

Cite this: *Chem. Sci.*, 2018, 9, 8850

All publication charges for this article have been paid for by the Royal Society of Chemistry

Structural studies suggest aggregation as one of the modes of action for teixobactin†

Carl Öster,^a Grzegorz P. Walkowiak,^{ab} Dallas E. Hughes,^c Amy L. Spoering,^c Aaron J. Peoples,^c Anita C. Catherwood,^b Julie A. Tod,^b Adrian J. Lloyd,^b Torsten Herrmann,^d Kim Lewis,^e Christopher G. Dowson^b and Józef R. Lewandowski^{*,a}

Teixobactin is a new promising antibiotic that targets cell wall biosynthesis by binding to lipid II and has no detectable resistance thanks to its unique but yet not fully understood mechanism of operation. To aid in the structure-based design of teixobactin analogues with improved pharmacological properties, we present a 3D structure of native teixobactin in membrane mimetics and characterise its binding to lipid II through a combination of solution NMR and fast (90 kHz) magic angle spinning solid state NMR. In NMR titrations, we observe a pattern strongly suggesting interactions between the backbone of the C-terminal "cage" and the pyrophosphate moiety in lipid II. We find that the N-terminal part of teixobactin does not only act as a membrane anchor, as previously thought, but is actively involved in binding. Moreover, teixobactin forms a well-structured and specific complex with lipid II, where the N-terminal part of teixobactin assumes a β conformation that is highly prone to aggregation, which likely contributes to the antibiotic's high bactericidal efficiency. Overall, our study provides several new clues to teixobactin's modes of action.

Received 16th August 2018
Accepted 19th September 2018

DOI: 10.1039/c8sc03655a

rsc.li/chemical-science

Introduction

The majority of antibiotics currently in clinical use have been discovered by screening cultivable soil bacteria. However, around 99% of microorganisms are uncultured, meaning that they do not grow under laboratory conditions. Recently, several methods to access potential antimicrobial compounds from uncultured microorganisms were developed,^{1–4} of which the iChip technology led to the discovery of teixobactin from a new species of β -proteobacteria, *Eleftheria terrae*.⁵ Teixobactin was shown to have very good activity against many difficult-to-treat bacterial pathogens, including *Mycobacterium tuberculosis*, and no resistant mutants were obtained *in vitro* with various bacteria. Teixobactin binds to both lipid II and lipid III and thus simultaneously inhibits the biosynthesis of peptidoglycan and teichoic acids, triggering synergistic effects leading to increased cell wall damage and delocalization of autolysins.^{5,6}

Additionally, teixobactin, in contrast to vancomycin, does not bind mature peptidoglycan, which enables the efficient killing of bacteria with increased cell wall density such as vancomycin-intermediate *Staphylococcus aureus* (VISA) against which vancomycin is ineffective.⁶ The biosyntheses of both peptidoglycan and teichoic acids rely on a common lipid carrier, bactoprenol, C₅₅-P, which is linked in both cases to a disaccharide unit *via* a pyrophosphate bridge. In this work we focus on the peptidoglycan precursor lipid II, which, apart from the lipid carrier and the pyrophosphate, contains the main building blocks of peptidoglycan; *N*-acetylmuramic acid, *N*-acetylglucosamine and a pentapeptide (L-Ala-D-Glu-L-Lys/DAP-D-Ala-D-Ala; see Fig. 1d). Lipid II biosynthesis varies, typically within the pentapeptide at position three between Gram-positive bacteria possessing lysine and Gram-negative bacteria diaminopimelic acid (DAP). *Mycobacteria* spp. possess DAP and can additionally modify *N*-acetyl- to *N*-glycolyl-muramic acid.⁷ Several other modifications to the peptide stem or glycan strands have been discovered and are discussed elsewhere.^{8,9} Teixobactin is expected to interact with the pyrophosphate of lipid II and it can thus bind both Gram-positive and Gram-negative variants of lipid II, regardless of any modifications to the sugars or peptides.⁵ Since the discovery of teixobactin, its biosynthetic pathway and the identification of lipid II as a primary target, a large number of studies have been conducted to gain more understanding on the mode of action of teixobactin, with the goal of developing analogues with better pharmacological properties. Several groups have synthesized

^aDepartment of Chemistry, University of Warwick, Coventry, CV4 7AL, UK. E-mail: j.r.lewandowski@warwick.ac.uk

^bSchool of Life Sciences, University of Warwick, Coventry, CV4 7AL, UK

^cNovoBiotic Pharmaceuticals, Cambridge, MA 02138, USA

^dUniv. Grenoble Alpes, CNRS, CEA, IBS, F-38000 Grenoble, France

^eAntimicrobial Discovery Center, Northeastern University, Department of Biology, Boston, MA 02115, USA

† Electronic supplementary information (ESI) available: Additional figures. NMR titration data. FRET assays. NMR spectra. NMR relaxation data. PDB of the structure of teixobactin in DPC micelles. See DOI: 10.1039/c8sc03655a



teixobactin analogues to investigate the roles of the different residues in teixobactin and potentially find active compounds that are easier to make and better suited for clinical use (see Fig. 1c for a summary of the effect on antimicrobial activity by substitution of amino acids). An NMR study of seven analogues showed the importance of the D-amino acids for activity.¹⁰ The residue in position 10, L-allo-enduracididine, is a non-proteinogenic amino acid, which has been difficult to synthesise and hence replacing it with a naturally available amino acid is an attractive approach. Arg₁₀teixobactin^{11,12} and Lys₁₀teixobactin¹² showed good activity and recently several other teixobactin analogues with different alternatives in position 10 were synthesised and found to have good activity against methicillin-resistant *Staphylococcus aureus* (MRSA), *Staphylococcus epidermidis* and *Bacillus subtilis*.¹³ Structural details of teixobactin analogues in isolation and some initial studies involving teixobactin analogues and mimics of lipid II in dimethyl sulfoxide (DMSO) and molecular dynamics simulations exploring the binding space were reported in the literature.^{10,13–15,19} However, to date no structure of wild type teixobactin and no high resolution structural information on the interactions with actual lipid II have been reported. Here we use a combination of solution and fast magic angle spinning (MAS) solid-state NMR complemented by a fluorescence based assay to obtain insights on the interactions of wild type teixobactin with both Gram-negative and Gram-positive lipid II and conformational changes of teixobactin induced by binding to lipid II. We find that the majority of the sites interacting with lipid II are located on the backbone of teixobactin and that teixobactin assumes an aggregation prone β -strand conformation for residues 2–6 upon binding lipid II. To mimic the natural membrane environment of lipid II we used dodecylphosphocholine (DPC) micelles but we also consider interactions of teixobactin with a water soluble lipid II in aqueous solution. Our data provide a rationale for understanding the behaviour of numerous analogues of teixobactin and valuable insights that should further guide efforts to engineer teixobactin analogues.

