*Pseudomonas aeruginosa* rhamnolipid micelles deliver toxic metabolites and antibiotics into *Staphylococcus aureus*

**Highlights**

- *Pseudomonas aeruginosa* rhamnolipids form micelles
- Rhamnolipid micelles delivery pyochelin into *S. aureus* cells
- Rhamnolipid micelles potentiate activity of lincosamide antibiotics against *S. aureus*
**Pseudomonas aeruginosa** rhamnolipid micelles deliver toxic metabolites and antibiotics into *Staphylococcus aureus*

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**SUMMARY**

Efficient delivery of toxic compounds to bacterial competitors is essential during interspecies microbial warfare. Rhamnolipids (RLPs) are glycolipids produced by *Pseudomonas* and *Burkholderia* species involved in solubilization and uptake of environmental aliphatic hydrocarbons and perform as biosurfactants for swarming motility. Here, we show that RLPs produced by *Pseudomonas aeruginosa* associate to form micelles. Using high-resolution microscopy, we found that RLP micelles serve as carriers for self-produced toxic compounds, which they deliver to *Staphylococcus aureus* cells, thereby enhancing and accelerating *S. aureus* killing. RLPs also potentiated the activity of lincosamide antibiotics, suggesting that RLP micelles may transport not only self-produced but also heterologous compounds to target competing bacterial species.

**INTRODUCTION**

*Pseudomonas aeruginosa* is an opportunistic pathogen causing acute infections in immunocompromised hosts and chronic infections in patients with cystic fibrosis. Owing to intrinsic and acquired antibiotic resistance determinants, *P. aeruginosa* remains a challenge when choosing optimal antimicrobial therapies. *P. aeruginosa* possesses a variety of host-directed virulence factors, including cell-contact dependent (T3SS, T6SS) and independent mechanisms (secreted toxins and secondary metabolites). Furthermore, its metabolic versatility and the production of bacteriocins (pyocins), confer a competitive advantage during inter- and intra-species competition, explaining its success as an opportunistic pathogen and niche colonizer (Atanaskovic et al., 2020; Gonzalez and Mavridou, 2019; Wood et al., 2019).

We have recently defined an antimicrobial cocktail consisting of proteins and several metabolites secreted by *P. aeruginosa*, which showed broad-spectrum killing activity against both Gram-positive and Gram-negative bacterial species (Gdaniec et al., 2020). An essential component of this cocktail was rhamnolipids (RLPs), which belong to the chemically diverse group of glycolipids. RLPs are composed of a hydrophilic rhamnose sugar moiety, which is linked through a \( \beta \)-glycosidic bond to a hydrophobic fatty acid moiety (Hauser and Karnovsky, 1957). RLPs have multiple roles in metabolite uptake, community and host interactions, and microbial competition (Chrzanski et al., 2012). The initially reported function of RLPs in *P. aeruginosa* was the solubilization and uptake of aliphatic hydrocarbons from the environment, that *P. aeruginosa* uses as a carbon and energy source (Noordman and Janssen, 2002). In the free form, their hydrophilic moiety can interact with the O-antigen component of lipopolysaccharides (LPS) of Gram-negative bacteria, thereby increasing the hydrophobicity of the outer membrane (Zhong et al., 2014).

One interesting feature of RLPs is their ability to form micelles above a critical micelle concentration (Haba et al., 2014). RLP micelles can remove LPS from the outer membrane of Gram-negative bacteria, which also leads to an increase in hydrophobicity due to the resulting lack of polar sugar residues on the cell surface (Al-Tahhan et al., 2000; Sotirova et al., 2009). However, the interactions between RLP micelles and Gram-positive bacteria are not well understood. Because *P. aeruginosa* is particularly efficient in killing *S. aureus* cells, we sought to get insight into the molecular interactions between *P. aeruginosa* RLPs and *S. aureus* using biochemical analysis and super-resolution microscopy. We found that RLP micelles are able to incorporate metabolites produced by *P. aeruginosa* and to deliver them to *S. aureus* cells. We further show that RLP micelles accelerate and potentiate the activity of lincosamide antibiotics directed against *S. aureus*.
RESULTS

Rhamnolipids secreted by *P. aeruginosa* form micelles

We have previously shown that supernatants of a wspF mutant of *P. aeruginosa* contain elevated concentrations of several secondary metabolites, including RLPs. RLPs were an essential ingredient of a synthetic cocktail, including the siderophores pyoverdine and pyochelin, as well as the alkylquinoline 2-heptyl-4-hydroxyquinoline-N-oxide (HQNO), able to cause a 5-log reduction of viable counts of *S. aureus* (Gdaniec et al., 2020) within 24 h. We wondered whether the RLPs concentration previously measured at 300–400 \( \mu \text{g/mL} \) in the \( \Delta \text{wspF} \) mutant supernatant (Gdaniec et al., 2020) would be sufficient to form micelles. We therefore analyzed by electron microscopy the supernatants of *P. aeruginosa* PA14 \( \Delta \text{wspF} \) and an isogenic \( \Delta \text{wspF} \Delta \text{rhlA} \) mutant, deficient in RLP production. As a control, we analyzed purified *P. aeruginosa* RLPs, containing a mix of mono- and di-RLPs with varying fatty acid chain lengths at a concentration of 250 \( \mu \text{g/mL} \). Electron microscopy imaging revealed the presence of round shaped, probably spherical structures in the supernatant of the \( \Delta \text{wspF} \) mutant with an average surface area of 0.15 \( \mu \text{m}^2 \) (Figure 1A, left panel), similar to those observed with the purified RLPs solution (Figure 1A, right panel), but absent in the supernatant of the \( \Delta \text{wspF} \Delta \text{rhlA} \) mutant (Figure 1A, middle panel). We therefore considered the structures observed in the \( \Delta \text{wspF} \) supernatant as RLP micelles. To further confirm their lipidic

