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Chapter 1

Introduction. Synergy by perturbing the Gramnegative outer membrane: opening the door for Gram-positive specific antibiotics

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Parts of this chapter have been submitted for publication

Abstract

New approaches to target antibacterial agents towards Gram-negative bacteria are key given the rise of antibiotic resistance. Since the discovery of polymyxin B nonapeptide as a potent Gram-negative outer membrane (OM) permeabilizing synergist in the early 1980s, a vast amount of literature on such synergists has been published. This review addresses a range of peptide-based and small organic compounds that disrupt the OM to elicit a synergistic effect with antibiotics that are otherwise inactive towards Gramnegative bacteria, with synergy defined as a fractional inhibitory concentration index of <0.5. Another requirement for the inclusion of the synergists here covered is their potentiation of a specific set of clinically used antibiotics: erythromycin, rifampicin, novobiocin, or vancomycin. In addition, we have focused on those synergists with reported activity against Gram-negative members of the ESKAPE family of pathogens namely, Escherichia coli, Pseudomonas aeruginosa, Klebsiella pneumoniae, and/or Acinetobacter baumannii. In cases where the FICI values were not directly reported in the primary literature but could be calculated from the published data we have done so, allowing for more direct comparison of potency with other synergists. We also address the hemolytic activity of the various OM disrupting synergists reported in the literature, an effect that is often downplayed but of key importance in assessing the selectivity of such compounds for Gram-negative bacteria.

1. Introduction

The increasing occurrence of antibiotic resistance among Gram-negative pathogens highlights the need for novel antibacterial agents and therapeutic strategies. It is well established that Gram-negative bacteria are inherently harder to kill with antibiotics than Gram-positives given the presence of the Gram-negative outer membrane (OM) as well as efflux pumps.¹⁻⁴ Among the limited number of clinically effective anti-Gram-negative agents, several are labeled as last resort further underscoring the urgent need for new treatments against Gram-negative pathogens.⁵⁻⁷ This troubling reality is further exacerbated by increasing accounts of emerging resistance mechanisms against Gramnegative antibiotics including: extended spectrum beta-lactamases (ESBLs) that can render even fifth generation cephalosporins and carbapenems innactive,⁸⁻¹¹ enzymes that structurally modify and deactivate aminoglycosides,¹²⁻¹⁵ and mcr-mediated polymyxin resistance.¹⁶⁻²⁷ In this context, the World Health Organization (WHO) recently listed (carbapenem-resistant), Acinetobacter baumannii Pseudomonas aeruginosa (carbapenem-resistant), and the Enterobacteriaceae (carbapenem-resistant and ESBLproducing strains) as the bacterial pathogens of highest priority for the development of new antibiotics.28

The Gram-negative OM functions as a barrier that prevents many antibiotics, that are otherwise active against Gram-positive species, from reaching their targets,^{3,29} The OM itself consists of an asymmetrical lipid bilayer (See Figure 1A).³⁰ The inner leaflet consist mostly of phospholipids and is similar to the cytoplasmic membrane,³¹ The outer leaflet is made up of an organized and fortified structure of densely packed lipopolysaccharides (LPS) and Mg^{2+}/Ca^{2+} cations that bridge the negatively charged phosphate groups of the lipid A component of LPS (See Figure 1B).^{3,32} Furthermore, the tightly packed saturated acyl chains result in a low level of membrane fluidity that limits the diffusion of hydrophobic compounds across the OM.^{2,3} The OM also contains porins which function as size exclusion channels across the OM that mediate the diffusion of small hydrophilic molecules between the periplasm and the extracellular environment while keeping large, hydrophobic molecules, including many antibiotics, out.^{1,2,29} Additionally, when lipophilic or amphiphilic antibiotics do manage to cross the OM, multidrug efflux pumps can transport these molecules back out.^{1-3,29} In many cases, the over-expression of efflux pumps provides an effective means for a Gram-negative pathogen to decrease its susceptibility to antibiotics.^{3,33} Taken together, their diverse resistance mechanisms and unique cellular features provide Gram-negative bacteria with a formidable range of defenses against antibacterial agents.

To address the specific challenges posed by Gram-negative bacteria a number of new and innovative approaches are currently under investigation. Such strategies include interfering with LPS biosynthesis,³⁴⁻³⁷ targeting OM proteins such as the BAM complex,^{34,38,39} developing siderophore-antibiotic conjugates as Trojan horse agents,⁴⁰⁻⁴² co-administering different antibiotics to restrict or reverse antibiotic resistance,^{43,44} and blocking efflux pumps.⁴⁵⁻⁴⁸ In addition to these promising strategies, the development of agents that can selectively disrupt the OM offers the possibility of sensitizing Gramnegative bacteria to antibiotics that otherwise function only against Gram-positive bacteria.^{3,7,32} The pursuit of such synergists continues to be a very active field of research and is the basis for this review.



Figure 1. A) Schematic depiction of the OM disruption required for potentiation of Grampositive specific antibiotics (created with BioRender.com); B) Lipid A (from Escherichia coli K-12), the hydrophobic anchor of LPS.

The best studied example of an OM disrupting synergist is polymyxin B nonapeptide (PMBN) which is obtained by enzymatic degradation of the clinically used lipopeptide polymyxin B (PMB).^{7,32} The potentiating effects of PMBN were first reported in the 1980s, and in the decades since a growing number of OM disrupting synergists have been discovered.^{7,32,49} To date, a number of reviews have been published on the general topic of antibiotic synergy,⁵⁰⁻⁵⁷ including compounds that potentiate Grampositive antibiotics through interactions with the OM⁵⁸ and OM disrupting synergists,^{32,59-} ⁶³ However, a comprehensive overview of OM disrupting synergists that also provides the reader with a direct comparison of both the potency and selectively of these compounds has, to date, been lacking. In this regard, the most widely accepted benchmark for synergistic activity is the so called fractional inhibitory concentration index (FICI, Box 1). In this review we discuss only those synergists for which FICI values are reported or could be calculated from published data. The other criterion we have also chosen to emphasize is the selectivity of OM disruption associated with these synergists. In this regard, we pay special attention to the hemolytic activity reported for the various OM disruptors as a means of assessing their membrane specificity.

Among the Gram-negative bacteria for which OM disrupting synergists have been reported, we have selected those pathogens noted on the WHO's priority list: A. *baumannii*, Escherichia coli, Klebsiella pneumoniae, or P. aeruginosa.²⁸ As for Grampositive specific antibiotics whose activity is potentiated by OM disrupting synergists, we have chosen to focus on clinically used agents that are most commonly evaluated for synergy with OM disruptors: erythromycin, rifampicin, vancomycin, and novobiocin.^{7,58} This criterion has, for example, led to the exclusion of OM disrupting agents for which synergy was reported with macrolide antibiotics other than erythromycin.⁶⁴⁻⁶⁷ Also, to further streamline the review, synergists for which an OM disrupting mechanism was not clearly demonstrated are not here discussed in detail.⁶⁸⁻⁷⁶ In addition, synergists that specifically engage with Gram-negative targets and subsequently cause OM disruption as a secondary effect are not discussed in this review.⁷⁷⁻⁸⁵

Box 1. An important formalism in the field of synergy is the fractional inhibitory concentration index (FICI). The FICI is calculated from experimental minimum inhibitory concentration (MIC) data as shown in Equation 1. A synergistic combination is generally defined as an FICI ≤ 0.5 . Additionally, it allows for a straightforward comparion of the potency of the synergistic combinations: the lower the FICI, the more potent the combination. Apart from the FICI, the minimum synergistic concentration (MSC) values are also relevant parameters. The MSCs represent the concentrations of each component required for synergy and are therefore also of clinical relevance.

$$FICI = \frac{MSC_{ant}}{MIC_{ant}} + \frac{MSC_{syn}}{MIC_{syn}}$$
(1)

Equation 1. Calculation of FICI. $MSC_{ant} = MIC$ of antibiotic in combination with synergist; $MIC_{ant} = MIC$ of antibiotic alone; $MSC_{syn} = MIC$ of synergist in combination with antibiotic; $MIC_{syn} = MIC$ of synergist alone.

The scope of the synergists included in this review ranges from peptides to synthetic small-molecules and small polymers of <1500 Da. In this regard, protein-based OM disruptors such as the membrane attack complex (MAC)⁸⁶, lactoferrin,⁸⁷ and the bactericidal/permeability-increasing protein (BPI)⁸⁸ or larger polymers or polymer-like agents^{89-92,92-96} will not be discussed. This review is further organized based on the chemical families of the synergists covered. We begin with cyclic peptides based on PMBN, followed by linear peptides, cationic steroids, peptide-steroids hybrids, and small molecules. For each subgroup of synergists a summary table has been assembled to provide a convenient comparative overview of FICI values. These tables also include the identity of the Gram-negative species and companion antibiotics employed in generating the FICIs. In addition, where possible, we have included the reported hemolytic activity of each synergist to provide an indication of their selectivity for Gram-negative cells.

2. Peptide-based potentiators

2.1. Polymyxin derived synergists

Polymyxin derived synergists have been extensively reviewed in the past and therefore only a concise summary of these analogues is here included.^{7,32,63} PMBN is a derivative of the parent lipopeptide PMB (see Figure 2A). Unlike its parent compound, PMBN has no inherent antimicrobial activity nor is it nephrotoxic.^{7,97} In their landmark 1983 paper, Martti and Timo Vaara demonstrated that the combination of PMBN with hydrophobic, generally Gram-positive specific, antibiotics results in a potent synergistic effect (See Table 1).^{32,49} In this regard, PMBN is often used as a benchmark for synergistic activity.⁷ Apart from PMBN, other truncated derivatives of PMB, like deacylpolymyxin B (DAPB), polymyxin B octapeptide (PMBO) and polymyxin B heptapeptide (PMBH) also display synergistic activity (Figure 1A and Table 1).³² The peptide macrocycle is of key importance for these synergists as linear PMBN variants lose their synergistic activity.⁹⁸



Figure 2. Molecular structures of A) polymyxin B (PMB), deacylpolymyxin B (DAPB), polymyxin B nonapeptide (PMBN), polymyxin B octapeptide (PMBO), and polymyxin B heptapeptide (PMBH); B) PMBN analogues SPR741, NAB739, and NAB7061.

