSY, total correlation spectroscopy; NOE, nuclear Overhauser enhancement; LPS, lipopolysaccharide; STD, saturation transfer difference; CPMG, Carr-Purcell-Meiboom-Gill; ITC, isothermal titration calorimetry; DLS, dynamic light scattering; FITC, fluorescein isothiocyanate; NPN, 1-N-phenylnaphthylamine; DPC, dodecyl phosphocholine.
activity of pardaxins, deduced from experiments in lipid vesicles, had been proposed to occur as a result of peptide aggregations (28–32). Solid-state NMR experiments pointed out the membrane orientation and interactions of pardaxins, hence mechanisms of action, are highly dependent on compositions of lipids in the bilayers (38, 39). The NMR structure of Pa4 in zwitterionic DPC micelles demonstrated a well defined C-terminal helix and a rather flexible turn/helix structure at the N-terminal (40).

In this study, tr-NOESY (44, 45), STD NMR (46, 47), and CPMG relaxation dispersion (48, 49) experiments were used to determine three-dimensional structure of pardaxin Pa4, and the localization of the amino acid residues in LPS and binding kinetics of the complex. Further, membrane permeability assay, CD, ITC, IR, dynamic light scattering (DLS), and 31P NMR studies of peptide/LPS revealed the binding and molecular mechanisms underlying the outer membrane disruption by Pa4.

EXPERIMENTAL PROCEDURES

Reagents—LPS of Escherichia coli 0111:B4 and fluorescein isothiocyanate (FITC)-conjugated LPS from E. coli 055:B5 were purchased from Sigma. Pa4 was synthesized by solid-phase peptide synthesis method containing a [15N-Leu-19] label and purified as described before (38).

NPN Uptake Assay—The outer membrane permeabilization activity of Pa4 was analyzed by using an 1-N-phenylphpenyl-amine (NPN) dye uptake assay (50). E. coli BL21(DE3) cells were grown in LB broth to mid-log phase (A600 of 0.38), centrifuged at 3000 × g for 10 min at 298 K, and resuspended in buffer containing 5 mM sodium HEPES buffer, pH 7.2, 5 mM glucose, and 5 mM NaN3. After washing once, the pellet was resuspended in the same buffer to a final A600 of 0.5. The basal fluorescence level of 500 µL of treated cells was acquired using a Cary Eclipse fluorescence spectrophotometer (Varian, Inc., Palo Alto, CA). Excitation and emission wavelengths used for this assay were 350 nm and 420 nm, employing bandwidths of 5 nm, respectively. NPN, at a concentration of 10 µM, was added to the cells, and a new basal fluorescence level was recorded after 5 s. Assay controls (5 µM polymyxin B as positive control) and increasing concentrations of pardaxin (3–15 µM) were added, and NPN fluorescence spectra were obtained in the presence of cells. Fluorescence intensity data were analyzed after subtracting the background level of fluorescence of NPN alone.

Secondary Structure by CD Spectroscopy—Secondary structure was determined for Pa4, 100 µM, in 10 mM sodium phosphate buffer at pH 4.5, at 298 K, in the absence and presence of 100 µM LPS from E. coli 0111:B4. CD data were collected using a Chirascan CD spectrometer (Applied Photophysics Ltd., UK). The far-UV CD spectra were scanned over the range of 180–235 nm wavelengths in a 0.01-cm path length cuvette, using a bandwidth of 1 nm, step size of 0.4 nm, a time of 5 s at each wavelength, and averaging over three scans. Baseline scan was obtained using the same parameters on buffer alone, and it was subtracted from the data scan of Pa4. The corrected data obtained in millidegrees (θ) was converted to molar ellipticity in deg.cm² dmol⁻¹.

ITC—All isothermal titration calorimetry (ITC) experiments were performed on a VP-ITC Micro-Calorimeter (MicroCal Inc., Northampton, MA) in 10 mM sodium phosphate buffer at pH 6.2. Prior to titration, LPS samples were vortexed for 15 min, heated at 60 °C for 5 min, and sonicated for 2 min. The molecular mass for E. coli 0111:B4 LPS was considered to be 10 kDa (51). A typical titration involved 30 injections of 1 mM pardaxin (10 µL per injection, except first injection of 3 µL) at 400-s intervals, into the sample cell (volume 1.4 mL) containing 50 µM LPS. The reaction cell was stirred continuously at 300 rpm. The high feedback level of mode/gain was selected. The titrations were carried out at two different temperatures, 298 K and 313 K. Raw data were corrected for the heat of dilution of Pa4 in buffer and integrated using a MicroCal Origin 5.0 supplied by the manufacturer. The first data point (3-µL injection) was not considered for the analysis. A single site binding model was used to fit the data using a non-linear least square method yielding the binding stoichiometry (n), equilibrium association constant (Kₐ), and enthalpy change (ΔH). The change of free energy (ΔG) and the change of entropy (ΔS) were calculated using the fundamental equations of thermodynamics, ΔG = −RT ln Kₐ and ΔS = (ΔH − ΔG)/T, respectively.

