

A Unique Tryptophan C-Prenyltransferase from the Kawaguchipectin Biosynthetic Pathway

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Abstract: Cyanobactins are a rapidly growing family of linear and cyclic peptides produced by cyanobacteria. Kawaguchipectins A and B, two macrocyclic undecapeptides reported earlier from *Microcystis aeruginosa* NIES-88, are shown to be products of the cyanobactin biosynthetic pathway. The 9 kb kawaguchipectin (*kgp*) gene cluster was identified in a 5.26 Mb draft genome of *Microcystis aeruginosa* NIES-88. We verified that this gene cluster is responsible for the production of the kawaguchipectins through heterologous expression of the *kgp* gene cluster in *Escherichia coli*. The *KgpF* prenyltransferase was overexpressed and was shown to prenylate C-3 of Trp residues in both linear and cyclic peptides *in vitro*. Our findings serve to further enhance the structural diversity of cyanobactins to include tryptophan-prenylated cyclic peptides.

Cyanobactins are a family of ribosomally-synthesized and posttranslationally modified peptides (RiPPs) produced by cyanobacteria.^[1–6] Cyanobactin posttranslational modifications include N-to-C macrocyclization; epimerization; heterocyclization to form thiazolines and oxazolines; oxidation of heterocycles to thiazoles and oxazoles; N-methylation of His; O-prenylation on Ser, Thr, and Tyr; and N-prenylation.^[2,7,8] Biological activities that have been reported for cyanobactins include anticancer, antimalarial, antibacterial, and protease inhibitory activity.^[2]

The posttranslational prenylation of cyanobactins is catalyzed by a prenyltransferase enzyme encoded within the cyanobactin biosynthetic gene cluster, which uses 3-methylbut-2-en-1-yl group derived from dimethylallyl pyrophosphate (DMAPP).^[9–12] Although a putative prenyltransferase gene is present in all known cyanobactin gene clusters, only a few of the cyanobactins are known to be prenylated, including prenylagaramides, aestuaramides, trunkamides, and anacyclamides.^[4,6,10,12] The known cyanobactin prenyltransferases are O-prenyltransferases that catalyze the O-prenylation of Tyr, Thr, and Ser in forward or reverse orientation. C-prenylated cyanobactins have been shown to be synthesized originally as O-prenylated peptides that later undergo a Claisen rearrangement to yield C-prenylated peptides.^[12]

Kawaguchipectins are macrocyclic undecapeptides produced by the cyanobacterial strain *Microcystis aeruginosa* NIES-88.^[13,14] Two variants of kawaguchipectin have been reported (Scheme 1). Kawaguchipectin A contains two C-3-prenylated tryptophan residues and a D-Leu residue.^[15] Kawaguchipectin B consists of solely unmodified amino acids and is reported to show antimicrobial activity against *Staphylococcus aureus*.^[14] Herein, we report a genome sequence for *Microcystis aeruginosa* NIES-88, identify the kawaguchipectin biosynthetic gene cluster, and confirm enzymatic prenylation activity.

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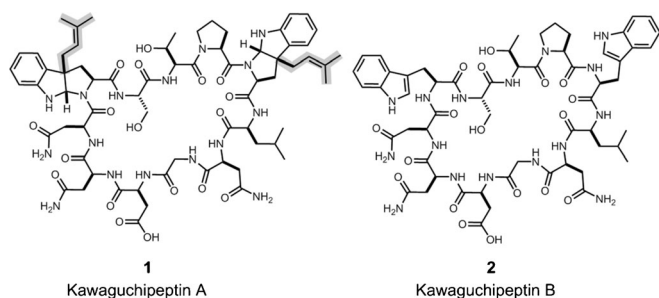
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Scheme 1. The cyclic undecapeptides kawaguchipeptin A (1) and kawaguchipeptin B (2) reported from *Microcystis aeruginosa* NIES-88. Kawaguchipeptin A contains two C-3-prenylated tryptophan residues (highlighted). Prenylation of C-3 is coupled to the formation of a new ring formation as a result of bond formation between C-2 and NH in the main chain.

We obtained a 5.26 Mb genome sequence for *Microcystis aeruginosa* NIES-88 by using a combination of 3 kb 454 sequencing and short-insert illumina Miseq paired-end data, which were subsequently assembled into 29 scaffolds. A total of 4,996 genes, including 4 ribosomal RNA operons and 41 tRNAs, were annotated from this genome. The kawaguchipeptin precursor gene was identified through tBLASTn using the predicted kawaguchipeptin peptide backbone (WLNGDNNWSTP). The KgpE precursor peptide was found to encode three exact copies of the WLNGDNNWSTP core (Figure 1). The KgpE precursor peptide was encoded in

