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## Re-evaluation of the N-terminal substitution and the D-residues of teixobactin†

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Teixobactin is a head to side-chain cyclic depsipeptide with a guanidino based residue within the cycle, three D-amino acids in the tail, and a N-methylated terminal residue. The synthesis of the first analogue, containing Arg, was recently described by our group. Herein, we demonstrated that analogues of Arg. Teixobactin bearing either (a) three L-amino acids in the tail and keeping the N-methyl at the N-terminal or (b) with three D-amino acids, but with acetylation of the N-terminal, are inactive against Gram(+) and Gram(−) bacteria. These results complement those published by the groups of Madder, Taylor, and Singh that have shown that both modifications: L-amino acids and N-acetylation also led to loss of biological activity.

The development of new antimicrobial drugs has become essential and crucial due the dramatic increase of bacterial resistance to traditional antibiotics.<sup>1–3</sup> Amongst these new antimicrobial compounds, antimicrobial peptides (AMP), especially cyclic peptides, have been shown to be very appealing due to the potent activities and stability.<sup>4–6</sup> Very recently (2015), a new head to side chain antimicrobial cyclodepsipeptides called teixobactin (**1**) was discovered by Ling *et al.* using an iChip multichannel device as a new discovery technique to simultaneously isolate and grow uncultured bacteria.<sup>7</sup> This new antimicrobial peptide exhibited higher activity against Gram positive than Gram negative bacteria. Furthermore, teixobactin has been reported to kill bacteria in absence of any detectable

resistance. The reason for that was ascribed to its mechanism of action; where it blocks one of the membrane-associated steps of peptidoglycan biosynthesis and then causes inhibition of lipid II, as a predominant target, and lipid III which plays a critical role in the synthesis of teichoic acid.<sup>7,8</sup>

Since the publication of isolation, characterization, and antimicrobial activity of teixobactin,<sup>7</sup> several groups started the development of synthetic strategies for a convenient synthesis that is suitable for running a medicinal chemistry program. Our group published in 2015, the first synthesis of an analogue of teixobactin (**2a**) in 2015 where the non-proteinogenic guanidine-based amino acid enduracididine was substituted with the proteinogenic arginine.<sup>9</sup> This analogue showed similar biological trends, being active against the Gram-positive bacteria and inactive against Gram-negative bacteria. However, teixobactin was one order of magnitude more active than the Arg analogue.<sup>9,10</sup>

At the beginning of 2016, the groups of Madder, Taylor, and Singh published the synthesis of the same analogue, Arg-teixobactin (**2a**),<sup>11</sup> using an identical strategy and reporting the same biological activity. In the same manuscript, the authors stated that “(they) established that the D-amino acids are critical for the antimicrobial activity”. They supported this statement with the synthesis of a second analogue (**3b**) where the three D-amino acids of the tail were substituted by the L-amino acid residues (Phe, Gln, Ile). However, the N-Me of the terminal Phe for this new analogue (**3b**) was substituted with N-Ac. Substitution of N-Me-Phe residue with Ac-Phe should have a significant impact in the biological activity. The impact of this Ac-Phe terminal could potentially be more critical than the change to L-amino acids, since the peptide (**3b**) has lost its cationic character (a normal requirement for antimicrobial peptides)<sup>12</sup> due to introduction of the acetyl group. In addition to that, the hydrogen bond map of the peptide is also dramatically altered.

Herein, we report the synthesis and biological evaluation of two unpublished analogues: (**2b**) that has three L-amino acids while keeping the N-Me group at the Phe terminal; and (**3a**) with D-amino acids, but where the D-Phe terminal is acetylated (Fig. 1).