Size Distribution of LPS by DLS—Particle size distributions of LPS micelles in free and in the presence of Pa4 were investigated using DLS measurements, performed on a BI-200SM instrument equipped with a He-Ne laser and a BI-900AT autocorrelation system (Brookhaven Instruments Corp., Holtsville, NY). E. coli 0111:B4 LPS, 1 µM, dissolved in a 10 mM sodium phosphate buffer, pH 6.2 at 298 K, was exposed to 90° light scattering for 3 min. DLS measurements were recorded for LPS in the presence of 2 µM Pa4, under similar conditions. All samples were filtered, degassed, and scanned using a 1-mm path length quartz cuvette. DLS data were analyzed by using the CONTIN method supplied by the manufacturer.

NMR Spectroscopy—All NMR spectra were recorded on a Bruker DRX 600 spectrometer, equipped with a cryo-probe and pulse field gradients. Data acquisition and processing were performed with the Topspin software (Bruker) running on a Linux workstation. Two-dimensional TOCSY (total correlation spectroscopy) and NOESY (nuclear Overhauser effect spectroscopy) spectra of Pa4 in free solution were acquired in an aqueous solution containing 10% D2O at pH 4.5 with 0.5 mM peptide concentration and mixing times of 80 ms and 150 ms, respectively. The measurements were performed at two different temperatures (288 and 298 K) using 80 scans and 2,2-dimethyl-2-silapentane-5-sulfonate sodium salt as an internal standard (0.0 ppm). However, due to severe signal overlap at 298 K, the resonance assignments of Pa4 in aqueous solution were done at 288 K. A 0.5 mM solution of Pa4 was titrated with various concentrations of LPS, ranging from 9 to 74 µM. The two-dimensional tr-NOESY experiments were performed at the molar...
ratio of [LPS]: [Pa4] = 1:12.5 with three different mixing times (100, 150, and 200 ms) and at two different temperatures (288 and 298 K). NOESY and tr-NOESY experiments were performed with 456 increments in t1 and 2K data points in t2 using the WATERGATE procedure for water suppression and States-TTPPI for the quadrature detection in the t2 dimension. The spectral width was typically 12 ppm in both dimensions. After zero filling along the t1 dimension, 4K (t2) × 1K (t1) data matrices were obtained. NMR data processing and analysis were carried out using Topspin (Bruker) and SPARKY4 programs, respectively.

For the one-dimensional STD experiments, LPS was saturated at −2.0 ppm (40.0 ppm for reference spectra) with a cascade of 40 selective Gaussian-shaped pulses (49 ms each) in an interval of 1 ms resulting in total saturation time of 2 s. The two-dimensional STD TOCSY spectra were recorded with 350 t1 increments and 80 transients using a MLEV-17 spin lock field of 80 ms. The relaxation delay was set to 2.1 s. Saturation transfer was achieved by using 40 selective Gaussian 270 pulses with a duration of 50 ms. The LPS resonances were irradiated at −2.0 ppm (on-resonance) and 40.0 ppm (off-resonance) for 2 s. Subtraction of the two spectra (on resonance off-resonance) by phase cycling (46) leads to the difference spectrum that contains signals arising from the saturation transfer. As a control, one-dimensional STD experiments were performed using an identical experimental setup with the same peptide concentration but in the absence of LPS.

Interactions of Pa4 with LPS were examined by recording a series of one-dimensional 31P NMR spectra of LPS whereby 0.2 mm of LPS in water at a pH of 4.5 was titrated with various concentrations, 0.05, 0.1, 0.2, and 0.4 mm, of Pa4 at 298 K. The 31P NMR spectra were recorded on a Bruker DRX 400 spectrometer.

**Relaxation Dispersion Measurements**—The binding kinetics parameters of Pa4 [15N-Leu-19] to LPS (molar ratio of [LPS]: [Pa4] = 1:12.5) were obtained from two dimensional 1H-15N relaxation dispersion measurements using the Carr-Purcell-Meiboom-Gill (CPMG) pulse sequence (48, 49) on a Bruker DRX 700 spectrometer, equipped with an actively shielded cryo-probe and pulse field gradients. The total CPMG period (T1,1s) was 60 ms in all the experiments, and the effective field strength was varied by changing the delay time (Tdp) between consecutive 15N 180° refocusing pulses, ranging from 7.76 and 0.67 ms in 9 experiments with two duplicates for error estimation. A reference experiment was also performed by omitting the CPMG sequence. Effective R2 rates were calculated as described elsewhere (52). To determine the exchange kinetics, effective relaxation rates (R2) as a function of inverse of the delay time (Tdp) were fitted to the Carver-Richards model (46) using an in-house C-program yielding kex and pb values. The grid-search algorithm was used to minimize the errors to the initial input values.

**NMR-derived Structure Calculation**—NMR structures were calculated using the DYANA program (version 1.5) (53) as described previously (54). Briefly, on the basis of cross-peak intensities from the tr-NOESY spectra recorded at a mixing time of 150 ms, the NOE intensities were qualitatively categorized to strong, medium, and weak and then translated to upper bound distance limits to 2.5, 3.5, and 5.0 Å, respectively. The LPS-bound conformation of Pa4 was calculated solely based on tr-NOE-derived distance constraints. To restrict the conformational search, the ϕ dihedral angles were set to −30° to −120° for all residues except for Gly and Pro. No hydrogen bond constraints were used for structure calculations. Several rounds of structure calculations were carried out, and, depending on NOE violations, the distance constraints were adjusted. Out of the 100 structures, the 20 lowest energy structures were used for further analysis.