**Figure 1.** *P. aeruginosa* \( \Delta \text{wspF} \) supernatant contains rhamnolipid micelles

(A) Electron microscopy imaging of RLP micelles present in the \( \Delta \text{wspF} \) supernatant (left panel) and formed by purified RLPs (right panel).

(B) Schematic representation of RLP micelles staining with Nile Red (NR).

(C) Critical micellar concentration (CMC) was determined by adding purified RLPs to either medium or to RLP-deficient \( \Delta \text{wspF} \Delta \text{rhlA} \) supernatant.

(D) Detection of RLP micelles by NR staining and fluorescence measurement in supernatants of *P. aeruginosa* wild type and mutants. Student test (***, p value <0.0001). The scale bar indicates 1 \( \mu \text{m} \). Six-thirteen images per condition, showing similar heterogeneous size distributions of RLP micelles, were analyzed.
nature, we performed selective fluorescent staining with Nile Red (NR). While NR is slightly fluorescent in water, its fluorescence signal is enhanced upon integration into a lipophilic environment like phospholipid membranes or vesicles (Greenspan et al., 1985) (Figure 1B). We measured the fluorescence emission of purified RLPs, added either to M14 medium or to RLP-deficient ΔwspFΔrhlA culture supernatant. This allowed us to determine the RLPs critical micelle concentration (CMC), which was comprised between 50 and 100 μg/ml (Figure 1C). Next, we measured NR fluorescence in bacterial supernatants of the PA14 wild type, ΔwspF, and ΔwspFΔrhlA mutants, as well as in the M14 growth medium. While fluorescence remained low for the wild type and the ΔwspFΔrhlA supernatants, indicating lack of micelles formation, the ΔwspF supernatant showed a strong fluorescence signal indicating micelle formation and an RLP concentration above the CMC (Figure 1D). Altogether, these data suggest that RLPs overproduced in a ΔwspF mutant supernatant form micelles, which have similar shape and NR fluorescence compared with purified RLPs.

**RLP micelles are essential for the killing activity of a ΔwspF mutant supernatant**

We next analyzed the role attributable to RLPs in the *S. aureus* killing activity of *P. aeruginosa* culture supernatants. For this, we investigated the *S. aureus* killing activity of supernatants produced by the PA14 wild type, the ΔwspF, and ΔwspFΔrhlA mutants. As expected, the ΔwspF supernatant showed a potent *S. aureus* killing activity, not found in supernatants produced by the other strains, confirming the role for RLPs promoting *S. aureus* killing (data not shown). To further characterize this killing activity, we separated the crude culture supernatants (crude-S) by ultracentrifugation into an ultracentrifuged pellet (ultra-P) and an ultracentrifuged supernatant (ultra-S). Determined by Orcinol assays, the ΔwspF ultra-P fraction contained 88% of the RLPs, while 12% remained in the ultra-S fraction (Figure 2A), suggesting enrichment of RLPs in the ultra-P fraction. We next compared the killing kinetics of the crude-S and ultra-S supernatants, with those of resuspended ultra-P, PA14 crude-S, ultra-S, and ultra-P resuspended in M14 medium to the initial supernatant volume (1x ultra-P) had no killing activity (Figure 2B). In contrast, ultra-P resuspended in M14 medium to 1/15 of the initial supernatant volume (15x ultra-P) presented moderate killing during the initial 1.5 h (1.5-log decrease in *S. aureus* CFU), which was followed by a further extended killing period (3.5-log reduction over the next 21 h). ΔwspF crude-S and ultra-S, as well as 15x ultra-P showed similar killing kinetics (4-log reduction during first 3 h and additional 2-log reduction at 24 h), while 1x resuspended ultra-P had no killing activity (Figure 2C). The ΔwspFΔrhlA crude-S and ultra-S, and resuspended ultra-P (both 1x and 15x) showed no killing during the first 3 h. However, at 24 h, the crude-S and ultra-S showed a 2-log, and the 15x ultra-P a 3-log reduction in *S. aureus* CFUs, respectively (Figure 2D). To quantitatively assess the effect of the individual mutations as well as RLP supplementation, we considered the log decrease per hour as the primary readout for killing activity. We restricted this analysis to the first 3 h as at longer timepoints many experiments reached the detection limit and could not be quantitatively interpreted (Figure S1A). We observe statistically highly significant enhancement of killing in the various supernatant fractions of the ΔwspF mutant (Figure S1B), which is again highly significantly abolished in the ΔwspFΔrhlA mutant (Figure S1C), and restored by addition of exogenous RLPs (Figure S1D). This confirms the importance of metabolites present in the ΔwspF supernatant and the specific requirement of RLPs for *S. aureus* killing.

