X-Ray Crystal Structures of Short Antimicrobial Peptides as *Pseudomonas aeruginosa* Lectin B Complexes

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Abstract

Herein we report X-ray crystal structures of 11-13 residue antimicrobial peptides (AMPs) active against *Pseudomonas aeruginosa* as complexes of fucosylated D-enantiomeric sequences with the *P. aeruginosa* lectin LecB. These represent the first crystal structures of short AMPs. In twenty-four individual structures of eight different peptides we found mostly α -helices assembled as two-helix or four-helix bundles with a hydrophobic core and cationic residues pointing outside. Two of the analogs formed an extended structure engaging in multiple contacts with the lectin. Molecular dynamics (MD) simulations showed that α -helices are stabilized by bundle formation and suggested that the N-terminal acyl group present in the linker to the fucosyl group can extend the helix by one additional H- bond and increase α -helix amphiphilicity. Investigating N-terminal acylation led to AMPs with equivalent and partly stronger antibacterial effects compared to the free peptide.

Introduction

The emergence of multidrug resistant (MDR) Gram-negative bacteria such as Pseudomonas aeruginosa and Acinetobacter baumannii in the hospital environment is calling for the development of new antibacterial agents.⁽¹⁻³⁾ Antimicrobial peptides (AMPs), which occur in microorganisms, plants and animals as components of innate immunity, offer a promising opportunity to address this challenge.⁽⁴⁻⁶⁾ Many of these AMPs are short peptides which fold into amphiphilic α -helices in contact with the bacterial membrane, (7-14) leading to aggregation inducing membrane disruption, morphological changes and cell lysis as documented by imaging studies.(15-18) Such short AMPs can exert potent effect against multidrug resistant bacteria and their biofilms, (19, 20) and show potent activities in vivo. (21)Interestingly, the experimental evidence for α -helix formation in these peptides is mostly based on circular dichroism (CD) and NMR studies.(22) Direct crystallographic evidence for α -helices in short AMPs(23) has only been reported for mastoparan MP-PJ (14 residues),(24) mellitin (26 residues), toxic to both prokaryotic and eukaryotic cells,(25) and an analog of magainin II (23 residues) with a crystal structure of the racemic form, (26) as well as for peptaibols, a special class of non-ribosomal peptides forming ion channels and which fold into an unusual 310-helix induced by amino-isobutyric acid residues.(27-29)

Herein we report X-ray crystal structures of analogs of the 13 residue peptide KYKKALKKLAKLL (**SB1**), initially reported by Hancock and coworkers to be active against *P. aeruginosa*,(*30*) in form of complexes of fucosylated D-enantiomeric sequences (*e.g.* **SB4**, Scheme 1) with *P. aeruginosa* lectin LecB.(*31*) These are the first X-ray crystal structures of short α -helical AMPs targeting Gram-negative bacteria. This report builds on our previous examples of using LecB complexes to access structural data for molecules otherwise difficult to crystallize such as oligonucleotides,(*32*) glycopeptide dendrimer biofilm inhibitors,(*33-39*) cyclic and bicyclic AMPs.(*40*, *41*)

The fucosylated peptides reported here act similarly to **SB1** by folding into amphiphilic and membrane disruptive helices. The crystal structures of their LecB complexes show that they mostly form α -helices assembled via hydrophobic contacts into two-helix or four-helix bundles, although two sequences present an unfolded extended structure in their LecB complexes. Molecular dynamics (MD) simulations show that α -helices are stabilized by bundle formation in the crystal but are not stable in pure water as single entities. They also suggest that the N-terminal acyl group added as a linker to the fucosyl group can extend the helix by one additional H-bond and thereby increase α -helix amphiphilicity. Investigating Nterminal acylation led to AMP analogs displaying equivalent and partly stronger antibacterial effects against different Gram-negative strains as well as *P. aeruginosa* biofilm inhibition compared to the free peptide.



Scheme 1. Synthesis of AMP **SB4**. Conditions: **a**. (i) Piperidine/DMF (1/4; v/v), RT, 2x 20 min, (ii) FmocAAOH, oxyma/DIC (3 eq), NMP/DCM (6/1, v/v); RT, 1.5 h. **b**. Peracetylated α -L-fucosyl-acetic acid, oxyma/DIC (5 eq), NMP/DCM (6/1, v/v), RT, overnight. **c**. MeOH/H₂O/25% NH₄OH (8:1:1, v/v/v), 12 h. **d**. TFA/TIS/H₂O (94:5:1, v/v/v), 3 h.