**Infrared Spectroscopy**—Fourier transform IR spectra were recorded on a Nicolet Nexus 560 spectrometer (Madison, WI) purged with N2 and equipped with an MCT/A detector cooled with liquid nitrogen. Attenuated total reflection spectra were measured with a 25-reflection attenuated total reflection accessory from Graseby Specac (Kent, UK) and a wire grid polarizer (0.25 mm, Graseby Specac). Approximately 200 μl of a D2O solution of LPS alone or in the presence of peptide in a 20:1 lipid/peptide molar ratio was applied onto a trapezoidal (50 mm × 2 mm × 20 mm) germanium internal reflection element. A dry, or D2O-saturated, N2 stream flowing through the attenuated total reflection compartment was used to remove the bulk water (low hydration) or to fully hydrate the sample (high hydration), respectively. A total of 200 scans was collected at a resolution of 4 cm−1, averaged, and processed with one-point zero filling and Happ-Genzel apodization. The secondary structure composition of the peptide was obtained by analyzing the amide I band of the non-polarized spectrum, using previous assignments (55). Main component bands were identified after mild Fourier deconvolution (full width half-height 20 cm−1 and k 2.5). Non-polarized spectra were obtained from the parallel ([∥]) and perpendicularly ([⊥]) attenuated total reflection polarized spectra, using the expression, 1([∥]) + 1.44([⊥]), as described previously (56).

**RESULTS**

**Disruption of Outer Membrane Integrity by Pa4**—We have examined whether Pa4 can permeabilize outer membrane by use of NPN uptake assay. NPN is a neutral, hydrophobic probe that fluoresces weakly in an aqueous environment, but strongly in the hydrophobic milieu of a membrane. When NPN was mixed with intact E. coli cells, it emitted a weak fluorescence at 515 nm, as it was unable to penetrate the outer membrane permeability barrier. Upon subsequent additions of Pa4, NPN partitioned into the hydrophobic region of the membrane and demonstrated an enhanced fluorescence intensity (Fig. 1). A dose-dependent increase in the fluorescence intensity was observed, indicating the loss of outer membrane integrity. A marked blue shift (~14 nm) in NPN fluorescence emission maxima (Fig. 1, inset) further indicates that NPN is well inserted into the non-polar environment of E. coli cell membranes. These data demonstrate that Pa4 efficiently permeabilizes the outer membrane as observed for the inner membrane.

**Secondary Structure of Pa4 in LPS Micelles**—In aqueous solution, the far-UV CD spectrum of Pa4 is characterized by a

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strong negative band at ~200 nm (Fig. 2), representative of random-coil like conformations. This observation is in accordance with previous CD studies, which reported that the pardaxins are mostly unstructured in aqueous solution (35, 37). But a drastic change in the CD spectrum of Pa4 in the presence of LPS indicates structural changes of Pa4 upon binding to LPS (Fig. 2). A very strong positive band at 192 nm and two negative maxima of approximately equal intensity at 222 nm and at 208 nm, typically observed for a protein rich in helical conformations. This observation clearly demonstrates that the Pa4 has random conformations in aqueous buffer but adopts predominantly helical structures in the context of LPS micelles.

**Binding Affinity of Pa4 to LPS**—ITC was used to determine the energetic of interactions and equilibrium binding constants involving Pa4 and LPS. The binding of Pa4 to LPS was determined by titrating Pa4 into LPS solutions at pH 6.2, at 298 K and 313 K. At 298 K, the interaction was found to be endothermic in nature, as evident by the upward position of titration peaks and the corresponding positive integrated heats (Fig. 3). In other words, binding is entropically driven and hence hydrophobic in nature. However, at 313 K, above the phase transition temperature of LPS, the interactions became exothermic, as seen with the downward position of titration peaks and the corresponding negative integrated heats ($\Delta H < 0$) (Table 1 and supplemental Fig. S1); these results suggest that the binding reaction is enthalpically driven. This temperature-dependent behavior (endothermic to exothermic transition) of LPS-peptide interactions has been attributed to the gel phase to liquid phase transition of LPS micelles (57, 58). However, irrespective of the phase transition thermodynamic effects, Pa4 binds to LPS with a similar equilibrium binding association constant ($K_a$) of $0.1\mu M^{-1}$ and $0.15\mu M^{-1}$ at 298 K and 313 K, respectively (Table 1).

**Conformations of Isolated Pa4**—A nearly complete sequence-specific proton resonance assignment of Pa4 in aqueous solution was achieved by the analysis of two-dimensional $^1$H-$^1$H NOESY and TOCSY (59) spectra (supplemental Table S1). The primary amino acid sequence of 33-residue Pa4 contains as many as 7 Ser, 5 Leu, 3 Phe, 3 Ile, and 3 Gly residues. In our experimental conditions (see “Experimental Procedures”) most of the resonances from these amino acids were unambiguously assigned (supplemental Table S1). The free-state
NOESY spectrum of Pa4 is characterized by the presence of intra-residue and sequential NOEs between the backbone protons and the side-chain protons of Pa4 (supplemental Fig. S2A). The lack of any diagnostic NOEs clearly shows that Pa4 does not adopt any unique conformation(s) in the free solution, which is in agreement with the CD data (Fig. 2) and other studies of pardaxins in aqueous solution (36, 37).

Binding Kinetics of Pa4 in Complex with LPS—Additions of aliquots of LPS into solutions containing Pa4 showed a concentration-dependent broadening of almost all the proton resonances of the peptide, without any significant changes in the chemical shifts (supplemental Fig. S2B). This observation demonstrates that the Pa4 undergoes a fast exchange between its free and LPS-bound states at the NMR time scale.