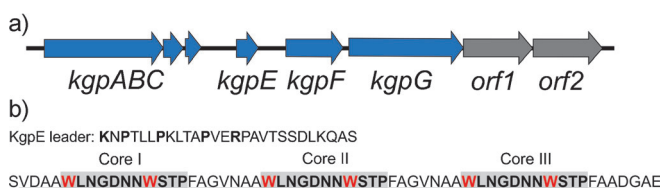


Figure 1. The kawaguchipeptin biosynthetic pathway in *Microcystis aeruginosa* NIES-88.^[23] A) The 9 kb *kgp* biosynthetic gene cluster consists of six biosynthetic genes (shown in blue), organized in a single operon, that show homology to genes present in other known cyanobactin gene clusters. The grey arrows indicate genes that encode proteins with unknown functions. B) The *kgpE* precursor gene encodes the 87 amino acid precursor peptide, which contains three identical copies of the undecapeptide core (highlighted in grey).

a 9 kb gene cluster (*kgp*) together with the KgpA and KgpG cyanobactin proteases, as well as the putative KgpF prenyltransferase (Figure 1). The KgpF prenyltransferase shows just 20–45% identity to known and putative cyanobactin prenyltransferases. A homologue of cyclodehydrase (PatD) was not detected in the gene cluster, which is consistent with the absence of heterocyclized amino acids in the cyclic peptides.

To demonstrate that the *kgpA-G* genes encode kawaguchipeptin production in vivo, we cloned the entire *kgp* operon into a broad host range yeast/bacteria shuttle vector pMQ123i^[15] and placed *kgpA* downstream of a pTac promoter to generate the expression plasmid pDK-kgp1 (Figure S1 in the Supporting Information). This construct allowed the regulated expression of the *kgpA-G* genes in *E. coli*

TOP10. Liquid chromatography with high-resolution mass spectrometry (LC–HRMS) guided metabolite profiling of *E. coli* cells transformed with pDK-kgp1 revealed the presence of 2, for which the LC retention time and HRMS profile matched authentic 2 isolated from *M. aeruginosa* NIES-88 (Figure 2, traces 2–3, and Figure S2). *E. coli* cells carrying pDK-kgp1 alone, however, did not produce 1 (Figure 2,

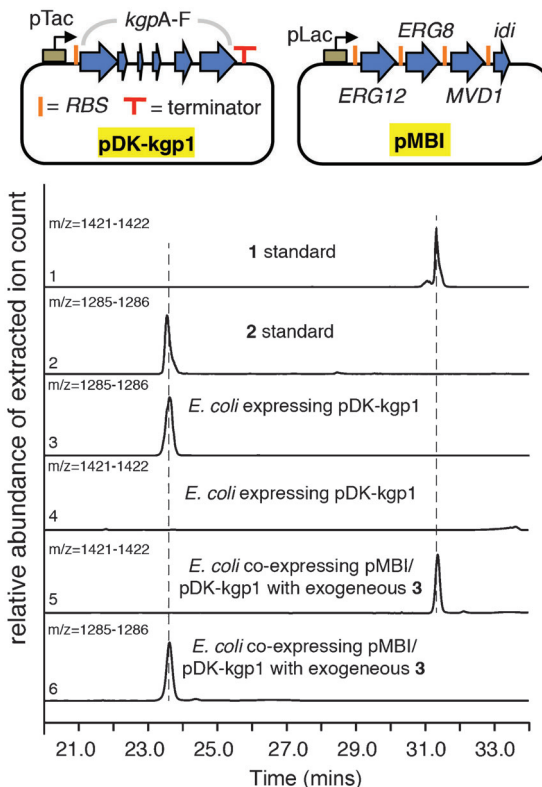


Figure 2. Heterologous expression of the *kgp* operon in *E. coli* demonstrates that the *kgpA-G* genes confer the production of 1 and 2 in vivo. Extracted ion chromatographs of LC–HRMS analysis of: Authentic 1 and 2 from *M. aeruginosa* NIES-88 (1, 2); 2 in *E. coli* transformed with pDK-kgp1 (3); 1 in *E. coli* transformed with pDK-kgp1 (4); 1 in *E. coli* co-transformed with pDK-kgp1 and pMBI and supplied with mevalonolactone 3 (5); and 2 in *E. coli* co-transformed with pDK-kgp1 and pMBI and supplied with mevalonolactone 3 (6).

trace 5), which is potentially derived from 2 by bisprenylation at C-3 of Trp by prenyltransferase KgpF. We hypothesized that this observation may have been due to the lack of sufficient endogenous prenyl donor dimethylallyl pyrophosphate (DMAPP) in *E. coli*. To overcome this problem, we co-transformed *E. coli* with pDK-kgp1 and the plasmid pMBI,^[16] which harbors four yeast mevalonate-dependent isoprenoid pathway biosynthetic genes that can convert mevalonate to isopentenyl pyrophosphate (IPP), a precursor to DMAPP. Co-expression of the pDK-kgp1 and pMBI genes in *E. coli* TOP10 supplied with exogenous mevalonolactone 3 (1.0 mM) led to the production of both 1 and 2 that matched their authentic standards from *M. aeruginosa* NIES-88 (Figure 2, traces 1–2, 6–7, and Figure S3), as assessed by LC–HRMS. These experiments establish that the *kgpA-G* genes confer

1 and **2** biogenesis in vivo. In addition, the coexistence of **1** and **2** in *E. coli* cells transformed with both pDK-kgp1 and pMBI and the lack of **1** in *E. coli* cells transformed with pDK-kgp1 alone strongly suggest that **2** is the direct biosynthetic precursor to **1** and bis-prenylation by KgpF is likely the last enzymatic step in the biosynthetic maturation of kawaguchi-peptins.