Results and Discussion

Synthesis of SB1 and analogs

Aiming to perform functional and structural studies we prepared the 13-mer AMP SB1 and its D-enantiomer SB2 as well as the corresponding *N*-terminally C-fucosylated derivatives SB3 and SB4 by N-terminal acylation using peracetylated α -C-fucosylacetic acid as described previously for peptide dendrimers (Scheme 1).(42) We also prepared shorter fucosylated analogs focusing on D-enantiomeric sequences because fucosylated L-sequences only generated few diffracting crystals where the peptides were not resolved. Specifically, we removed one or two residues at each end of the peptide to test the dependence of folding on sequence length (SB5-SB9). Furthermore, we synthesized the retro-inverso sequence (SB10) and its shortened analogs (SB11-SB15) to investigate folding in a modified context. Following our structural studies, we finally prepared N-terminally acylated analogs SB16-SB18 to test the influence of this modification on folding and activity (Table 1).

SB1 is a typical membrane disruptive AMP

The 13-residue AMP **SB1** is predicted to fold into an amphiphilic α -helix clustering cationic and hydrophobic side chains on opposite sides of the helix (Figure 1a). Analysis of CD spectra using Dichroweb(*43*) showed a transition from a disordered conformation in water to an α -helical conformation upon addition of up to 20% v/v trifluoroethanol (TFE), which partly mimics the reduced polarity environment of the bacterial membrane. This behavior is typical of short amphiphilic and membrane disruptive AMPs (Figure 1b).(*5*, *44*) As expected from this folding behavior, **SB1** exhibited membrane disruptive activity in vesicle leakage experiments with vesicles consisting of phosphatidyl glycerol (PG) mimicking the bacterial membrane (Figure 1c). Vesicles consisting of phosphatidyl choline (PC) similar to the membrane of eukaryotic cells were also affected (Figure 1d), although **SB1** showed negligible hemolysis (Table 1). The D-enantiomeric sequence **SB2** showed similar folding and membrane disruptive properties.

Cpd.	Sequence ^{a)}	N ^{b)}	% α-helix ^{c)}		Vesicle leakage (%) ^{d)}		MHC	MIC
-	-		0% TFE	20% TFE	PG	PC	(µg/mL) ^{e)}	(µg/mL) ^{f)}
SB1	KYKKALKKLAKLL	13	12	63	60	45	>2000	2
SB2	kykkalkklakll	13	9	69	70	30	>2000	1
SB3	*KYKKALKKLAKLL	13	n.d.	n.d.	n.d.	n.d.	n.d.	n.d.
SB4	*kykkalkklakll	13	11	68	41	10	1000	4
SB5	*ykkalkklakll	12	11	66	76	11	500	8
SB6	*kykkalkklakl	12	12	48	60	0	>2000	64
SB7	*kykkalkklak	11	11	33	6	2	>2000	>64
SB8	*kkalkklakll	11	13	53	57	3	>2000	32
SB9	*ykkalkklakl	11	11	47	70	1	>2000	64
SB10	*llkalkklakkyk	13	9	60	37	4	1000	n.d.
SB11	*llkalkklakky	12	11	64	49	10	1000	16
SB12	*lkalkklakkyk	12	9	42	25	0	>2000	64
SB13	*kalkklakkyk	11	10	29	3	0	>2000	64
SB14	*llkalkklakk	11	11	56	22	1	>2000	32
SB15	*lkalkklakky	11	11	50	61	1	>2000	32
SB16	Ac-kykkalkklakll	13	13	63	75	32	500	2
SB17	Pr-kykkalkklakll	13	15	68	71	35	500	2
SB18	C ₈ -kykkalkklakll	13	14	60	31	99	16	2

Table 1. Sequences, α -helical content and activity of **SB1** and C-fucosylated analogs on *P. aeruginosa*.