CPMG relaxation dispersion experiments are useful to determine millisecond to microsecond dynamics of 15N-labeled proteins utilizing 1H-15N heteronuclear single quantum coherence correlations (48, 49). Recent studies on transient binding of peptide ligands with the target proteins demonstrated that the kinetic parameters of the exchange processes can be extracted from relaxation dispersion measurements (52, 60). To understand the dynamics of Pa4 in the context of LPS binding, CPMG relaxation dispersion experiments were carried out on the complex at a Pa4:LPS molar ratio of 12:5:1. Fig. 4 shows relaxation dispersion profile of the 15N resonance of residue Leu-19 in the presence of LPS micelles. In the absence of LPS, Pa4 did not respond to the CPMG pulse trains showing no significant relaxation dispersion (Fig. 4). The dispersion data of the Pa4-LPS complex was fitted into a two-state exchange model (50) yielding the exchange rate \( k_{ex} \), the absolute value of the chemical shift difference \( \Delta \text{ppm} \), and bound-state population \( p_b \) (Fig. 4, inset). The data suggest that the rate of exchange of the complex to be fast (~1725 s\(^{-1}\)), typically suitable for tr-NOESY experiments (44, 45).

Analysis of tr-NOESY Spectra—tr-NOESY is a useful tool to determine three-dimensional structures of ligands bound to the macromolecules (45). LPS forms high molecular weight micelles at low concentrations (≤0.1 μM) (51, 61), permitting observation of tr-NOEs from the bound peptides with a dissociation constant \( K_d \) in the micromolar to millimolar range (63–66). Fig. 5 shows the two-dimensional 1H-1H tr-NOESY spectra of Pa4, correlating fingerprint and aromatic regions in the presence of 40 μM LPS. By contrast to the NOESY spectra of Pa4 in free solution (supplemental Fig. S2A), there was a dramatic improvement, in terms of number and intensity of NOE cross-peaks, in the NOESY spectrum of Pa4 in complex with LPS micelles. In addition to the backbone \( \alpha \text{N} \) (i,i+1) and sequential NN (i,i+1) tr-NOEs of Pa4 in LPS, a large number of medium range \( \alpha \text{N} \) (i,i+3/i+4) tr-NOEs was detected, indicating a helical structure (Fig. 5A). Medium range NOEs were unambiguously identified for residues Ala\(^8\)–Ser\(^{12}\) and Phe\(^{15}\)–Ser\(^{28}\) (Fig. 6A). Based on the tr-NOEs alone, two helical segments (helix 1, Ala\(^{8}\)–Ser\(^{12}\); helix 2, Phe\(^{15}\)–Ser\(^{29}\)) could be qualitatively assigned for Pa4 in LPS micelles. The peptide bonds between residues Ile\(^6\) and Pro\(^7\) and residues Ser\(^{12}\) and Pro\(^{13}\) adopt only a trans configuration as judged by the strong tr-NOEs between Ile\(^6\)C\(^{\alpha}\)H-Pro\(^7\)C\(^\beta\)Hs and Ser\(^{12}\)C\(^\beta\)H-Pro\(^{13}\)C\(^\alpha\)Hs, respectively. The chemical shifts of the aromatic ring protons and the amide proton of Phe\(^{15}\) are well separated from residues Phe\(^2\) and Phe\(^3\) (Fig. 5 and supplemental Table S1). There were a number of unambiguous long range tr-NOE contacts involving aromatic ring protons of Phe\(^{15}\) with CßH\(^5\) of residue Ala\(^{21}\) and also with the side-chain C\(^\alpha\)H\(^2\)/C\(^\alpha\)H\(^4\) groups of Ile\(^9\) (Fig. 5, B–D). The aromatic ring protons of Phe\(^{15}\) also showed an ample number of medium range tr-NOEs with the C\(^\alpha\)H\(^2\) of Pro\(^{13}\), C\(^\alpha\)Hs of Leu\(^{19}\), C\(^\alpha\)H\(^5\) of T17 and C\(^\beta\)Hs of Leu\(^{18}\) and Leu\(^{19}\) (Figs. 5D and 6A). By contrast, the aromatic ring protons of Phe\(^2\) and Phe\(^3\) appeared to be lacking any medium or long range NOE contacts in the Pa4-LPS complex. These data suggest that residue Phe\(^{15}\) is engaged by a large number of NOE contacts, ≥40 NOEs, with the neighboring residues in complex with LPS (Fig. 6B).