Results

Teixobactin in solution

Since teixobactin binds lipid II in the cytoplasmic membrane of bacteria, we investigated its conformation in a simulated membrane using DPC micelles. The 20 lowest energy structures of teixobactin in DPC micelles in phosphate buffer determined by solution NMR are shown in Fig. 1a (see ESI Table 1† for details). The structures are overlaid with a crystal structure of the teixobactin analogue Ac- Δ_{1-5} Arg₁₀-teixobactin as a hydrochloride salt¹⁴ (Fig. 1b). The proposed pyrophosphate binding cage formed by residues 8–11 is nearly identical in the NMR and crystal structures. The most obvious differences between the structures are that the bond between Ser₇C' and D-Thr₈N points in different directions and the side-chain of residue 10, where in the analogue the allo-enduracididine has been replaced by arginine. Additionally, the N-termini are different in the

structures since residues 1–5 were replaced by an acetyl group in the crystal structure construct.¹⁴

Structures of several analogues including teixobactin-Arg₁₀ were also solved in DMSO by solution NMR.¹⁰ In the structure of teixobactin-Arg₁₀ and in contrast to the structure of native teixobactin in DPC micelles, presented here, residues 1–7 are largely unstructured. This discrepancy is likely due to DMSO being used as a solvent for the teixobactin analogues, as DMSO is known to destabilise the secondary structure and promote disorder.²⁰ Since, similar to other lipid II binding antibiotics,²¹ the solvent environment appears to have a potentially large effect on the structure of teixobactin, we have also performed solution NMR experiments on teixobactin in a phosphate buffer in the absence of micelles. A comparison of the chemical shifts in the two media (*i.e.* aqueous solution in the presence and absence of DPC micelles) suggests that no large conformation change (except a slight change near residues 1 & 6) occurs between them with most changes being attributable to hydrophobic interactions between teixobactin and DPC micelles (with residues 2, 6, 7 and 11 being most likely to be involved in or influenced by binding to the micelle; see Fig. ESI1†).

It has been previously suggested that teixobactin in aqueous solution binds lipid II in a 2 : 1 ratio,⁵ which could be potentially explained if teixobactin dimerizes under these conditions. To investigate the oligomeric state of teixobactin we measured ¹⁵N NMR relaxation in aqueous solution (see Fig. ESI3, Table ESI2†). The correlation time (τ_c) for the overall tumbling and subsequently the approximate size of the molecule were estimated from ¹⁵N relaxation times. The relaxation measurements indicate that teixobactin is not monomeric in aqueous solution. The correlation time calculated from the relaxation times corresponds much better to a dimer or a trimer than to a monomer (see Fig. ESI3, Table ESI3†). The relaxation rates are also consistent with considerable local flexibility of the N-terminal residues (order parameter, S^2 , of 0.4–0.6). Since the N-terminus appears quite mobile we have also considered whether the increased apparent hydrodynamic radius of teixobactin in solution could be potentially explained by a monomeric teixobactin where the molecule assumes a more extended conformation as observed for some teixobactin analogues in DMSO (though minimal changes in chemical shifts between teixobactin in DPC micelles and in aqueous solution do not support such interpretation). However, all of the extended monomers we generated resulted in underestimated values for τ_c (see Fig. ESI3, Table ESI3†). Because DPC micelles dominate the tumbling a similar method cannot be used to estimate the oligomeric state of teixobactin in DPC micelles (see Fig. ESI4 and Table ESI2† for relaxation data). Due to the lack of any plausible intermolecular NOEs, teixobactin in DPC micelles is assumed to be monomeric for structure calculations, which is a general assumption made for peptides in DPC micelles.

Teixobactin–lipid II interactions

Whereas teixobactin did not exhibit any particular propensity for aggregation in DPC micelles, we found that addition of lipid



II resulted in rapid aggregation with peaks disappearing in the solution NMR spectra due to the large size of the aggregates. We observed the same behaviour when a water soluble variant of lipid II, C₁₅ lipid II as opposed to the native C₅₅ lipid II, was added to teixobactin in aqueous solution in the absence of DPC micelles. This suggests that the aggregation is induced by interactions between teixobactin and lipid II and does not require a membrane environment. Previously, adding various amounts of DMSO was demonstrated to facilitate obtaining soluble antibiotic–lipid II complexes and to prevent their aggregation and precipitation.^{22,23} However, we observe strong aggregation in the teixobactin–lipid II complex even in solutions containing 80% DMSO. Since in our case DMSO does not alleviate the aggregation and because of its high potential for artificially introducing disorder and suppressing potentially important interactions, we have avoided it in further experiments.

To gain insights into the aggregation process we performed a Fluorescence Resonance Energy Transfer (FRET) assay on a series of lipid II binders. Among several tested antibiotics including teixobactin, ramoplanin, deoxyactagardin B, mersacidin and vancomycin, only ramoplanin and teixobactin promoted aggregation of lipid II molecules (see Fig. ESI5†).

In order to determine whether the aggregation occurs due to nonspecific or specific interactions we have also sedimented the soluble aggregates formed between teixobactin and lipid II in DPC micelles using ultracentrifugation and performed fast magic angle spinning (MAS) solid-state NMR experiments. We observed only one set of resonances in the solid state NMR spectra, consistent with the presence of a single dominant conformer in the sedimented complex (see Fig. 2e and ESI6, 7†). The narrow, under the applied conditions, linewidths (at 90 kHz spinning) ¹H 0.2–0.4 ppm (120–240 Hz), ¹³C 0.4–0.6 ppm (60–90 Hz) and ¹⁵N ~0.7 ppm (45 Hz) indicate unambiguously that the aggregates are homogeneous in nature and are driven by specific rather than nonspecific interactions.²⁴ Note that even moderate local disorder would be reflected in substantial broadening.

Interestingly, in a study by Parmar *et al.*¹⁵ no aggregation was observed for teixobactin analogues in DMSO in the presence of geranyl pyrophosphate, a lipid II mimic possessing a pyrophosphate and C₁₀ isoprenyl chain but no sugar moieties or the peptide stem. This result combined with our observations suggests that the interaction with the sugar moieties of lipid II may play a role in the aggregation process for the teixobactin : lipid II complex.

To obtain a site-specific view of the effect of the binding of teixobactin to lipid II we have performed NMR titrations in solution, where changes in teixobactin spectra were followed as a function of lipid II concentration. Upon addition of lipid II we did not observe changes in chemical shifts for any peaks but rather a decrease in intensities of the peaks, which is behaviour characteristic of slow exchange with dissociation constants, K_d , in the low μ M to nM range. To determine binding affinities in the presence of slow exchange, typically, one follows the build-up of peak intensities of the bound form.²⁵ This was not possible in the case of lipid II titrations because the resonances

for the bound form of teixobactin were broadened beyond detection due to aggregation. Instead, we fitted intensities of the peaks from free teixobactin as a function of lipid II concentration to extract apparent association constants²⁶ (K_a is the inverse of K_d ; see eqn (1) in methods, see Fig. ESI8–11, Tables ESI4, 5†). It should be noted that the K_a values obtained by these fits may be higher than the actual K_a values due to the effect aggregation has on the NMR spectra. However, even if the calculated K_a values may not be quantitative, they yield qualitative information on which sites of teixobactin are the most affected by the binding of lipid II either due to interaction or conformational change upon interaction. We performed the experiments using both native Gram-negative lipid II (diaminopimelic acid in position 3 of the peptide stem, see Fig. 1d) with the full C₅₅ lipid tail in DPC micelles (in the following we refer to this sample as TXB_{DPC}; see Fig. 2e) and a water soluble version of Gram-positive lipid II (lysine in position 3 of the peptide stem, see Fig. 1d) with a C₁₅ lipid tail in aqueous solution (in the following we refer to this sample as TXB_{aqueous}; see Fig. 2c). To facilitate the analysis, in Fig. 2b–e we have highlighted in light blue all the sites for which the apparent K_a is larger than the average apparent K_a and in dark blue all the sites for which apparent K_a is larger than the average apparent K_a plus one standard deviation.

