Discovery and Heterologous Expression of Microginins from *Microcystis aeruginosa* LEGE 91341

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**ABSTRACT:** Microginins are a large family of cyanobacterial lipopeptide protease inhibitors. A hybrid polyketide synthase/non-ribosomal peptide synthetase biosynthetic gene cluster (BGC) found in several microginin-producing strains—mic—was proposed to encode the production of microginins, based on bioinformatic analysis. Here, we explored a cyanobacterium, *Microcystis aeruginosa* LEGE 91341, which contains a mic BGC, to discover 12 new microginin variants. The new compounds contain uncommon amino acids, namely, homophenylalanine (Hphe), homotyrosine (Htyr), or methylproline, as well as a 3-aminodecanoic acid (Ada) residue, which in some variants was chlorinated at its terminal methyl group. We have used direct pathway cloning (DiPaC) to heterologously express the mic BGC from *M. aeruginosa* LEGE 91341 in *Escherichia coli*, which led to the production of several microginins. This proved that the mic BGC is, in fact, responsible for the biosynthesis of microginins and paves the way to accessing new variants from (meta)genome data or through pathway engineering.

**KEYWORDS:** cyanobacteria, natural products, direct pathway cloning (DiPaC), heterologous expression, microginins, biosynthesis

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**INTRODUCTION**

Cyanobacteria are rich secondary metabolite producers. This is evidenced by the large number of natural products reported from these organisms and by the presence of multiple biosynthetic gene clusters (BGCs) in their genomes. In line with what is observed for most bacteria, cyanobacterial BGCs greatly outnumber the reported natural products for this phylum. These observations highlight the potential for the discovery of new natural products from cyanobacteria, but they also underscore the lack of effective tools for revealing the natural products associated with orphan BGCs (i.e., those for which no compound can be ascribed). Deorphanizing BGCs from bacteria has been one of the main challenges in natural products discovery ever since next-generation sequencing became available at large. For many natural products-rich organisms, genetic tools and appropriate heterologous hosts are readily available to reach this goal. However, modifying cyanobacterial genomes, in particular of non-model strains, is a slow and complicated process, even if possible. The heterologous expression of whole cyanobacterial BGCs in a non-cyanobacterial host has long been considered as a valid strategy but has progressed slowly, in particular due to challenges in finding suitable hosts. Nevertheless, it has been possible to express over a dozen cyanobacterial BGCs, in particular in small-sized clusters, in *Escherichia coli* or cyanobacterial hosts. Direct pathway cloning (DiPaC) is a recently developed strategy for BGC capture and heterologous expression that has shown promise in the expression of cyanobacterial BGCs, including complex polyketide synthase/non-ribosomal peptide synthetase pathways.

Microginins are a large group of over 80 cyanobacterial secondary metabolites. Their structures can be summarized as linear lipopeptides featuring a 3-amino fatty acyl residue connected to a peptide (typically three or four amino acid residues). Modifications of the fatty acyl residue such as terminal halogenation, hydroxylation, or N-methylation are common. Several microginins were shown to inhibit a variety of proteases with considerable potency. Rouge et al. proposed that microginin production is carried out by enzymes encoded in the mic BGC (Figure 1a), found in the genome of the oscillaginin (a type of microginin)-producing strain *Planktothrix prolifica* NIVA-CYA 98. This proposal was grounded on the predicted functions of the mic genes, which were consistent with the generation of a metabolite with a structure similar to that of oscillaginins A and B. Specifically, the presence of NRPS adenylation domains, whose predicted specificity matched the amino acids found in the oscillaginins,
and co-linearity between such adenylation and the sequence of amino acids in these peptides formed the basis for the proposal. However, the mic BGC reported by Rounge et al. with a single PKS module and three NRPS modules could not explain the origin of the Ahda moiety in oscillaginins. Later, Kramer reported a fatty acyl AMP-ligase (FAAL) enzyme associated with a Microcystis aeruginosa mic BGC and proposed that it would activate and load octanoic acid onto an acyl carrier protein (ACP), which would then interact with the PKS module to generate the Ahda residue. The biosynthetic origin of the halogenations that are often observed in microginins was proposed only after the discovery of dimetal carboxylate halogenases—Nakamura et al. identified one putative halogenase of this type within a mic BGC from M. aeruginosa PCC 9432. In line with these observations, the updated genome data for the oscillaginin-producing P. prolifica NIVA-CYA 98 (NCBI: AVFZ00000000) include a longer contig harboring the mic BGC, extending its 5′ region, which features additional genes encoding FAAL, ACP, a dimetal-carboxylate halogenase, and a SAM-dependent methyltransferase (Figure 1a). In addition, a luciferase-like monooxygenase domain is annotated in its micD gene (Figure 1a) by the NCBI.