Three-dimensional Structure of Pa4 Bound to LPS—An ensemble of conformations of Pa4 in complex with LPS micelles was determined based on 242 distance constraints derived from 104 sequential, 72 medium range, and 16 long range tr-NOEs (Table 2). Fig. 6C delineates root mean square deviation (r.m.s.d.) for each residue. The LPS-bound structure of Pa4 is well defined with an average backbone and heavy atoms r.m.s.d. of 0.6 and 1.0 Å, respectively (Table 2). At the central region of the peptide, especially for the residues Ala\(^8\)–Ser\(^{29}\) except for Lys\(^3\) (Fig. 7A). However, both the termini of Pa4, e.g. residues Gly\(^{1}\)–Phe\(^{3}\) and Gly\(^{30}\)–Glu\(^{33}\), appear to be experiencing somewhat a larger conformational space (Fig. 7A). Fig. 7B represents the disposition of backbone topology and...
side-chain orientation of a selected structure of Pa4 bound to LPS. The LPS-induced conformation of Pa4 is defined by a short helix at the N-terminal (residues Leu5–Ser12) and a comparatively longer helix at the C-terminal (residues Lys16–Ser28), connected by a short loop consisted by residues Pro13, Leu14, and Phe15 (Fig. 7B). Other residues, Phe2–Ala4 and Ser29–Glu33, of Pa4 are found to be assuming extended conformations in LPS micelles (Fig. 7B). The N and C termini helices of Pa4 assume a defined orientation dictated by the aromatic residue Phe15 from the loop, giving rise to a helix-loop-helix or helical hairpin structure (Fig. 7B). The longer C-terminal helix shows an approximate tilt, moved toward the shorter helix, of 25° from the helical axis.

The hydrophobic face of the N-terminal helix is maintained by the residues Leu5, Ile6, Ile9, and Ile10, whereas residues Lys8, Ser11, and Ser12 are positioned at the opposite hydrophilic face of Pa4 (Fig. 7B). In the longer helix, residues Leu19, Val22, and Leu26 are located in the non-polar face, whereas residues Thr27, Ser28, and Ser29 are in the polar region (Fig. 7B). However, the polar or hydrophilic face of the C-terminal helix is interrupted by the presence of non-polar residues Leu18, Ala21, and Ala25 (Fig. 7B). The aromatic residue Phe15, from the intervening loop, plays an important role in bringing the non-polar faces of two helices, consisting of residues Ile9, Ile10, Pro13, Leu18, Leu19, and Ala21, to form a hydrophobic core (Fig. 7C). Interestingly, this structural arrangement appears to place the two positively charged residues (Lys8 and Lys16) across the two helices maintaining a distance of ~13 Å (Fig. 7D). This distance separation is compatible with the inter-phosphate distance of lipid A moiety of LPS (see below). The LPS-induced structure of Pa4 essentially demonstrates a “horseshoe”-type fold (Fig. 7E). A substantial part of the horseshoe, including the “head or bend” is non-polar, sustained by residues
The ring proton resonances of Phe2, Phe3, Leu5, Ile6, Ile9, Ile10, Pro13, Phe15, Leu19, Val22, and Leu26 (Fig. 7E). The end regions of the two arms of the horsehoe are maintained by oppositely charged residues (Fig. 7E). The LPS-bound structure of Pa4 shows striking differences with the structure obtained in DPC micelles (Fig. 7F) (see below).

**Interactions between Pa4 and LPS by STD NMR**—To understand the localization of Pa4 in LPS micelles, STD NMR experiments were performed. We have carried out one-dimensional STD and two-dimensional STD-TOCSY to examine the interacting residues of Pa4 with LPS. Overall, the one-dimensional STD spectra show aromatic ring protons, and a number of aliphatic side-chain proton resonances are in direct contact with LPS micelles (data not shown). The two-dimensional STD-TOCSY spectra are of high resolution facilitating unambiguous identification of LPS-binding residues (Fig. 8). Fig. 8 (A and B) shows off-resonance (or reference) TOCSY spectra and corresponding STD-TOCSY spectra of Pa4 in LPS, respectively, for the downfield-shifted aromatic ring protons (left panels) and upfield-shifted aliphatic side-chain proton resonances (right panels). STD-TOCSY spectra are interpreted following a previous study (47), whereby resonances showing cross-peaks in the STD-TOCSY spectra are considered to be strongly saturated owing to their close contacts, ≤5 Å, with macromolecules. As can be seen, the ring proton resonances of Phe2, Phe3, and Phe15 residues of Pa4 delineate intense STD effects, indicating their close contacts with LPS (Fig. 8B, left panel). The two β-proton resonances of Phe15 are non-degenerate, resonating at 3.13 and 3.04 ppm, showing TOCSY correlations in the off-resonance spectrum (Fig. 8A, right panel). However, in the STD-TOCSY spectra, there are no correlations for the β-proton resonances of Phe13 (Fig. 8B, left panel), suggesting these protons are not in close proximity with LPS micelles and, hence, had received less or a lower degree of saturations. Similarly, a low degree of saturations or lack of contact with LPS were observed for the CαHs of Pro7 and Pro13 and C-terminal residues Gln32 and Glu33 of Pa4 (Fig. 8, A and B). On the other hand, strong STD effects were detected for the side-chain resonances of non-polar residues Leu5, Ile6, Ile9, Ile10, Leu14, Leu18, and Val22, indicating their close proximity to LPS micelles (Fig. 8B, left panel). Most importantly, two-dimensional STD (Fig. 8B, right panel) spectra clearly show the cross-peaks from the side chains (CβHs, CαHs, and CαHs) of two basic residues Lys8 and Lys16, establishing their direct contacts with LPS. Taken together, STD-TOCSY data demonstrate critical residues, including aromatic, aliphatic, and two cationic, that are in close contact with LPS micelles.