A new generation of PMBN analogues containing only three positive charges was developed more recently.^{99,100} SPR741, previously named NAB741, has passed the Phase I clinical trials (See Figure 2B).⁷ Like PMBN, SPR741 has no lipophilic tail resulting in improved renal clearance compared to PMB and other analogues including a lipophilic tail such as NAB739 and NAB7061.¹⁰⁰ NAB7061 has little inherent antimicrobial activity, but is a very potent synergist, while NAB739 has very potent antimicrobial activity (Table 1).¹⁰¹ Remarkably, this difference in activity between NAB739 and NAB7061 is attributed to the absence of one hydroxyl group in NAB7061 (See Figure 2B).⁹⁹ NAB739 has been reported to be exhibit generally moderate synergistic activity against wild-type strains with the exception of the A. *baumannii* strain indicated in Table 1.^{99,102} Interestingly, against *mcr*-positive strains, the loss of antimicrobial activity for NAB739 is accompanied by a significant increase in its synergistic activity, an effect also noted for colistin.^{102,103}

Name	Ref.	FICI	Pathogen	Antibiotic
PMBN	104	0.013*	E. coli	rifampicin
РМВО	104	0.013*	E. coli	rifampicin
РМВН	104	0.020*	E. coli	rifampicin
DAPB	104	0.043*	E. coli	rifampicin
SPR741	105	0.06	E. coli	rifampicin
NAB739	99	0.126	A. baumannii	rifampicin
NAB7061	99	0.055	E. coli	rifampicin

Table 1. Synergistic activity of polymyxin analogues.

*FICI calculated from MSC and MIC values reported in the cited reference.

2.2. Dilipidated polymyxins

Polymyxin analogues bearing an additional lipid tail have also been explored to test the hypothesis that additional hydrophobicity might enhance membrane interactions.¹⁰⁶ To generate these variants a variety of acyl tails were added to both amino groups of the N-terminal 2,4-diaminobutyric acid (Dab) residue of PMB (Figure 3).^{106,107} The introduction of simple propyl lipids as in analogue **1** led to a complete loss of inherent activity (\leq 128 µg/mL), while the analogues **2** and **5**, bearing larger, more hydrophobic groups, maintained moderate activity with MICs of 4-64 µg/mL against most Gram-negative bacteria.¹⁰⁶ Notably, the reduced inherent activity was accompanied by a higher synergistic potential (Table 2), indicating that these dilipidated analogues have an increased capacity to disrupt the OM.¹⁰⁶ Also, of note is the reported activity of analogues **2** and **5** against Gram-positive bacteria (MICs of 8-32 µg/mL) compared to colistin, which has no such activity (MICs of \leq 128 µg/mL).¹⁰⁶



Figure 3. Molecular structures of the dilipidated polymyxin analogues.

Name	Ref.	FICI	Pathogen	Antibiotic	Hemolytic activity ^a
Dilipid polymyxin 1	106	0.02	P. aeruginosa	rifampicin	<10% (1h)
Dilipid polymyxin 2	106	0.26	P. aeruginosa	novobiocin	<10% (1h)
Dilipid polymyxin 5	106	0.31	P. aeruginosa	rifampicin	<10% (1h)

Table 2. Synergistic activities of dilipidated polymyxin analogues.

^aNon-hemolytic is defined as <10% hemolysis compared to positive control with incubation times denoted in parentheses

2.3. Linear peptide-based synergists

In most reviews published on the topic of OM-targeting synergists, relatively little attention has been paid to linear peptides. Peptides have several drawbacks including poor metabolic stability, low bioavailability, potential immunogenicity, and high production costs.¹⁰⁸⁻¹¹⁰ To improve their metabolic stability, the structures of peptides can be adapted by a number of approaches including: peptidomimetics, lipidation, head-to-tail cyclization, N- and C-terminus modifications, backbone stereochemistry changes, and incorporation of unnatural amino acids.^{108,109,111-115} Improvements to the bioavailability of peptides have also been explored by applying formulation techniques, adjusting the properties of peptides, or linking them to a moiety to improve passage over the blood-brain barrier.¹⁰⁸⁻¹¹⁰ These advances, combined with the development of more economical methods for peptide synthesis support a future role for peptide-based therapeutics with a number of antimicrobial peptides already in (pre)clinical development.¹¹⁶⁻¹²⁰

In the literature an increasing number of peptides synergists that function through OM disruption have been reported (see Table 3). In some studies, panels of structurally similar peptides are screened, resulting in the identification of multiple hits with FICIs lower than 0.5. In such cases we have opted to select up to four of the most potent synergists to limit the number of peptides. Given that most peptide-based synergists are derived from specific lead proteins or antimicrobial peptides (AMPs), we have divided the linear peptide synergists accordingly, both in the discussion below and in the overview Table 3.

2.3.1. Cathelicidin antimicrobial peptides

The cathelicidins are AMPs that play an important role in the innate immune defense system of mammals and function by binding to bacterial membranes resulting in their destabilization and lysis.¹²¹⁻¹²⁴ In addition to their direct antibacterial activity, cathelicidins have also been found to play a role in recruiting immune cells to the site of infection as well as in LPS neutralization.^{56,121,125} The sole human cathelicidin-AMP gene encodes for hCAP-18 which cleaved by proteases into the active LL-37.¹²²⁻¹²⁴ The mature LL-37 peptide forms an amphipathic α -helix that upon interaction with bacterial cell surfaces is associated with a detergent-like antimicrobial activity.¹²⁶⁻¹²⁸ Recently, a truncated version of LL-37, termed FK16, was reported to potentiate the activity of vancomycin against P. aeruginosa (Table 3).129 Similarly, the Kuipers group showed that another LL-37 derived sequence termed KR-12-2, is able to synergize with azithromycin (and erythromycin, Table 3).¹³⁰ Further optimization of the peptide sequence resulted in peptide L11 which was also synthetized as the D-amino acid variant (D11) as a means of improving serum stability (Table 3).^{130,131} These peptides were screened in combination with multiple antibiotics against different Gram-negative strains and OM disruption assays verified their mode of action.130-132

In addition to the human cathelicidins, derivatives of cathelicidins from other mammals have also been screened for synergistic activity including novicidin (sheep), bactenectin (bovine), and indolicidine (bovine).^{121,133,134} Among these, only novicidin was reported to display potent synergy (Table 3).¹³³ In the case of bactenectin, which normally contains a disulfide bridge, a number of linear analogues have been prepared, including peptides G2, R2, amd DP7 which were found to exhibit OM disruption and exhibit moderate synergy (Table 3).^{134–136,137} In the case of indolicidin, structure-activity relationship (SAR) studies have led to the discovery of the synergists Indopt 10 and

CLS001 (Table 3). CLS001 is particularly effective and displays synergy with both vancomycin and azithromycin against multiple Gram-negative pathogens.^{134,137} Marketed under the name Omiganan, CLS001 is also much less hemolytic than indolicidin and is currently in clinical trials for the treatment of skin-related infections.^{101,138,139}

2.3.2. Lactoferrin-derived peptides

Lactoferrin is a multifunctional protein found in mammals and plays key roles in the human immune system. Lactoferrin has inherent activity against a range of bacterial, fungal, and viral pathogens and in the case of Gram-negative bacteria, it can disrupt the OM.⁸⁷ Based on the LPS binding region of lactoferrin known as LF11, the Martínez-de-Tejada group synthesized a series of LF11 homologues (Table 3) which were screened in combination with novobiocin for synergistic activity.¹⁴⁰ Based these findings a new generation of peptide synergists was designed using PEptide DEscriptors from Sequence (PEDES) software to predict OM permeabilizing sequences.¹⁴¹ The peptides thus obtained (i.e. peptide P2-16, Table 3) generally showed synergistic activity on par with the original series.¹⁴¹ Given the abundance of lactoferrins in other mammals, Svendsen and coworkers also investigated a series of peptides derived from bovine lactoferrin, both for antimicrobial activity and synergistic activity.¹⁴²⁻¹⁴⁵ This led to the identification of a 12-mer peptide termed P12, along with P15, a 15-mer containing biphenylalanine (Bip), and a longer 18-mer termed P18 all of which were found to exhibit moderate synergy with erythromycin when tested against E. *coli* (Table 3).

2.3.3. Thrombin-derived peptides

Thrombin is an enzyme that plays a critical role in coagulation and recent studies have also shown that certain thrombin-derived C-terminal peptides are capable of binding to LPS and neutralizing its toxic and inflammatory effects.¹⁴⁶ Given the capacity of PMB to also bind and neutralize LPS, our group was interested in assessing whether these thrombin-derived peptides might also exhibit the synergistic behavior of PMB. To this end we prepared a series of 12-mer thrombin-derived peptides and showed that a number of them are indeed potent synergists.¹⁴⁷ The most active synergist thus identified (Peptide **6**, Table 3) was further investigated by means of an alanine scan, leading to the discovery of more potent variants (Peptides **14** and **19**, Table 3). Notably, these peptides were found to be non-hemolytic and their synergistic activity was shown to extend to rifampicin, erythromycin, and novobiocin against multiple Gram-negative strains including those with *mcr*-mediated resistance.¹⁴⁷

2.3.4. Histatins

The histatins are a unique group of histidine-rich peptides found in human saliva that play roles in both defending against infection as well as in aiding wound-healing.¹⁴⁸ Among the most common histatins, the 24 amino acid Histatin **5** has been shown to bind Lipid A and has endotoxin neutralizing properties.¹⁴⁹ SAR studies with Histatin **5** led to the identification of a 12-mer sub region termed P-113 that exhibits antimicrobial activity against Gram-positive and Gram-negative bacteria.^{148,150-152} Further structural optimization to enhance the stability of P-113 led to analogues incorporating β -naphthylalanine (Nal) and Bip residues to yield Nal-P-113 and Bip-P-113 and wherein the 4th, 5th, and 12th histidine resides were replaced by Nal or Bip respectively (Table 3).¹⁵² Bip-P-113 and Nal-P-113 exhibit antimicrobial activity, improved serum proteolytic

stability, and were also found to permeabilize LPS containing large unilamellar vesicles used to model the Gram-negative OM.^{152,153} These findings prompted investigation of vancomycin potentiation by Bip-P-113 and Nal-P-113 revealing both to exhibit moderate synergy.¹⁵⁴ However, a notable drawback of Bip-P-113 and Nal-P-113 is their significantly increased hemolytic activity relative to P-113.¹⁵²