The apparent K_a values for TXB_{aqueous} are approximately one order of magnitude larger than in TXB_{DPC}, which might indicate that the peptide stem of lipid II plays a role in the binding or that the addition of DPC has an effect on the binding strength. However, due to the above-mentioned ambiguity of the K_a values obtained from fitting of disappearing peaks we limit ourselves to qualitative comparisons only.

Although we see an overall difference between the binding strength in TXB_{DPC} and in TXB_{aqueous}, the binding mode seems to be very similar in both cases. The strongest effect appears in the same parts of teixobactin in both media. The majority of the sites strongly affected by the titration of lipid II are backbone atoms. In fact, there are backbone atoms showing strong affinity in all residues except 2 and 3; for residue 2 data are missing for N-H (TXB_{aqueous}) and C^α-H^α (TXB_{aqueous} and TXB_{DPC}), for residue 3 N-H and C^α-H^α show apparent K_a above average (light blue in Fig. 2a–d) but not significantly above average (dark blue in Fig. 2b–e) for TXB_{DPC}. For the side chains there are no strongly affected sites in residues 2–5, the most strongly affected side-chains include: *N*-Me-D-Phe₁C^{Me}, *N*-Me-D-Phe₁C^β (TXB_{aqueous} only), *L*-Ile₆C^β, *L*-Ser₇C^β (TXB_{DPC} only, above average K_a for TXB_{aqueous}), *D*-Thr₈C^β, *D*-Thr₈C^{γ2}, *L*-End₁₀C^β, *L*-End₁₀C^δ (TXB_{aqueous} only, above average apparent K_a for TXB_{DPC}), *L*-Ile₁₁C^{γ2} (TXB_{DPC} only), and *L*-Ile₁₁C^δ (TXB_{DPC} only). Most sites that show strong apparent affinity (apparent K_a above average plus standard deviation) in either TXB_{aqueous} or TXB_{DPC} show at least apparent affinity above average in the other. The only exceptions are the *N*-Me-D-Phe₁C^β (strong apparent affinity in TXB_{aqueous} only) and the side chains of residue 11, where C^{γ2} and C^δ show strong affinity in TXB_{DPC} but not in TXB_{aqueous}. In the backbone of *L*-Ile₄ C^α shows strong affinity in TXB_{aqueous} but not in TXB_{DPC} and the NH shows strong affinity in TXB_{DPC} but not in TXB_{aqueous}. The deviations for the C^β of *N*-Me-D-Phe₁



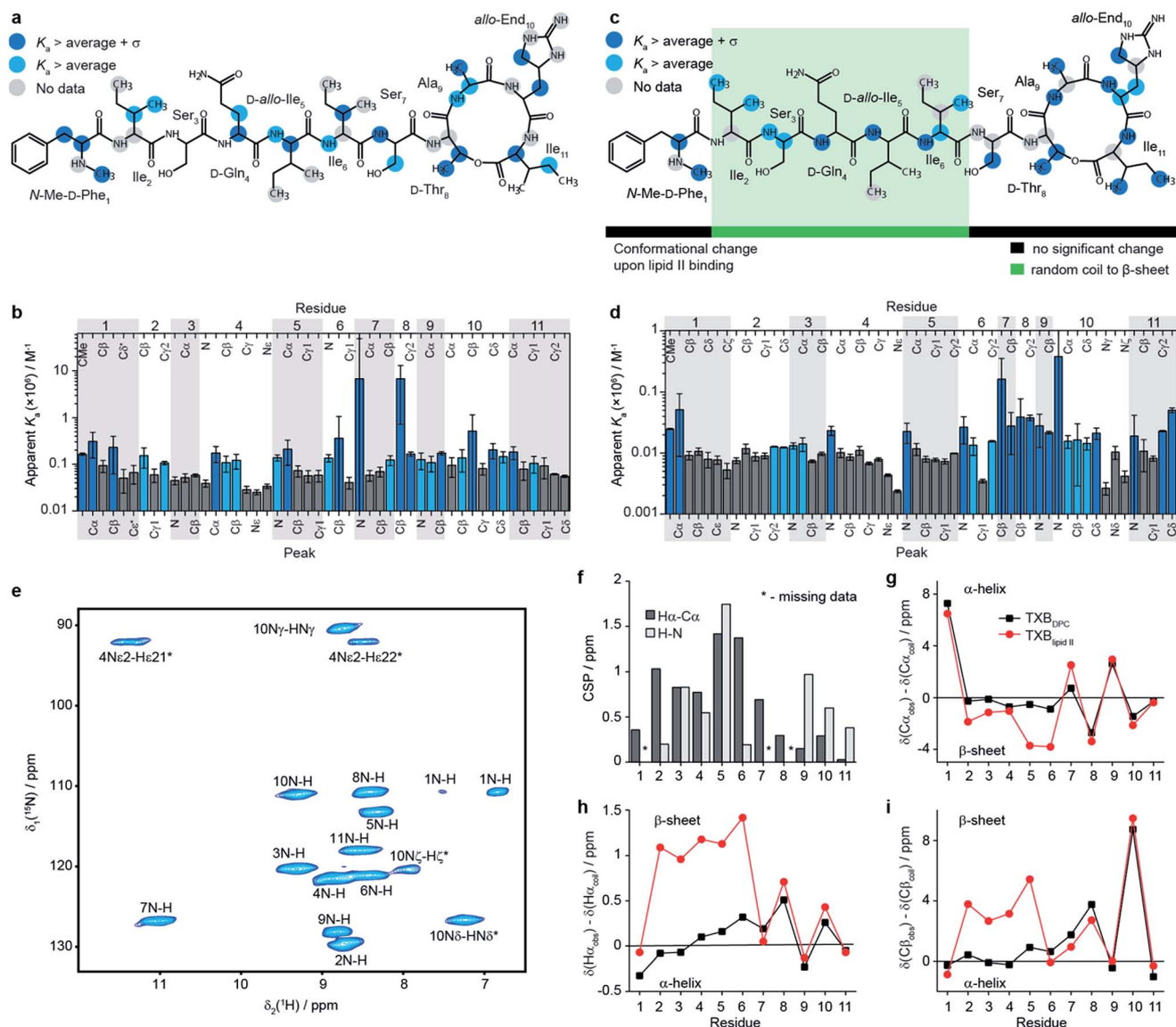


Fig. 2 Characterisation of the teixobactin–lipid II complex. (a–d) Results of solution NMR titrations of teixobactin with a water soluble C_{15} variant of Gram-positive lipid II in aqueous solution (a, b) and with native C_{55} Gram-negative lipid II in DPC micelles (c, d). Panels b and d show the measured apparent K_a values and plotted onto the chemical structure of teixobactin in a and c. Dark blue indicates sites with apparent K_a values higher than the average plus one standard deviation and light blue indicates sites with apparent K_a values higher than the average. Dark grey indicates sites with a below average apparent K_a (b and d). In a and c grey circles indicate sites for which data are not available. (e) 2D 1H - ^{15}N solid state NMR correlation spectrum of sedimented [$U^{13}C$ - ^{15}N]teixobactin in complex with natural abundance Gram-negative lipid II in DPC micelles acquired at 600 MHz 1H Larmor frequency and 90 kHz magic angle spinning frequency. In panel e star indicates peaks folded in the ^{15}N dimension. The two resonances for residue 1 are due to the zwitterionic form of *N*-Me-D-Phe₁ being the major form at pH 6.5. (f) 1H - ^{15}N (light grey) and 1H - ^{13}C (dark grey) chemical shift perturbations between TXB_{DPC} and TXB_{lipid II}. Comparisons of secondary $^{13}C\alpha$ (g), $^1H\alpha$ (h) and $^{13}C\beta$ (i) chemical shifts between TXB_{DPC} (black squares) and TXB_{lipid II} (red circles).

and for the side-chains of *L*-Ile₁₁ can possibly be explained by hydrophobic interactions with the DPC micelles which are affected due to conformational changes occurring in teixobactin when lipid II is added. The phenyl group of *N*-Me-D-Phe₁ has previously been suggested to interact through hydrophobic interactions with the membrane and the lipid tail of lipid II.²⁷ The deviations for C^α and NH for residue 4 could possibly indicate slightly different backbone conformations in the two samples: NH of residue 4 is among those peaks showing largest chemical shift changes between TXB_{aqueous}

and TXB_{DPC} (see Fig. ESI1[†]). It is also possible that the different peptide stems between the Gram-negative lipid II used in the titrations to TXB_{DPC} and the Gram-positive lipid II used in the titrations to TXB_{aqueous} have some effect on which sites are mostly affected but the available data are inconclusive in this respect.

