INVESTIGATIONS INTO THE MICROCYSTIN GENE CLUSTER

FROM HAPALOSIPHON HIBERNICUS BZ-3-1 AND

PLANKTOTHRIX AGARDHII CYA 126/8

A THESIS SUBMITTED TO THE GRADUATE DIVISION OF THE UNIVERSITY OF HAWAII IN PARTIAL FULFILLMENT OF THE REQUIREMENTS FOR THE DEGREE

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By

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Abstract

Hapalosiphon hibernicus BZ-3-1 is a terrestrial cyanobacterium isolated from a soil sample from Maui, Hawaii. It was shown to produce microcystin-LA during screening for protein phosphatase inhibitors. Using a PCR approach involving degenerate primer mixes, which were derived from comparison of the three known gene clusters previously sequenced from Microcystis aeruginosa, Planktothrix agardhii and Anabaena circinalis. Using the primer mixes the genes mcyA, -B, -C, -G, -H, and -J were amplified in almost their entirety. The genes mcyD and -E did not give amplicons under similar conditions and therefore needed to be amplified in a slightly different manner. Fragments from the 5' and 3' end of each gene (1 kb) were amplified with the degenerate primer mixes and then the known fragments were connected together using the gathered sequence. After sequencing the genes, the organization of the gene cluster was established by connecting PCR. The resulting gene cluster yielded two operons, mcyABC and mcyGDJEFIH, which are transcribed in opposite directions from a putative bi-directional promoter. The H. hibernicus cluster shows the greatest similarity to the Anabaena cluster (mcyA, -B, -D, -G, -J) and the nodularin gene cluster from Nodularia (mcy-E, F, -I).

It was noticed that the Planktothrix cluster was the only one to contain mcyT and be deficient in mcyI and -F. Genetic studies were then undertaken to disrupt mcyT using homologous recombination. The DNA was introduced into the cyanobacterial cells via electroporation and resistance to chloramphenicol was used to select for the mutant. The mutant was analyzed for peptide production and it was discovered that microcystin biosynthesis had been reduced to 5-10% of that found in the wild type. During
characterization of the mutant peptide production a new anabeanopeptin derivative was found. Structure determination was accomplished using NMR and HR-MS techniques. Stereochemistry of the amino acid residues was determined by the advanced Marfey analysis using LC-MS.
Table of Contents

Acknowledgments

Abstract

List of Tables

List of Figures

List of Abbreviations

1. Sequencing of the mcy gene cluster from Hapalosiphon hibernicus

1.1 INTRODUCTION

1.1.1 Natural products

1.1.2 Natural products from microorganisms

1.1.3 Microcystin

1.1.4 Polyketide synthetases and non-ribosomal peptide synthetases

1.1.5 Microcystin toxicity

1.1.6 Hapalosiphon hibernicus

1.1.7 Research objectives

1.2 EXPERIMENTAL SECTION

1.2.1 General

1.2.1.1 General laboratory procedures

1.2.1.2 Preparation of electro-competent E. coli cells

1.2.1.3 Transformation of electro-competent cells

1.2.1.4 Preparation of chemical competent cells

1.2.1.5 Transformation of chemically competent cells
1.2.1.6 Colony PCR using *Taq* polymerase from Qiagen

1.2.1.7 Plasmid preparation by hand

1.2.1.8 Cyanobacterial DNA isolation

1.2.2 Cyanobacterial culture

1.2.2.1 Z+G media

1.2.2.2 BG-11 media

1.2.2.3 Cultivation of alga

1.2.3 *E. coli* culturing

1.2.3.1 Antibiotics

1.2.3.2 Agar plate preparation

1.2.4 Recombinant DNA procedures

1.2.4.1 Ligation using T4 DNA ligase

1.2.4.2 Ligation into pDrive vectors

1.2.4.3 PCR using *Taq* DNA polymerase from Qiagen

1.2.4.4 PCR using Advantage 2 PCR kit

1.2.4.5 Gel electrophoresis for DNA analysis

1.2.5 DNA sequencing

1.2.6 Sequencing of the *mcy* cluster

1.2.6.1 Gene amplification of *mcyA*, -B, -C, and -G

1.2.6.2 Amplification of *mcyD* and -E

1.2.6.3 Amplification of *mcyH* and -I

1.2.6.4 Sequencing of *mcyA*, -B, -C, -D, -E, and -G

1.2.6.5 Orientation of the *mcy* gene cluster
1.2.6.6 Completing *mcyH* and -C ........................................................................ 32

1.3 RESULTS AND DISCUSSION ........................................................................ 35

1.3.1 Amplification/Identification of the *mcy* gene cluster ............................ 35

1.3.2 Organization of the microcystin gene cluster ......................................... 36

1.3.3 Characterization of the microcystin genes ............................................. 38

1.3.4 NRPS adenylation domain specificities .................................................. 40

1.3.5 Phylogenetic analysis of adenylation domains ........................................ 41

1.4 CONCLUSION ................................................................................................. 43

2 Genetic knockout of *mcyT* in *P. agardhii* CYA 126/8 ................................. 44

2.1 INTRODUCTION ............................................................................................ 44

2.1.1 McyI ........................................................................................................... 44.