2.3.5. Other Natural AMPs, their hybrids, and derivatives

A number of other naturally occurring AMPs have been reported to potentiate antibiotics that are otherwise excluded by the OM. These AMPs are all polycationic and include: buforrin II, esculentin 1b, sphistin, HE2α, HE2β2, anoplin, magainin II, and cecropin A (Table 3).¹⁵⁵⁻¹⁵⁹ The sources of these AMPs are diverse and include toads, wasp venom, or even the human male reproductive tract.^{157,158,160} The AMPs here discussed have all been reported to disrupt the OM,^{156,158,161-163} bind to LPS, and/or show endotoxin neutralizing activity.^{155,159,164} In general, these AMPs exhibit modest FICIs (0.2-0.36) which has also led to interest in hybrids and derivatives with enhanced synergistic activity. For example, Park and coworkers developed a series of hybrid peptide synergists, termed CAME, CAMA, and HPMA containing sequences derived from crecopin A, magainin II, and melittin (Table 3).^{164,165} Other approaches include truncation as in the case of the lipopeptide AMP Tridecaptin A₁ (TriA₁), which exhibits potent inherent anti-Gramnegative activity, were found to be effective synergists. Notably, removal of the TriA₁ Nterminal lipid vielded H-TriA₁ which was found to be much less active as an antibiotic but exhibited very potent synergism when combined with rifampicin resulting in an FICI of 0.002 against E. coli (Table 3).^{166,167} Like the tridecaptins, the recently discovered paenipeptins contain a number of Dab residues and have been subject to SAR studies.¹⁶⁸ These efforts led to the discovery of a potent paenipeptin inspired synergist termed SLAP-S25 which effectively potentiates the activity of rifampicin and vancomycin against E. coli (Table 3).¹⁶⁹ In addition to OM disruption, the binding of SLAP-S25 to LPS and phosphatidylglycerol (PG) was established, suggesting that SLAP-S25 is also an inner membrane disruptor.¹⁶⁹ This was confirmed by dose-dependent uptake of propidium iodide and release of cellular contents in cells treated with SLAP-S25.169 Notably, SLAP-S25 was also demonstrated to effectively enhance the in vivo activity of colistin against a colistin-resistant strain of E. coli in both G. mellonella and mouse infection models.¹⁶⁹

Originally isolated from wasp venom, anoplin is one of the smallest known amphipathic, α -helical AMPs.^{158,160} Multiple SAR investigations have been performed to improve its antimicrobial activity and stability.^{170–174} A recent study with anoplin reported the systematic introduction of tryptophan and lysine residues to determine the optimal hydrophobicity, amphipathicity, and number of positive charges required for antibacterial activity and minimal cytotoxicity.¹⁵⁸ A number of these analogues were also found to be synergistic when combined with rifampicin (see peptides A13, A17, and A21 in Table 3) via a mechanism involving OM disruption.¹⁵⁸ A similar study with Mastoparan-C, a peptide found in the venom of the European hornet, led to the identification of an analogue termed L7A (Table 3) which also displays synergy via OM perturbation.¹³⁸ Another example of a synergist derived from a toxic peptide is myotoxin II which is isolated from certain snake venoms. Studies with peptide sequences based on the Cterminus of myotoxin II resulted in the generation of peptide S1 (Table 3) which showed a good balance of synergy with vancomycin and low hemolytic activity.^{175,176} Attempts at further improving the S1 peptide involved the introduction of Nal residues at the Cterminus to generate S1-Nal which exhibited enhanced synergistic activity and S1-NalNal which also exhibited enhanced synergistic activity but at the expense of increased hemolytic activity (Table 3).^{177–180}

2.3.6. Peptide synergists discovered via library screening

Guardabassi and coworkers recently reported the development and validation of an assay meant to enable high throughput screens for identifying OM disruption agents.¹⁸¹ To this end they applied a whole-cell screening platform that allows for detection of OM permeabilization in E. *coli* based on the signal generated by a chromogenic substrate reporter for a cytoplasmic β -galactosidase. To validate the assay, a library of peptides and peptidomimetics was screened which generated a notable hit termed peptide 79 that showed potentiation of various antibiotics at therapeutically relevant levels (Table 3).¹⁸¹ In a follow-up study the same group went on to develop two improved synergists termed Peptides **1** and **2** along with the all D-amino acid variants which were also found to effectively potentiate rifampicin against K. *pneumoniae* (Table 3).^{70,181}

2.3.7. Peptide synergists from phage display

Phage display techniques have also been applied to identify novel peptides capable of interaction with the OM. In one such investigation, a phage library displaying random 12mer peptides was screened for the ability to bind to the cell surface of Gram-negative bacteria.¹⁸² Specificity for the Gram-negative OM was ensured by removal of peptides binding to Gram-positive bacteria by pre-incubation of the library with *Staphylococcus aureus*.¹⁸² This approach led to the identification of a peptide termed EC5, that exhibits moderate antibacterial activity against *E. coli* and *P. aeruginosa*, with MICs in the range of 8-16 µg/mL against both.¹⁸² The EC5 peptide was shown to cause OM disruption and cytoplasmic membrane depolarization while exhibiting very little hemolytic activity.¹⁸² Subsequent synergy studies showed that the peptide was also capable of potentiating the activity of erythromycin, clarithromycin, and telithromycin against P. *aeruginosa*.¹³⁰

2.3.8. Rationally designed peptide synergists

Inspired by the structure of DAPB (see Figure 2), Vaara and coworkers designed a series of linear and cyclic peptides for evaluation as synergists.¹⁸³ The sequences of these peptides were based on an ABB_n motif in which A is a basic amino acid and B a hydrophobic residue (see Peptides 4 and 5 in Table 3).¹⁸³ Cyclic peptides were also prepared bearing a similar AB_n motif (see Peptide 7, Table 3).¹⁸³ All peptides were screened for synergistic activity with erythromycin, rifampicin, novobiocin, and fusidic acid with the rifampicin combinations being the most potent (Table 3).¹⁸³ While the synergistic activity of these peptides could be correlated to their OM disrupting activity, the effect was not specific given their high hemolytic activity.¹⁸³

De novo-designed peptides have also been explored as a means of generating novel synergists. To this end the Sahal group developed a number of peptides incorporating key elements found in AMPs and synergists including amphipathicity, positive charge, and helical conformation.^{184,185} Of note was the introduction of α , β -didehydrophenylalanine (Δ F) into the peptides as a means of constraining the helical conformation of the peptides.¹⁸⁶⁻¹⁸⁸ Using this approach two peptides termed Δ Fm and Δ Fmscr were identified as effective synergists with low toxicity towards mammalian cells (Table 3).¹⁸⁷

In another recent approach to identifying novel peptide synergists, Yu and colleagues reported the construction of a small library wherein amphipathic peptides where subjected to a proline-scanning strategy to generate novel hinged peptides.¹⁹⁴ Such proline hinged peptides are reported to have lower toxicity towards mammalian cells given that their membrane binding is reduced compared to conventional AMPs with a high α -helical conformation.¹⁸⁹ Proline scanning of two model peptides, LK (LKKLLKLLKLLKL) and KL (KLLKLLKKLLKL), provided a set of peptides that were screened for synergistic activity with the four most potent peptides displayed in Table 3. The peptides were also screened for hemolysis which led to identification of peptide KL-L9P as the most promising hit. This peptide was subsequently shown to permeabilize the OM, as evidenced by uptake of N-phenylnaphthalen-1-amine (NPN), and was also found to bind LPS without disturbing the inner membrane.¹⁹⁰ Mouse sepsis studies were also performed to evaluate the *in vivo* synergistic effect of KL-L9P, which displayed a significant potentiation of a number of clinically used antibiotics and resulted in improved overall survival.¹⁹⁰

In another recently reported study, Zeng et al. described the application of rational design approaches to generate novel helix-forming AMPs based on cytolytic peptide toxins produced by highly virulent strains of S. *aureus*.^{191,192} The peptides thus obtained were shown to have improved physicochemical properties and antibacterial activity, while maintaining low hemolytic activity and cytotoxicity. Among the 16-mers thus generated, two peptides, termed zp12 and zp16, were also found to exhibit potent synergy (Table 3). Notable in this regard is the finding that peptide zp16 specifically potentiates the effect of the glycopeptide antibiotics vancomycin and teicoplanin against highly pathogenic *K. pneumonia*.¹⁹² The vancomycin-zp16 combination exhibits negligible toxicity *in vitro* and *in vivo* and mechanistic studies indicate that zp16 enhances vancomycin's cell permeability, leading to markedly reduced biofilm formation and rapid bactericidal effect.¹⁹²

In 2022 the group of Ni reported the potentiation of multiple antibiotics, including rifampicin, by two rationally designed peptides named K4 and K5 (Table 3).¹⁹³ These peptides were selected from a library of variants all containing a repeating motif (WRX)_n wherein X represents I, K, L, F, and W.¹⁹⁴ Hemolysis and cytotoxicity assays led to the selection of peptides K4 and K5 as leads.¹⁹⁴ The finding that these peptides permeabilize the OM resulted in follow-up studies on the potentiation of antibiotics against Gram-negative bacteria.¹⁹³ Apart from synergy, a 15-day resistance assay was also performed for the K4 and K5 peptides, with or without antibiotics, showing no significant resistance development.^{193,194} Also of note, while the inherent activity of K4 was found to be comparable to PMB, K4 was reported to display significantly less toxicity.¹⁹⁴

2.4. Lipopeptide synergists

In addition to the exclusively peptide-based synergists described above, lipopeptides have also been explored as synergists. We here cover examples of lipopeptides that do not possess potent inherent antibacterial activity but rather have the capacity to effectively potentiate the activity of other antibiotics. A recent example are the synthetic paenipeptins developed by Huang and coworkers.¹⁹⁵ The design of these lipopeptides is based on peptides produced by *Paenibacillus sp.* strain OSY-N that contain a number of unnatural and D-amino acids. Using low hemolytic activity as a selection criterion, a subset of these lipopeptides were selected and screened for synergistic activity. This led

to the identification of paenipeptins 1, 9, 15, and 16 which exhibit potent synergy (Table 3).^{195,196} These lipopeptides were further shown to have OM disrupting activity as indicated by the NPN assay. Furthermore, in an murine thigh infection model, paenipeptin **1** was shown to effectively potentiate the *in vivo* activity of both clarithromycin and rifampin against polymyxin-resistant E. coli.¹⁹⁶