In our titrations we see a strong effect on the backbone for residues 4–6, which is not seen in the titration of the teixobactin-Ala₁₀ analogue with geranyl pyrophosphate¹⁵ (lacking the sugars and peptide chain compared to the native lipid II).



Absence of indication of any specific interaction with geranyl pyrophosphate for these residues suggests that they might be involved in interactions with sugar moieties and possibly parts of the peptide stem in native lipid II.

As mentioned previously due to aggregation we could not detect the bound form of teixobactin by solution NMR but we could study it by fast MAS solid state NMR, which is not limited by the large size of the aggregates.^{24,28–30} Fig. 2e shows a cross-polarization (CP) based ^1H - ^{15}N correlation spectrum of [^{13}C , ^{15}N]teixobactin in a sedimented complex with unlabelled Gram-negative lipid II in the presence of DPC (in the following we refer to this sample as $\text{TXB}_{\text{lipid II}}$). The high quality spectra obtained by solid state NMR allowed for complete backbone assignments of $\text{TXB}_{\text{lipid II}}$ and almost complete side-chain assignments (the aromatic group of residue 1 was not completely assigned).

To gain some insights into the intermolecular interactions and conformation of teixobactin in the complex we calculated chemical shift perturbations (CSPs, eqn (4) in methods) between TXB_{DPC} (solution) and $\text{TXB}_{\text{lipid II}}$ (solid; see Fig. 2f) and secondary chemical shifts (Fig. 2g–i). The ^1H - ^{15}N CSPs are sensitive to both binding and conformational changes and $^1\text{H}^\alpha$ - $^{13}\text{C}^\alpha$ CSPs are mostly sensitive to conformational changes. The large ^1H - ^{15}N CSPs for residues 3–5 and 9–10 confirm the view available from the titrations in solution that the binding of teixobactin to lipid II extends beyond the pyrophosphate binding cage formed by residues 8–11. The $^1\text{H}^\alpha$ - $^{13}\text{C}^\alpha$ CSPs suggest that notable conformational changes take place for residues 2–7 with the conformation of residues 8–11 being very similar to what we observe in solution for isolated teixobactin. The conformational changes are more evident from the secondary chemical shifts, which report on secondary structure deviations from random coil. Combined analysis of $^{13}\text{C}^\alpha$, $^1\text{H}^\alpha$ and $^{13}\text{C}^\beta$ secondary chemical shifts suggests that residues 2–6 rearrange from a random coil like conformation to a more extended β -strand like conformation upon binding to lipid II. Again, the back-bone conformation for residues 8–11 appears largely unchanged upon complex formation.

Discussion

The charged side chain of the L-allo-enduracididine amino acid at position 10 was initially thought to be important for the antimicrobial activity and hence most of the initial work focused on analogues of teixobactin with positively charged side chains: $\text{End}_{10}\text{Arg}$ and $\text{End}_{10}\text{Lys}$ variants both displayed antimicrobial activity but not as good as the wild type teixobactin.^{12,13} In the X-ray structure of $\text{Ac-}\Delta_{1-5}\text{Arg}_{10}$ -teixobactin analogue the guanidinium side chain of Arg_{10} formed a hydrogen bond to the chloride ion¹⁴ (mimicking the pyrophosphate of lipid II). Interestingly, according to our titrations there is no strong interaction between the side chain nitrogens of residue 10 and lipid II. However if we look at chemical shift changes between TXB_{DPC} and $\text{TXB}_{\text{lipid II}}$ the CSPs are large for the side-chain NH groups; 0.44, 1.36 and 0.74 ppm for N^γ , N^δ and N^ϵ respectively, which, compared to the CSPs for the backbone amides, would suggest that N^δ is one of the most

affected nitrogen sites. The side-chains of D-Gln_4 show even larger chemical shift changes (the chemical shift in TXB_{DPC} is 112 ppm for the $\text{N}^{\epsilon 2}$ and 41.1 ppm in $\text{TXB}_{\text{lipid II}}$ and the ^1H chemical shifts for $\text{H}^{\epsilon 21}$ and $\text{H}^{\epsilon 22}$ are 6.8 and 7.5 ppm in TXB_{DPC} and 8.5 and 11.4 ppm in $\text{TXB}_{\text{lipid II}}$). The large chemical shift changes for side-chain nitrogen can be explained by conformational changes due to interactions with the carbon side-chains in the case of End_{10} . The unusual chemical shift for D-Gln_4 side-chain nitrogen suggests interactions with charged moieties, e.g. D-Glu in the pentapeptide of lipid II: the only sites with similar chemical shifts for the systems deposited in BMRB occur for glutamines in close contact with charged, mostly negatively charged, side chains. The fact that a charged side chain for residue 10 is not required was recently demonstrated by creating teixobactin analogues where L-allo- End_{10} was replaced by residues with hydrophobic side chains that exhibited antimicrobial activity^{15,16} even at the level matching the activity of wild type teixobactin.¹⁵ Interestingly, the best performing analogues are the ones where the side chain can assume a similar conformation to the one found for interacting sites in our titrations, $\text{End}_{10}\text{Ile}$ and $\text{End}_{10}\text{Leu}$,¹⁵ suggesting that for the side chain of residue 10 the shape is more important than the charge or hydrophobicity. The other residues of teixobactin whose replacement in analogues did not lead to an overly detrimental effect on the antimicrobial activity include Ser_3 and Ala_9 . Residue 3 did not show any significant interaction according to our titrations but backbone conformational changes of residue 3 appear important for the activity. According to our titrations the side-chain of residue 9 is strongly involved in the binding to lipid II, however it has been shown that the alanine in position 9 can be exchanged to a lysine or arginine without significant loss of antimicrobial effect¹⁴ (see Fig. 1c). This would suggest that the backbone of residue 9 is involved in the interactions but it is not important that the side-chain is hydrophobic. The N-terminus of teixobactin has been suggested to function primarily as an anchoring point to the cell membrane, however our data suggest that it could have a more involved role in the bactericidal mechanism of teixobactin by promoting aggregation, which may explain the lower activity seen in analogues where residues 1–5 were replaced by a dodecanoyl group¹⁴ and why substitution of residues 1, 2, 5 and 6 is not well tolerated.¹⁶ Our data suggest that most of the sites except the aromatic ring in the first residue are involved in the interactions with lipid II both in the presence and absence of DPC micelles. The removal of the methyl group interacting with lipid II leads only to a modest reduction of the antibacterial activity¹⁷ but loss of the positive charge or replacement of the methyl group with a bulkier hydrophobic group has much greater negative impact.³¹ In the context of our study this could suggest that bulky substituents interfere with the interaction with lipid II. The removal of the aromatic ring, that is not interacting with lipid II according to our data, leads to loss of antibacterial activity altogether but substitution with larger hydrophobic groups has a positive effect.^{16,27} That larger hydrophobic groups do not decrease the activity is consistent with the side chain of residue 1 not being involved in binding to lipid II but rather contributing to anchoring to the membrane.