2.1.2 McyT ......................................................................................................... 46

2.2 EXPERIMENTAL SECTION .......................................................................... 48

2.2.1 Insertional mutagenesis of *mcyT* ............................................................ 48

2.2.1.1 Construction of the homologous recombination plasmid.................. 48

2.2.1.2 Creating the chloramphenicol disrupted *mcyT* gene ...................... 48

2.2.1.3 Preparation of the transformation construct ...................................... 49

2.2.1.4 Transformation of *P. agardhii* CYA 126/8 ....................................... 50

2.2.1.5 Confirmation of mutant identity ......................................................... 51

2.2.2 Peptide analysis ........................................................................................ 51

2.2.2.1 Peptide extraction of *P. agardhii* CYA126/8 Δ*mcyT* .................... 51

2.2.2.2 HPLC analysis of CYA 126/8 Δ*mcyT* peptide content ............... 52

2.2.2.3 LC-MS analysis of CYA 126/8 Δ*mcyT* peptide content .............. 53
List of Tables

Table

1. Known microcystin variants .................................................................................................................. 6

2. Degenerate primer sequences used in the amplification of \( mcyA, -B, -C, -D, -E, -G, -H, \) and \(-J.\) ................................................................................................................................. 29

3. Primers used during the orientation PCR reactions. ........................................................................ 31

4. Primers used during the completion of \( mcyC \) and \(-H.\) ........................................................................ 34

5. Percent homology of genes from \( H. hibernicus \) to known \( mcy \) genes........................................ 38

6. Amino acid specificity conferring sequences of adenylation domains as
   Stachelhaus et al. .................................................................................................................................. 40

7. NMR data for Anabaenopeptin I (14). ............................................................................................. 63
List of Figures

1. Starter units for fatty acid synthetase and polyketide synthetase systems. 7
2. Conversion of apo protein into holo protein by nucleophilic attack of the conserved serine residue on coenzyme A. 9
3. Schematic diagram of the microcystin clusters from M. aeruginosa, P. agardhii, A. circinalis and the nodularin gene cluster from N. spumigena. 11
4. Biosynthesis of microcystin-LA. 12
5. Schematic diagram of the mcy cluster sequenced from H. hibernicus as well as the known clusters from M. aeruginosa, P. agardhii, A. circinalis and the nodularin gene cluster from N. spumigena. 37
6. (a) Evolutionary tree diagram of the adenylation domains found in the microcystin clusters and the nodularin cluster using GrsA-A1 as an out-group.
   (b) Evolutionary tree diagram of the condensation domains found in the microcystin clusters as well as the nodularin cluster. 42
7. Multalign of 3-D-phosphoglycerate dehydrogenases from E. coli W3110, E. coli K-12, Yersinia pestis, Mycobacterium tuberculosis, Synechocystis sp. PCC 6803, Homo sapiens and McyI from H. hibernicus BZ-3-1. 45
8. 2.0% agarose gel verifying the correct mutant had been created. PCR performed with primers mcyT5’seq and mcyT3’seq. 60
9. Synthesis of enantiomerically enriched N-Me homotyrosine. 65
# List of Abbreviations