Small cationic lipopeptides have also been explored as synergists with the aim of identifying smaller, less hemolytic agents. To this end Schweizer and coworkers recently reported a series of "dilipid ultrashort cationic lipopeptides" (dUSCLs) capable of enhancing the activity of clinically used antibiotics against Gram-negative bacteria. The design of these dUSCLs consists of lysine rich tetrapeptides bearing various lipids at the N-terminal residue as illustrated in Figure 4A. It was found that dUSCLs bearing lipids of \geq 11 carbon atoms caused significant hemolysis. However, analogues with slightly shorter lipid were found to achieve an acceptable balance of low hemolytic activity and synergistic activity. This led to the identification of dUSCLs **2** and **6** as the most promising synergists (Table 3) capable of sensitizing a range of Gram-negative strains to various antibiotics. The authors also noted that in addition to permeabilizing the OM, the dUSCLs may also function by indirectly disrupting antibiotic efflux.¹⁹⁷

The Schweizer group also recently reported a series of ultrashort tetrabasic lipopeptides (UTBLPs) synergists.¹⁹⁸ These compounds were specifically prepared to assess the effect of lysine N-ζ-methylation on the potentiation of antibiotics and was inspired by reports suggesting N-methylation can lead to reduced hemolysis, increased proteolytic stability, and improved antibacterial activity.¹⁹⁹⁻²⁰¹ Compared to the dUSCLs, UTBLP **5** and **6** contain an extra lysine while an octanoyl group was employed as the lipophilic moiety (Figure 4B).^{197,198} Methylation of the lysine side-chain resulted in a reduction of potentiation for rifampicin and novobiocin in both wild-type and resistant Gramnegative strains.¹⁹⁸ A correlation between the number of methyl groups and loss of activity was seen, while the increase in NPN fluorescence of the tri-methylated UTBLP were on par their un- or mono-methylated analogues.¹⁹⁸

2.5. Lipopeptidomimetic synergists

The Schweizer group also expanded the scope of their dUSCLs by exploring a series of dilipid ultrashort tetrabasic peptidomimetics (dUSTBPs) as a proteolytically stable alternative.²⁰² In a focused SAR study they prepared dUSTBPs consisting of three basic amino acids separated by a molecular scaffold, bis(3-aminopropyl)glycine, along with ligation to simple fatty acids (see Figure 4C).²⁰² This led to identification of a number of dUSTBPs capable of potentiating the activity of several antibiotics against pathogenic Gram-negative bacteria while exhibiting low hemolytic activity (Table 3). In particular, dUSTBP **8**, consisting of three L-arginine units and a dilipid of 8 carbons long, was found to potentiate novobiocin and rifampicin against multidrug-resistant (MDR) clinical isolates of P. *aeruginosa*, A, *baumannii*, and *Enterobacteriaceae* species.²⁰²

In 2007 Mor and coworkers introduced the oligo-acyl-lysyls (OAKs) as peptidomimetics of the antimalarial peptide dermseptin S3 (Figure 4D) that were initially evaluated primarily for antimicrobial activity.²⁰³⁻²⁰⁵ Among the first series of analogues prepared, OAK $C_{12(\omega7)}$ was found to adhere to the OM with minimal insertion and its antibacterial activity against Gram-negative bacteria improved in combination with ethylenediaminetetraacetate (EDTA).²⁰⁵⁻²⁰⁷ The introduction of a double bond in OAK $C_{12(\omega7)}$ resulted in significant reduction of hemolytic activity compared to OAK C_{12} while 20

the slightly less hydrophobic OAK C_{10} and OAK C_8 analogues also showed no hemolytic activity.^{205,208} In 2013 these four OAKs, as well as the more recently described OAK $C_{14(\omega5)}OOc_{10}O$ containing ornithine instead of lysine (Figure 4D), were reported to potentiate rifampicin against Gram-negative bacteria (Table 3).^{208,209} Interestingly, the synergistic activity of the OAKs was maintained in human plasma but was suppressed by addition of anti-complement antibodies, suggesting that these compounds sensitize Gram-negative bacteria to the action of antibacterial innate immune mechanisms.²⁵²



Figure 4. Lipopeptide and lipopeptidomimetic synergists. Representative structures of A) dilipid ultrashort cationic lipopeptides (dUSCLs); B) Ultrashort tetrabasic lipopeptides (UTBLPs); C) dilipid ultrashort tetrabasic peptidomimetics (dUSTBPs); and D) oligo-acyl-lysyls (OAKs).

Name	Ref	Peptide sequence ^a	FICI	Pathogen	Antibiotic	Hemolytic activity ^b
		Cathelic	idin deriv	ed peptides		
FK16	129	FKRIVQRIKDFLRNLV	0.25	P. aeruginosa	vancomycin	<10% (1h)
KR-12-a2	130,210	KRIVQRIKKWLR-NH2	0.156	P. aeruginosa	erythromycin	<10% (1h)
L-11	131	RIVQRIKKWLR-NH2	0.070	A. baumannii	vancomycin	NR
D-11	131,132	rivqrikkwlr-NH2	0.032	A. baumannii	rifampicin	<10% (1h)
Novicidin	133	KNLRRIIRKGIHIIKKY F	0.018	E. coli	rifampicin	<10% (1h)
G2	134	RGARIVVIRVAR-NH2	0.38	P. aeruginosa	erythromycin	NR
R2	134	RRARIVVIRVAR-NH2	0.27	P. aeruginosa	erythromycin	NR
DP7	137,211	VQWRIRVAVIRK	0.25	P. aeruginosa	vancomycin	<10% (1h)
Indopt 10	134	ILKWKIFKWKWFR-NH2	0.38	P. aeruginosa	erythromycin	NR
CLS001	137,139	ILRWPWWPWRRK-NH2	0.28	P. aeruginosa	vancomycin	10% (30 min)
		Lactofer	rin deriv	ed peptides		
P10	140	FWQRNIRKVKKK-NH2	0.113	P. aeruginosa	novobiocin	<10% (1h)
P14	140	FWQRNIRKVKKKI-NH2	0.113	P. aeruginosa	novobiocin	<10% (1h)
P22	140	RFWQRNIRKYRR-NH2	0.431	P. aeruginosa	novobiocin	<10% (1h)
P2-16	141	FWRNIRIWRR-NH2	0.116	P. aeruginosa	novobiocin	NR
P12	144,212	RRWQWRMKKLGA	0.43	E. coli	ery thromycin	<10% (2h)
P15	144	FK-Bip- RRWQWRMKKLGA°	0.38	E. coli	erythromycin	NR
P18	144	PAWFKARRWAWRMLKKA A	0.38	E. coli	erythromycin	NR
		Thromb	oin derive	d peptides		
Peptide 6	147	VFRLKKWIQKVI-NH2	0.094	E. coli	rifampicin	<10% (20h)
Peptide 14	147	VFRLKKAIQKVI-NH2	0.078	E. coli	erythromycin	<10% (20h)
Peptide 19	147	VFRLKKWIQKVA-NH2	0.078	E. coli	rifampicin	<10% (20h)
		Histati	in derived	l peptides		
Nal-P-113	152,154	Ac-AKR-Nal-Nal- GYKRKF-Nal-NH2 ^d	0.38	E. coli	vancomycin	>10% (1h)
Bip-P-113	152,154	Ac-AKR-Bip-Bip- GYKRKF-Bip-NH2°	0.38	E. coli	vancomycin	>10% (1h)
		Other Natural AMP	s, their h	ybrids, and der	ivatives	
Buforin II	155,213	TRSSRAGLQFPVGRVHR LLRK	0.312	A. baumannii	rifampicin	<10% (1h)
Esculentin 1b	156,214	GIFSKLAGKKLKNLLIS G-NH2	0.36	E. coli	erythromycin	>10% (1h)
ΗΕ2α	157,161	VHISHREARGPSFRICV GFLGPRWARGCSTGN	0.3	E. coli	rifampicin	<10% (1h)
ΗΕ2β2	157,161	ICRLFFCHSGTGQQHRQ RCG	0.2	E. coli	rifampicin	<10% (1h)
Anoplin	158	GLLKRIKTLL	0.3125	P. aeruginosa	rifampicin	<10% (1h)

Table 3. Overview of linear peptide-based synergists (compound names provided as given in the cited literature references).