Figure 1. Microginin BGCs (mic) and discovery of microginins 1–12. Architecture of the mic BGC from M. aeruginosa LEGE 91341 and of the initially proposed and now reannotated mic BGC from P. prolifica NIVA-CYA 98 (asterisks indicate genes/domain annotated in this study), putatively encoding the production of oscillaginins A and B (a). Structures of the herein reported microginins 1–12 (b), colors of the circle for each group correspond to those in panel c. The compound isolated and structurally characterized using NMR is labeled with “NMR”. LC–HRESIMS analysis of an organic extract of M. aeruginosa LEGE 91341 cells indicates the presence of four groups of microginins, each consisting of non-chlorinated, mono- and di-chlorinated variants, as illustrated by the represented extracted ion chromatograms for the [M + H]+ ion of each of these compounds (c).
Conserved Domain Search tool. Despite these bioinformatics-based data, direct biochemical evidence connecting microginins with their putative BGC (mic) is lacking. Our group recently detected a putative mic BGC in the cyanobacterium M. aeruginosa LEGE 91341 (NCBI: JADEXY0000000000) encoding a dimetal carboxylate halogenase homolog (Figure 1a). Here, we report the discovery of 12 new microginins (compounds 1−12, Figure 1b) from this strain, including mono- and di-chlorinated as well as non-halogenated congeners. The mic BGC from M. aeruginosa LEGE 91341 was heterologously expressed in E. coli using DiPaC, leading to the production of several of the microginins found in the cyanobacterium but also of new variants. Our work confirms that mic is the microginin BGC and opens the door for accessing novel microginin diversity, namely, through the expression of additional variants of mic BGCs captured from (meta)genomic data or by engineering the mic pathway. Furthermore, this study reinforces the usefulness of the DiPaC strategy for cyanobacterial NRPS/PKS BGC deorphanization.

## RESULTS AND DISCUSSION

### Discovery of New Microginins from M. aeruginosa LEGE 91341

Our previous observation of a putative microginin BGC (mic) in M. aeruginosa LEGE 91341...
motivated a search for known or new microginins in this strain. AntiSMASH-based annotation of this BGC allowed for predicting the specificity of three out of the four NRPS modules: two would incorporate Pro and one would incorporate Tyr. Manual inspection of liquid chromatography high-resolution electrospray ionization mass spectrometry (LC−HRESIMS) data for a methanolic extract of *M. aeruginosa* LEGE 91341 revealed 12 features corresponding to compounds with isotope patterns characteristic of one or two chlorinations or to their non-halogenated counterparts (Figure 1c). The most abundant features were in the m/z 600–800 range ([M + H]+), which would be consistent with microginin-like metabolites. Overall, these mass features could be considered as making up four groups of analogues, each group containing variants with different degrees of halogenation (absent, one and two chlorine atoms). Searches for the accurate masses (±5 ppm) of the compounds in the NP Atlas and Dictionary of Natural Products returned no hits for cyanobacteria, suggesting that these were novel compounds.

**Isolation and Structure Elucidation of Microginin 789 (1).** Given the apparent abundance of some microginins in the small-scale organic extract of *M. aeruginosa* LEGE 91341 (Figure 1c), we focused next on the isolation of sufficient amounts of these metabolites to enable NMR-based structure elucidation. Fermentation of *M. aeruginosa* LEGE 91341 in 120 L of the culture medium yielded 42 g of biomass (d.w.), which was extracted by repeated percolation with methanol. The resulting crude extract (7.8 g) was subjected to reversed-phase (C18) vacuum liquid chromatography (VLC) with a stepwise gradient of decreasing polarity from water to MeOH, to give seven fractions (A−G). LC−HRESIMS analysis indicated that fraction E, eluting with 100% MeOH, contained dihalogenated 1 in high abundance. Global Natural Products Social Molecular Networking (GNPS) analysis of this fraction revealed what could be additional minor microginins (Figure S1), but their identification was not pursued in this study.