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Definition</th>
</tr>
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<tbody>
<tr>
<td>4' -PP</td>
<td>4' -phosphopantetheinyl arm</td>
</tr>
<tr>
<td>A</td>
<td>adenylation</td>
</tr>
<tr>
<td>Ac</td>
<td>acetyl</td>
</tr>
<tr>
<td>AcCN</td>
<td>acetonitrile</td>
</tr>
<tr>
<td>ACP</td>
<td>acetyl carrier protein</td>
</tr>
<tr>
<td>Adda</td>
<td>(2S,3S,8S,9S)-3-amino-9-methoxy-2,6,8-trimethyl-10-phenyl-(4E, 6E)-decadienoic acid</td>
</tr>
<tr>
<td>ADP</td>
<td>adenosine diphosphate</td>
</tr>
<tr>
<td>Amp</td>
<td>ampicillin</td>
</tr>
<tr>
<td>AT</td>
<td>acetyl transferase</td>
</tr>
<tr>
<td>ATP</td>
<td>adenosine triphosphate</td>
</tr>
<tr>
<td>C</td>
<td>condensation</td>
</tr>
<tr>
<td>Cm</td>
<td>chloramphenicol</td>
</tr>
<tr>
<td>CMT</td>
<td>carbon methyl transferase</td>
</tr>
<tr>
<td>CoA</td>
<td>Coenzyme A</td>
</tr>
<tr>
<td>dH₂O</td>
<td>deionized water</td>
</tr>
<tr>
<td>DNA</td>
<td>deoxyribonucleic acid</td>
</tr>
<tr>
<td>Ep</td>
<td>epimerization</td>
</tr>
<tr>
<td>ESI</td>
<td>electrospray ionization</td>
</tr>
<tr>
<td>EtBr</td>
<td>ethidium bromide</td>
</tr>
<tr>
<td>EtOAc</td>
<td>ethyl acetate</td>
</tr>
<tr>
<td>EtOH</td>
<td>ethanol</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Description</td>
</tr>
<tr>
<td>--------------</td>
<td>-------------</td>
</tr>
<tr>
<td>FABMS</td>
<td>fast atom bombardment mass spectrometry</td>
</tr>
<tr>
<td>FDAA</td>
<td>1-fluoro-2-4-dinitrophenyl-5-L-alanine amide</td>
</tr>
<tr>
<td>HMBC</td>
<td>heteronuclear multiple bond correlation</td>
</tr>
<tr>
<td>HMQC</td>
<td>heteronuclear multiple quantum coherence</td>
</tr>
<tr>
<td>Hty</td>
<td>homotyrosine</td>
</tr>
<tr>
<td>IPTG</td>
<td>isopropyl-beta-D-thiogalactopyranoside</td>
</tr>
<tr>
<td>KR</td>
<td>ketoreductase</td>
</tr>
<tr>
<td>KS</td>
<td>ketosynthase</td>
</tr>
<tr>
<td>LB_Amp</td>
<td>LB media containing 100 μg/mL ampicillin</td>
</tr>
<tr>
<td>LC-MS</td>
<td>liquid chromatography-mass spectrometry</td>
</tr>
<tr>
<td>Masp</td>
<td>methyl aspartate</td>
</tr>
<tr>
<td>MC</td>
<td>microcystin</td>
</tr>
<tr>
<td>Mdha</td>
<td>N-methyl dehydroalanine</td>
</tr>
<tr>
<td>MeOH</td>
<td>methanol</td>
</tr>
<tr>
<td>MOPS</td>
<td>3-(N-Morpholino)propanesulfonic acid</td>
</tr>
<tr>
<td>MSD</td>
<td>mass selective detector</td>
</tr>
<tr>
<td>Na$_2$EDTA</td>
<td>ethylenediaminetetraacetate, disodium salt</td>
</tr>
<tr>
<td>NAD</td>
<td>nicotinamide adenine dinucleotide</td>
</tr>
<tr>
<td>NMR</td>
<td>nuclear magnetic resonance</td>
</tr>
<tr>
<td>NMT</td>
<td>N-methyl transferase</td>
</tr>
<tr>
<td>NRPS</td>
<td>nonribosomal peptide synthetase</td>
</tr>
<tr>
<td>ORF</td>
<td>open reading frame</td>
</tr>
<tr>
<td>PCR</td>
<td>polymerase chain reaction</td>
</tr>
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<td>Abbreviation</td>
<td>Definition</td>
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<tr>
<td>--------------</td>
<td>------------</td>
</tr>
<tr>
<td>PKS</td>
<td>polyketide synthase</td>
</tr>
<tr>
<td>PP 1/PP 2a</td>
<td>protein phosphatase 1/protein phosphatase 2a</td>
</tr>
<tr>
<td>RT</td>
<td>room temperature</td>
</tr>
<tr>
<td>SAM</td>
<td>S-adenosyl methionine</td>
</tr>
<tr>
<td>T</td>
<td>thiolation</td>
</tr>
<tr>
<td>Te</td>
<td>thioesterase</td>
</tr>
<tr>
<td>Tm</td>
<td>melting temperature</td>
</tr>
<tr>
<td>TMS-Cl</td>
<td>chlorotrimethylsilane</td>
</tr>
<tr>
<td>TOCSY</td>
<td>total correlated spectroscopy</td>
</tr>
<tr>
<td>TOF</td>
<td>time of flight</td>
</tr>
<tr>
<td>XGal</td>
<td>5-bromo-4-chloro-3-indoly-β-D-galactoside</td>
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</table>
1. Sequencing of the mcy gene cluster from *Hapalosiphon hibernicus*

1.1 INTRODUCTION

1.1.1 Natural products

Throughout the history of human civilization, humans and natural products have had a close, intertwining relationship. From the ancient Incas chewing on coca leaves for the energizing effect of cocaine to the use of willow bark as a source of pain and fever relief, to the discovery of penicillin by Sir Alexander Fleming, nature has provided a tremendous number of bioactive metabolites that have influenced civilizations.