Magainin II	159,213	GIGKFLHAAKKFAKAFV AEIMNS-NH2	0.312	P. aeruginosa	rifampicin	>10% (1h)			
Cecropin A	159,164	GIIKAGPAVAVVGQATQ IAK-NH2	0.312	P. aeruginosa	rifampicin	<10% (1h)			
CAME	215,216	KWKLFKKIGIGAVLKVL TTG-NH2	0.375	A. baumannii	erythromycin	<10% (1h)			
CAMA	215,216	KWKLFKKIGIGKFLHSA	0.25	A. baumannii	erythromycin	<10% (1h)			
НРМА	215,217	AKKVFKRLGIGKFLHSA KKF-NH2	0.313	A. baumannii	erythromcyin	<10% (1h)#			
H-TriA1	166,167	v-dab-Gsw-Dab- dab-FEI-alle-A ^{e,f}	0.002	E. coli	rifampicin	<10% (30 min)#			
SLAP-S25	169	Dab-fL-Dab-vLA- NH2	0.031	E. coli	rifampicin	<10% (1h)			
A13	158	GWWKRIKTWW	0.375	K. pneumoniae	rifampicin	<10% (1h)			
A17	158	KWWKRWKKWW	0.3125	P. aeruginosa	rifampicin	>10% (1h)			
A21	158	KWWKKWKKWW	0.3125	K. pneumoniae	rifampicin	<10% (1h)			
L7A	138	LNLKALAAVAKKIL- NH2	0.31	E. coli	rifampicin	<10% (1h)			
S1	177,180	Ac-KKWRKWLAKK-NH2	0.38	A. baumannii	vancomycin	<10% (1h)#			
S1-Nal	177,180	Ac-KKWRKWLAKK- Nal-NH2	0.27	A. baumannii	vancomycin	<10% (1h)#			
S1-Nal-Nal	177,180	Ac-KKWRKWLAKK- Nal-Nal-NH2	0.27	A. baumannii	vancomycin	>10% (1h)			
Peptide synergists via library screening									
Peptide 79	176,181	KKWRKWLKWLAKK-NH2	0.14	E. coli	rifampicin	<10% (1h)			
Peptide 1	70,218	KLWKKWKKWLK-NH2	0.02	K. pneumoniae	rifampicin	<10% (1h)			
Peptide 2	70,184	GKWKKILGKLIR-NH2	0.04	K. pneumoniae	rifampicin	<10% (1h)			
Peptide D1	70	klwkkwkkwlk-NH2	≤0.03	K. pneumoniae	rifampicin	NR			
Peptide D2	70	gkwkkilgklir-NH2	≤0.04	K. pneumoniae	rifampicin	NR			
		Peptide syne	rgists fro	m phage displa	у				
EC5	130,182	RLLFRKIRRLKR	0.266	P. aeruginosa	erythromycin	<10% (24h)			
		Des	signed pe	ptides					
Peptide 4	183	KFFKFFKFF	0.03	E. coli	rifampicin	>10% (30 min)			
Peptide 5	183	IKFLKFLKFL	0.06	E. coli	rifampicin	NR			
Peptide 7	183	C KFKFKFKF C	0.20	E. coli	rifampicin	NR			
ΔFm	187	Ac-G Δ FRK Δ FHK Δ FWA-NH2 ^g	0.3	E. coli	rifampicin	<10% (1h)			
∆Fmscr	187	Ac-GAFRKAFKAAFWH- NH2 ^g	0.14	E. coli	rifampicin	<10% (1h)			
LK-L8P	219	LKKLLKLPKKLLKL- NH2	0.18	E. coli	erythromycin	<10% (4h)			
LK-L11P	219	Ac- LKKLLKLLKKPLKL- NH2 Ac-	0.47	E. coli	erythromycin	<10% (4h)			
KL-L6P	219	LKKLLPLLKKLLKL- NH2	0.33	E. coli	erythromycin	>10% (4h)			

		Ac-				
KL-L9P	219	LKKLLKLLPKLLKL- NH2	0.12	E. coli	erythromycin	<10% (4h)
zp12	192	GIKRGIIKIIKRIKRI- NH2	0.25	K. pneumoniae	vancomycin	NR
zp16	192	GIKRGIIKIIRRIKRI- NH2	0.06	K. pneumoniae	vancomycin	<10% (1h)
K4	193,194	WRKWRKWRKWRK-NH2	0.2	K. pneumoniae	rifampicin	<10% (1h)
К5	193,194	WRKWRKWRKWRKWRK- NH2	0.2	E. coli	rifampicin	<10% (1h)
		Lipop	eptide Sy	nergists		
Paenipeptin 1	195,196	C ₆ -Dab-I-Dab-fL- Dab-vLS-NH2 ^h	0.125*	E. coli	rifampicin	<10% (30 min)
Paenipeptin 9	195	C ₈ -Dab-I-Dab-fL- Dab-vL-Dab-NH2 ⁱ	≤0.03*	K. pneumoniae	rifampicin	<10% (30 min)
Paenipeptin 15	5 ¹⁹⁵	Cbz-Dab-I-Dab-fL- Dab-vLS-NH2 ^j	≤0.03*	K. pneumoniae	rifampicin	<10% (30 min)
Paenipeptin 16	B ¹⁹⁵	Cha-Dab-I-Dab-fL- Dab-vLS-NH2 ^k	0.06*	K. pneumoniae	rifampicin	<10% (30 min)
dUSCL 2	197	$C_{10}-K(C_{10}) KKK-NH2^1$ (Figure 4A)	0.07	P. aeruginosa	rifampicin	<10% (1h)
dUSCL 6	197	$C_{10}-K(C_{10}) KGK-NH2^1$ (Figure 4A)	0.25	P. aeruginosa	rifampicin	<10% (1h)
UTBLP 5	198	C ₈ -K(C ₈)KKKK-NH2 ⁱ (<i>Figure 4B</i>)	≥0.016	P. aeruginosa	novobiocin	NR
UTBLP 6	198	C_8 - K (C ₈) K (Me) K (Me) K (Me) K (Me) - NH2 ⁱ (Figure 4B)	¹ 0.047	A. baumannii	rifampicin	NR
		Lipopepti	domimet	ic Synergists		
dUSTBP 2	202	Figure 4C	≥0.250	P. aeruginosa	rifampicin	<10% (1h)
dUSTBP 5	202	Figure 4C	≥0.125	P. aeruginosa	rifampicin	<10% (1h)
dUSTBP 8	202	Figure 4C	≥0.002	A. baumannii	novobiocin	<10% (1h)
OAK C ₁₂ (ω7)	208	Figure 4D	≤0.073*	E. coli	rifampicin	>10% (3h)
OAK C ₁₂	208	Figure 4D	≤0.211*	E. coli	rifampicin	>10% (3h)
OAK C ₁₀	208	Figure 4D	≤0.036*	E. coli	rifampicin	<10% (3h)#
OAK C ₈	208	Figure 4D	≤0.078*	E. coli	rifampicin	<10% (3h)#
OAK C14(ω5)OOc10O	209	Figure 4D	0.20*	K. pneumoniae	rifampicin	<10% (3h)#

^aLower case letters indicate D-amino acids; ^bNon-hemolytic is defined as <10% hemolysis compared to positive control with incubation times denoted in parentheses, NR denotes no data reported; ^cBip, biphenylalanine; ^dNal, β-naphthylalanine; ^eDab, 2,4-diaminobutyric acid; ^falle, D-*allo*-isoleucine; ^gΔF, α ,β-didehydrophenylalanine; ^hC₆, hexanoyl; ⁱC₈, octanoyl; ^jCbz, benzyloxycarbonyl; ^kCha, cyclohexylalanyl; ^lC₁₀, decanoyl; [#]denotes that the concentration tested was lower than 100 µg/mL; ^{*}FICI calculated from MSC and MIC values reported in the cited literature reference.

3. Cationic steroids

In 1993 the isolation of squalamine from tissues of the dogfish shark *Squalus acanthias* was reported.²²⁰ Squalamine consists of a steroid core linked to a spermidine moiety (Figure 5A) and was found to exhibit broad antimicrobial activity.²²⁰ Later, it was established that squalamine disrupts membranes and is also hemolytic. Notably, investigations into its synergistic activity showed that it was unable to potentiate erythromycin against wild-type strains, showing an effect only against a P. *aeruginosa* strain overproducing MexAB-OprM efflux pumps (See Table 4).^{221,222} A few years after its discovery, novel squalamine mimics (SMs) were synthesized in an attempt to enhance antibacterial activities (Figure 5B).²²³ These synthetic analogues consist of cholic and deoxycholic acid as the steroid backbone to which a spermidine chain is appended. This approach resulted in the identification of analogue SM-7, which was found to potentiate rifampicin against multiple Gram-negative bacteria (Table 4).²²³ However, like squalamine, SM-7 also possesses significant hemolytic activity limiting its potential for systemic use.²²³

In another approach, the Savage group also employed the cholic acid backbone but with the aim of mimicking polymyxins through the amphiphilic positioning of positive charges (Figure 5C and 5D).^{224,225} In doing so, a variety of cationic steroids were developed and screened both for inherent antimicrobial activity as well as the capacity to potentiate antibiotics against Gram-negative bacteria.²²⁵⁻²³³ The orientation of the hydroxyl groups of cholic acid backbone provide convenient functionalities for the incorporation of positively charged moieties via formation of ether (Figure 5C) or ester (Figure 5D) linkages. Among the ether-linked series, an analogue bearing three carbon atom spacers between the steroid and the primary amine groups, along with an N-benzylated tertiary amino group at the C24 position (analogue I, Figure 5C), was found to exhibit both inherent antimicrobial and synergistic activity.²²⁵ Interestingly, replacement of the lipophilic N-benzyl moiety with a hydroxyl group led to analogue II which showed a significant reduction of inherent activity while maintaining a strong ability to potentiate the activity of erythromycin against E. coli.^{224,225} The decreased lipophilicity of analogue II also reduced the hemolytic activity seen with analogue I (Table 4). Follow-up studies revealed that conversion of the free hydroxyl group at the C24 position to the propyl ether as in analogue III significantly increased hemolytic activity.^{226,227} Notably, addition of a terminal amino group to the propyl ether moiety provided analogue IV which exhibited significantly reduced hemolysis relative to analogue III while maintaining effective synergistic activity (Table 4).²²⁸ A series of ester linked analogues were also prepared by the Savage group (Figure 5D), wherein compounds V,VI, and VII exhibited synergistic activity comparable to the corresponding ether variants (Table 4).^{229,230} Amide analogues were also explored, however, they exhibited a significant lower potentiation of erythromycin, presumably due to conformational constraints relative to the more active esters.229

In addition to the polycationic steroids described above, steroid-peptide hybrids have also been explored as synergists.²³³⁻²³⁵ In a one case, Bavikar *et. al* reported a series of hybrids wherein simple tetrapeptides were coupled to cholic acid in an attempt to mimic the squalamine tail (Figure 5E).²³⁵ As indicated in Table 4, these steroid-peptide hybrids exhibit potent synergy with erythromycin against *E. coli*. While the hemolytic activity of these compounds was not reported, they were described as having low cytotoxicity towards HEK293 and MCF-7 cells.²³⁵



Figure 5. Overview of the synergistic steroids A) Squalamine; B) squalamine mimic SM-7; C) polycationic cholic acid ether linked steroid synergists; D) polycationic cholic acid ester linked steroid synergists; and E) steroid-peptide hybrids.