![Figure 2. Structural elucidation of microginins. Key NMR correlations supporting the proposed structure of microginin 789 (1) (a); HRESIMS/MS-based deduction of the structures of the remaining microginins, as illustrated for the monochlorinated 2, 5, 8, and 11 (b).](https://doi.org/10.1021/acssynbio.2c00389)
Subsequent reversed-phase flash chromatography and high-performance liquid chromatography (HPLC) steps afforded pure microginin 789 (1, 2.8 mg). This compound was subjected to 1D and 2D NMR-based structure elucidation. The $^1$H NMR data for 1 (Table 1) showed the presence of two broad singlets resonating at approx. 9 ppm, two exchangeable amide protons doublets ($\delta^1_H 8.28$ and 8.12), several protons in the aromatic/olefin region, and resonances around $\delta^1_H \sim 5.0$–3.5 that could correspond to alpha protons. In addition, a methylene envelope was observable ($\delta^1_H \sim 1.4$–1.2), and a

Figure 3. Heterologous expression of the mic BGC in E. coli leads to the production of microginins. Two-step SLIC-based DiPaC strategy used to clone and express the mic BGC from *M. aeruginosa* LEGE 91341 in *E. coli* (a). LC–HRESIMS analysis of *E. coli* culture extracts reveals the production of microginins in strains carrying the mic BGC that are not found in the producing cyanobacterium. Shown are EICs and proposed peptide sequences for novel compounds 13–16 (b). Supplementation with Hphe leads to the production of microginins 5 and 6 by *E. coli* harboring the mic BGC genes (c), which are identical to the ones produced by *M. aeruginosa* LEGE 91341, as confirmed by HRESIMS/MS analysis (d).
The terminal methyl triplet group resonance for a saturated alkyl chain was not present. The $^{13}$C NMR data (Table 1) notably indicated the presence of five carbonyls ($\delta_C \sim 175-169$) and two aromatic, likely phenol, moieties (eight resonances in the $\delta_C \sim 160-110$ range). These $^1$H and $^{13}$C NMR data would thus be consistent with a lipopeptide containing two aromatic residues. Additional structural information was obtained by analysis of the 2D NMR data. A combination of heteronuclear single quantum coherence (HSQC), heteronuclear multiple bond correlation (HMBC) and correlation spectroscopy (COSY) data (Table 1) was used to establish several substructures of 1, as follows. Two very similar aromatic spin systems could be readily constructed. The first corresponded to a para-substituted phenol that further connected to an alpha proton (H-32, $\delta_H$ 3.96) through a diastereotopic methylene (CH$_2$-33, $\delta_{Ha}$ 3.04, $\delta_{ab}$ 2.68) and was attributed to a Tyr residue. The second also corresponded to a para-substituted phenol moiety, which was HMBC-correlated to a COSY-derived spin system composed of two methylene groups (CH$_2$-19 and CH$_2$-18) and an alpha position (CH-17), thereby establishing a homotyrosine (Htyr) moiety. The two amide protons were COSY-correlated to each of the aromatic residue alpha protons (Table 1 and Figure 2a). Two proline residues could also be established (Table 1 and Figure 2a) and these explained the absence of additional amide protons. A conspicuous methine (CH-10, $\delta_C$ 74.8, $\delta_H$ 6.25) was assigned as a dichloromethyl group, taking into account the particular stereochemistry of all the proteinogenic amino acid residues in 1 and was considered to be the bridging position between the two alkyl chain spin systems (CH$_2$-8). This established that 1 contains a dichlorinated 3-amino decanoic acid (Ada) residue (Table 1 and Figure 2a). Inter-residue HMBC correlations supported that 1 corresponds to a linear peptide of sequence [Cl$_2$-Ada]-Pro-Htyr-Pro-Tyr (Table 1 and Figure 2a). The stereochernistry of all the proteinogenic amino acid residues in 1 was deduced, from the absence of epimerase domains, as being of the L configuration, while the stereochernistry of the Htyr residue and of the $\beta$-amino group were not elucidated. While angiotensin-converting enzyme (ACE) inhibitory activity has been reported for several microginins, compound 1 did not show ACE inhibitory activity at a concentration of up to 1 $\mu$M (Figure S2).

MS/MS-Based Structure Elucidation of Additional Microginins (3–12). To obtain further structural information, we acquired MS/MS data for the 12 new microginins, (Figures S3–S14). To establish a comparative framework, we looked in detail at the MS/MS data for the monochlorinated representative of each of the four different groups (compounds 2, 5, 8, and 11, Figure 2b). Extensive analysis of microginin fragmentation under ESI conditions is available in the literature (Table S1) and was used as the reference. Notably, the MS/MS spectra of 2, 5, 8, and 11 contained an m/z 162 ion corresponding to fragmentation of a chlorinated 3-amino decanoic acid-derived moiety. The three larger metabolites (2, 5, and 8) were composed of a C-terminal Pro--Tyr moiety, giving rise to a characteristic m/z 279 fragment ion. The smaller compound (microginin 592) did not show this ion and instead showed a m/z 116 fragment ion corresponding to a C-terminal Pro. From this observation and because the mass difference between 2 and 11 is consistent with a Tyr residue, we propose that 10, 11, and 12 are truncated versions of 1, 2, and 3, respectively, lacking the C-terminal Tyr.