On the other hand, the aromatic ring could also aid the aggregation process. The general chemical shift changes indicative of β -strand formation for residues 2–6 and the assembly of large soluble aggregates upon interaction of teixobactin and lipid II suggest that these residues may aid in the aggregation process (exposed β -strands are involved in an important mode for protein–protein interactions linked to aggregation³²), perhaps by fibril formation, as was previously reported for the ramoplanin : lipid II complex²² or the type of aggregation observed for the nisin : lipid II complex^{33,34} (though the precise nature of this specific aggregate is unclear and requires further studies). This highlights that flexibility of the N-terminal part of teixobactin is important for its biological activity, as has been suggested based on studies of several analogues of teixobactin involving substitutions of amino acids in this part.¹⁰ However, our data suggest that the conclusion of the mentioned study should be slightly modified: it is not the disorder that is the key (the N-terminus is mobile but well-ordered in our structure of teixobactin in DPC micelles) but rather the range of accessible conformations promoting intermolecular rather than intramolecular β -sheet formation.

Aggregation makes it difficult to study structural aspects of the teixobactin : lipid II complex using conventional methods such as X-ray crystallography and solution NMR. However, we have shown that ultra-fast MAS solid state NMR can be used to gain valuable site-specific information in this system. More investigations into the teixobactin : lipid II complex using isotope labelled lipid II are required to resolve ambiguities in intermolecular cross-peak assignments and to gain further understanding into the structural arrangement of these large aggregates.

Conclusions

The 3D structure of teixobactin in DPC micelles confirms the presence of a C-terminal pyrophosphate binding cage previously suggested in truncated teixobactin analogues. The N-terminal part of the peptide is well-structured in DPC micelles, in contrast to structures of teixobactin analogues in DMSO where it appears mostly disordered. A combination of titrations in solution NMR and solid state NMR experiments reveals that most of the teixobactin is involved in binding lipid II, either *via* interactions or conformational changes upon binding. The pattern of the strongest affected residues is overall very similar in both titrations using water soluble Gram-positive lipid II in the absence of DPC micelles and native Gram-negative lipid II in the presence of DPC micelles, with mostly backbone sites being affected. Residues 2–6 undergo a change from coil to β conformation upon binding of lipid II. We suggest that residues 2–6 are important for the activity due to their involvement in aggregation of teixobactin–lipid II complexes, which in the cell would lead to accumulation of lipid II and inhibition of cell wall synthesis. Importantly, using solid state NMR we have for the first time been able to study the full complex formed between native teixobactin and native lipid II, something that would not be possible with other biophysical techniques such as solution NMR and X-ray crystallography. This study paves the

way for more detailed structural analysis of aggregating antibiotics–substrate complexes.

Experimental

Sample preparation

Lipid II was synthesized as described previously.^{34,35} The biotinylated variant of lipid II used in the FRET experiments was obtained by biotinylation of the *N*-acetylmuramyl-pentapeptide precursor prior to the lipid II synthesis.

Natural abundance teixobactin and [U - ^{13}C , ^{15}N]teixobactin were produced the same way by growing *Eleftheria terrae* in R4 growth medium⁵ without (for natural abundance teixobactin) and with (for labeled teixobactin) replacement of labeled materials (purchased from Cambridge Isotope Laboratories, Inc., Tewksbury, MA, USA). The labeled Modified Celtone R4 was prepared as 10 g D-glucose U - $^{13}C_6$, 1.1 g Celtone base U - ^{13}C U - ^{15}N , 0.5 g L-proline U - $^{13}C_5$ ^{15}N , 10 g $MgCl_2 \cdot 6H_2O$, 4 g $CaCl_2 \cdot 2H_2O$, 0.2 g K_2SO_4 , 5.6 g *N*-[tris(hydroxymethyl)methyl]-2-aminoethanesulfonic acid (TES), and deionized H_2O to 1 L and pH adjusted to 7.0 using KOH. 1 L of labelled media was inoculated with biomass from an agar plate and grown at 28 °C for 7 days with shaking. The fermentation was centrifuged to separate the cell biomass from the supernatant. The supernatant was decanted into a new container and extracted with an equal volume of butanol, while the cell pellet was extracted with acetone and centrifuged. The acetone and butanol extracts were combined and evaporated to dryness. The resulting residue was dissolved in DMSO and separated by HPLC (water/acetonitrile/0.1% TFA). Of the forty eight fractions generated, fraction 23 contained the majority of the target compound. This fraction was then examined by LC-MS and the expected mass for fully ^{13}C and ^{15}N labelled teixobactin was the major ion peak ($[M + H] = 1315.9$). This fraction was dried down, resulting in 10 mg of the desired product.

Solution NMR

For preparation of TXB_{DPC} [U - ^{13}C , ^{15}N]teixobactin was dissolved to a concentration of 2 or 3 mM in 10 mM NaP buffer, pH 6.5, with 100 or 150 mM d_{38} DPC (Eurisotop). For preparation of $TXB_{aqueous}$ [U - ^{13}C , ^{15}N]teixobactin was dissolved to a concentration of 0.3 mM in 10 mM NaP buffer, pH 6.5. 2,2-Dimethyl-2-silapentane-5-sulfonate (DSS) was used as internal reference. Lipid II was dissolved in 10 mM NaP buffer, pH 6.5, with 150 mM d_{38} DPC. Water soluble lipid II was dissolved in 10 mM NaP buffer, pH 6.5. All solution NMR experiments were performed on a Bruker Avance II 700 MHz spectrometer equipped with a cryo-probe, using 3 mm NMR tubes. Experiments were performed at 25 °C, additional 1H - ^{15}N and 1H - ^{13}C correlation experiments were performed at 37 °C so that assignments could be compared with assignments obtained in solid state NMR experiments. Initial assignments for TXB_{DPC} were obtained from a natural abundance teixobactin (prepared in the same way as labelled teixobactin) using 2D experiments: 1H - ^{13}C HSQC, 1H - ^{15}N SOFAST HMQC (0.3 s recycle delay), 1H - 1H TOCSY (70 ms mixing time), 1H - 1H COSY. Assignments were