a) Single letter code for amino acids, * indicates a C-fucosyl group as shown in Scheme 1. Ac=acetyl, Pr=propyl, C₈=octanoyl. b) Number of amino acids. c) CD spectra were recorded at 0.1 µg/mL in aq. 10 mM phosphate buffer pH 7.2 with addition of 0, 5, 10, 15 and 20 % v/v trifluoroethanol (TFE). The primary CD spectra were analyzed using DichroWeb, and the percentages of α -helical signals were extracted. The full table is shown in the SI (Table S2). The Contin-LL method and reference set 4 were used.(*43*) n.d. = not determined. d) Membrane disruption was assessed on phosphatidyl glycerol (PG) or phosphatidyl choline (PC) lipid vesicles loaded with 50 mM 5(6)-carboxyfluorescein buffer and treated with peptide solutions (45 s) and 1.2% Triton X-100 (285 s). The percentage of lysis at 10 µg/mL was determined as I_t/I_∞ where I_t is the fluorescence intensity at t = 250 s and I_∞ is the fluorescence at t = 350 s. e) Minimum hemolytic concentration (MHC, µg/mL) was measured on human red blood cells in phosphate buffered saline at room temperature for 4 hours. All experiments were done in at least two independent duplicates. f) Minimum inhibitory concentration (MIC, µg/mL) was measured in Mueller-Hinton medium on *P. aeruginosa* PAO1, after incubation for 16-20 h at 37 °C. Values were obtained from at least 2 independent duplicate experiments giving the same results.



Figure 1. Evidence for α -helical folding and membrane disruption by the 13-residue AMP **SB1**. **a**) Prediction of **SB1** with α -helix wheel. **b**) CD spectra of **SB1** in phosphate buffer pH7.2 with 0, 5, 10, 15 and 20% v/v 2,2,2-trifluoroethanol. 5(6)-Carboxyfluorescein leakage from **c**) phosphatidylglycerol (PG) and **d**) phosphatidylcholine (PC) lipid vesicles. Addition of **SB1** to a lipid vesicle solution in buffer (10 mM TRIS, 107 mM NaCl, pH 7.4) at 45 s and addition of 1.2% Triton X-100 at 285 s.

Fucosylated D-peptide SB4 is a membrane disruptive AMP

The N-terminally fucosylated D-enantiomeric sequence **SB4** and the fucosylated retro-inverso sequence **SB10** showed a folding and membrane disruptive behavior similar to the parent sequence **SB1** (Table 1, Figure S1, Supplementary Table S2 and Figure S3). α -Helical folding was significantly reduced upon sequential removal of leucine residues at the C-terminus (**SB4** \rightarrow **SB6** \rightarrow **SB7**) respectively at the N-terminus of the retro-inverso sequence (**SB10** \rightarrow **SB12** \rightarrow **SB13**), in agreement with the α -helix propensity of leucine. Furthermore,

membrane disruption as measured by vesicle leakage was strongly reduced for SB7 and SB13

lacking both terminal leucine residues and showing negligible α-helical folding with TFE. None of the fucosylated sequences showed any significant hemolysis. Remarkably, the fucosylated analog **SB4** of the parent AMP showed the same minimum inhibitory concentration (MIC) against *P. aeruginosa* PAO1 as the parent AMP, however the activity was strongly reduced in all modified sequences. Taken together, these data showed that the fucosylated AMP **SB4** behaved as a membrane disruptive AMP similarly to the parent sequence.

SB4 forms a four-helix bundle as LecB complex

Crystallization attempts with the full sequence **SB1** or its D-enantiomer **SB2**, the racemic mixture of both, or with the fucosylated L-enantiomer **SB3** in complex with LecB were unsuccessful. By contrast, we obtained good quality diffraction data of LecB complexes with the fucosylated analog **SB4** (2.01 Å, PDB 6Q86, Table S4.1 and Figure 2, and 2.01 Å, PDB 6Q79, Supplementary Table S3, Table S4.2 and Figure S5.2). In these two structures, **SB4** peptides are bound to LecB monomers at their fucose binding sites via the N-terminal fucose and folded into fully or partially resolved amphiphilic α -helices, with two (PDB 6Q86) respectively four (PDB 6Q79) non-equivalent **SB4** peptides in the asymmetric unit. In both structures, there are no contacts between the peptide portion of the ligands and the protein, suggesting that folding occurs independently of the lectin. The two respectively four non-equivalent α -helices form four-helix bundles in which hydrophobic residues are packed on the inside and cationic, hydrophilic residues face outwards. In the first structure (PDB 6Q86), the four peptides forming the bundle are complete. However, in the second case (PDB 6Q79), only two of the α -helices are entirely resolved and well-formed, while the other two are only

observed for the six N-terminal residues preceding the fucosyl group, which can be interpreted in terms of a partially folded α -helix with a disordered C-terminus.