Prominent MS/MS ions of m/z 112 and 140, observed for 2, 5, and 11 and corresponding to Ada-Pro fragments were shifted by 14.015 a.m.u. (CH$_3$) in compound 8. Given the absence of methyltransferase genes/domains in the mic BGC of M. aeruginosa LEGE 91341, which could lead to N-methylation, we propose that methyl proline (mPro) is present in 7, 8, and 9, as observed in other cyanobacterial natural products. It is unclear how this modification takes place in the biosynthesis of 7–9, and thus, we refrain from ascribing the methylation to a specific Pro carbon.

The MS/MS spectrum for compound 5 shows a prominent m/z 462 peak that is not observed in 2, while the m/z 478 peak corresponding to the neutral loss of Pro--Tyr in 2 is not observed in 5. The difference between these two fragments (15.994 a.m.u.) corresponds to an oxygen atom. Considering that, as described above, the [Cl-Ada] and Ada-Pro fragmentations in 2 and 5 are identical, we ascribe the oxygen atom difference to an Hphe residue being present in 4, 5, and 6, instead of the Htyr residue found in 1, 2, and 3. Analogues of cyanobacterial secondary metabolites with this type of modification have been described previously and are related to the relaxed specificity of the corresponding NRPS adenylation domain.

Overall, our findings confirmed that M. aeruginosa LEGE 91341 produces 12 new microginins. These congeners contain the less common Ada fatty acyl moiety and a new combination of amino acids when compared to previously reported microginins. In addition, neither Hphe nor mPro had previously been encountered in this large group of cyanobacterial peptides.

Heterologous Expression of the mic BGC. To obtain biochemical evidence for the involvement of mic genes in the biosynthesis of microginins, we sought to heterologously express the mic BGC from M. aeruginosa LEGE 91341 in E. coli. A few NRPS/PKS pathways from cyanobacteria have been successfully expressed heterologously (using cyanobacterial or E. coli hosts). Using a Sequence- and Ligation-Independent Cloning (SLIC) DiPaC strategy, we assembled the roughly 25 kb micro gene cluster from M. aeruginosa LEGE 91341 into a modified pET-28 vector backbone, which includes a PctO promoter and generates a C-terminal gfp fusion for the last gene in the BGC (pET28-pctO:gfp2, Figure 3a). The cloning strategy was achieved through SLIC ligation of two PCR-derived ampiclons (Figure 3a). The resulting construct was cloned into the E. coli BAP1 strain, which has a chromosomal copy of the gfp gene, encoding a promiscuous Ppant transferase. This enables the post-translational phosphopanthetheinylation of the several thiolation domains encoded in the mic BGC. Comparison of the LC--HRESIMS profiles of E. coli pET28-pctO:mircABCDEFG-gfp2 transformants with those of empty vector transformants, both grown in the minimal M9 medium supplemented with a vitamin mix (see the Methods section and Table S1), revealed MS features that were only present in cells containing the mic BGC (Figure 3b). Full MS spectra for these peaks showed [M + H]$^+$ ions for non-halogenated compounds of m/z 658.417 (13) and 672.433...
Table 2. Strains and Plasmids Used in the Present Study

| strain                  | description                          | reference or source          |
|-------------------------|--------------------------------------|------------------------------|
| E. coli TOP10           | cloning host strain                  | ThermoFisher Scientific      |
| E. coli BAP1            | strain for heterologous expression   | Kerafast                     |
| M. aeruginosa LEGE 91341| microginins native producer          | LEGEcc culture collection    |
| pET28-pterO::gfpv2 (6552 bp) | tetracycline inducible expression plasmid, ColE1, KanR, gfp reporter gene downstream of the promoter | Duell et al.28 |
| pET28-pterO::micABCDE-gfpv2 (22,033 bp) | pET28b-pterO::gfpv2 with micABCDE cloned as the first fragment between PetO and gfp construct built using pET28-pterO::micABCDE-gfpv2 as a vector and miFG as a single piece nucleotide insert between the ligation site and gfp | this study |
| pET28-pterO::micABCDEFG-gfpv2 (31,587 bp) | | this study |

*Isolation of 1 from large-scale culturing of M. aeruginosa LEGE 91341.