**Perturbation of LPS Micelles by Pa4**—We have probed plausible structural changes of LPS micelles upon binding to Pa4 using three different methods, namely FITC fluorescence of FITC-conjugated LPS, dynamics light scattering, and 31P NMR experiments. Fluorescence intensity of FITC of Pa4 had been found to be sensitive to the aggregation states of FITC-LPS (67). A quenched fluorescence intensity of FITC-LPS could be observed in the absence of Pa4 (Fig. 9A). However, the binding of Pa4 to LPS has caused an enhancement of fluorescence intensity of FITC (Fig. 9A). A 5-fold increase in the fluorescence intensities of FITC has been observed upon additions of Pa4 from nanomolar (100 nM) to micromolar (25 μM) concentrations. This quenching effect results from the dissociation of LPS aggregates, indicating destabilization of LPS assembly. Several host defense AMPs, including magainin and human cathelicidin LL37, had demonstrated a similar effect on the LPS structure (18, 68). The effect of Pa4 on the apparent size distribution of LPS micelles was determined by DLS using photon correlation spectroscopy at a scattering angle of 90°. Fig. 9B shows particle size analysis of LPS aggregates in the absence (upper panel) and in the presence of Pa4 (lower panel). The hydrodynamic diameter of LPS aggregates is shown along the x-axis. The vertical bars represent the relative percentage of the LPS particle size distribution, with the larger aggregates having a normalized frequency of 100. As can be seen, in the presence of Pa4, high molecular weight clusters of LPS aggregates with a diameter of >500 nm are completely absent. In addition, the polydispersity of LPS aggregates with a diameter centered at ~1347 nm were shifted with concomitant 80% decrease in mean hydrodynamic diameter of LPS to 271 nm in the presence of Pa4. This observation once again explains the dissociation of LPS micelles into smaller sizes upon interactions with Pa4.

Binding of Pa4 with the phosphate head groups of LPS were further examined using 31P NMR experiments. The 31P NMR spectrum of LPS micelle is characterized by two well resolved resonances (Fig. 9C). By the comparison of 31P NMR spectra of LPS dissolved in SDS detergent micelle (66, 69), we assumed that the intense peak ~2.00 ppm corresponds to the diphasate groups of LPS, and the downfield-shifted peak at ~0.72 ppm belongs to the monophosphate group of LPS, attached to the O4′ of GlcN I. There was a drastic change of 31P resonances of the LPS micelles upon additions of increasing concentrations of Pa4, as monitored by progressive chemical shift changes as well as broadening of 31P resonances (Fig. 9C). The 31P resonance of LPS at ~2.00 ppm was shifted significantly upfield in complex with Pa4. On the other hand, the 31P resonance of LPS at ~0.72 ppm undergoes a significant broadening upon binding to Pa4 (Fig. 9C). Such line broadening of 31P resonances of LPS upon binding to Pa4 suggests that plausible conformational exchanges among the different states of LPS-peptide complexes. The upfield shift of the 31P resonance of LPS may occur due to a close proximity of the one or more Phe residues (Phe2, Phe3, and Phe15) of Pa4 with the head groups of LPS, as demonstrated by the STD experiments (Fig. 8). Taken together, these results (Fig. 9) confirm that the LPS micelles undergo structural rearrangements or a dissociation of larger LPS aggregates into smaller sizes in complex with Pa4.
IR Studies—In the presence of hydrated LPS, Pa4 showed two main bands in the amide I region of the IR spectrum (Fig. 10, blue). These bands become more apparent after Fourier self-deconvolution and are centered at 1653 and 1637 cm⁻¹. The first band is clearly assigned to α-helix, whereas the second band is usually assigned to β-structure, although it has also been assigned to the 3₁₀-helix (70). The amide I also has a shoulder at ~1670 cm⁻¹ that is usually assigned to turns (loop regions). The bands at 1732 and 1586 cm⁻¹ probably correspond to a protonated and a deprotonated side chain of residue Glu³³ (71), respectively. The band at 1543 cm⁻¹ corresponds to the amide II band of the peptide, which therefore is at least partially protected from D₂O exchange in these conditions. Although a quantitative estimate of the secondary structure is not possible, because the areas of the two main bands are not similar. The structural characterization of Pa4 in LPS micelles by IR is in good agreement with the NMR-derived structure showing the presence of helices, loop/turn, and extended conformations at the termini.

DISCUSSION

Pardaxins are originally identified as pore-forming membrane lytic peptides (30, 31). Later on, antibacterial activities, against both Gram-negative and Gram-positive organisms, of pardaxins were discovered (33, 35). The antibacterial activities of AMPs are often positively correlated with the net content of cationic residues in their amino acid sequences (5). Analysis of charge distribution in the AMPs had revealed average positive charges of +4 to +6 for the most potent AMPs (5). A high cationicity and amphipathic structure have been attributed to AMPs membrane interactions and cell lysis (1–3).
particular, a higher positive charge content has been thought to be necessary to overcome the LPS-mediated barrier in Gram-negative organisms (11–13). Recent studies with amphibian AMPs temporins demonstrated the role of LPS in preventing the outer membrane permeabilization by low cationic (+1 to +2) antimicrobial peptides (20, 21). Temporins A and B are largely inactive against Gram-negative bacteria as a result of the lack of their outer membrane permeabilization. However, an interesting synergistic effect has been demonstrated among these temporins to disrupt outer membranes (20, 21). The structural basis of the synergistic mechanism is not known. Although it has been demonstrated that single temporins might aggregate in the LPS layer, thus preventing transversal through the outer membrane (20, 21). In contrast to these, Pa4 contains a net positive charge of +1, disregarding terminal charges, and is found to be independently active. A considerably large body of work has been carried out for Pa4 and other members of the family in synthetic lipids of different compositions. However, no studies on pardaxins embedded in outer membranes or LPS have been reported. Taken into consideration temporin synergistic mechanisms, it becomes important to determine structural consequences of this particular class of AMPs in the outer membrane or LPS.