confirmed and completed from 3D experiments using [$U\text{-}^{13}\text{C},^{15}\text{N}$]teixobactin: BEST HNCACB (0.3 s recycle delay), CCH-TOCSY (16.3 ms DIPSI-3 mixing time), CBCA(CO)NH (25% non-uniform sampling (NUS) reconstructed in TopSpin using the MDD algorithm³⁶). The assignments were referenced to DSS using an external reference. A 3D ^{15}N HSQC-NOESY (200 ms mixing time) and a 2D ^1H - ^1H NOESY (200 ms mixing time) were used for distance restraints in the structure calculations. ^{15}N T_1 and T_2 relaxation times were measured using interleaved pseudo 3D experiments. Relaxation delays for ^{15}N T_1 experiments (s): 0.005, 0.075, 0.1, 0.2, 0.3, 0.4, 0.5, 0.65, 0.8, 1, 1.25, 1.5. Relaxation delays for ^{15}N T_2 experiments (s): 0.017, 0.034, 0.051, 0.068, 0.085, 0.102, 0.136, 0.170, 0.204, 0.254, 0.339. Titrations with native lipid II to [$U\text{-}^{13}\text{C},^{15}\text{N}$]teixobactin were measured using ^1H - ^{15}N SOFAST HMQC (0.3 s recycle delay) and ^1H - ^{13}C HSQC. Lipid II with a concentration of 15 mM was titrated to the following concentrations (mM): 0.33, 0.79, 1.5, 2, 2.7, 3, 3.4, 3.7, 4, 4.3, 4, leading to a dilution of teixobactin to a final concentration of 2 mM from an initial concentration of 3 mM. Assignments for $\text{TXB}_{\text{aqueous}}$ were initially transferred from TXB_{DPC} using 2D experiments: ^1H - ^{15}N SOFAST HMQC (0.3 s recycle delay) and ^1H - ^{13}C HSQC. The assignments were completed using 3D experiments: HNCA and CCH-TOCSY (16.3 ms DIPSI-3 mixing time). The assignments were referenced internally to DSS. ^{15}N T_1 and T_2 relaxation times were measured using interleaved pseudo 3D experiments. Relaxation delays for ^{15}N T_1 experiments (s): 0.075, 0.1, 0.2, 0.3, 0.4, 0.6, 0.8, 1.1, 1.4. Relaxation delays for ^{15}N T_2 experiments (s): 0.017, 0.034, 0.068, 0.102, 0.136, 0.170, 0.237, 0.305. Titrations of water soluble lipid II to [$U\text{-}^{13}\text{C},^{15}\text{N}$]teixobactin were measured using ^1H - ^{15}N SOFAST HMQC and ^1H - ^{13}C HSQC. Water soluble lipid II with a concentration of 2.4 mM was titrated to the following concentrations (mM): 0.05, 0.1, 0.14, 0.19, 0.23, leading to a dilution of teixobactin from 0.3 to 0.27 mM.

Solid state NMR

After titrations of lipid II to [$U\text{-}^{13}\text{C},^{15}\text{N}$]teixobactin (in DPC micelles) the sample was transferred from the NMR tube, 10 mM NaP buffer pH 6.5, with 2% DSS was added up to 500 μL leading to a final concentration of approximately: 0.6 mM [$U\text{-}^{13}\text{C},^{15}\text{N}$]teixobactin, 1.4 mM lipid II, 45 mM d_{38} DPC. The sample was sedimented by ultracentrifugation (Beckmann Coulter Optima MAX-XP Ultracentrifuge) for 46 hours at $700\,000 \times g$, forming a solid paste. Most of the liquid was removed and a small amount of 10 mM NaP buffer containing 2 mM gadolinium diethylenetriaminepentaacetic acid bismethylamide (Gd(DTPA-BMA)) and 2% DSS was added to the sediment. The sediment was packed into a 0.81 mm Samoson rotor. To keep the sample hydrated during packing small amounts of buffer with Gd(DTPA-BMA) and DSS were added to the rotor. All experiments were recorded at a 600 MHz Bruker Avance II spectrometer using a Samoson HXY 0.81 mm probe at 90 kHz magic angle spinning and a sample temperature of 39 ± 2 °C measured from the water peak referenced to DSS. Proton detection with 30 ms acquisition time was used for all

experiments. The addition of Gd(DTPA-BMA) enabled a recycle delay of 0.5 s. Water suppression was achieved by 100–150 ms slpTPPM³⁷ at 22.5 kHz nutation frequency ($\frac{1}{4}$ of the spinning speed). The following spectra were acquired: 2D ^1H - ^{13}C inverse CP (0.4 ms CP between ^1H - ^{13}C and ^{13}C - ^1H) 2D ^1H - ^{15}N inverse CP (1 ms CP between ^1H - ^{15}N and 0.9 ms between ^{15}N - ^1H), 3D hCANH (0.4 ms CP between ^1H - ^{13}C , 11 ms between ^{13}C - ^{15}N , 0.9 ms between ^{15}N - ^1H), 3D hCONH (2.5 ms CP between ^1H - ^{13}C , 11 ms between ^{13}C - ^{15}N , 0.9 ms between ^{15}N - ^1H), 3D hCCH TOCSY (0.4 ms CP between ^1H - ^{13}C and ^{13}C - ^1H , 16.3 ms DIPSI-3 mixing time at 10 kHz nutation frequency) 3D hCOCAHA DREAM (2.5 ms CP between ^1H - ^{13}CO , 0.4 ms CP between ^{13}CA - ^1HA , 7 ms DREAM between ^{13}CO - ^{13}CA at 45 kHz nutation frequency, $\frac{1}{2}$ of the spinning speed).

NMR data analysis

TopSpin 3.2 was used to process all spectra. The spectra were assigned in NMRFAM Sparky.³⁸ Structure calculation of TXB_{DPC} was performed using UNIO 10 with Cyana 2.1³⁹ as molecular dynamics software using NOEs as structural restraints. Raw spectra (^1H - ^1H 2D NOESY and ^{15}N 3D HSQC-NOESY) were used as input into the structure calculation. Automatic peak picking and NOE assignments were achieved by UNIO ATNOS-CANDID.^{40,41} Cyana library entries for D -amino acids were produced using CyLib:⁴² D -glutamine (converted from DGN.cif), D -allo-isoleucine (converted from 28J.cif), D -threonine (converted from DTH.cif). A Cyana library entry for N -methylated- D -phenylalanine was initially converted by Cylib from ZAE.cif and slightly modified by producing a .cor file in Cyana containing the new ZAE residue as the first residue and a peptide bond to another amino acid. The amide proton was added in UCSF Chimera⁴³ and a pdb file was exported and read in MOLMOL⁴⁴ where coordinates were exported as a library file. The library file was edited manually in a text editor to fit to the Cyana library format. The .cif files were obtained from http://www.bpc.uni-frankfurt.de/guentert/wiki/index.php/Cyana_Residue_Library_Entries, except for D -allo-isoleucine (28J.cif), which was obtained from <http://www.ebi.ac.uk/pdbe-srv/pdbechem/chemicalCompound/show/28J>. A Cyana library entry for allo-enduracididine was produced from a drawing of the chemical structure in ChemSketch (ACDLabs Freeware 2012) including peptide bonds to residues before and after. The chemical structure was exported as a .mol file and converted to PDB in UCSF Chimera.⁴³ MOLMOL⁴⁴ was used to write coordinate files and the Cyana library entry was finalized by manually rearranging the atoms, adding pseudo atoms and torsion angles in a text editor (PSPAD editor).

Peak integrals from TopSpin were exported to MatLab where a single decaying exponential function was used to fit the R_1 and R_2 relaxation data. Errors were calculated by a Monte Carlo error estimation. A random number between 0 and 1 was multiplied with the integral error, which was based on the noise level compared to the signal for each peak, and added to the recalculated integrals. The fitting was then repeated 2000 times with a new random number between 0 and 1 generated each time. Two times the standard deviations of the R_1 or R_2 values received from the fits for each residue were used as errors.