Name	Ref.	FICI	Pathogen	Antibiotic	Hemolytic activity ^a
Squalamine	220,222	0.35*	P. aeruginosa	erythromycin	>10% (10 min)
SM-7	223	0.063	K. pneumoniae	rifampicin	<10% (24h)
		Polycationi	c cholic acid analo	gues	
Ether linked					
I	225,226	0.035	K. pneumoniae	rifampicin	>10% (24h)
II	226	0.029	K. pneumoniae	novobiocin	<10% (24h)
III	226	0.022	K. pneumoniae	novobiocin	>10% (24h)
IV	228	0.13	K. pneumoniae	rifampicin	<10% (24h)
Ester linked					
v	229	0.057*	E. coli	erythromycin	NR
VI	229	0.064*	E. coli	erythromycin	NR
VII	230	0.176*	E. coli	erythromycin	<10% (24h)
Steroid-peptide	hybrids				
VIII	235	0.099	E .coli	erythromycin	NR
IX	235	0.093	E .coli	erythromycin	NR
x	235	0.078	E .coli	erythromycin	NR

Table 4. Overview of synergists based on cationic steroids.

^aNon-hemolytic is defined as <10% hemolysis compared to positive control with incubation times denoted in parentheses, NR denotes no data reported; *FICI calculated from MSC and MIC values reported in the cited literature reference.

4. Non-steroid small molecule synergists

4.1. Synergists based on approved drugs

Recently, Brown and coworkers reported an innovative screening platform for the identification of non-lethal, OM-active compounds with potential as adjuvants for conventional antibiotics.²³⁶ They applied their screen to a library of 1,440 previously approved drugs which resulted in the identification of three hits. Among the three hits identified, the antiprotozoal agent pentamidine (Figure 6A), was subsequently found to display the highest synergistic potency (Table 5).²³⁶ Notably, while pentamidine's OM targeting mechanism was found to be driven by interaction with LPS, *mcr*-resistance did not affect its synergistic potential.²³⁶ The potentiation of novobiocin by pentamidine was also established *in vivo* against wild-type and resistant A. *baumannii*.²³⁶ Subsequently, a focused SAR study using commercially available bis-amidines similar in structure to pentamidine led to the identification of compound **9** as an even more potent synergist (Figure 6A and Table 5).²³⁶

Inspired by these findings, our group recently undertook a broad SAR investigation wherein a number of structurally unique bis-amidines were synthesized and evaluated as synergists.²³⁷ Specifically we focused our attention on the length and rigidity of the linker motif as well as the geometry of the amidine groups on the aromatic rings. In addition to assessing the synergistic activity of the new bis-amidines prepared, we also performed hemolysis assays with each compound to ascertain OM selectivity. Given the potent synergy previously reported for bis-amidine **9**²³⁶ we also synthesized it to use as a benchmark. Among the compounds prepared in our study, bis-amidine **21**, containing an *ortho*-substituted benzene linker, was found to be significantly more synergistic than pentamidine and displayed no hemolytic activity (Figure 6A and Table 5).²³⁷ We also found that the introduction of additional aromatic groups to the linker, such as in compound **38**, led to further enhancement of synergy, however, this came at the costs of increased hemolytic activity (Table 5). Interesting, our studies also revealed benchmark bis-amidine **9** to be hemolytic. These findings further highlight the importance of assessing OM selectivity when pursuing synergists.²³⁷

The Brown group also recently reported a follow-up SAR study aimed at further enhancing the therapeutic potential of bis-amidines synergists.²³⁸ Similar to our own SAR study, the rigidity, conformation flexibility, and lipophilicity were further explored. In addition, the role of chirality and charge were also investigated.²³⁸ A key focus of this study was to identify bis-amidine synergists with improved off-target effects relative to pentamidine, especially the QT prolongation resulting from its effect on the hERG ion channel.²³⁸⁻²⁴⁰ This led to compound **P35** which was shown to have the same synergistic mode of action as pentamidine, displayed a strong potentiation of novobiocin, and no hemolytic activity (Table 5). Furthermore, compound **P35** outperformed pentamidine on multiple levels: an improvement in cytoxicity, a higher efficacy in a mouse infection model, and reduced hERG inhibition.²³⁸

Wang and coworkers also recently reported a study wherein the Prestwick Chemical Library, comprising 158 FDA-approved drugs, was assessed for compounds exhibiting synergy with doxycycline.²⁴⁰ This led to the finding that metformin, a commonly prescribed anti-diabetic agent (Figure 6B), effectively potentiates vancomycin as well as tetracycline antibiotics, particularly doxycycline and minocycline, against MDR S. *aureus*, *Enterococcus faecalis*, E. *coli*, and *Salmonella enteritidis*.²⁴⁰ The capacity for 28

metformin to disturb the OM was assessed using the NPN assay, revealing an increase in *E. coli* OM permeability in a dose-dependent manner. Of particular note was the finding that metformin was also able to fully restore the activity of doxycycline in animal infection models.²⁴¹



Figure 6. Representative structures of reported A) bis-amidine synergists; and B) metformin.

4.2. Small molecule synergists via high throughput screening

Following the success in applying their OM perturbation reporter assay to identify pentamidine as a potent synergist, the Brown group applied the same approach in a much larger high throughput screening (HTS) campaign with a library of ca. 140 000 synthetic compounds.^{236,242} This in turn led to the identification of 39 hits that were subsequently screened for synergistic activity with rifampicin.²⁴² Among these hits MAC-0568743 and liproxstatin-1 and (Figure 7A) were found to be particularly active synergists (Table 5).²⁴² Both compounds were found to potentiate the activity of the Gram-positive-targeting antibiotics rifampicin, novobiocin, erythromycin, and linezolid. This potentiation was further shown to be due to selective disruption of the OM, driven by interactions with LPS, and neither compound impacted the inner membrane.²⁴²

In another recently reported campaign, Datta and coworkers screened a focused library of 3000 drug-like compounds for antibiotic synergy using a whole-cell-based phenotypic assay.²⁴³ This led to the identification of a series of azaindoles that potentiate the MICs of macrolides, novobiocin, and rifampicin, by 100–1000-fold vs. Gram-negative bacteria. Optimization studies led to compounds BWC-Aza1 and BWC-Aza2 (See Figure 7B) both of which were screened for synergistic activity with an extensive panel of antibiotics against *E. coli* (Table 5). The OM permeabilizing activity of the azaindoles was also probed using the NPN assay revealing dose-dependent disruption.²⁴³

4.3. Small molecule polyamine synergists

In recent years the polyamines norspermine and norspermidine have been explored as starting points for the development of antibacterial and antibiofilm agents.^{244,245} Building on this work, the Haldar group recently reported the development of D-LANA-14 comprised of a norspermidine core linked to two D-lysine along with conjugation to a tetradecanoyl chain at the central secondary amine (Figure 7C).²⁴⁶ D-LANA-14 showed potent synergy with tetracycline or rifampicin against meropenem-resistant A. *baumannii* and P. *aeruginosa* clinical isolates (Table 5) and importantly was also found to disrupt established biofilms formed by these pathogens.²⁴⁶ D-LANA-14 was shown to perturb the OM by means of the NPN assay and importantly also found to exhibit potent *in vivo* activity when combined with rifampicin resulting in a significant reduction of bacterial burden in a mouse model of burn-wound infection.²⁴⁶

In another study involving small molecule polyamines, Katsu and coworkers investigated synthetic analogues of the joro spider toxin as OM disrupting agents leading to the identification of napthylacetylspermine (Figure 7D) which was found to potentiate the activity of novobiocin against E. coli (Table 5).247 Mechanistic studies revealed that administration of napthylacetylspermine causes OM disruption, which was attributed to displacement of LPS-associated Ca²⁺. In addition, napthylacetylspermine was found to promote cellular uptake of the tetraphenylphosphonium (TPP⁺), indicating membrane permeabilization, a finding similar to that obtained with PMBN.^{247,248} Interestingly, spermidine and spermine were also found to induce loss of Ca²⁺ but did not cause uptake of TPP⁺, pointing to the importance of the napthyl mojety for membrane permeabilization.248 Given that hemolysis data no was reported for napthylacetylspermine, it is not possible to assess the selectively of its OM activity.

The David group also reported the development of acylated polyamines as LPS neutralizing agents capable of functioning as OM disrupting synergists.^{249–251} A series of monoacyl- and bisacyl-homospermines were prepared and evaluated as potentiators of rifampicin resulting in the identification of two potent synergists, compounds **8a** and **8b** (see Figure 7E and Table 5).²⁴⁹ A clear correlation between length of the lipophilic tails and hemolytic activity was seen with compound **8a** appearing to strike an optimal balance.²⁴⁹ Using a similar approach, Copp and coworkers introduced the indole-3-acrylamido-spermine conjugates inspired by a class of indole spermidine alkaloid natural products.^{252,253} A SAR study led to the development of spermidine analogues like **14** and **17** which exhibited effective synergy with various antibiotics (Figure 7F and Table 5).^{252,254} These compounds affect bacterial membrane integrity, show low cytotoxicity and hemolytic activity. Interestingly, compound **14** was also found to inhibit bacterial efflux pumps suggesting that the potentiation of antibiotics by these compounds may be attributed to a dual mechanism of action.^{252,254}

Given the inclusion criteria noted in the introduction, only small molecules synergists (MW under 1500 kDa) are included in this review and as such we do not discuss larger polycationic polymers even though some have been shown to exhibit synergistic activity.^{89–95,25256} It is noteworthy, however, that branched polyethylenimine (BPEI) with a MW of 600 Da shows synergistic activity (Figure 7G, Table 5) and can also eradicate biofilms when co-administered with a variety of antibiotics.²⁵⁷ Mechanistic studies using isothermal titration calorimetry and fluorescence spectroscopy indicate that at the concentration required for antibiotic potentiation, 600 Da BPEI reduces diffusion barriers from LPS without disrupting the OM itself.²⁵⁷



Figure 7. Non-steroid small molecule synergists A) synergists identified via HTS; B) azaindole synergists: C) D-LANA-14 based on a norspermidine core linked to two D-lysine residues and a central tetradecanoyl moiety; D) joro spider toxin inspired napthylacetylspermine; E) bisacyl-homospermines; F) indole-3-acrylamido-spermine conjugates; and G) representation of 600 Da branched polyethylenimine (BPEI).