In this context, we have determined the atomic level resolution three-dimensional structure of Pa4 as a complex with LPS for the first time and studied its interactions with LPS by various methods. Pa4 induces uptake of NPN in a concentration-dependent manner, indicating its ability to disrupt the outer membrane barrier. Energetics of interactions between LPS and Pa4 were investigated at 298 K and 313 K, corresponding to the gel and liquid crystalline phases of LPS, respectively. The binding affinity or equilibrium association constant between Pa4 and LPS has been estimated to be very similar at both temperatures (Table 1). Pa4 binding to the gel phase of LPS is primarily driven by the positive change of entropy or hydrophobic interactions. At 313 K, enthalpy change appears to be slightly negative with positive entropy change contributing to the interactions. It is noteworthy that ITC studies on polycationic antimicrobial peptide polymyxins and low charged temporins showed very similar temperature-dependent binding behavior with LPS (21, 58). Taken together, these results suggest that Pa4 also interacts similarly with LPS. CD studies indicate that LPS-bound Pa4 acquires helical con- formations, as demonstrated for pardaxins in lipid vesicles and membrane mimetic solvents (29, 33). tr-NOESY demonstrates a well defined structure of Pa4 in complex with LPS micelles. The tertiary fold, resembling a horseshoe, of Pa4 is determined by two helices, a short N-terminal helix and a longer C-terminal helix, connected by a tight turn. The aromatic ring of Phe15 from the turn acts as a pivot that interconnects the two helices by non-polar packing interactions. Few residues, Phe2–Ala4 and Ser29–Glu31 appear to be extended in LPS-bound Pa4 structure. The horseshoe structure of Pa4 in LPS determined in this work can be con-

### TABLE 2
A summary of structural statistics for the twenty final structures of Pa4 bound to LPS micelles

| Distance restraints                          | 50 | 104 | 72 | 16 | 242 | 29 |
|---------------------------------------------|----|-----|----|----|-----|----|
| Intra-residue (|j| | 0)   |    |    |    |    |
| Sequential (|j| | 1)   |    |    |    |    |
| Medium range (|j| | 4)   |    |    |    |    |
| Long range (|j| | 5)   |    |    |    |    |
| Total                                         |    |    |    |    |    |    |
| Angular restraints (φ)                       |    |    |    |    |    |    |

| Distance restraints violations (Å)            |   ≤0.21 |   ≤0.09 |
|---------------------------------------------|---------|---------|
| Average violations                           | 162.6   | 162.6   |
| Maximum violations                           | 0.01    | 0.01    |

| Deviation from mean structure                | 3.3%    | 1.06%   |
|---------------------------------------------|---------|---------|
| All residues (N, Cα, and C')                  | 0.62    | 1.06    |
| Heavy atoms                                   |         |         |

| Ramachandran plot for the mean structure*     |         |         |
|---------------------------------------------|---------|---------|
| % residues in the most favorable and additionally allowed region | 100 |
| % residues in the generously allowed region  | 0 |
| % residues in the disallowed region           | 0 |

* Based on Procheck-NMR.

**FIGURE 7.** Three-dimensional structure of Pa4 in LPS micelles. A, superposition of backbone atoms (N, Cα, and C') of the twenty lowest energy structures of Pa4 bound to LPS, obtained from DYANA 1.5. B, a representative conformation of LPS-bound Pa4, showing side-chain orientation and backbone topology. This figure was produced using PyMOL. C, a space-filling representation of the side-chain to side-chain contacts of Pa4, involving residues Ile8, Ile10, Pro14, Phe15, Leu16, Leu17, and Ala21, in complex with LPS. The figure was prepared using InsightII. D and E, electrostatic surface potential of Pa4 structure in LPS micelles in two different orientations. The dotted line indicates the distance between ammonium groups of Lys8 and Lys16 residues. F, superposition of the LPS-bound structure (in blue ribbon) and the DPC-bound structure (in red ribbon) of Pa4. These images were produced using MOLMOL software.
sidered as novel, because three-dimensional structures of AMPs determined to date either in LPS (64–66) or in synthetic lipid micelles (1, 5) do not resemble this conformation. Recently, we have determined a helical hairpin structure of a potent antimicrobial peptide MSI-594 in LPS micelles (72). Although MSI-594 adopts a helix-loop-helix structure in LPS, the topology of Pa4 and MSI-594 is largely different (Fig. 11A).

A T-shaped conformation and boomerang-type folds were observed for an 11-residue antimicrobial fragment of protein lactoferrin (64) and de novo designed peptides (66), respectively, in complex with LPS (Fig. 11A).