Currently 60% of anticancer and 70% of anti-infective antibiotics are natural products or semisynthetic derivatives. For example the anticancer chemotherapeutic agent paclitaxel (1) is obtained in commercially relevant amounts by synthetic derivation of baccatin III (2), which is isolated from the needles of the English yew tree (*Taxus baccata*). Cyclosporin (3), the immunosuppressant used to suppress graft rejections after organ transplants, is isolated from cultures of the fungus *Tolypocladium inflatum* Gams.

Microorganisms have been a tremendous source of bioactive molecules; the actinomycetes are especially valuable in the search for and production of clinically useful drugs, while more recently other groups of microorganisms such as cyanobacteria have been investigated. Marine invertebrates, particularly from the phylum *Porifera* (sponges) and *Coelenterata* (bryozoans, corals, etc.) have also been heavily explored for novel natural products. The natural products discovered from these macroscopic organisms have been hypothesized to originate, at least in part, from symbiotic microorganisms. With the recent advances in microbial culture and genetic techniques, additional evidence for the endosymbiosis production theory has been obtained. An example is swinholide A
(4), which was originally isolated from the marine sponge *Theonella swinhoei* by Kobayashi and co-workers.\(^5\) Bewley *et al.* found swinholide A to be localized within the symbiotic unicellular bacteria within this sponge, which were separated using centrifugation.\(^6\) More recently swinholide A and a derivative were isolated from a collection of marine cyanobacteria.\(^7\)

The isolation of bioactive compounds from microorganisms offers unique advantages over isolation from larger organisms. First, microorganisms typically contain higher percentages of metabolite per kilogram dry mass, and microorganisms can be easier to culture and maintain in the laboratory. However there are numerous problems associated with culturing of microorganisms, as it has been estimated from genetic data
that fewer than 1% of known microorganisms have been cultured and examined in the lab. To make things more complicated some bacterial cultures have been known to only produce certain metabolites in the presence of a specific stimulus.8

The possibility for the discovery of novel structural skeletons from microbes is immense considering the large unexplored microbial genetic diversity and the low percentage of microbes that have been cultured. To this date over 7,000 natural products have been isolated from the actinomycetes alone with most produced by the genus *Streptomyces* (80%) and a few other well-known soil dwelling genera.4 The diversity of known structural backbones and biological activities of compounds isolated from microorganisms range from vancomycin (5), a cyclic peptide antibiotic isolated from *Amycolatopsis orientalis* to barbamide (6), a linear polyketide-peptide hybrid with potent molluscicidal activity9 to the linear peptide dolastatin 10 (7), which disrupts microfilament depolymerization.10 Both barbamide and dolastatin 10 were isolated from the marine cyanobacterium *Lyngbya majuscula.*
1.1.2 Natural products from microorganisms

Large multi-modular enzymes called polyketide synthases (PKSs) and non-ribosomal peptide synthetases (NRPSs) catalyze the formation of the two largest families of natural products including the compounds mentioned above. However in silico analysis of *Streptomyces coelicolor* and *S. avermitilis* showed that only 12% (14 of 118) of putative PKS and NRPS clusters had been correlated with the production of a specific natural product. This data coupled with the estimate that 1% of microorganisms have been cultured suggests that a wide variety of unknown structures could/should exist.

Cyanobacteria have recently become a hot area of study because of their ability to produce prolific amounts of secondary metabolites, typically cyclic peptides and depsipeptides, which are produced by PKS and NRPS systems. As is the case with *S. coelicolor* and *S. avermitilis*, the sequenced genome of *Nostoc punctiforme* contains 22 putative PKS and NRPS clusters, however no natural product has been definitively associated with any of these clusters. To date, ten different cyanobacterial gene clusters
have been sequenced including the clusters responsible for nostocyclopeptide A1 (8), microcystin-RR (9), jamaicamide (10), and barbamid (6) biosynthesis.