In the ultra-P fraction, the situation is more subtle because both in the wild type and the ΔwspF mutant the 15x concentrated ultra-P shows significant killing activity, whereas the 1x ultra-P does not. Nevertheless, abrogation of RLP production prevents killing also in the 15x ultra-P (Figure S1C), with restoration by exogenous RLPs (Figure S1D).

To further substantiate the specific role of RLPs in *S. aureus* killing, we supplemented the ultra-S and 15x ultra-P of the ΔwspFΔrhlA mutant with commercial RLPs. Our previous metabolomic analysis of ΔwspF supernatant had shown a large diversity of RLPs, dominated by the C10-C12 di-RLP species (Gdaniec et al., 2020). We therefore used a mix of purified mono- and di-RLPs (Figure 2E), but also C10-C10 (Figure 2F) or C12-C12 mono-RLPs (Figure 2G) alone for the supplementation experiments. ΔwspFΔrhlA ultra-S and 15x ultra-P supplemented with RLPs showed strong *S. aureus* killing. The killing activities differed significantly between the RLP types (p<0.001), but not between the ultra-S and 15x ultra-P fractions (p = 0.155) (Figure S1D). The effect of purified RLPs was most pronounced with the C10-C10 mono-RLPs, suggesting that the shorter acyl chains provide a more efficient killing (Figure S1E). Altogether, these results suggest that RLPs are crucial for the initial killing of *S. aureus* and that a single species of RLPs is sufficient for this activity.
Rhamnolipids interact with *S. aureus* cell membranes and cross their peptidoglycan

To further explore the mechanisms underlying the killing activity and to identify the localization of RLPs in *S. aureus* cells, we covalently labeled a mix of purified commercial RLPs with the fluorescent dye Abberior STAR 555 (Schermelleh et al., 2019). We performed comparative confocal microscopy of unlabeled and Abberior STAR 555 labeled RLP micelles, which showed a similar micelle

Figure 2. RLP micelles are essential for initial *S. aureus* killing activity of ΔwspF mutant supernatants

(A) Scheme of isolation for crude and ultracentrifuged supernatants, and ultracentrifuged pellets.

(B) Killing curves of *S. aureus* exposed to crude or ultracentrifuged supernatants, or resuspended ultracentrifuged pellets produced by PA14 wild type (B), ΔwspF mutant (C), and ΔwspFΔrhlA mutant (D). ΔwspFΔrhlA mutant crude or ultracentrifuged supernatants, or resuspended ultracentrifuged pellets complemented with commercial rhamnolipids (E), RLP C_{10}C_{10} (F) and RLP C_{12}C_{12} (G). Parts of the data (E–G) are imported from (C) for comparison. Additional data and statistical details are given in Figure S1.
area for the most abundant micelle fractions, confirming that labeled RLPs still form micelles (Figure S2).

We then labeled the *S. aureus* peptidoglycan layer using vancomycin coupled with the Abberior STAR 488 NHS ester dye and visualized the cells using stimulated emission depletion (STED) nanoscopy to study the interaction between the labeled RLPs and *S. aureus* cells. As expected, we observed a fluorescence signal corresponding to the *S. aureus* cell wall, visualized by the incorporation of fluorescent vancomycin into the peptidoglycan layer (Watanakunakorn, 1984) (Figures 3A–3F). For up to 66.5% of the cells, the maximum RLPs signal formed a circular shape on the cytosolic side of the peptidoglycan, while the signal intensity decreased toward the cell center (Figures 3A–3F). This result suggests that the RLPs cross the peptidoglycan layer of *S. aureus*, accumulating in or next to the cell membrane.

**Rhamnolipid micelles serve as a carrier for *P. aeruginosa* metabolites**

To decipher the role of RLP micelles as potential carriers for *P. aeruginosa* metabolites into *S. aureus* cells, we labeled the *P. aeruginosa* siderophore pyochelin with the fluorescent dye Abberior 580, and performed confocal microscopy of *S. aureus* cells with peptidoglycan visualized with Abberior 488 labeled vancomycin, in the absence or presence of commercial RLPs (Figure 4). To distinguish delivery of labeled pyochelin between free RLP molecules and micelles, we choose two RLPs concentrations: 50 μg/mL (below CMC) and 250 μg/mL (above CMC). In the presence of RLPs below CMC or without RLPs (Figure 4, second and third rows, respectively), uptake of labeled pyochelin reached 25% and 4%, respectively. When RLPs were added at a concentration above the CMC, labeled pyochelin was visible in the cytosol of 66% of *S. aureus* cells (Figure 4, first row).
Chemically inactivated Abberior 580 dye itself did not enter the cells even at the highest RLPs concentration (Figure S3). These data suggest that RLP micelles serve as a carrier for pyochelin in vitro. They also highlight the specificity of translocation of pyochelin in S. aureus, as opposed to non-specific increase in cellular permeability.