The three-dimensional structures of Pa4 and Pa1 have been determined in DPC micelles and in 2,2,2-trifluoroethanol/water mixture, respectively (40, 41). Under these conditions, independently folded helical structures were stabilized with a kink at residue Pro\(^{13}\) (Fig. 7F) (40). Most importantly, the DPC-bound structure of Pa4 lacks any interhelical packing interactions as observed for the LPS-bound structure (Fig. 7F). Recent NMR studies of AMPs in LPS micelles demonstrated constellations of non-polar/aromatic residues defining compact structures (54, 64–66, 72). A more recent work on the designed β-sheet peptide conformations in LPS had delineated functional significances, in terms of antimicrobial and anti-endotoxic activities, of packing interaction (66). STD experiments indicate a close proximity of the three aromatic ring protons (Phe\(^2\), Phe\(^3\), and Phe\(^{15}\)) and aliphatic side-chain protons with LPS micelles. It is interesting to note that, although the N-terminal aromatic residues Phe\(^2\) and Phe\(^3\) do not assume a folded structure in LPS, these residues are found to be in contacts with the micelles. The side chains of two cationic residues, Lys\(^8\) and Lys\(^{16}\), of Pa4 demonstrate strong STD effects for C\(^{\alpha}\)Hs, C\(^{\beta}\)Hs, and C\(^{\gamma}\)Hs, indicating their localization at the LPS micelles. Pa4 appears to bring significant perturbations of LPS micelle organization. FITC fluorescence and DLS data demonstrate disaggregation of LPS micelles into smaller sizes. In addition, the \(^{31}\)P NMR resonances of LPS are highly affected upon additions of Pa4. The chemical shift perturbations as well as the line broadening effect of phosphate groups of LPS in the presence of Pa4 suggest that an increase is plausible in the dynamics at the phosphate head groups of LPS. Taken together, these results elucidate a disorderliness of the LPS micelle and dissociation of LPS aggregates into smaller size complexes in complex with Pa4. Interestingly, similar structural perturbations of the phosphate head groups of LPS were observed for a highly active magainin- and melittin-derived AMP MSI-594 (72) and designed peptides (66). Collectively, these results indicate that overall effects on LPS micelle structure of Pa4 are similar to other highly cationic AMPs.

The three-dimensional structure of Pa4 in LPS and its interactions presented in this work provide mechanistic insight(s) at the outer membrane. In the horseshoe-shaped structure of Pa4 obtained in LPS micelles two cationic residues Lys\(^8\) and Lys\(^{16}\) located at the middle of the N terminus helix and at the beginning of the C terminus helix can interact with the bis-phosphate groups of lipid A moiety of LPS via salt bridge or hydrogen bonds.

**FIGURE 8. Localization of Pa4 in LPS micelles by STD NMR.** A, the reference or off-resonance TOCSY spectrum and (B) STD TOCSY spectrum of Pa4 in the presence of LPS in D\(_2\)O at pH 4.5 and 288 K, showing the downfield-shifted aromatic resonances (left panels) and upfield-shifted side-chain resonances (right panels). Spectra were acquired using spin-lock MLEV17 sequence with a mixing time of 80 ms. Saturation of LPS was achieved by applying a cascade of 40 Gaussian pulses (50 ms each), resulting in a total saturation of \(-2\) s (on-resonance, \(-2.0\) ppm; off-resonance, 40 ppm).
bonding. The close proximity of these residues is deduced from the STD studies. The three-dimensional structure of Pa4 in LPS demonstrates that the two charged H3N/H11001 groups of Lys8 and Lys16 are ~12–13 Å apart. This distance is geometrically compatible with the inter-phosphate distance of the phosphate groups of lipid A domain of LPS. Hydrophilic residues, e.g. Ser11 and Ser12 from the N-terminal helix, located at the same helical face of residue Lys8 and residue Thr17 from the C-terminal helix may complement the phosphate head group interactions by forming additional hydrogen bonds with the sugar moieties of LPS. In other words, this mode of binding of Pa4 to LPS may orient two of its helices providing facile contacts with the hydrophobic residues along the acyl chains of LPS (Fig. 11B). These interactions between Pa4 and LPS may lead to a plausible disruption or fluidization of LPS structures facilitating traversal of the peptide through the outer membrane. The compact structure of Pa4 will certainly aid in the process of permeation. The mode of LPS recognition by Pa4 has also partly been supported by functional and vesicle binding studies, whereby replacements of charged residues Lys8/Lys16 or the aromatic residue Phe15 had significantly reduced membrane interactions and antibacterial activities (32, 33).

Although our NMR studies have provided atomic level insights into the outer membrane permeation of pardaxin, it may be pointed out that the LPS micelles lack the contributions from other components of the bacterial outer membrane and the related constrained, planar surface of the membrane. In this context, the development of more physiologically relevant model membranes to closely mimic the bacterial outer membrane particularly for high resolution NMR studies could be valuable. Such model membranes are bound to be immobile in the NMR timescale and therefore could be utilized to understand the role of individual components on the function of antimicrobial peptides using solid state NMR experiments (73, 74). It is also worth mentioning that pardaxins have been demonstrated to form pores in the model lipid vesicles or liposome (28, 29). Such pore formation may require oligomerization of pardaxins in the lipid membranes. The LPS-bound structure of Pa4 determined here does not appear to show any detectable inter-peptide interactions as judged by the paucity of inter-peptide NOEs. Therefore, Pa4 may be active as a monomeric form in the outer membrane lipid like LPS as described for other highly active AMPs (66, 72). Recent biophysical studies had also demonstrated that oligomerization of AMPs in LPS caused a reduction of antimicrobial activities (18–21). However, self-association of
NMR Structure of Pardaxin in LPS

[Graph showing NMR spectrum and molecular model]

plausible mechanism of the outer membrane disruption. We believe that the LPS-bound structure of Pa4 and other finding in this study will be useful in the development of new antimicrobials with enhanced binding affinity for LPS (62, 75).

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