4.4. Plant derived synergists

A number of plant-derived compounds have also been reported to potentiate the activity of antibiotics against Gram-negative bacteria (Table 5). These include natural products like eugenol, a major component of cloves oil, linalool which can be isolated from coriander, thymol which is extracted from thyme, and cinnamaldehyde and cinnamic acid which are found in the bark and leaves of the cinnamon tree (Figure 8).²⁵⁸⁻²⁶⁴ Important to note is that only pure compounds derived from plants are included in our assessment. We refer the reader to other reviews on the synergistic activity of essential oils or crude extracts.^{265,266} Notably, most plant-derived compounds reported to potentiate antibiotics against Gram-negative bacteria are not cationic, setting them apart from most other synergists. Despite their lack of positive charge, a number of investigations have shown that the synergy associated with these compounds is a function of their ability induce OM permeabilization (Table 6).^{258,259,267-269} The broad range of biological activities associated with cinnamic acid and its derivatives, including ferulic acid, 3,4dimethoxycinnamic acid, and 2,4,5-trimethoxy cinnamic acid (Figure 9), have been recently reviewed including synergistic effects associated with OM disruption.²⁷⁰ Interestingly, despite its clear structural similarities with cinnamic acid, studies with cinnamaldehyde suggest it may operate via a different synergist mechanism. Unlike cinnamic acid, cinnamaldehyde does not increase OM permeabilization based on the NPN assay, but does exhibit synergistic effects with erythromycin and novobiocin (Table 5).267,269



Figure 8. Plant-derived natural products reported to potential the activity of antibiotics against Gram-negative bacteria.

Table 5. Overview of non-steroid small molecule synergists (compound names provided as given in the cited literature references).

Name	Ref	FICI	Pathogen	Antibiotic	Hemolytic activity ^a	
Synergists based on approved drugs						
Pentamidine	236,237	0.25	E. coli	rifampicin	<10% (20h)	
Compound 9	236,237	< 0.047	E. coli	rifampicin	>10% (20h)	
Compound 21	237	≤0.094	E. coli	rifampicin	<10% (20h)	
Compound 38	237	≤0.039	E. coli	rifampicin	>10% (20h)	
Compound P35	238	0.094	A. baumannii	novobiocin	<10% (45 min)#	
Metformin	241	0.375	E coli	vancomycin	<10% (1h)	

High throughput screening (HTS)-hits									
MAC-0568743	242	≤0.16	E. coli	rifampicin	NR				
Liproxstatin-1	242	0.25*	E. coli	rifampicin	NR				
BWC-Aza1	243	0.258	E. coli	rifampicin	<10% (45 min)				
BWC-Aza2	243	0.06	A. baumannii	rifampicin	<10% (45 min)				
Peptidomimetics									
OAK C ₁₂ (ω7)	208	≤0.073*	E. coli	rifampicin	>10% (3h)				
OAK C ₁₂	208	≤0.211*	E. coli	rifampicin	>10% (3h)				
OAK C ₁₀	208	≤0.036*	E. coli	rifampicin	<10% (3h)#				
OAK C ₈	208	≤0.078*	E. coli	rifampicin	<10% (3h)#				
C _{14(ω5)} OOc ₁₀ O	209	0.20*	K. pneumoniae	rifampicin	<10% (3h)#				
dUSTBP 2	202	≥0.250	P. aeruginosa	rifampicin	<10% (1h)				
dUSTBP 5	202	≥0.125	P. aeruginosa	rifampicin	<10% (1h)				
dUSTBP 8	202	≥0.002	A. baumannii	novobiocin	<10% (1h)				
Synergists with a polyamine motif									
D-LANA-14	245,246	0.09	P aeruginosa	rifampicin	<10% (1h)				
Naphthylacetylspermine	247	0.125*	E. coli	novobiocin	NR				
Bisacyl-homospermine 8a	249	0.304*	E. coli	rifampicin	<10% (30min)				
Bisacyl-homospermine 8b	249	0.297*	E. coli	rifampicin	>10% (30min)				
Spermidine analogue 14	254	0.255*	E. coli	erythromycin	<10% (1h)#				
Spermidine analogue 17	254	0.255*	P. aeruginosa	erythromycin	<10% (1h)#				
600-Da BPEI	257,271	0.26	P. aeruginosa	erythromycin	<10% (1h)				
		Plant der	ived synergists						
Eugenol	258,272	≤0.2*	P. aeruginosa	rifampicin	<10% (24h)				
Linalool	259,273	0.37	E. coli	erythromycin	<10% (4h)				
Thymol	267,274	0.25	E. coli	erythromycin	<10% (1h)				
Cinnamaldehyde	267,275	0.24	E. coli	erythromycin	<10% (48h)				
trans-Cinnamic acid	268,276	0.36	E. coli	erythromycin	<50% (1h)				
Ferulic acid	268,276	0.48	E. coli	erythromycin	<50% (1h)				
3,4-dimethoxycinnamic acid	268,276	0.42	E. coli	erythromycin	<50% (1h)				
2,4,5-trimethoxycinnamic acid	268,276	0.22	E. coli	erythromycin	<50% (1h)				

^aNon-hemolytic is defined as <10% hemolysis compared to positive control with incubation times denoted in parentheses, NR denotes no data reported; [#]denotes that the concentration tested was lower than 100 μ g/mL; *FICI calculated from MSC and MIC values reported in the cited literature reference.

5. Antibiotic-derived synergists

In general, the antibiotic potentiators discussed above show little-to-no inherent antibacterial activity. There are, however, a number of reports describing antibacterial compounds that also exhibit OM disrupting effects and in doing so synergize with antibiotics that are otherwise inactive towards Gram-negative bacteria. The synergists described in this section are specifically included based upon their OM disrupting activity rather than a contribution of their inherent activity to synergy. We therefore do not include the combination of rifampicin with imipenem or trimethoprim which is solely based on functional synergy.^{277,278} In addition, we also do not cover reports describing systems where an OM perturbing motif like PMBN is covalently linked to another antibiotic as a means of enhancing anti-Gram-negative activity.^{39,279-281}

5.1. Tobramycin-derived synergists

Tobramycin (Figure 9A) belongs to the aminoglycoside class of antibiotics that function by inhibiting ribosomal protein synthesis in bacteria. Recent studies have also revealed that aminoglycosides like tobramycin also interact with bacterial membranes by specifically binding to LPS and in doing so cause membrane depolarization.²⁸²⁻²⁸⁶ Building on these insights Schweizer and coworkers have prepared and assessed a number of conjugates wherein one tobramycin molecule is linked to a second antibiotic providing hybrid systems that possess both inherent antibacterial activity as well as potent synergy with other antibiotics (Figure 9A).^{287-290,279,291-297} Among the first hybrids prepared was a series tobramycin-fluoroquinolone conjugates.^{287,288} Both the optimal sites of conjugation and linker lengths between the two antibiotics were investigated revealing TOB-MOX, a tobramycin-moxifloxacin hybrid, and tobramycin-ciprofloxacin conjugate 1e to be potent synergists (Table 6).²⁸⁸ Notably, the conjugates generally showed lower inherent antibacterial activity than the parent antibiotics indicating that their synergistic activity comes at the price of inherent activity.^{287,288} OM disruption was confirmed for both hybrids using the NPN assay and both were found to potentiate multiple antibiotics including rifampicin, erythromycin, novobiocin, and vancomycin.^{287,288} Also of note was the finding that these hybrids exhibited a significantly reduced capacity to inhibit of protein translation compared to that of tobramycin.^{287,288} Conversely, the hybrids were found to maintain, and some cases exceed, the gyrase inhibiting activity of the parent fluoroquinolones.^{287,288} Another series of hybrids were prepared by coupling tobramycin with rifampicin, which targets the bacterial RNA polymerase.²⁸⁹ As for the fluoroquinolone conjugates, the inherent activity of the tobramycin-rifampicin conjugates was significantly reduced compared to the parent antibiotics. Again, however, some hybrids were found to exhibit synergy via an OM-disrupting mechanism (see tobramycin-rifampicin 1, 2, 3, Figure 9A).^{288-290,298}

A number of other hybrids have also been reported by the Schweizer group wherein tobramycin was coupled to various other small molecules known to engage with different bacterial targets. In one case, tobramycin was coupled to a lysine-based amphiphile known to function as membrane permeabilizer (see tobramycin-lysine **3**, Figure 9A).^{290,299} This conjugate was found to effectively potentiate the activity of novobiocin, erythromycin, and vancomycin (Table 6).^{290,300} The same group also explored hybrids wherein tobramycin was coupled to small molecule efflux pumps inhibitors such as 1-(1-naphthylmethyl)-piperazine (NMP) and paroxetine (PAR) (Figure 9A).^{45,291,301-303} Along with potent synergy against P. *aeruginosa* (Table 6), these hybrids were also found

to cause OM disruption and inner membrane depolarization.^{315,316} Two additional generations of tobramycin conjugates were also reported: tobramycin homodimers and tobramycin coupled to chelating cyclams (Figure 9A).^{293,294} The dimerization of tobramycin was conveniently achieved by means of copper catalyzed azide-alkyne click chemistry, resulting in potent synergists that also exhibit enhanced OM disruption relative to tobramycin itself (Table 6).²⁹³ A combination of novobiocin and tobramycin homodimer **1** (both administered at 50 μ g/mL) was further shown to have *in vivo* efficacy against A. baumannii in a wax worm larvae model.²⁹³ Studies with the corresponding monomeric tobramycin azide and alkyne precursors revealed neither to be synergistic, underscoring the need for dimerization to achieve synergy.²⁹³ In the case of the tobramycin-cyclam conjugates, the introduction of the cyclam chelating group was hypothesized to aid in the OM permeabilization by sequestration of divalent cations bridging the Lipid A phosphate groups.^{294,304-306} While tobramycin-cyclam hybrids 1, 2, and **3** effectively potentiated novobiocin, rifampicin, vancomycin and erythromycin (Table 6), it is also particularly noteworthy that they also enhanced the activity of meropenem against both carbapenem-resistant and -sensitive strains.²⁹⁴ This effect was abrogated by the addition of excess MgCl₂ further supporting a mode of action driven by OM disruption.294