Figure 2. Details of the first X-ray structure of the **SB4**·LecB complex (PBD 6Q86) (**a-b**) The two nonequivalent peptides with electron density map as blue mesh, Ca^{2+} atoms shown as magenta spheres and the bound LecB monomer as green cartoon. (**c**) Charge distribution within the 4-helix bundle. Hydrophobic side chains are colored brown and cationic side chains in blue. Backbone atoms are shown as cartoon. (**d**) Overview of the 4-helix bundle in the crystal. Each of the four peptide (spheres) - LecB monomer (cartoon) complexes is independently color-coded.

SB4 analogs form helix-bundles or extended structures as LecB complexes

We next investigated shortened and retro-inverso sequences **SB5-SB15** to test if additional peptide conformations might occur in LecB complexes besides the fully and partially folded α -helices observed in the two structures of **SB4**·LecB. While we did not obtain any crystals for the least helical and inactive peptides **SB7** and **SB13**, or for **SB9** and **SB14** which also

belonged to the shortest and least helical sequences, the remaining seven fucosylated Dpeptides **SB5**, **SB6**, **SB8**, **SB10**, **SB11**, **SB12** and **SB15** yielded well diffracting crystals as complexes with LecB (Supplementary Table S3, Tables S4.3-4.10). These crystals revealed one to four different individual peptide structures each depending on crystal symmetry. In most cases the entire peptide sequence was well resolved (Supplementary Figures S5.3-5.10, Figure 3).

A detailed analysis of peptide conformations in the different structures showed that almost all residues were in an α -helical conformation as indicated by their Ramachandran angles (Supplementary Table S5). LecB complexes with **SB5**, **SB6**, **SB8**, **SB10** and **SB11** were globally comparable to the two **SB4**·LecB structures and showed α -helical peptides engaging in two-helix or four-helix bundles with a hydrophobic core and hydrophilic residues on the outside (Supplementary Figures S5.3-5.8). As observed with **SB4** the helices did not form significant contacts with LecB. In fact, for two of the four α -helices in the bundle formed by **SB10**, the fucosyl group did not even bind to the lectin and the corresponding helix was only held in place by hydrophobic contacts within the bundle (Figure 3a, b, c). On the other hand, peptides **SB12** and **SB15** were observed in an extended, β -sheet like conformation engaging in four H-bonding contacts with LecB, providing a direct picture of an alternative, unfolded conformation (Figure 3d, e, f and Supplementary Figures S5.9-5.10). Both of these sequences belonged to the retro-inverso series and showed a less complete α -helical folding in 20% TFE compared to the parent AMP, which might explain the absence of α -helical folding in the crystal (Table 1).



Figure 3. Details of the X-ray structures of α -helical AMPs in complex with LecB. (a) Electron density map of **SB10**, with Ca²⁺ atoms shown as magenta spheres and the bound LecB monomer as green cartoon. (b) Charge distribution within the 4-helix bundle formed by **SB10**. Backbone atoms are shown as cartoon. Hydrophobic side chains are colored brown and cationic side chains in blue. (c) 4-helix bundle of peptide **SB10**. Peptides are shown as spheres and bound lectin monomers are displayed as cartoon of the same color. (d) Electron density map of **SB15** as in (a). (e) Details of the four stabilizing hydrogen bonds (black dashes) engaged on a LecB monomer by peptide **SB15**. Hydrophobic side chains are colored brown and cationic side chains in blue. (f) β -sheet engagement of peptide **SB15** bound to LecB. Peptides are shown as spheres and bound lectin monomers are displayed as cartoon of the same color.