Rhamnolipids potentiate the activity of lincosamide antimicrobials
As RLPs are essential for killing of S. aureus in combination with other metabolites present in the P. aeruginosa ΔwspF mutant supernatants (Gdaniec et al., 2020), we investigated whether RLP micelles might potentiate the effect of antibiotics. We tested the effect of azithromycin, novobiocin, rifampicin, nalidixic acid, ciprofloxacin, ampicillin, and sulfamethoxazole on S. aureus in the presence/absence of RLPs, as representative members of the main antibiotic classes. The antimicrobial activity of these compounds on S. aureus Newman (MSSA) and COL (MRSA) strains were not affected by the presence of RLPs at
concentrations above the CMC (250 \( \text{mg/mL} \)) (data not shown). In contrast, the minimal inhibitory concentrations (MICs) of lincomycin and clindamycin, two members of the lincosamide antibiotic family, decreased by 4–8 times in the presence of RLPs at concentrations above the CMC, whereas the MICs were not affected by RLPs at concentrations below the CMC (25 \( \text{mg/mL} \)) (Table 1).

Table 1. Lincosamide MICs for \( S. \text{ aureus} \) strains Newman and COL in the presence of RLPs

| Antibiotics/compound | MIC (\( \text{mg/mL} \)) | \( S. \text{ aureus Newman} \) | \( S. \text{ aureus COL} \) |
|----------------------|--------------------------|-----------------------------|---------------------------|
| Rhamnolipids (RLPs)  | 1000                     | 1000                        |
| Clindamycin          | 0.25                     | 0.25                        |
| Clindamycin + RLPs 25 \( \text{mg/mL} \) | 0.125 | 0.125                      |
| Clindamycin + RLPs 250 \( \text{mg/mL} \) | 0.032 | 0.032                      |
| Lincomycin           | 4                        | 4                           |
| Lincomycin + RLPs 25 \( \text{mg/mL} \) | 2    | 2                           |
| Lincomycin + RLPs 250 \( \text{mg/mL} \) | 0.5  | 0.5                         |

To evaluate the effect of RLPs on antibiotic activity, we performed checkerboard MIC assays and calculated the fractional inhibitory concentration (FIC) (Table S1). For the \( S. \text{ aureus} \) Newman strain, the combination of RLPs with lincomycin revealed an additive effect (FIC 0.5–1.0), while combination of RLPs with clindamycin showed a synergistic effect (FIC \( \leq 0.5 \)). For the \( S. \text{ aureus} \) COL strain, the combination of RLPs with clindamycin or lincomycin also resulted in a synergistic effect (FIC \( \leq 0.5 \)).

We then investigated the killing kinetics of clindamycin and lincomycin in the presence of RLPs mix. RLPs accelerated and increased killing by both lincosamides as shown by a 2–3 log decrease in \( S. \text{ aureus} \) CFUs (Figure 5). Of note, the effect of RLPs was more pronounced on the MSSA strain Newman than on the MRSA strain COL.

**DISCUSSION**

Antibiotic resistance is a worldwide threat to human health that requires major efforts to develop new antimicrobial strategies and/or increase the efficacy of already existing antibiotics (ref: https://www.who.int/news-room/fact-sheets/detail/antimicrobial-resistance). Consequently, drug delivery systems enhancing the efficacy of antibiotics, while reducing off-target toxicity, have attracted attention. While synthetic liposomes (Chen et al., 2018) or siderophore-drug conjugates have been studied to increase antibiotic delivery to their targets (Schalk, 2018), little is known about natural bacterial delivery tools. Here, we demonstrate that \( P. \text{ aeruginosa} \) uses naturally produced rhamnolipid micelles to enhance the killing of \( S. \text{ aureus} \). Our results reveal new properties of rhamnolipids acting as carriers for toxic metabolites, increasing their delivery to their targets in competing microorganisms. Therefore, our study provides evidence for natural antimicrobial delivery systems used by pathogenic bacteria in interspecies competition.

The properties of biosurfactants were intensively investigated over the last 50 years since the first biosurfactant “surfactin” was purified and characterized from \( B. \text{ subtilis} \) (Arima et al., 1968). Among the various categories of biosurfactants, rhamnolipids belong to the glycolipids group and are studied not only as a virulence factor of \( P. \text{ aeruginosa} \) but also as an interesting compound for industrial processes (Sekhon Randhawa and Rahman, 2014). \( P. \text{ aeruginosa} \) synthesizes a variety of RLPs composed of mono- and di-RLP species, which carry either one or two acyl chains of variable chain length (C\(_8\) to C\(_{14}\)), respectively. The dominant species in \( P. \text{ aeruginosa} \) are the di-RLPs C\(_{10}\)-C\(_{10}\) and C\(_{12}\)-C\(_{12}\) (Abdel-Mawgoud et al., 2010; Déziel et al., 1999). RLPs play an essential role as biosurfactant for swarming motility (Köhler et al., 2000), and help maintain the biofilm architecture (Davey et al., 2003; Tremblay et al., 2007). RLPs also play a role in protecting \( P. \text{ aeruginosa} \) from the host defense by inhibiting phagocytosis by macrophages (Alhede et al., 2009; McClure and Schiller, 1996; Van Gennip et al., 2009b) and by lysing polymorphonuclear neutrophils (Jensen et al., 2007; Van Gennip et al., 2009a). At high concentrations, RLPs possess modest intrinsic antimicrobial activity against both Gram-positive and Gram-negative bacteria (Haba et al., 2003; Nitschke et al., 2010; Samadi et al., 2012), as well as ameba (Cosson et al., 2002) and fungi (Goswami et al., 2002).
et al., 2015). Moreover, it has been shown that RLPs increase the fluidity of fungal membranes (Monnier et al., 2019).