1.1.3 Microcystin

The microcystins (MCs) are a class of compounds with the general structure \( \text{cyclo-(D-Ala-L-X-D-erythro-\beta-Masp-L-Z-Adda-D-isoGlu-Mdha)} \), where X and Z are variable amino acids, \( D-\text{erythro-\beta-Masp} \) is \((3S)-3\)-methyl aspartate, Mdha is N-methyl dehydroalanine and Adda is \((2S,3S,8S,9S)-3\)-amino-9-methoxy-2,6,8-trimethyl-10-phenyl-(4E, 6E)-decadienoic acid. Over 70 variants have been described (Table 1) with the majority of the substitutions taking place at the X and Z variable positions. The naming of the microcystins is based on the above base formula with the variable position amino acids listed afterwards in one letter code. For example, microcystin-LA (MC-LA) has the structure \( \text{cyclo-(D-Ala-L-Leu-D-erythro-\beta-Masp-L-Ala-Adda-D-isoGlu-Mdha)} \).
While some characterized variants can be viewed as artifacts of isolation, thereby reducing the actual number of naturally occurring variants, the structural diversity remains impressive. Interestingly, most strains that have been surveyed for microcystin production contain only a few of these multiple variants, albeit at different levels (i.e. Planktothrix agardhii CYA 126/8 produces (D-Asp\(^3\))-MC-RR (90%), (D-Asp\(^3\))-MC-LR (9%) and (D-Asp\(^3\))-MC-YR (1%)). The production of multiple peptide variants is not a unique occurrence found only in microcystin producing strains. Moore and co-workers have isolated 24 cryptophycin variants from the strain Nostoc sp. GSV 224.\(^\text{12}\)

Table 1: Known microcystin variants

<table>
<thead>
<tr>
<th>D-Ala(^1)</th>
<th>X(^2)</th>
<th>D-Masp(^3)</th>
<th>Z(^4)</th>
<th>Adda(^5)</th>
<th>D-Glu(^6)</th>
<th>Mdha(^7)</th>
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<tbody>
<tr>
<td>D-Ser</td>
<td>Leu</td>
<td>D-Asp</td>
<td>Ala</td>
<td>DMAAdda</td>
<td>D-Glu(OMe)</td>
<td>Dha</td>
</tr>
<tr>
<td>D-Leu</td>
<td>Ala</td>
<td></td>
<td>Aba</td>
<td>ADMAAdda</td>
<td>Ser</td>
<td>Dhh</td>
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<tr>
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<td>Tyr</td>
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<td>Leu</td>
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<td>D-Me-Ser</td>
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<td></td>
<td>Glu</td>
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<td>Me-Lan</td>
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<tr>
<td></td>
<td>Val</td>
<td></td>
<td>Glu</td>
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<td></td>
<td>Glu(OMe)</td>
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<td>(H(_2))Tyr</td>
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</tbody>
</table>

1.1.4 Polyketide synthetases and non-ribosomal peptide synthetases

Polyketides are synthesized by polyketide synthases, while peptides (linear and cyclic) and depsipeptides are synthesized by non-ribosomal peptide synthetases. There are three types of PKS systems, Type I, II and III, and they are classified by their mechanistic characteristics and resulting products. Type I PKS systems are the aforementioned large multifunctional enzymes found in the microcystin gene cluster. Type I polyketide synthases (from hereon out simply referred to as PKS) are related to fatty acid synthases (FAS) in that they both contain conserved domains for selection, coupling and modification of carbon extender units and both rely on the thio-template mechanism for product construction. Unlike FAS, which can only incorporate acetyl-CoA or malonyl-CoA building blocks, PKSs can utilize a wide variety of starting materials (Figure 1). After the Claisen reaction, the FAS system reduces the resulting β-keto thioester to the β-hydroxy thioester, followed by dehydration to form the α,β-unsaturated thioester and lastly

\[
\text{Figure 1:} \text{ Starter units for fatty acid synthase systems (red) and polyketide synthase systems (red and blue). Adapted from http://203.90.127.50/~pksdb/polyketide.htm}
\]
reduction of the resulting alkene to an alkane. Done repetitively, this results in a long saturated alkyl chain. In contrast PKS systems can do all or none of these reactions following the condensation. Moreover they may introduce methyl groups at the C1 and C2 position of the incorporated acetate unit. This flexibility, along with the diverse pool of extender units contributes to the large structural diversity of polyketide natural products. NRPSs operate by a similar means only using amino acids instead. NRPSs also modify the incorporated amino acids by N-methylation, epimerization, cyclization (of Thr, Ser and Cys residues) and oxidation or reduction of the cyclized amino acid.

NRPS and PKS contain conserved domains that perform different aspects of the biosynthesis. Adenylation or A domains (NRPS) and ketosynthase or KS domains (PKS) are analogous in function. They select and activate the correct building block for incorporation. Adenylation domains activate the amino acid as the AMP mixed anhydride with the formation of pyrophosphate while ketosynthase domains utilize Co-enzyme A bound substrates. Condensation or C domains (NRPS) and acyl transferase or AT domains (PKS) perform the coupling of neighboring units by amide bond formation or a thio-Claisen reaction respectively. Thiolation or T domains (NRPS) and acyl carrier proteins or ACPs (PKS) are responsible for tethering the growing chain to the enzyme through a thioester linkage.