SB4 α-helices are stabilized by bundle formation in the crystal

The ten crystal structures from eight different peptides described above revealed twenty-four individual structures of short (11 to 13 residues) peptides. Assuming that all unresolved residues reflect disordered conformations, the overall α -helix content across all twenty-four resolved structures amounts to 76% on a per residue basis, which is significantly higher than the average α -helical content calculated from CD spectra showing unfolded conformations in water, which corresponds to the crystal environment (~10% α -helix content), and partially α -helical conformations in the presence of TFE (~58% α -helix content in 20% TFE). Considering that there were almost no contacts between the peptides and the protein outside the fucose binding site in these structures, we hypothesized that α -helices might be stabilized by helix bundle formation in the crystal.

To test this hypothesis, we performed MD simulations using GROMACS(45) in explicit water for 1000 ns at 300 K starting from the 4-helix bundle of peptide **SB4** as observed in the X-ray structure (PDB 6Q86), or from a single isolated α -helix. The 4-helix bundle was stable throughout the entire simulation as shown by overlays across the whole trajectory (Figure 4a), the stable number of hydrogen bonds and the low RMSD variance (Figure 4c/d, orange line). By contrast, the isolated α -helix rapidly unfolded under the same conditions and transitioned to a conformation containing a disordered N-terminus, a turn and a short intramolecular β -sheet at the C-terminus (Figure 4b and Figure 4c/d, blue line), which is consistent with CD data in water (Table 1 and Supplementary Table S2). The MD data thus clearly showed that the 4-helix bundle observed in the LecB crystal was stable on its own, and that its formation could explain the occurrence of α -helical conformations in the aqueous environment of the protein crystal despite the fact that the sequence does not fold significantly in pure water.



Figure 4. MD simulations of **SB4** as 4-helix bundle and as single peptide in water. (**a**) Overlays of 100 states over the whole 1000 ns trajectory of a **SB4** bundle. The average structure is shown as sticks, with the 100 overlayed states as thin lines. Hydrophobic side chains are colored brown and cationic side chains blue. Backbone atoms are represented as cartoon. (**b**) Overlay of the major conformations from the whole 1000 ns trajectory of **SB4** as single peptide in water. The average structure is shown as large sticks, with overlayed states as thin lines. Hydrophobic side chains are colored brown and cationic side chains blue. (**c**) Intramolecular Hydrogen bonds of a single helix within the bundle against the single peptide in water. (**d**) Root Mean Square Distances of the backbone atoms relative to the backbone atoms of the starting model.

H-bonding to the N-acyl group enhances α -helix amphiphilicity

The presence of the C-fucosyl group might affect folding by enabling an additional α -helical backbone H-bond interaction involving its linker amide carbonyl group, which is absent in the free peptide. The crystallographic data showed that this additional H-bond was indeed present in 10 out of 18 individual structures of α -helical peptides, as illustrated here with the structure of LecB·**SB6** which features both the H-bonded (Figure 5a) and the non H-bonded conformations (Figure 5b).

To test if this additional α -helical H-bond might also participate in folding in an isolated α -helix, we performed MD simulations with **SB2** and the C-fucosylated version **SB4**. We also investigated the N-acetyl (**SB16**), N-propanoyl (**SB17**) and N-octanoyl (**SB18**) derivatives, which we synthesized as new analogs of potential interest as antimicrobial peptides (Table 1). Indeed, although most AMPs have free N-termini, N-acylated AMPs can also show potent antimicrobial effects.(*7*, *46*) We ran our MD simulations in explicit water solvent with 20% v/v TFE for 1000 ns starting from a model of each peptide in a pre-folded α -helix conformation derived from the structure of **SB4** (PDB 6Q86) including the additional H-bond with the N-acyl group.

The N-fucosylated peptide **SB4** rapidly lost its acyl H-bond in the course of the simulation, leading to N-terminal conformations related to those of the free peptide **SB2**. In this conformation the N-terminal lysine side chain points to the opposite side of the α -helix relative to the cluster of cationic side chains (Figure 5c, d, f). By contrast, The N-acyl H-bond was preserved across the entire MD trajectory in **SB16** (Figure 5e, f), **SB17** and **SB18** (Supplementary Figures S6 and S7). These more stable trajectories resulted in an N-terminal conformation in which the lysine side chain at position 1 is rotated by 180° compared to **SB2** and **SB4** and is located on the same side of the α -helix as the other cationic side chains. This arrangement of the side chains corresponded to an increased α -helix ampliphilicity compared

to **SB2** and **SB4**. The conservation of the N-acyl H-bond in **SB16-SB18** also induced a more rigid conformation of the N-terminal lysine side chain compared to **SB2** and **SB4** (Figure 5g).