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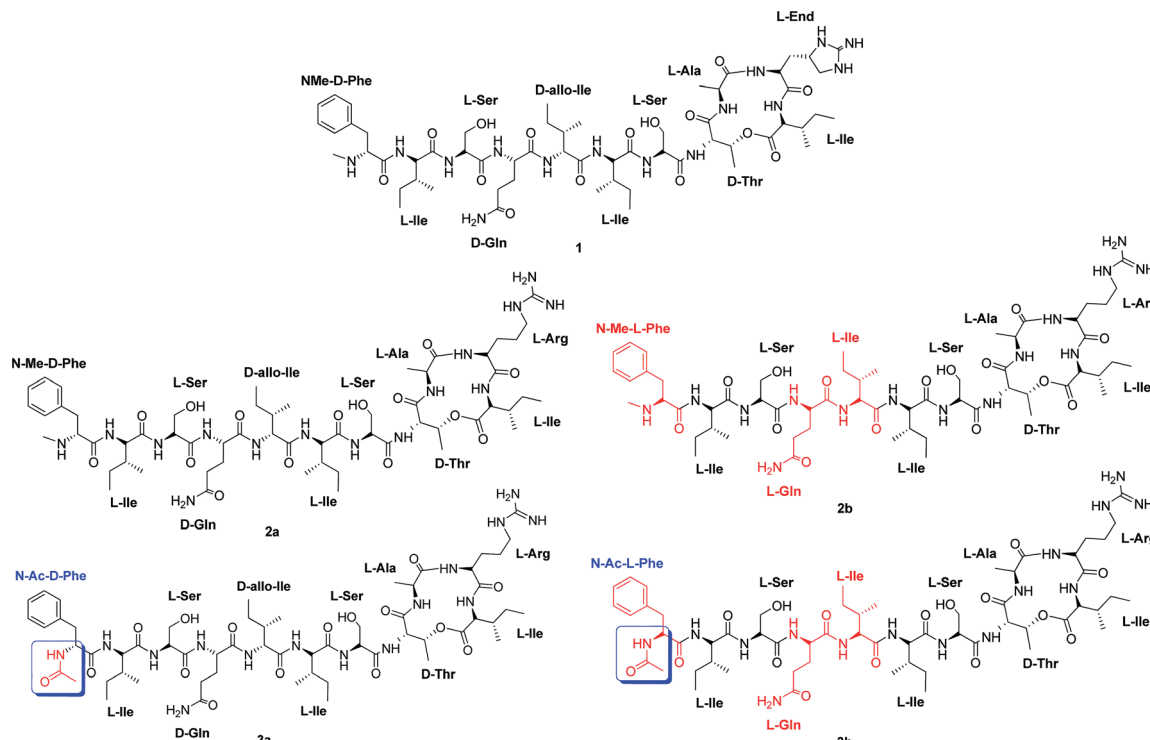
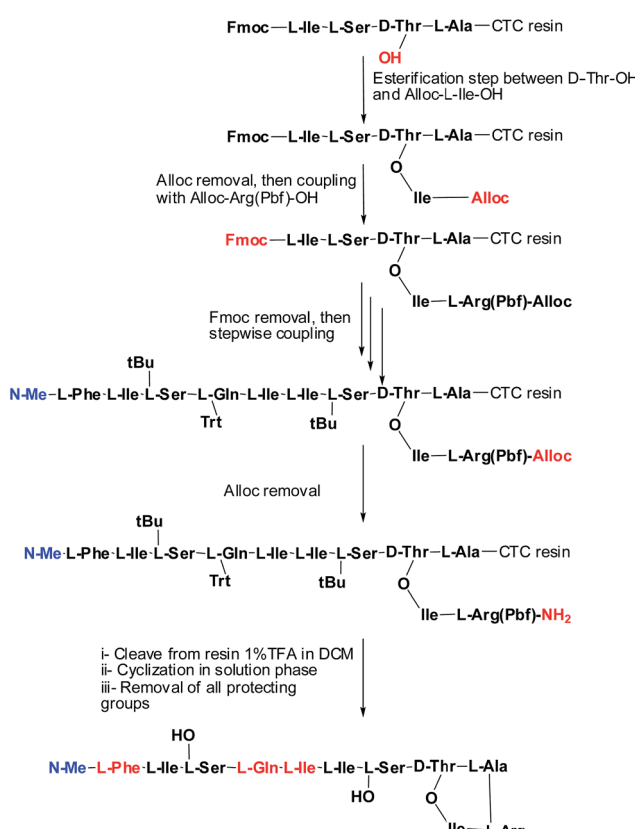
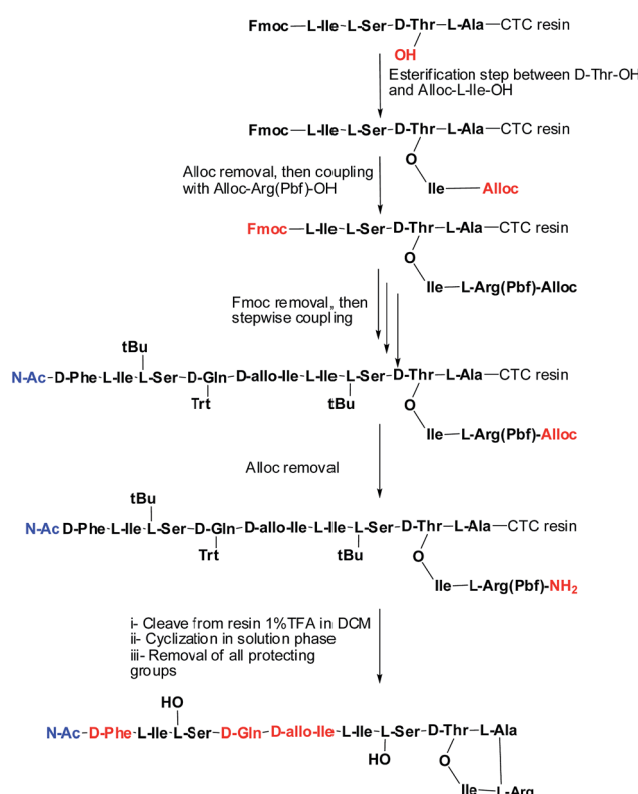