There are two forms of each enzyme, the apo or inactive form and the holo or active form. An enzyme known as a phosphopantetheinyl transferase mediates the conversion of the apo to the holo form. This enzyme transfers a 4'-phosphopantethieryl arm (4'-PP) onto a conserved serine residue of the T or ACP domain. The holo form of
the enzyme is established by nucleophilic attack of the β-phosphate of Co-enzyme A by the serine hydroxyl group (Figure 2).

![Figure 2: Conversion of apo protein into holo protein by nucleophilic attack of the conserved serine residue on coenzyme A](image)

Type I thioesterase domains (NRPS/PKS) are responsible for cleavage of the final product from the enzyme either by hydrolysis to form the linear product or by cyclization to yield a lactone or lactam type product. Alternatively, a reductase domain can also cause the release of the peptide as the aldehyde, as in the case of nostocyclopeptide biosynthesis. Exclusive to PKS systems are the C-methyl transferase (CMT), the ketoreductase (KR), the dehydratase (DH), and the enoyl reductase (ER) domains. These domains are responsible for methylating the C2 of an acetate extender unit (the methyl is derived from S-adenosyl methionine (SAM)), reducing the ketone to the β-hydroxy-
thioester, dehydration to generate the α,β-unsaturated thioester and finally reduction to
the saturated thioester respectively. NRPSs have their own associated domains including
the N-methyl transferase (NMT) and the epimerization (Ep) domains. The NMT
catalyzes the addition of a methyl group to the –NH₂ group prior to amide bond
formation while the epimerization domain converts the naturally occurring L-amino acid
to the unnatural D-amino acid.

The microcystins are synthesized by a PKS-NRPS hybrid system. The gene
clusters from three limnic cyanobacterial strains, Microcystis aeruginosa PCC 7806,²⁶
Planktothrix agardhii CYA 126/8 ᵃ and Anabaena circinalis strain 90²⁷ have been
sequenced in addition to the gene cluster for the related natural product nodularin. The
gene clusters contain either nine or ten genes involved in microcystin biosynthesis. All
three contain mcyA, -B, -C, -D, -E and -G which are NRPS/PKSs that are responsible for
the assembly of the structural backbone of microcystin, mcyJ, which encodes for an O-
methyl transferase and mcyH which codes for a putative ABC transporter. McyJ is
responsible for the methylation of the hydroxyl group to form the methoxy group found
on the Adda side chain. The clusters obtained from M. aeruginosa and A. circinalis
contained two modifying genes named mcyF, an aspartate racemase and mcyI, a D-3-
phosphoglycerate dehydrogenase homolog. P. agardhii was discovered to be deficient in
these genes but did contain a putative type II thioesterase gene, mcyT (Figure 3). Figure 4
shows the domains present in each gene and what aspect of the biosynthesis each gene is
responsible for. The Te domain found at the 3’ end of mcyC is responsible for cyclization
by forming the amide bond between the carboxyl group of the variable amino acid Z and
the amino group found on the Adda chain.
1.1.5 Microcystin toxicity

The toxicity of the microcystins has been studied intensively because of the danger toxic blooms pose to humans and livestock. The microcystins have recently been blamed for the deaths of approximately 50 dialysis patients in a Brazilian hospital,\textsuperscript{19} and currently microcystins are being investigated as the cause of high levels of liver cancer in China.\textsuperscript{20,21} The World Health Organization (WHO) has set the limit for microcystin-LR at 1\(\mu\)g per liter of water as safe for human consumption.\textsuperscript{22} Microcystin-LR was found to be the most toxic variant with an LD\textsubscript{50} in mice of 43 \(\mu\)g per kg body weight while microcystin-RR was found to be less toxic with a LD\textsubscript{50} of 235 \(\mu\)g per kg body weight.

The mechanism of action of the MCs is a result of their high affinity for the eukaryotic PP1 and 2a. Most microcystin variants described to date have some level of inhibition of eukaryotic PP1 and PP2A. Protein phosphatases 1 and 2A regulate a wide...
Figure 4: Biosynthesis of microcystin-LA
variety of cellular functions by working in concert with protein kinases to regulate the activity of enzymes. Processes such as cell signaling, glycogen storage and cell division are regulated by PP1 and 2A. Goldberg and coworkers co-crystallized human PP1 and MC-LR and therefore were able to deduce the key interactions by X-ray crystallography.