We overexpressed and purified the recombinant KgpF from *E. coli* and assessed the enzymes ability to process a range of cyclic and linear peptides and to use isopentenyl pyrophosphate (IPP), DMAPP, and geranyl pyrophosphate (GPP; Table 1 and Figures S4–S24).

Table 1: Substrates and cofactors used for the in vitro reactions.

| Substrate | Cofactor | Mono-prenylation | Di-prenylation |
|---------------------------------------|----------|------------------|----------------|
| Cyclic [WLNGDNNWSTP] (2) | DMAPP | + | + |
| Cyclic [WLNGDNNWSTP] (2) | IPP | + | – |
| Cyclic [WLNGDNNWSTP] (2) | GPP | – | – |
| Cyclic [TSQIWGSPVP] (4) | DMAPP | + | NA |
| Cyclic [SAQWQNFVGP] (5) | DMAPP | + | NA |
| Cyclic [HAFIGYDQDPTGKYP] (6) | DMAPP | – | – |
| Cyclic [RERFVYP] (7) | DMAPP | – | – |
| Cyclic [LIGIMHP] (8) | DMAPP | – | – |
| WLNGDNNWSTP (9) | DMAPP | + | – |
| WLNGDNNWSTPAYDG (10) | DMAPP | + | – |
| EDWYFDHPAYDG (11) | DMAPP | – | – |
| VPWFPAYDG (12) | DMAPP | – | – |
| Boc-Trp (13) | DMAPP | – | – |
| Boc-Trp (13) | IPP | – | – |
| Boc-Tyr (14) | DMAPP | – | – |
| Boc-Tyr (14) | IPP | – | – |

[+] Product detected, [–] Product not detected.

Our results show that the enzyme processes a second Trp residue within macrocyclic peptide substrate **2**, whereas in the linear peptide **9**, despite 40 h of incubation, only a single modification was observed with DMAPP as the cofactor (Table 1; Figures S5–S7, S18–S20). The catalytic activity of the enzyme decreased when IPP was used as a cofactor instead of DMAPP, as seen in the processing of one Trp residue out of two in kawaguchi-peptin B (**2**) when IPP instead of DMAPP was used (Table 1; Figures S8). We did not detect any processing of other residues in the linear or macrocyclic peptide substrates. The enzyme did not process Boc-Trp (**13**) in presence of DMAPP or IPP. Interestingly, the enzyme could not use GPP as a cofactor. According to these results, the selectivity of prenylation in linear substrates can be explained by the necessity of the Trp residue to be sandwiched between two residues. Terminal Trp residues would presumably be too strongly solvated to bind efficiently the enzymatic site and undergo transformation (Figures S20, S22).

O- or C-prenylation of Tyr, Ser, and Thr in forward or reverse orientation have been observed for cyclic cyanobactins.^[2,5,12] C-prenylated peptides like the C-prenylated Tyr in aestuaramides have been reported.^[12] However, the latter is the result of reverse O-prenylation on the oxygen atom of Tyr followed by Claisen rearrangement.^[12] The biochemical characterization of C-3 Trp prenylation^[13] and demonstration

of the presence of a homologue of the prenyltransferase gene *kgpF* in the gene cluster confirm that this is a direct posttranslational modification, which is rare in cyanobactins. To our knowledge, C-3 prenylation of Trp through posttranslational modification of a peptide has been demonstrated only once before for the ComX peptide, a pheromone produced by *Bacillus subtilis* and related bacilli.^[17,18]

Trp prenylation is common in some plants and bacteria, but mostly in fungi, and the respective prenyltransferases catalyze the addition of a dimethylallyl group to Trp during the synthesis of secondary metabolites.^[19,20,21] The synthesis and biosynthesis of these compounds, particularly indole alkaloids that contain prenylated Trp at their core, have been the subject of considerable interest.^[22] Our findings therefore expand the chemical diversity of cyanobactins and confirm the existence of a rare Trp prenyltransferase. The prenyltransferases of the cyanobactin family are now known to catalyze the O-, C-, and N-prenylation of amino acids in cyclic peptides.

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- [23] The draft genome of *Microcystis aeruginosa* NIES-88 has been deposited in GenBank under the accession number JXYX00000000.

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Communications

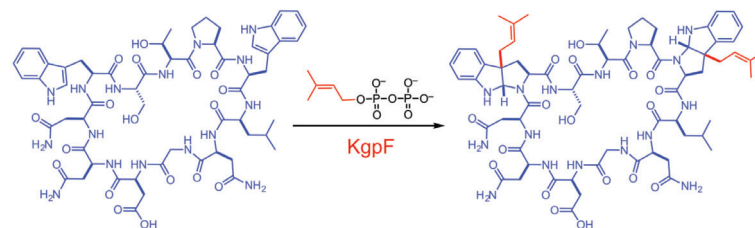


Biosynthesis

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