Fig. 1 Structure of teixobactin. (1) Structure of teixobactin analogue (2a) with D-amino acids at the tail and a N-(Me)-terminal.<sup>6,9</sup> Structure of teixobactin analogue (2b) with L-amino acids at the tail and a N-(Me)-terminal (new analogue). Structure of teixobactin analogue (3a) with D-amino acids at the tail and a N-(Ac)-terminal (new analogue). Structure of teixobactin analogue (3b) with L-amino acids at the tail and a N-(Ac)-terminal.<sup>11</sup>



Scheme 1 Synthetic strategy of (2b).



Scheme 2 Synthetic strategy of (3a).

Table 1 MIC ( $\mu\text{g ml}^{-1}$ ) for (2a) (2b) (3a), and (3b)

Entry	MIC $\mu\text{g ml}^{-1}$ (ATCC strains)			
	Gram(+) <i>S. aureus</i> <i>B. subtilis</i>		Gram(−) <i>E. coli</i> <i>P. aeruginosa</i>	
	<i>S. aureus</i>	<i>B. subtilis</i>	<i>E. coli</i>	<i>P. aeruginosa</i>
Teixobactin <sup>7</sup>	0.25	0.06	25	>32
2a <sup>9</sup>	1.99	0.49	63.9	NI
2a <sup>11</sup>	2	—	64	—
2b	NI <sup>a</sup>	NI	NI	NI
3a	NI	256	NI	NI
3b <sup>11</sup>	128	—	GAW <sup>b</sup>	—
Meropenem <sup>c</sup>	0.0625	0.0625	0.03125	0.25

<sup>a</sup> NI-no inhibition. <sup>b</sup> GAW: growth in all wells. <sup>c</sup> Use as control in our experiments.

Syntheses of analogues (2b) and (3b) were carried following basically the same synthetic scheme used by our group for the synthesis of Arg-teixobactin (2a),<sup>9</sup> (Schemes 1 and 2), with the only modification for the analogue (3a) where Fmoc protection was used for the D-Phe, instead of Boc. Before cleavage of the latter from the resin, the Fmoc group was removed and the free amine was acetylated with HOAc in presence of K-Oxyma/DIC. Both analogues were obtained with an overall non-optimized yield of 50% for (2b) and 56% for (3a) before purification. The respective yields after purification were 3.8% for (2b) and 3.9% for (3a), and these peptides were characterized by HPLC and HRMS.

Analogues (2b) and (3a) were evaluated against two ATCC Gram positive bacterial strains (*Staphylococcus aureus* 29213 and *Bacillus subtilis* 6051) and two ATCC Gram negative bacterial strains (*Escherichia coli* 25822 and *Pseudomonas aeruginosa* 27853) resulting in a total lack of activity as shown in (Table 1).

In conclusion, the synthesis of two new analogues of 2a was achieved. Peptide 2b is different to 2a in that three D-amino acids have been substituted by the L-residues (Phe, Gln, Ile), while 3a is different to the teixobactin analogue 2a only in that the terminal N-Me group was replaced with a N-Ac group. Both of these compounds lost totally the bacterial activity. These results complement those reported by the groups of Madder, Taylor, and Singh that have found that the double change (3b),

L-amino acids and acetyl at the N-terminal acetylated is also inactive.

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