One observation was that the solution and bound conformations of MC-LR did not differ significantly which results in a low activation barrier for binding. The Adda side chain fits snuggly in the hydrophobic groove of PP1 where it interacts with the hydrophobic residues. The carboxylate group and the amide carbonyl of the D-isoGlu residue hydrogen bonds to two water molecules found in the active site of the enzyme. The water molecules are in turn coordinated to two metal ions (Fe$^{3+}$, Mn$^{2+}$) found in the active site. The carboxylate of the D-erythro-Masp hydrogen bonds to two residues (Arg-96 and Tyr-134) while the Mdha unit was covalently bound to Cys-273. However this last interaction was found to be non-essential as mutation of the cysteine did not reduce the binding affinity of MC-LR.

1.1.6 *Hapalosiphon hibernicus*

*Hapalosiphon hibernicus* W & GS West BZ-3-1 was isolated from a soil sample collected on Maui, HI. It was shown to produce microcystin-LA during screens for protein phosphatase inhibition by Moore and co-workers. This became an important discovery because it was the first terrestrial strain as well as first subsection V cyanobacterium to be identified as a microcystin producer. Cyanobacteria in subsection I (*Microcystis*), III (*Planktothrix*), and IV (*Anabaena*) had previously been shown to produce microcystin. More recently another terrestrial, lichen-associated cyanobacterium (*Nostoc* sp. strain IO-102-I) was shown to produce a few microcystin variants.
1.1.7 Research objectives

These studies were undertaken to 1) study the gene cluster of a terrestrial strain in the hopes that new proteins involved in the biosynthesis of microcystin could be located. As well as examine whether the biosynthesis of microcystin was conserved between limnic and terrestrial strains. 2) to develop a PCR based method for sequencing microcystin gene clusters from any cyanobacterial source.
1.2 EXPERIMENTAL SECTION

1.2.1 General

1.2.1.1 General laboratory procedures

All cell handling and transfer was performed using sterile technique under a laminar flow hood (Labconco, Kansas City, MO). All centrifugations were performed in a Sorvall RC 5B Plus centrifuge (Thermo Electron Corporation, Asheville, NC). All DNA primers were obtained from Integrated DNA Technologies (Coralville, IA) with standard desalting purification and were resuspended in dH₂O to a stock concentration of 100 pmol/μl and used without further purification. Growth media used for *Escherichia coli* were prepared and sterilized according to standard protocols. All PCR reactions were performed in a GeneAmp PCR system 2700® (Applied Biosciences). All OD₆₀₀ measurements were taken in a SP-830 spectrophotometer (Barnstead International, Dubuque, IA) using disposable polypropylene cuvettes (Fisher Scientific, Pittsburgh, PA). *E. coli* cells in liquid media were grown in an Environ shaker (Lab-line) at 37°C with shaking at 220 rpm. Agar plates were grown in an IsoTemp incubator (Fisher Scientific, Pittsburgh, PA) at 37°C for 15-20 h.

1.2.1.2 Preparation of electro-competent *E. coli* cells

An individual colony was inoculated into 5 ml of LB broth and grown to saturation (OD₆₀₀ > 2). The following day 2.5 ml of the culture was inoculated into 500 ml YENB media. The cells were grown to an OD₆₀₀ between 0.5 and 0.9 at which time the cells were transferred to sterile 250 ml collection bottles and pelleted in a SLA-1500 rotor pre-cooled to 4°C (5000 rpm/15 min/4°C). The supernatant was decanted and the cells were resuspended in 25 ml of sterile reverse osmosis water (pre-cooled to 4°C) and
then diluted up to 200 ml. The cells were then pelleted (5000 rpm/15 min/4°C) and the washing and centrifugation repeated. The cells were then resuspended in 50 ml total volume of 10% glycerol (pre-cooled to 4°C, sterile filtered). The cells were centrifuged again in a SH-3000 rotor pre-cooled to 4°C (4500 rpm/15 min/4°C). The cells were resuspended in 1-1.5 ml 10% glycerol and allocated in sterile 1.5 ml micro-centrifuge tubes in 40 µl aliquots, which were then flash frozen in liquid nitrogen and stored at -80°C until use.

1.2.1.3 Transformation of electro-competent cells

Previously aliquoted electro-competent cells were defrosted on ice while ligation mixtures were dialyzed against reverse osmosis water using a 0.2 µ nylon membrane (Phenomenex, Torrance, CA) for 5 minutes. The ligation mixture was added to the thawed cells and the mixture was transferred to a pre-cooled (-20°C) 0.2 cm Gene Pulser® cuvette (Bio-Rad, Hercules, CA). The electroporation was performed using a Gene Pulser II® (Bio-Rad) (1.7 kV, 200 Ω, 25 µF) resulting in time constants of 4.65 - 4.91 msec. Immediately after electroporation the cells were resuspended in 250 µL room temperature SOC media and transferred to a sterile 1.5 ml micro-centrifuge tube. The cells were then grown in an orbital shaker for 1 hour. The cells were then spread on LB agar plates containing the appropriate antibiotics using a glass spreader.