(14) as well as minor amounts of 495.354 (15) (Figure 3b). A monochlorinated version of 13 (m/z 692.378, [M + H]**+) could also be detected (16). LC−HRESIMS/MS analysis of the most abundant compounds 13, 14, and 16 (Figure S15) indicated that these new metabolites are microginins related to 1−12, in which Leu or Ile replaces Htyr/Hphe. The formation of 13−16 can be explained by the fact that the mic BGC does not include the hphpABCD operon necessary for the production of Htyr/Hphe.24 In fact, this set of genes is found elsewhere in the genome of LEGE 91341, as part of an anabaenopeptin BGC (Table S2). Supplementation of the mic BGC transformant cultures with exogenous Hphe led to the detectable production of 6 and 8 (Figure 3c)—as confirmed by MS/MS analysis (Figure 3d)—but did not lead to the production of dichlorinated 4 (Figure 3c). Dichlorinated microginin variants were also not observed in the absence of exogenously supplied Hphe. Increasing NaCl levels boosted monochlorinated but not dichlorinated microginin production (Figure S16). Interestingly, the addition of NaBr to the culture medium led to the production of mono- and di-brominated microginin variants (Figure S16), which prompts future investigations of the MicC halogenase. Using an LC−HRESIMS calibration curve for compound 1 (Figure S17) and assuming similar ionization behavior, we estimated that pellet and supernatant levels of the major microginin in E. coli (13) amount to 1.3 mg L**−1 in yield, compared to 2.8 mg L**−1 of compound 1 in the cyanobacterial cultures, as estimated from crude extracts. We tested the influence of induction and growth period on microginin production in E. coli. We found that tetracycline is required for expression (Figure S16) and that higher yields are obtained with a 2 day growth period (Figure S16, roughly corresponding to a 2-fold increase in supernatant titers of 13). Overall, the heterologous expression data confirm experimentally that the mic BGC is responsible for microginin production and that all the structural features (including halogenation) of the natural microginins detected in M. aeruginosa LEGE 91341 can be generated in E. coli.

## METHODS

**General Experimental Procedures.** Flash chromatography was performed in a Pure C-850 Flash Prep chromatography system (Buchi). HPLC separations were performed in a Waters Alliance e2695 Separations Module instrument with a 2998 PDA module and an automatic Waters Fraction Collector III or on a Waters 1525 binary pump, coupled to a Waters 2487 UV/vis detector (for the last step of the purification of compound 1). LC−HRESIMS and LC−HRESIMS/MS analyses were performed on an UltiMate 3000 UHPLC (Thermo Fisher Scientific) system composed of a LPG-3400SD pump, a WPS-3000SL autosampler, and a VWD-3100 UV/VIS detector coupled to a Q Exactive Focus Hybrid Quadrupole Orbitrap mass spectrometer controlled by Q Exactive Focus Tune 2.9 and Xcalibur 4.1 (Thermo Fisher Scientific). For LC−HRESIMS data, the full scan mode was used with a resolution of 70,000 full width at half-maximum (fwhm), a capillary voltage of −3.8 kV, a capillary temperature of 300 °C, and a sheath gas flow rate of 35 units. The LC−HRESIMS/MS analysis was performed on the same system, and fragmentation was carried out at 17,500 fwhm, using higher energy collision dissociation (HCD), a 3.0 a.m.u. isolation window, and a normalized collision energy of 35. Optical rotations were measured using a JASCO P-2000 polarimeter (JASCO) using SpectraManager 2.14.02 software. The UV spectra were acquired on a UV-1600PC spectrometer (VWR) controlled using MWAVE 1.0.20 software. 1D and 2D NMR data were obtained in the Materials Center of the University of Porto (CEMUP) on a Bruker Avance III, 400 MHz controlled using TopSpin 3.2. Compounds were analyzed in DMSO-d_6 (Sigma). Chemical shifts (δ H and δ C) are expressed in δ (ppm), referenced to the residual non-deuterated solvent (δ_H 2.5000, δ_C 39.520). NMR data were visualized in MestreNova 12.0.4. All solvents used were MS grade for MS-based experiments, HPLC gradient grade for...
HPLC analysis/purification, and ACS grade for extraction, VLC, and flash chromatography. i-homophenylalanine was purchased from TCI Europe.

**Strains and Plasmids.** The strains and plasmids used in this work and their characteristics are summarized in Table 2.