While low concentrations of rhamnolipids (below critical micellar concentration) were shown to enhance the action of aminoglycoside antibiotics (Radlinski et al., 2019), the mode of action of RLPs above the CMC has not been investigated. Importantly, the molecular details of how RLP micelles interact with the bacterial membrane are poorly understood. The amphiphilic nature of free rhamnolipids points to the membrane as their hypothetical site of action. Ortiz et al. showed that di-RLPs intercalate into artificial phosphatidylcholine bilayers and produce structural perturbations, which might affect the function of the membrane (Ortiz et al., 2006). Our data support the notion that the affinity of RLPs micelles for bacterial membranes allows them to interact with the S. aureus cell surface and to release toxic compounds inside the cells. Notably, RLPs interacted with various Gram-positive pathogens, whose membranes contain a range of diverse phospholipid compounds and molecules (Sohlenkamp and Geiger, 2015), suggesting a “broad-spectrum” activity. Fluorescent pyochelin labeling revealed the RLPs' ability to deliver pyochelin to S. aureus cells, where it generates reactive oxygen species and participates in cell killing (Adler et al., 2012). However, the physical nature of S. aureus membrane and RLP micelle interactions will require further investigations. We further demonstrate the synergistic effect of RLPs with lincosamide antimicrobials, accelerating the killing activity of clindamycin and lincomycin against S. aureus by enhancing their delivery to their intracellular targets. The origin of the apparent selectiveness of RLP micelles for some cargo compounds remains to be elucidated.

The RLPs cargo function described in our study yields novel insights on the pathogenesis of P. aeruginosa infections and the remarkable capacities of this pathogen during interspecies competition. We previously described the selection of ΔwspF mutants of P. aeruginosa during a co-evolution experiment with S. aureus (Tognon et al., 2017). Notably, supernatants of ΔwspF mutants had the ability to kill not only S. aureus but also other Gram-positive as well as Gram-negative bacterial species. The killing activity requires the combined action of four different molecules, namely alkyl quinoline N-oxides...
(AQNOs), pyoverdine, pyochelin, and rhamnolipids (Gdaniec et al., 2020). Previous studies reported that outer membrane vesicles (OMVs) of P. aeruginosa could fuse to the membranes of other Gram-negative bacteria, as well as to S. aureus cells (Kadurugamuwa and Beveridge, 1995). OMVs incorporate hydrophobic molecules, like AQNOs and the QS signal molecule 3-oxo-C12-HSL (Calfee et al., 2005; Mashburn-Warren et al., 2008a, 2008b). While we confirmed high levels of AQNO molecules in ΔwspF mutant supernatants, the concentrations of 3-oxo-C12-HSL were lower when compared with PA14 wild type supernatants (Gdaniec et al., 2020). Notably, synthesis of AQNOs and 3-oxo-C12-HSL were strongly reduced in a ΔwspFΔrhlA mutant. Thus, in the absence of rhamnolipids, AQNOs and 3-oxo-C12-HSL may remain attached to the P. aeruginosa membrane, leading to reduced release into culture supernatants. However, whether RLP micelles can also deliver AQNOs, phenazines, or QS molecules to target bacteria remains to be determined. Our data suggest that RLPs, provided as a mixture or as single species (C10-C10, C12-C12), exert their cargo effect at concentrations above the CMC, suggesting the importance of micelles formation for the carrier function.

Altogether, our findings support a novel role for RLPs as a remarkable strategy of P. aeruginosa to transport self-produced or environmentally available metabolites to target competitor microorganisms. Our data highlight the potential for enhancing antimicrobial activity via well-designed drug delivery strategies.

Limitations of the study
We focused here on the interaction of RLP micelles with S. aureus and did not investigate other Gram-positive bacteria. Our work was not designed to identify all endogenous molecules, which could be transported by RLP micelles from P. aeruginosa. However, this could be achieved by mass-spectroscopy analyses of purified RLP micelles.

STAR METHODS
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SUPPLEMENTAL INFORMATION
Supplemental information can be found online at https://doi.org/10.1016/j.isci.2021.103669.

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AUTHOR CONTRIBUTIONS

Conceptualization and methodology, B.G.G., F.B., T.K., T.B., C.v.D., and F.P.; Investigation, B.G.G., F.B., and F.P.; Writing, B.G.G., F.B., T.K., C.v.D., and T.B.; Visualization, B.G.G. and F.B.; Funding Acquisition, B.G.G, C.v.D., and T.B.; Supervision, T.K., T.B., and C.v.D.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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**STAR METHODS**