The correlation times (τ_c) for TXB_{aqueous} and TXB_{DPC} were calculated from the ^{15}N T_1 and T_2 relaxation times and the ^{15}N resonance frequency (ν_N), 71 MHz at a 16.4 T NMR magnet (700 MHz ^1H Larmor frequency) using eqn (1):

$$\tau_c \approx \frac{1}{4\pi\nu_N} \sqrt{6 \frac{T_1}{T_2} - 7} \quad (1)$$

which is generally used for rigid protein molecules in the limit of slow molecular motion ($\tau_c \gg 0.5$ ns), using high magnetic field (^1H Larmor frequency ≥ 500 MHz).⁴⁵

To determine binding affinities the peak intensities of unbound teixobactin (TXB_{DPC} or TXB_{aqueous}) were measured in the titration spectra and fitted to eqn (2) (based on eqn (6) in ref. 25):

$$P = \frac{m}{2n[P_t]} (-K_d - [L_t] + n[P_t]) + \sqrt{4K_d n[P_t] + (-K_d + [L_t] - n[P_t])^2} \quad (2)$$

where P is the normalized peak intensity, $m = 1$ if the point without an added ligand is included and m is fitted if the point without an added ligand is not included, n is the molar ratio for binding (lipid II/teixobactin), $[L_t]$ is ligand concentration (lipid II), $[P_t]$ is protein concentration (teixobactin). The data points used for the fitting were: 0.33, 0.79, 1.5, 2, 2.7, 3 (mM) lipid II for TXB_{DPC} and 0.05, 0.1, 0.14, 0.19, 0.23 (mM) water soluble lipid II for TXB_{aqueous}. The fitting of the data was done by minimization of the χ^2 target function (eqn (3)):

$$\chi^2 = \sum \frac{(X_{i,\text{calc}} - X_{i,\text{exp}})^2}{\sigma_{i,\text{exp}}^2} \quad (3)$$

where X_i are the data sets and σ_i the corresponding error. Errors for the K_d fits were calculated in the same way as for the relaxation data using 500 steps in the Monte Carlo error estimation.

Chemical shift perturbations were calculated as Euclidian distances²⁵ (eqn (4)):

$$\text{CSP} = \sqrt{\frac{1}{2} [\delta_H^2 + (\alpha\delta_X^2)]} \quad (4)$$

where δ is the difference in chemical shift between TXB_{aqueous} and TXB_{DPC} or between TXB_{DPC} and TXB_{lipid II}, $\alpha = 0.14$ if $x = ^{15}\text{N}$ and $\alpha = 0.3$ if $x = ^{13}\text{C}$.

FRET. 1 μM biotinylated lipid II was labelled with streptavidin-conjugated FRET dyes (HTRF, CisBio) and incubated with ramoplanin, teixobactin, deoxyactagardine B, mercacidin and vancomycin at concentrations ranging between 0.5 and 32 μM . Samples were incubated for 30 min at ambient temperature and FRET magnitude was measured in a time-resolved manner, using the Clariostar plate reader (BMG Labtech). The experiments were performed in triplicates. Full reaction composition: 1 μM biotin-DAP-lipid II, 1 \times HTRF labelling mixture (terbium cryptate, d2), 50 mM bis-Tris propane pH 8.5, 10 mM MgCl_2 , 20 mM NaCl, 0.05% Triton X-100, 1% glycerol.

Author contributions

C. Ö. and J. R. L. designed and analysed the NMR experiments and wrote the paper, C. Ö. performed the NMR experiments, C. Ö. and T. H. performed the NMR structure calculation. G. P. W. and C. G. D. designed and analysed the FRET experiments, G. P. W. performed the FRET experiments. C. G. D. and A. J. L. optimized lipid II synthesis and analysis, C. Ö., G. P. W., A. C. C., J. A. T. and A. J. L. produced and characterised lipid II. D. E. H., A. L. S. and A. J. P. produced and characterised the natural abundance teixobactin and [^{13}C , ^{15}N]teixobactin. The manuscript was written through contributions from all authors.

Conflicts of interest

The following authors, A. J. Peoples, A. L. Spoering, D. E. Hughes, and K. Lewis declare competing financial interests as they are employees and consultants of NovoBiotic Pharmaceuticals.

Data availability

NMR assignments were deposited in the BMRB under accession numbers 27478 (TXB_{DPC}), 27479 (TXB_{aqueous}) and 27480 (TXB_{lipid II}). Raw NMR data are available from <http://wrap.warwick.ac.uk/108554/>.

Acknowledgements

The research leading to these results has received funding from the Bill & Melinda Gates Foundation (grant OPP1160394, and OPP1132809), European Research Council under the European Union's Seventh Framework Programme (FP/2007–2013)/ERC Grant Agreement 639907. J. R. L. also acknowledges funding from EPSRC Grant EP/L025906/1, BBSRC Grant BB/L022761/1 & BB/R010218/1. A. J. L., J. A. T., A. C. C. and C. G. D. acknowledge funding from AMR Cross Council funding MR/N002679/1. C. Ö. and G. P. W. acknowledge funding from the European Union under a Marie Curie Initial Training Network FP7-PEOPLE-2012-ITN Grant Agreement Number 316630 CAS-IDP. We thank Peter Güntert for providing the CyLib software and Tobias Madl for providing Gd(DTPA-BMA). Molecular graphics and analyses were performed with the UCSF Chimera package. Chimera is developed by the Resource for Biocomputing, Visualization, and Informatics at the University of California, San Francisco (supported by NIGMS P41-GM103311).

Notes and references

- 1 T. Kaerberlein, K. Lewis and S. S. Epstein, *Science*, 2002, **296**, 1127–1129.
- 2 D. Nichols, N. Cahoon, E. M. Trakhtenberg, L. Pham, A. Mehta, A. Belanger, T. Kanigan, K. Lewis and S. S. Epstein, *Appl. Environ. Microbiol.*, 2010, **76**, 2445–2450.
- 3 A. D'Onofrio, J. M. Crawford, E. J. Stewart, K. Witt, E. Gavrish, S. Epstein, J. Clardy and K. Lewis, *Chem. Biol.*, 2010, **17**, 254–264.