Figure 5. MD simulations with **SB2**, **SB4** and **SB16** in 20% TFE. The two conformations of peptide **SB6** (PDB 6Q6X) with (**a**) and without (**b**) the additional H-bond induced by the C-fucoside carbonyl moiety. Overlays of 100 states over the whole 1000 ns trajectories of **SB2** (**c**), **SB4** (**d**) and **SB16** (**e**) respectively. The average structure is shown as large sticks, with the 100 overlayed states as thin lines. Hydrophobic side chains are colored brown and cationic side chains blue. (**f**) H-bond counts. The N-terminal acetyl group is taking part in an additional H-bond compared to the fucosylated and non-fucosylated peptides. (**g**) The Root Mean Square fluctuations per atom show the high flexibility of the C-fucosyl group (cFuc) and the first lysine residue (k¹) compared to the acetylated peptide.

N-acylated analogs of SB2 are potent antimicrobial and antibiofilm agents

Our initial screening had shown that fucosylated **SB4** and the N-acylated **SB16-SB18** were as active as **SB2** in terms of membrane disruption and as AMPs against *P. aeruginosa* PAO1 (Table 1). This suggested that the N-terminal amino group in **SB2** did not contribute significantly to activity, possibly because reduction by one positive charge was compensated by an increase in α -helix amphiphilicity induced by the additional H-bond as discussed above and pointed to the possibility to use N-acylation to modulate AMP activity.

A survey of antimicrobial activity across additional pathogenic bacteria showed that N-acylated AMPs **SB4** and **SB16-SB18** were as active as **SB2** across the different strains, and even slightly more active in the case of *Klebsiella pneumoniae-S*, a pathogen which is generally difficult to control. Increasing peptide hydrophobicity by an N-terminal octanoyl group resulted in AMP **SB18** showing the strongest antimicrobial activity (Table 2), however this sequence was unselective and strongly hemolytic (Table 1). Note that the free, nonfucosylated D-peptide **SB2** was slightly more active than its L-enantiomer **SB1**, which might reflect higher stability of the D-enantiomer in the culture medium where the L-enantiomer undergoes proteolysis (Supplementary Figure S2).

Strain	SB1	SB2	SB4	SB16	SB17	SB18
PAO1	2	1	4	2	2	2
A. baumannii 19606	2	1	1	1	1	2
K. pneumoniae-S	32	16	16	8	8	4
E. coli (W3110)	1	0.5	1	1	1	2
S. Epidermiditis	n.d.	2	4	2	2	1
MRSA*	n.d.	4	8	4	4	2

Table 2. Minimal inhibitory concentration data.

a) MIC (minimum inhibitory concentration) in μ g/mL against wild type Gram-negative, Gram-positive and multidrug resistant (MDR, marked*) were measured by 1/2 serial dilutions on 96 well plates in Mueller-Hinton (MH) broth after incubation overnight at 37 °C. MIC values were measured in independent duplicates with at least two experiments giving the same value.

We further investigated inhibition of *P. aeruginosa* biofilms by **SB4** since this peptide binds lectin LecB, and related multivalent(*33*, *39*, *47*) and monovalent LecB ligands(*48-50*) have been shown to inhibit biofilms from this bacterium. Indeed, **SB4** inhibited *P. aeruginosa* biofilm formation, however the effect was comparable to the free D-enantiomeric peptide **SB2** (Figure 6). The L-enantiomer **SB1** was again significantly less active against biofilms, probably due to proteolysis in the medium. We propose that the observed biofilm inhibition effect by **SB2** and **SB4** primarily results from a direct antibacterial effect mediated by membrane disruption. This hypothesis is supported by vesicle leakage experiments showing that both peptides induce significant membrane perturbations (Table 1).