**KEY RESOURCES TABLE**

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Bacterial and virus strains** | | |
| *Pseudomonas aeruginosa PA14* | (He et al., 2004) | N/A |
| *Pseudomonas aeruginosa PA14 Δ wspF* | (Tognon et al., 2017) | N/A |
| *Pseudomonas aeruginosa PA14 Δ wspFΔrhlA* | (Gdaniec et al., 2020) | N/A |
| *Staphylococcus aureus Newman* | (Baba et al., 2008) | N/A |
| *Staphylococcus aureus COL (MRSA)* | A. Renzoni, University of Geneva | N/A |
| **Chemicals, peptides, and recombinant proteins** | | |
| Rhamnolipids 90% | AGAE Technologies | Corvallis, OR 97333, USA |
| Rhamnolipids C10-C10 | GlycoSurf | Salt Lake City, UT 84103, USA |
| Rhamnolipids C12-C12 | GlycoSurf | Salt Lake City, UT 84103, USA |
| Pyochelin | G. Mislin, I. Schalk; University of Strasbourg, France | N/A |
| Lincomycin | Merck-Sigma-Aldrich | Cat#62143 |
| Clindamycin | Chemie Brunschwig | Cat#ACR44351-0100 |
| Vancomycin | Merck-Sigma-Aldrich | Cat#P91732 |
| Abberior STAR 580 (RED-NHS) | Merck-Sigma-Aldrich | Cat#38377 |
| Abberior STAR 488 | Merck-Sigma-Aldrich | Cat#61408 |
| Nile Red | Merck-Sigma-Aldrich | Cat#72485 |
| Oracinol | Merck-Sigma-Aldrich | Cat#447420 |
| Adipic acid dihydrazide | Merck-Sigma-Aldrich | Cat#A0638 |
| N-(3-Dimethylaminopropyl)-N'-ethylcarbodiimid-hydrochlorid | Merck-Sigma-Aldrich | Cat#BCB28350 |
| 2-(N-morpholinol)ethanesulfonic acid | Merck-Sigma-Aldrich | Cat#M3671 |

**Deposited data**

| RAW AND ANALYZED DATA | IDENTIFIER |
|-----------------------|------------|
| This paper | Zenodo; https://doi.org/10.5281/zenodo.5159447 |

**Software and algorithms**

| Fiji open-source software | https://imagej.net/loci | https://fiji.sc/ |

**RESOURCE AVAILABILITY**

**Lead contact**

Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact Thilo Köhler (thilo.kohler@unige.ch).

**Materials availability**

This study did not generate any new biological material.

**Data and code availability**

- Microscopy data reported in this paper will be shared by the lead contact upon request.
- All original code and raw data supporting the conclusions of this study have been deposited at Zenodo and are publicly available from the lead contact upon request (https://doi.org/10.5281/zenodo.5159447).
- Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.
**EXPERIMENTAL MODEL AND SUBJECT DETAILS**

**Bacterial strains, growth conditions and supernatant preparation**

Bacterial strains, plasmids and primers used in this study are listed in the key resources Table. M14 medium was adapted from the literature (Rudin et al., 1974) and is based on M9 salts (Na2HPO4 6 g/L; KH2PO4 3 g/L; NaCl 0.5 g/L; NH4Cl 1 g/L) supplemented with casamino acids (BD™, USA) 10 g/L, magnesium sulfate (MgSO4·7H2O) 1 mM, thiamine (vitamin B1) 2 mg/L, niacin (vitamin B3) 2 mg/L, calcium pantothenate (vitamin B5) 2 mg/L, biotin (vitamin B9) 0.1 mg/L and glucose 2 g/L. Casamino acids, vitamins and glucose solutions were sterilized by filtration and stored separately at 4°C. M9 salts and magnesium sulfate were sterilized by autoclaving at 121°C for 15 min (Tognon et al., 2017). Conditioned medium of bacterial cultures were recovered after 24 h of static growth at 37°C in microtiter plates (TPP, Switzerland). The cultures were pooled and centrifuged at 8,000 rpm for 5 min. Supernatants were sterilized by filtration (0.22 µm filters, Millipore, Switzerland) and stored at –20°C. Medium conditioned in this way is also referred to as crude supernatant. To determine the critical micellar concentration (CMC) of commercial RLPs or in crude supernatants, sterilized supernatants of ΔwspFΔrhlA mutant or M14 medium were supplemented with RLPs (90% purity, AGAE Technologies, USA) to the given concentrations and stained with Nile Red (Sigma Aldrich) at a final concentration of 1 mg/mL for 30 min. Crude supernatants were directly stained with Nile Red at a final concentration of 1 mg/mL for 30 min. Next, 200 µL of solution was deposited in a 96 well plate in triplicates. Fluorescence of Nile Red (ex 559 nm/em 635 nm) was measured in a plate reader (Synergy 1, Bio Tek®, USA). To concentrate the micelles and obtain the RLP pellet, 50 mL of supernatant was stained with Nile Red at a final concentration of 1 mg/mL for 30 min. Samples were centrifuged at 150,000 x g for 4 h at 4°C (Optima XPN, Beckman Coulter ROTOR TYPE 45 Ti). After ultracentrifugation supernatants were collected and the pellets resuspended in 1 mL of M14 medium yielding the 50x concentrated pellet sample, while 15x and 1x concentrated samples were prepared by dilution of 50x sample in M14 medium.