### Small-Scale Extraction of *M. aeruginosa* LEGE 91341 and LC−HRESIMS Analysis.** The strain *M. aeruginosa* LEGE 91341 was grown for 20 days in 50 mL of Z8 medium, at 25 °C, under a 14 h/10 h light (10–30 μmol photons s \(^{-1}\) m \(^{-2}\))/dark cycle. Biomass from the stationary phase was harvested by centrifugation (5000 g, 10 min, 4 °C, Gyrozen 2236R) and frozen. For small-scale extraction, 30 mg of the frozen biomass was homogenized using liquid nitrogen and extracted using methanol (2 × 20 mL) at room temperature with shaking. The supernatant was obtained by centrifugation, and the solvent was evaporated in a rotary evaporator. The resulting extract was weighted, resuspended in MeOH (1 mg mL \(^{-1}\)), and filtered through a 0.2 μm regenerated cellulose syringe filter. Ten microliters (10 μL) of the extract were injected into the LC−HRESIMS system fitted with an ACE UltraCore 2.5 Super C18 (75 mm × 2.1 mm) column. Samples were eluted at 0.35 mL min \(^{-1}\) using a linear gradient from 99.5% solution A (95% H\(_2\)O, 5% MeOH, and 0.1% HCOOH) v/v/v and 0.5% solution B (95% isopropanol, 5% MeOH, and 0.1% HCOOH, v/v) to reach 10% solution B over 0.5 min, followed by an increase to 60% solution B in 8 min and then to 90% in 1 min; these conditions were maintained for 6 min before returning to the initial conditions. The column oven was set at 40 °C, and UV monitoring was carried out at 215 and 254 nm. These chromatographic conditions were used in subsequent analyses.

In order to obtain sufficient amounts of microginins for NMR-based structure elucidation from the biomass of *M. aeruginosa* LEGE 91341, a large-scale cultivation was carried out. A total of 120 L of *M. aeruginosa* cultures in Z8 medium was carried out in 20 L of polycarbonate carboys (Nalgene). Biomass from stationary-phase batch cultures was harvested by centrifugation at 8500 g, lyophilized, and stored at −20 °C until extraction. The freeze-dried biomass (42 g) was extracted with MeOH (6 × 500 mL). The resulting crude extract was filtered through a 0.2 μm regenerated cellulose syringe filter. Ten microliters (10 μL) of the extract were injected into the LC−HRESIMS system fitted with an ACE UltraCore 2.5 Super C18 (75 mm × 2.1 mm) column. Samples were eluted at 0.35 mL min \(^{-1}\) using a linear gradient from 99.5% solution A (95% H\(_2\)O, 5% MeOH, and 0.1% HCOOH) v/v/v and 0.5% solution B (95% isopropanol, 5% MeOH, and 0.1% HCOOH, v/v) to reach 10% solution B over 0.5 min, followed by an increase to 60% solution B in 8 min and then to 90% in 1 min; these conditions were maintained for 6 min before returning to the initial conditions. The column oven was set at 40 °C, and UV monitoring was carried out at 215 and 254 nm. These chromatographic conditions were used in subsequent analyses.

**Microginin 789 (1) (White Amorphous Solid).** \([\alpha]_D^{27} \pm 8^\circ (c 0.1, \text{MeOH}); \text{UV (MeOH)} \lambda_{\text{max}} (log ε) 228.5 (3.26), 275.5 (3.18) \] (H and Cl NMR data, see Table 1 and Figures S18–S22; HRESIMS m/z: 790.3333 [M + H\(^+\)) \(= 30.5 \text{ min).}

**Molecular Networking.** To investigate if other microginins are being produced by *M. aeruginosa* LEGE 91341, each VLC fraction was analyzed by LC−HRESIMS/MS. Raw data files of each fraction but also the blanks of the run were converted to the mzML format using MSConvert, using the parameters recommended for GNPS molecular networking. The appropriate files were uploaded to the GNPS web platform using the default parameters recommended for high-resolution MS data. The molecular network was created using feature-based molecular networking analysis on GNPS and visualized in Cytoscape 3.9.0.