- 4 E. Gavriš, C. S. Sit, S. Cao, O. Kandrór, A. Spoering, A. Peoples, L. Ling, A. Fetterman, D. Hughes, A. Bissell, H. Torrey, T. Akopian, A. Mueller, S. Epstein, A. Goldberg, J. Clardy and K. Lewis, *Chem. Biol.*, 2014, **21**, 509–518.
- 5 L. L. Ling, T. Schneider, A. J. Peoples, A. L. Spoering, I. Engels, B. P. Conlon, A. Mueller, T. F. Schäberle, D. E. Hughes, S. Epstein, M. Jones, L. Lazarides, V. a Steadman, D. R. Cohen, C. R. Felix, K. A. Fetterman, W. P. Millett, A. G. Nitti, A. M. Zullo, C. Chen and K. Lewis, *Nature*, 2015, **517**, 455–459.
- 6 T. Homma, A. Nuxoll, A. B. Gandt, P. Ebner, I. Engels, T. Schneider, F. Götz, K. Lewis and B. P. Conlon, *Antimicrob. Agents Chemother.*, 2016, **60**, 6510–6517.
- 7 D. Münch and H. G. Sahl, *Biochim. Biophys. Acta, Biomembr.*, 2015, **1848**, 3062–3071.
- 8 W. Vollmer, D. Blanot and M. a. de Pedro, *FEMS Microbiol. Rev.*, 2008, **32**, 149–167.
- 9 W. Vollmer, *FEMS Microbiol. Rev.*, 2008, **32**, 287–306.
- 10 A. Parmar, S. H. Prior, A. Iyer, C. S. Vincent, D. Van Lysebetten, E. Breukink, A. Madder, E. J. Taylor and I. Singh, *Chem. Commun.*, 2017, **53**, 2016–2019.
- 11 Y. E. Jad, G. A. Acosta, T. Naicker, M. Ramtahal, A. El-Faham, T. Govender, H. G. Kruger, B. G. de la Torre and F. Albericio, *Org. Lett.*, 2015, **17**, 6182–6185.
- 12 H. Yang, K. H. Chen and J. S. Nowick, *ACS Chem. Biol.*, 2016, **11**, 1823–1826.
- 13 A. Parmar, A. Iyer, D. G. Lloyd, C. S. Vincent, S. H. Prior, A. Madder, E. J. Taylor and I. Singh, *Chem. Commun.*, 2017, **53**, 7788–7791.
- 14 H. Yang, D. R. Du Bois, J. W. Ziller and J. S. Nowick, *Chem. Commun.*, 2017, **53**, 2772–2775.
- 15 A. Parmar, A. Iyer, S. H. Prior, D. G. Lloyd, E. T. Leng Goh, C. S. Vincent, T. Palmal-Pallag, C. Z. Bachrati, E. Breukink, A. Madder, R. Lakshminarayanan, E. J. Taylor and I. Singh, *Chem. Sci.*, 2017, **8**, 8183–8192.
- 16 K. H. Chen, S. P. Le, X. Han, J. M. Frias and J. S. Nowick, *Chem. Commun.*, 2017, **53**, 11357–11359.
- 17 C. Wu, Z. Pan, G. Yao, W. Wang, L. Fang and W. Su, *RSC Adv.*, 2017, **7**, 1923–1926.
- 18 S. A. H. Abdel Monaim, S. Noki, E. J. Ramchuran, A. El-Faham, F. Albericio and B. G. de la Torre, *Molecules*, 2017, **22**, 1–9.
- 19 Y. Liu, Y. Liu, M. B. Chan-Park and Y. Mu, *Sci. Rep.*, 2017, **7**, 17197.
- 20 M. Jackson and H. H. Mantsch, *Biochim. Biophys. Acta*, 1991, **1078**, 231–235.
- 21 S.-T. D. Hsu, E. Breukink, G. Bierbaum, H.-G. Sahl, B. de Kruijff, R. Kaptein, N. A. J. van Nuland and A. M. J. J. Bonvin, *J. Biol. Chem.*, 2003, **278**, 13110–13117.
- 22 P. Cudic, J. K. Kranz, D. C. Behenna, R. G. Kruger, H. Tadesse, A. J. Wand, Y. I. Veklich, J. W. Weisel and D. G. McCafferty, *Proc. Natl. Acad. Sci. U. S. A.*, 2002, **99**, 7384–7389.
- 23 S.-T. D. Hsu, E. Breukink, E. Tischenko, M. A. G. Lutters, B. de Kruijff, R. Kaptein, A. M. J. J. Bonvin and N. A. J. van Nuland, *Nat. Struct. Mol. Biol.*, 2004, **11**, 963–967.
- 24 J. M. Lamley, D. Iuga, C. Öster, H.-J. Sass, M. Rogowski, A. Oss, J. Past, A. Reinhold, S. Grzesiek, A. Samoson and J. R. Lewandowski, *J. Am. Chem. Soc.*, 2014, **136**, 16800–16806.
- 25 M. P. Williamson, *Prog. Nucl. Magn. Reson. Spectrosc.*, 2013, **73**, 1–16.
- 26 M. Jenner, S. Kosol, D. Griffiths, P. Prasongpholchai, L. Manzi, A. S. Barrow, J. E. Moses, N. J. Oldham, J. R. Lewandowski and G. L. Challis, *Nat. Chem. Biol.*, 2018, **14**, 270–275.
- 27 Y. Zong, X. Sun, H. Gao, K. J. Meyer, K. Lewis and Y. Rao, *J. Med. Chem.*, 2018, **61**, 3409–3421.
- 28 A. Mainz, S. Jehle, B.-J. J. van Rossum, H. Oschkinat and B. Reif, *J. Am. Chem. Soc.*, 2009, **131**, 15968–15969.
- 29 A. Loquet, N. G. Sgourakis, R. Gupta, K. Giller, D. Riedel, C. Goosmann, C. Griesinger, M. Kolbe, D. Baker, S. Becker and A. Lange, *Nature*, 2012, **486**, 276–279.
- 30 C. Gardiennet, A. K. Schütz, A. Hunkeler, B. Kunert, L. Terradot, A. Böckmann and B. H. Meier, *Angew. Chem., Int. Ed.*, 2012, **51**, 7855–7858.
- 31 S. A. H. Abdel Monaim, Y. E. Jad, G. A. Acosta, T. Naicker, E. J. Ramchuran, A. El-Faham, T. Govender, H. G. Kruger, B. G. de la Torre and F. Albericio, *RSC Adv.*, 2016, **6**, 73827–73829.
- 32 J. a. Siepen, S. E. Radford and D. R. Westhead, *Protein Sci.*, 2003, **12**, 2348–2359.
- 33 K. M. Scherer, J. H. Spille, H. G. Sahl, F. Grein and U. Kubitschek, *Biophys. J.*, 2015, **108**, 1114–1124.
- 34 E. Breukink, H. E. Van Heusden, P. J. Vollmerhaus, E. Swiezewska, L. Brunner, S. Walker, A. J. R. Heck and B. De Kruijff, *J. Biol. Chem.*, 2003, **278**, 19898–19903.
- 35 A. J. Lloyd, A. M. Gilbey, A. M. Blewett, G. De Pascale, A. El Zoeiby, R. C. Levesque, A. C. Catherwood, A. Tomasz, T. D. H. Bugg, D. I. Roper and C. G. Dowson, *J. Biol. Chem.*, 2008, **283**, 6402–6417.
- 36 V. Y. Orekhov and V. A. Jaravine, *Prog. Nucl. Magn. Reson. Spectrosc.*, 2011, **59**, 271–292.
- 37 J. R. Lewandowski, J. Sein, H. J. Sass, S. Grzesiek, M. Blackledge and L. Emsley, *J. Am. Chem. Soc.*, 2010, **132**, 8252–8254.
- 38 W. Lee, M. Tonelli and J. L. Markley, *Bioinformatics*, 2015, **31**, 1325–1327.
- 39 P. Güntert, C. Mumenthaler and K. Wüthrich, *J. Mol. Biol.*, 1997, **273**, 283–298.
- 40 T. Herrmann, P. Güntert and K. Wüthrich, *J. Biomol. NMR*, 2002, **24**, 171–189.
- 41 T. Herrmann, P. Güntert and K. Wüthrich, *J. Mol. Biol.*, 2002, **319**, 209–227.
- 42 E. M. Yilmaz and P. Güntert, *J. Biomol. NMR*, 2015, **63**, 21–37.
- 43 E. F. Pettersen, T. D. Goddard, C. C. Huang, G. S. Couch, D. M. Greenblatt, E. C. Meng and T. E. Ferrin, *J. Comput. Chem.*, 2004, **25**, 1605–1612.
- 44 R. Koradi, M. Billeter and K. Wüthrich, *J. Mol. Graphics*, 1996, **14**, 51–55.
- 45 J. Cavanagh, J. W. Fairbrother, A. G. Palmer, M. Rance and N. J. Skelton, *Protein NMR Spectroscopy: Principles and Practice*, Elsevier, 2nd edn, 2007.