**METHOD DETAILS**

**Orcinol assay for rhamnolipids quantification**

Rhamnolipid quantification was performed as described (Wittgens et al., 2011) with slight modifications. Samples containing 50 mL of filtered supernatant before and after ultracentrifugation were treated with equal volumes of ethyl acetate. A standard curve was established by dissolving a mix of mono and di-rhamnolipids (R90-10G, Sigma Aldrich, Switzerland) in 2 mL ddH2O and subsequent extraction with equal volumes of ethyl acetate. After vortexing for 1 min, samples were centrifuged for 1 min at 5000 rpm to separate the phases. The organic solvent was evaporated in a SpeedVac (Thermo Scientific) and residues containing rhamnolipids were resuspended in 100 µL of ddH2O and mixed with an equal volume of 1.6% orcinol (Sigma Aldrich, Switzerland) in ddH2O. Subsequently, 800 µL of 60% sulfuric acid (vol/vol) were added, and samples were incubated for 45 min at 80°C. A 100 µL aliquot of the reaction mix was transferred to a microtitre plate, and OD420 was measured in a Synergy H1 Multi-Mode plate reader (BioTek®).

**Transmission electron microscopy**

Supernatant samples and RLP suspensions were loaded onto 2 mm single slot copper grids (Electron Microscopy Sciences) coated with 1% Pioloform plastic support film. Grids were dried for 30 min and then stained with 1% aqueous uranyl acetate solution for 3 min and examined using a Tecnai 20 TEM (FEI) electron microscope operating at an acceleration voltage of 80 kV and equipped with a side-mounted MegaView III CCD camera (Olympus Soft-Imaging Systems) controlled by iTEM acquisition software (Olympus Soft-Imaging Systems). Grids for EM were prepared twice independently and multiple pictures were acquired.

**Killing assays and kinetics**

Killing assays were performed on S. aureus cells grown for 6 h in M14 medium under static growth conditions in microtiter plates as described (Tognon et al., 2017). After the 6 h incubation, 100 µL of S. aureus culture was removed and replaced with either 100 µL M14 medium or crude or fractionated P. aeruginosa supernatant. Growth (OD600) was monitored in a plate reader (BioTek®, USA) for 24 h. At the end of the killing assay, viable plate counts were performed to determine S. aureus survival. For killing kinetics, samples were taken at different time intervals during growth and surviving S. aureus cells determined by viable plate counts. Other bacterial strains were grown on LB-plates or specific growth media. Cells were scraped from the plate and a suspension was prepared and adjusted to obtain 10⁸ CFU/mL.

**Soft-Imaging Systems**

Killing assays and kinetics were performed on S. aureus cells grown for 6 h in M14 medium under static growth conditions in microtiter plates as described (Tognon et al., 2017). After the 6 h incubation, 100 µL of S. aureus culture was removed and replaced with either 100 µL M14 medium or crude or fractionated P. aeruginosa supernatant. Growth (OD600) was monitored in a plate reader (BioTek®, USA) for 24 h. At the end of the killing assay, viable plate counts were performed to determine S. aureus survival. For killing kinetics, samples were taken at different time intervals during growth and surviving S. aureus cells determined by viable plate counts. Other bacterial strains were grown on LB-plates or specific growth media. Cells were scraped from the plate and a suspension was prepared and adjusted to obtain 10⁸ CFU/mL.
After 24 h incubation in the presence of culture supernatants, surviving cells were determined by plate counts.

**Minimal inhibitory concentration determinations**

To determine antibiotic susceptibility, minimal inhibitory concentrations were performed in triplicates by two-fold serial dilutions in Müller-Hinton broth (Becton Dickinson, Franklin Lakes, NJ, USA) according to Clinical and Laboratory Standard Institute (CLSI) guidelines.

**Fractional inhibitory concentration (FIC)**

For all of the wells of the 96 wells plates that corresponded to an MIC, the sum of the FICs ($\Sigma$FIC) was calculated for each well with the equation $\Sigma$FIC = FIC$_A$ + FIC$_B$ = ($C_A$/MIC$_A$) + ($C_B$/MIC$_B$), where MIC$_A$ and MIC$_B$ are the MICs of drugs A and B alone, respectively, and $C_A$ and $C_B$ are the concentrations of the drugs in combination, respectively, in all of the wells corresponding to an MIC (isoeffective combinations) (Hall et al., 1983).

**Covalent labeling of rhamnolipids, pyochelin and vancomycin with Abberior STARNHS ester dye**

A stock solution of rhamnolipids (RLP) (90% purity, AGAE Technologies, USA) was prepared at 100 mg/mL in DMSO. When necessary, sonication was performed until complete dissolution of the RLP aggregates. When required, the stock solution was further diluted in DMSO. Sequentially 800 µL of a solution of adipic acid dihydrazide (AAD, Sigma Aldrich, A0638, Switzerland) at 31.25 mg/mL in ddH$_2$O and 50 µL of Abberior STAR 488 or 580 NHS-ester (depending on the experiment, respectively Sigma Aldrich, 61,048 and 38,377, Switzerland) at a concentration of 2 mg/mL in DMSO were added. After 1 h incubation at room temperature under constant agitation (vortex on the lowest speed), 200 µL of N-(3-Dimethylaminopropyl)-N’-ethylcarbodiimide hydrochloride (EDC, Sigma Aldrich, E1769, Switzerland) at 6 mg/mL (final concentration of EDC at 0.8 mg/mL) in MES buffer 0.5 M (pH 5.5) was added. Reaction was incubated for 20 min at room temperature under constant agitation.