**Cloning of the mic BGC from *M. aeruginosa* LEGE 91341.** DiPaC-SLIC is based on long amplification PCR combined with homologous nucleotide overhangs, which allows for in vitro DNA assembly via SLIC of the BGC directly into the selected expression vector, in this case pET28-iptetO:gfpv2, but a restriction site for PmlI (NEB). The cloning of the *mic* BGC was achieved via pET28-iptetO:gfpv2, the first step of the cloning strategy was to identify putative terminators in the *mic* BGC using the ARNold tool,\(^{29,30}\) but none were detected. Because after several attempts, we were unable to amplify the entire 25 kb *mic* BGC, it was split in two fragments (Figure 3a) to facilitate their insertion into pET28-iptetO:gfpv2. The primer pairs used in the cloning strategy are detailed in Table S3. The vector backbone (pET28-iptetO:gfpv2) was linearized by PCR amplification. The *micABCDE* fragment was PCR-amplified with homology tails for pET28-iptetO:gfpv2, but a restriction site for PmlI (NEB) was added to the 3’ end of the amplicon to allow linearization of the large pET28-iptetO:micABCDE-gfpv2 construct through endonuclease digestion. The *micFG* fragment was obtained...
through PCR with homology tails to ptetO:micABCDEF-gfpv2, which enabled assembly through SLIC. The ptetO:micABCDEF-gfpv2 construct was then used for heterologous expression (Figure S3a). Amplification of all fragments was carried out as standard 50 μL PCR reactions with a final concentration of 1× Q5 High-Fidelity 2× Master Mix (NEB), 0.5 μM of forward and reverse primers, and 50–100 ng of DNA template using a Veriti Thermal Cycler. Thermal cycling programs varied according to the template and were as follows: initial denaturation at 98 °C for 30 s; denaturation at 98 °C for 10 s; annealing at 62 °C for linearization of vector backbone pET28-p tetO:gfpv2, 63 °C for amplification of the micABCDF amplicon, or 57 °C for amplification of the micEF amplicon; and extension at 72 °C, 6 min for pET28-p tetO:gfpv2 or 8 min for both micABCDF and micFG, followed by a final extension of 10 min at 72 °C. The PCR product of vector pET28-p tetO:gfpv2 was first treated with DpnI (NEB) (37 °C 1 h and 65 °C for 20 min) in order to remove the template plasmid that could act as the transformation background. Prior to cloning, all linear fragments were purified by gel excision using the NZYGelpure kit (NZYTech) and eluted in 30 μL. For the construction of ptetO:micABCDEF-gfpv2, the SLIC method was performed using 1× buffer 2.1 (NEB), 0.5 μL of T4 polymerase (NEB), a variable volume of linear DNA fragments for assembly (concentrations calculated as described by Greunke et al.9 ranging from 0.02 to 0.5 pmol), and molecular biology-grade water in a total reaction volume of 10 μL. The SLIC reaction mixture was incubated for 30 s at 50 °C and then for 10 min on ice. A total of 5 μL of the reaction mixture was transferred by heat shock into chemically competent E. coli TOP10 cells. The presence of positive clones was evaluated by colony PCR (Table S3), using a primer pair amplifying a portion of the ptetO:micA region. Clones leading to amplicons of the expected size were grown overnight at 37 °C in liquid LB medium supplemented with 50 μg mL⁻¹ of kanamycin, with shaking at 180 rpm. Plasmids were isolated from overnight cultures using the NZYMiniprep kit (NZYTech) and sent for sequencing to confirm the integration site. Next, the vector backbone ptetO:micABCDEF-gfpv2 (22 Kbp) was treated with PmlI (NEB) (ca. 1 μg of DNA of ptetO:micABCDEF-gfpv2 at 37 °C for 1 h and 65 °C for 20 min). The micABCDF amplicon was purified by gel excision, as indicated above, and the construction of ptetO:micABCDEF-gfpv2 (31 Kbp) was performed by the SLIC reaction using the same conditions described above. To confirm the positive transformants, colony PCR was performed using two primer pairs amplifying micEF and micG-gfpv2 regions. Purified plasmids were obtained and sequenced as detailed above. The positive transformants confirmed by sequencing were then cryopreserved and used for heterologous expression. The entire BGC in the construct was sequenced by primer walking, and no mutations were detected (Figure S23).

**Heterologous Expression.** For heterologous expression experiments, E. coli BAP1 cells were transformed by electroporation with pET28-p tetO:gfpv2 (empty vector) or with ptetO:micABCDEF-gfpv2. Expression cultures were carried out using M9 minimal medium [17.1 g L⁻¹ Na₂HPO₄, 12H₂O, 3.0 g L⁻¹KH₂PO₄, 1.0 g L⁻¹NH₄Cl, 0.5 g L⁻¹NaCl, 1.0 mL⁻¹MgSO₄ (2 M), and 0.2 mL⁻¹CaCl₂ (0.5 M), pH 7.0] to which 10.0 mL of glucose solution (40% [w/v] and 2 mL of Vitamin mix (500X) were added after autoclaving, and which was then filtered (0.2 μm) and supplemented with 50 μg mL⁻¹ kanamycin. For cultures additionally supplemented with 1-homophenylalanine (Hphe), this amino acid was dissolved in the culture medium at a concentration of 1.0 mM prior to inoculation. For cultures additionally supplemented with NaCl (final concentration of 2 g L⁻¹) or NaBr at equivalent concentrations, the salts were dissolved in the culture medium prior to inoculation.