5.2. Nebramine-derived synergists

Following on their work with tobramycin hybrids, the Schweizer group also prepared a number of analogous nebramine conjugates (Figure 9B). Nebramine (NEB) is a disaccharide subunit of tobramycin that interestingly displays activity against tobramycin resistant strains and also interacts with the OM.^{283,307-313} The NEB hybrids synthesized included conjugates with moxifloxacin (MOX), ciprofloxacin (CIP), NMP, and cyclam (Figure 9B).^{295,296} These hybrids were all found to effectively potentiate the activity of multiple classes of antibiotics against a range of Gram-negative bacteria (Table 6). Furthermore, NEB-MOX **1a**, NEB-CIP **1b**, and NEB-NMP **2** were also reported to dissipate proton motive force and proposed to cause OM disruption as for the corresponding tobramycin conjugates.^{287,290,291,295,296}

5.3. Levofloxacin derived synergists

Schweizer and coworkers also recently reported another class of antibiotic based synergists based on polybasic peptide–levofloxacin conjugates (Figure 9C).²⁹⁷ While these levofloxacin-peptide hybrids were found to be non-hemolytic, they were also shown to be essentially devoid of inherent antimicrobial activity (MICs typically > 128 μ g/mL). They did however, exhibit strong potentiation of numerous antibiotics against MDR clinical isolates of *P. aeruginosa*, *E. coli*, *K. pneumoniae* and to a lesser extent, *A. baumannii* (Table 6).²⁹⁷ Preliminary mechanistic studies indicate that these conjugates potentiate other antibiotics by both blocking active efflux and by permeabilization of the OM.²⁹⁷

A) Tobramycin conjugates



Figure 9. Synergists based on clinically used antibiotics. A) Tobramycin (TOB) conjugates; B) Nebramine (NEB) analogues; and C) polybasic conjugated levofloxacin hybrids.

Name	Ref	FICI	Pathogen	Antibiotic	Hemolytic activity ^a			
		Tobramy	cin derivatives					
TOB-MOX 1	287	0.125	P. aeruginosa	novobiocin	<10% (30 min)			
Tobramycin-Ciprofloxacin 1e	288	< 0.04	P. aeruginosa	rifampicin	<10% (30 min)			
Tobramycin-Rifampicin 1	289	0.28	P. aeruginosa	rifampicin	<10% (1h)			
Tobramycin-Rifampicin 2	289	0.15	P. aeruginosa	erythromycin	<10% (1h)			
Tobramycin-Rifampicin 3	289	0.06	P. aeruginosa	erythromycin	<10% (1h)			
Tobramycin-Lysine 3	290	0.008	P. aeruginosa	novobiocin	<10% (1 h)			
TOB-NMP 1	292	≥0.008	P. aeruginosa	rifampicin	<10% (30 min)			
TOB-PAR 2	292	≥0.008	P. aeruginosa	rifampicin	<10% (30 min)			
Tobramycin homodimer 1	293	0.07	P. aeruginosa	novobiocin	<10% (1h)			
Tobramycin homodimer 2	293	0.08	P. aeruginosa	novobiocin	<10% (1h)			
Tobramycin homodimer 3	293	0.05	P. aeruginosa	novobiocin	<10% (1h)			
Tobramycin-Cyclam 1	294	0.13	P. aeruginosa	novobiocin	<10% (30 min)			
Tobramycin-Cyclam 2	294	0.13	P. aeruginoa	novobiocin	<10% (30 min)			
Tobramycin-Cyclam 3	294	0.08	P. aeruginosa	novobiocin	<10% (30 min)			
		Nebram	ine derivatives					
NEB-MOX 1a	295	≥0.002	K. pneumoniae	rifampicin	NR			
NEB-CIP 1b	295	≥0.008	P. aeruginosa	rifampicin	<10% (1h)			
NEB-NMP 2	295	≥0.004	P. aeruginosa	rifampicin	NR			
Nebramine-cyclam	296	0.25	P. aeruginosa	rifampicin	<10% (1h)			
Levofloxacin derivatives								
Levofloxacin conjugate 10	297	0.10	P. aeruginosa	rifampicin	<10% (1h)			
Levofloxacin conjugate 11	297	0.10	P. aeruginosa	novobiocin	<10% (1h)			
Levofloxacin conjugate 12	297	0.08	P. aeruginosa	novobiocin	<10% (1h)			
^a Non-hemolytic is defined as <	10% h	emolysis o	compared to posi	tive control wit	h incubation times			

Table 6. Overview of synergists based on clinically used antibiotics (compound names provided as given in the cited literature references).

denoted in parentheses, NR denotes no data reported.

6. Chelating agents as OM disrupting synergists

The activity of antibiotics can also be potentiated by chelating agents that disturb the integrity of the OM by sequestering the divalent cations Mg²⁺ or Ca²⁺ coordinated by the phosphate groups of the lipid A core of LPS (Figure 1B).³² The preeminent chelating agent, EDTA (Figure 10) is a well described synergist and its reported ability to potentiate antibiotics actually predates the reported synergistic activity of PMBN.^{49,314-317} Exposure of Gram-negative bacteria to EDTA is accompanied by the significant release of LPS and, as for treatment with PMBN, also results in the increased uptake of NPN.³¹⁸⁻³²⁰ While the potentiating effects of EDTA on antibiotics such as novobiocin and rifampicin are well documented, FICI values have not been reported in literature and cannot be readily calculated from published data.^{316,317,319,321} Similarly, for the other chelating here discussed, no FICI values could be found in the literature and as such we do not provide a summary table as done for the other synergists discussed in this review.

In additional to his seminal work with PMBN, Vaara also reported the potentiation of hydrophobic antibiotics by sodium hexametaphosphate (HMP, Figure 10) against Gramnegative bacteria as well as the increase in NPN uptake in cells treated with this potent Ca²⁺ binding agent.³²² In a similar study, Ayres and Russell also described sodium polyphosphates as potent synergist with several antibiotics (structures not shown).³²³ In the same study, citric acid (Figure 10) was also demonstrated to exhibit synergistic activity with erythromycin, novobiocin, rifampicin, methicillin, and gentamicin.³²³ In addition, 2,3-dimercaptosuccinic acid (Figure 10), clinically used in the treatment of lead intoxication, was also found to potentiate the activity of hydrophobic antibiotics.³¹⁹ The synergistic activity of 2,3-dimercaptosuccinic acid was attributed to an OM permeabilizing mechanism as evidenced by increased NPN uptake in bacterial cells treated with the compound.³¹⁹



Figure 10. Chelating agents with demonstrated synergistic activity.

7. Concluding remarks

New strategies are required to address the growing threat posed by MDR Gram-negative pathogens. To this end, a large and growing number of synergists capable of potentiating Gram-positive specific antibiotics against Gram-negative bacteria have been described in literature to date. Within this review we provide the reader with a comprehensive and up-to-date overview of those synergists reported to have a demonstrated OM targeting mechanism. We also draw attention to the importance of selective OM disruption, a factor that has often been overlooked by researchers when characterizing their synergists. In this regard, and based on our assessment of the literature, the majority of hemolysis studies reported for such synergists use relatively short incubation times compared to the incubation times actually used in assessing synergy (i.e. in checkerboard assays). Based on our own experience, not only is the concentration at which hemolysis is assessed relevant, but incubation time can also make a significant difference in describing a compound as hemolytic or not. For example, in cases where 5% hemolysis is reported after one hour, it is our experience that such compounds are often much more hemolytic after overnight incubation. For this reason we have included both the concentrations and incubation times of the synergists described in this review. Doing so provides for a more honest and accurate assessment of the OM specificity of these synergists.

To provide a means of comparing the relative activity of the synergists here summarized, we have emphasized their FICI values, a descriptor broadly applied as a scale to quantify synergistic potency. However, another important consideration that is not directly revealed by the FICI is of course the concentration at which a synergist actually potentiates the companion antibiotic. Related to this is the importance of the pharmacokinetic/pharmacodynamic profile of the synergist and how well it matches that of the antibiotic it potentiates. Given that the vast majority of synergists covered in this review have only been characterized using cell-based *in vitro* and biochemical assays, we have not touched on this. It is clear, however, that establishing and optimizing such parameters will be essential to the (pre)clinical development of any such synergist.

8. Thesis outline

This thesis describes the development of novel synergists designed to selectively disrupt the outer membrane of Gram-negative bacteria.

Chapter 2 reports the optimization of bis-amidines as outer membrane disrupting agents that can potentiate Gram-positive specific antibiotics. The synthesis of a number of unique bis-amidines was followed by an initial screening with checkerboard assays revealing the most potent synergists. The compounds were also evaluated for hemolytic activity to provide a rough measure of their selectivity. The most potent, non-hemolytic compounds were then evaluated in combination with rifampicin against multiple strains of Gram-negative bacteria. Lastly, their outer membrane disrupting activity was compared to the well-known synergist PMBN.

Chapter 3 describes the development of peptide-based synergists with the capacity to enhance the activity of antibiotics against Gram-negative bacteria. The approach taken was inspired by recent reports of LPS-binding activity by thrombin-derived peptides. This prompted us to further evaluate these peptides as outer membrane disrupting synergists. The structures of the peptides were optimized by adjusting the C- and N-termini as well as by applying an alanine scan. In addition, hemolysis and outer membrane disrupting assays were performed to establish the selectivity of the peptides for the outer membrane of Gram-negative bacteria. The synergistic potential of the lead peptides was evaluated with several Gram-positive antibiotics and for multiple Gram-negative bacterial strains.

In **Chapter 4** the focus was shifted from outer membrane disrupting synergists based on synthetic small molecules and peptides, to the synergistic activity of the complement system found in with human serum. A broad range of Gram-positive specific antibiotics was evaluated with serum in two assays: an inner membrane permeability assay serving as a screen, followed by a bacterial viability assay allowing for a validation of the hits. In addition, four antibiotics of the glycopeptide class were also evaluated to allow for an inclass comparison of synergy with the complement system.

Chapter 5 diverges from the synergy theme of this thesis: in this chapter the inherent anti-Gram-positive activity of the bis-amidines described in Chapter 2 is described. In addition to the previously synthesized bis-amidines, four new bis-amidines were prepared and evaluated for hemolytic and antimicrobial activity. In addition to the screening of several Gram-positive bacteria, the effects of different media were also evaluated.

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