Finally, RLP extraction was performed with ethyl acetate as described previously with slight modifications (Loiseau et al., 2018). Briefly, 2 mL of ethyl acetate (Sigma Aldrich, 270,989, Switzerland) was added to the solution and vortexed for 30 s. After complete separation, the upper phase containing the RLP was collected and 20 µL of DMSO was added before drying for at least 2 h in a SpeedVac Vacuum (Thermofisher, Germany). RLP were suspended in the 100 µL of DI and sonication was performed until complete dissolution. Pyochelin covalent modification with Abberior STAR RED-NHS ester dye was performed as follows: 16 µg of pyochelin were incubated with 5 mg of AAD and 40 µg of Abberior STAR red-NHS ester in 200 µL of ddH$_2$O for 1 h at room temperature. Then, 300 µg of EDC solubilized in MES buffer (0.5 M, pH 5.5) were added to the reactive mix and incubated for 20 min. Pyochelin was then extracted as previously described (Hoegy et al., 2014). Briefly, adjustment to pH 3 was performed by adding 5 mg of citric acid monohydrate (Sigma Aldrich, G1909, Switzerland) before pyochelin extraction using 1 mL of dichloromethane. After vortexing and complete phase separation, the lower phase was collected. A second extraction was carried out when necessary. Extracted pyochelin was dried for at least 2 h under nitrogen flow. Labeling of vancomycin was performed by resuspending 0.2 mg of vancomycin in 500 µL of ddH$_2$O, addition of 0.1 mg of Abberior STAR 488 NHS-ester and incubation at room temperature for 2 h.

**Confocal imaging and pyochelin RLP-assisted delivery**

Inactivation of the Abberior STAR RED-NHS dye was performed to prevent any reaction with cell surface compounds. For this, Abberior STAR RED-NHS ester dye was incubated with NaOH 0.1 M for 10 min prior incubation with or without RLP. S. aureus cells were incubated in absence or presence of RLPs at different concentration 25 µg/mL (below the CMC) or 250 µg/mL (above the CMC) either with labeled pyochelin (12 ng/condition) or with inactivated dye (20 ng/conditions) in 200 µL of PBS, for 10 min at room temperature. Then cells were centrifuged and washed with 200 µL of PBS. After centrifugation, cells were incubated with 0.01 µg of labeled vancomycin in 200 µL of PBS for 5 min. Cells were centrifuged and washed again in 200 µL of PBS. 1 µL of the cell suspension was spotted onto a microscopy glass slide with 1 µL of mounting buffer. Coverslips were sealed using nail polish. A Zeiss confocal microscope LSM800 was used for all observations.
STED nanoscopy

2D-STED dual color imaging was performed with a Leica TCS SP8 STED 3X microscope in a thermostatic chamber at 21 °C and equipped with a STED motorized glycerol immersion objective (HC PL Apo 93X/N.A. 1.30 motCOR). Fluorescently-labelled samples were mounted in Prolong Antifade Gold (Thermofisher Scientific) between a coverslip (0.170 ± 0.01 mm thick, Hecht-Assistent) sealed on a microscope slide with nail polish. Excitation was performed with a White Light Laser (WLL), depletion with either a continuous 592 nm laser (STED 592) or a 775 nm pulsed laser (STED 775). Excitation and depletion lasers were calibrated with the STED Expert Alignment Mode and Abberior gold nanoparticles (80 nm in diameter) before starting each imaging session, or with the STED Auto Beam Alignment tool during imaging sessions (Leica LAS X software, Leica Microsystems CMS GmbH). 2D-STED dual color was made using an excitation at 559 nm (WLL) and a STED 775 depletion laser line for Nile Red, followed by an excitation at 488 nm (WLL) and a STED 592 depletion laser line for Abberior STAR 488. Detection signals were collected from 650 nm to 740 nm for Nile Red and from 500 nm to 564 nm for Abberior STAR 488 using highly sensitive Leica Hybrid Detectors (HyDs) with a fixed gain and offset (100 mV and 0, respectively). Time-gated detection was used for Nile Red (0.50–5.92 ns). Acquisitions were performed sequentially with a line average of 4, a speed of 400 Hz and an optimized pixel size (20 nm). Images were deconvolved using the Leica Lightning Mode (LAS X software) and analyzed with Fiji open-source software (https://fiji.sc/).

QUANTIFICATION AND STATISTICAL ANALYSIS

All statistical analysis was performed with GraphPad Prism version 8.0 and R package. To compare killing activities of crude-S, ultra-S and ultra-P fractions of PA14 wild type and mutants, we extracted the slope in a log-diagram up to 3 h by linear regression. Saturation due to detection limits did not allow evaluation after 3 h. The data consisted of three replica per condition (Figure S1A). All triplets showed similar variance (Bartlett test failed to reject the hypothesis of identical variance in all groups of three killing curves, p = 0.17), therefore, unpaired t-tests assuming equal sample variance were used for individual comparisons (Figure S1). To compare the rescue efficiency of different rhamnolipid supplements (Figure S1E), rescue efficiency was calculated as the enhancement of killing activity over the corresponding unsupplemented controls. Given the presence of both supernatant and pellet fractions, ANOVA was then performed, followed by Tukey’s test for honest significant differences to analyze the effect of different RLP supplemenations. Experiments were performed with three biological replicates from at least two independent experiments when possible. Statistical significance is reported in Figure Legends, and data are presented as mean +/- SD as indicated.