For heterologous expression, 500 μL of a 37 °C LB overnight starter culture (supplemented with 50 μg mL⁻¹ kanamycin) was inoculated in 50 mL of M9 minimal medium in 100 mL Erlenmeyer flasks. Cultures were grown at 37 °C with shaking (180 rpm) until reaching an OD₆₀₀ of 0.25–0.3, at which point the cultures were placed at 4 °C for ~1 h. The cultures were then induced with 50 μL of 0.5 mg mL⁻¹ tetracycline. Expression was carried out for 4 days at 20 °C with shaking (180 rpm). To test the microginin production over time, cultures were harvested on the 2nd, 3rd, 4th, and 5th day of growth. In experiments testing the effect of tetracycline induction in microginin production, four conditions were tested: no induction and final concentrations of 0.2, 0.5, and 1 μg mL⁻¹ of tetracycline in 50 mL of culture. The cultures were harvested by centrifugation (5000g, 10 min at 4 °C). Cell pellets were extracted with EtOAc (20 mL) for 1–2 h with shaking and then centrifuged. The supernatants were transferred to a previously weighed vial and dried in a rotary evaporator. A second extraction was performed using 100% MeOH for 1–2 h with shaking and then centrifuged, and the supernatants were transferred to the same vial. In the case of culture supernatants, a liquid–liquid extraction was performed using 3 × 3 volumes of EtOAc. The organic phases were transferred to a round-bottom flask, dried in a rotary evaporator, and then transferred to a previously weighed vial. The extracts were weighted, dissolved in MeOH to a concentration of 5 mg mL⁻¹, filtered (0.2 μm), and used for LC–HRESIMS (or LC–HRESIMS/MS) analysis. Differences between the control (E. coli carrying the empty vector pET28-p tetO:gfpv2) and E. coli carrying the pET28-p tetO:micABCDEF-gfpv2 plasmid were detected by manual inspection of the resulting data.

**Quantification of Microginins in M. aeruginosa and E. coli.** To quantify the production of microginins in M. aeruginosa LEGE 91341 and E. coli, a calibration curve for microginin 789 (1) was obtained from LC–HRESIMS analysis. Briefly, standards of 1 were prepared at concentrations of 0.05, 0.01, 0.005, and 0.001 mg mL⁻¹. Five microliters (5 μL) of each standard solution were injected under the same conditions as those used for the analysis of cyanobacterial and E. coli extracts, as detailed above. The area under the curve (AUC) from the extracted ion chromatograms (EIC) for 1 was plotted (0.05 mg mL⁻¹ to 0.001 mg mL⁻¹), and a linear regression was fitted. This was then used to extrapolate (rough estimation assuming a similar ionization behavior) the concentration of each microginin in cyanobacteria and E. coli extracts.

**ACE Inhibition Assay.** ACE inhibitory activity was evaluated using the ACE activity assay test (Sigma-Aldrich CS0002). This assay is based on the hydrolysis of angiotensin I by the ACE to yield active angiotensin II. This test uses a synthetic fluorogenic peptide as the substrate, and the measured fluorescence is directly proportional to the ACE activity present. Briefly, all reagents were diluted in the assay buffer, according to the kit instructions. A standard curve was prepared in 100 μL of assay buffer per well ranging from 0 to 8 nmol for calculating the enzymatic activity. A total of 10 μL of
1 (final concentration of 1 μM), an inhibitor of ACE activity (lisinopril, Cayman Chemical) (a final concentration of 12 nM), a positive control of the ACE (provided by the kit), and a negative control (assay buffer) were pipetted into a 96-well black opaque plate. Then, 40 μL of the ACE, provided in the kit, were added to the sample and control wells and incubated at 37 °C for 5 min. The reaction was initiated by adding 50 μL of the fluorogenic substrate to experimental, control, and blank sample wells. The reaction was carried out at 37 °C, and fluorescence was read every minute for 5 min using a microplate reader (BioTek Cytation 5 Cell Imaging Multimode Reader, Agilent). The absorbance wavelengths of excitation and emission were set at 320 and 405 nm.

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