Figure 6. Relative absorbance readouts for Biofilm Inhibition on *Pseudomonas aeruginosa* PAO1 biofilm in M63 medium compared to untreated biofilms. Compound-containing wells were inoculated with the bacteria and the biofilms were allowed to grow 24-25 hours at 37° under static conditions. After washing, the biofilms were revealed with 0.5 mM WST-8 and 20 μ M phenazine ethosulfate for 3 hours and the absorbance measured. The experiments were performed as triplicates.

The membrane perturbation effects induced by **SB2** and **SB4** were further investigated by transmission electron microscopy (TEM). Images of *P. aeruginosa*, *E. coli* and *A. baumannii* cells showed that **SB4** and **SB2** induced similar morphological changes (Figure 7). After

treatment of PAO1 and *E. coli* with the peptides, vesicle-like structures and blebs were observed around the cells (blue arrows), consistent with a membrane disruptive mechanism. Peptides **SB2** and **SB4** also similarly affected the pili of *A. baumannii* (black arrows), one of its key virulence factors. In PAO1 and *A. baumannii*, dense aggregates were formed within the bacteria (orange arrows), which can be explained by peptide accumulation inside the cell and possibly aggregation with DNA.



Figure 7. TEM images on *Pseudomonas aeruginosa* PAO1, *E. coli* and *A. baumannii* treated with 40 µg/mL **SB2** and 40 µg/mL **SB4** after 60 min. Arrows indicate effects of the peptides on bacterial membranes (blue arrows: vesicle like structures and blebs), cell contents (orange arrows: dense aggregates), and pili (black arrows: pili well present in control but less so in peptide treated cells). See Figure S4 for enlarged versions of the images.

Conclusion

Here we reported the first X-ray crystal structures of short α -helical AMPs at the example of analogs of the 13-residue AMP **SB1**. The structures were obtained in form of complexes of the fucosylated D-enantiomeric sequence **SB4** with the bacterial lectin LecB. We also investigated shortened and retro-inverso sequences, resulting in a total of twenty-four

individual structures of eight different peptides. These structures show that these short AMPs form α -helices which assemble as two-helix or four-helix bundles displaying hydrophobic residues on the inside and cationic residues on the outside, without significant interaction with the lectin except for the anchoring fucosyl group bound at the fucose binding site. In two cases the peptide is observed in an extended β -sheet like conformation making four H-bonding interaction with residues on the lectin surface. Overall 76% of the residues are observed within an α -helix, while the remaining residues are either not visible or in a different conformation. Molecular dynamics (MD) simulations show that α -helices are stabilized by bundle formation and suggest that the N-terminal acyl group present in the linker to the fucosyl group can extend the helix by one additional H-bond and increase α -helix amphiphilicity. Indeed, several N-acylated AMP displayed equivalent or stronger antibacterial effects compared to the free peptide.

Methods

Procedures for peptide synthesis, CD-spectroscopy, vesicle leakage experiments, microbiology, crystallization, X-ray data collection and refinement statistics, molecular dynamics, transmission electron microscopy, and analytical data for all peptides, are described in the supporting information.

Associated Content

Supporting Information. The supporting information is available free of charge on the ACS Publication website. Synthesis and characterization of all peptides, X-ray data collection tables, detailed procedures for CD spectroscopy, vesicle leakage experiments, molecular dynamics, microbiology (MIC and biofilm assays) and transition electron microscopy.

Accession Codes. Coordinates and structure factors of refined LecB complexed with fucosylated peptides are available from the Protein Data Bank with accession codes 6Q86 and 6Q79 (SB4), 6Q6W (SB5), 6Q6X (SB6), 6Q8G (SB8), 6Q8H and 6Q87 (SB10), 6Q85 (SB11), 6Q77 (SB12) and 6Q8D (SB15).

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Author Contributions. SB synthesized peptides, performed X-ray crystallography and molecular dynamics, interpreted data and wrote the paper. BHG performed TEM studies and interpreted data. TNS performed vesicle leakage and antimicrobial activity assays. RV performed X-ray crystallography. MR synthesized peptides. SJ designed and supervised MD studies, interpreted data and wrote the paper. TD and JLR designed and supervised the study, interpreted data and wrote the paper.

Notes. The authors declare no competing financial interest.

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