1.2.1.4 Preparation of chemical competent cells

Transformation Buffer A (100 ml):

<table>
<thead>
<tr>
<th>Chemical</th>
<th>Amount</th>
</tr>
</thead>
<tbody>
<tr>
<td>RbCl</td>
<td>1.2 g</td>
</tr>
<tr>
<td>MnCl$_2$·4 H$_2$O</td>
<td>0.99 g</td>
</tr>
</tbody>
</table>
KOAc (1M, pH 7.5) 3 ml
CaCl₂·2H₂O 0.15 g
Glycerol 15 g

-Adjust pH to 5.8 with 0.2M acetic acid

-Sterilize with 0.2 μm filter

Transformation Buffer B (25 ml):

<table>
<thead>
<tr>
<th>Component</th>
<th>Amount</th>
</tr>
</thead>
<tbody>
<tr>
<td>MOPS (0.5M, pH 6.8)</td>
<td>0.5 ml</td>
</tr>
<tr>
<td>RbCl</td>
<td>0.03 g</td>
</tr>
<tr>
<td>CaCl₂·2H₂O</td>
<td>0.275 g</td>
</tr>
<tr>
<td>Glycerol</td>
<td>3.75 g</td>
</tr>
</tbody>
</table>

-Adjust pH 6.8 with NaOH

-Sterilize with 0.2 μm filter

An individual colony was inoculated into a 3 ml overnight culture in LB media. The following day 500 μl of the overnight culture was inoculated in 50 ml SOB broth containing the appropriate antibiotics. The cells were grown in an orbital shaker until the OD₆₀₀ was between 0.3 and 0.5. The cells were transferred to a 50 ml Falcon tube, placed on ice for 15 min and pelleted using a SH-3000 rotor pre-cooled to 4°C (4500 rpm/15 min/4°C). The supernatant was decanted, the pellet resuspended in 16 ml of Transformation Buffer A, and allowed to sit on ice for 15 min. The cells were pelleted in a SH-3000 rotor (4500 rpm/15 min/4°C), the supernatant drained and the pellet resuspended in 4 ml Transformation Buffer B. The cells were then aliquoted in sterile 1.5 ml micro-centrifuge tubes in 200 μl aliquots and flash frozen in liquid nitrogen and stored at -80°C until use.
1.2.1.5 Transformation of chemically competent cells

Chemically competent cells were thawed on ice, DNA was then added and the tube mixed by quickly vortexing. The cells were incubated on ice for 30 min followed by heat shock at 42°C for 40-45 sec and then placed back on ice for 1 min. 500 µl SOC media (pre-warmed to 37°C) was added to the cells and they were incubated in an orbital shaker for 1 hour. Using a glass spreader the cells were then plated out on LB agar plates containing the appropriate antibiotic.

1.2.1.6 Colony PCR using Taq polymerase kit from Qiagen (Valencia, CA)

Master mix

36.25 µl reverse osmosis water
5 µl 5X PCR buffer 3 µl 25mM MgCl₂ solution
1.5 µl dNTP mixture (10 µM each)
1.5 µl forward primer (10pmol/µl)
1.5 µl reverse primer (10pmol/µl)
0.25 µl Taq DNA polymerase

Individual colonies were suspended in 10 µl of reverse osmosis water. The master mix was mixed and aliquots of 9 µl were placed in 0.2 µl thin walled PCR tubes (Corning, Acton, MA) and 1 µl of the cell suspension in water was added to each PCR tube. The PCR reaction was performed using the following program. 95°C for 3 min followed by 32 cycles of 95°C for 30 sec, 66°C for 30 sec then 72°C for a variable amount of time (1 min per kb of DNA to be amplified) followed by cooling to 25°C. The PCR products were then observed by gel electrophoresis.

1.2.1.7 Plasmid preparation by hand

Plasmid Isolation Buffer (PIB) 1: 50 mM Tris•HCl, 10mM Na₂EDTA, pH 8.0
Plasmid Isolation Buffer (PIB) 2: 0.2 M NaOH, 1% SDS

Plasmid Isolation Buffer (PIB) 3: 4 M guanidine-HCl, 0.5 M KOAc, pH 4.2

The cell pellet was resuspended in 250 µl PIB 1 by vortexing. PIB 2 (250 µl) was added and mixed by inverting gently 4-5 times. After 5 min at RT, PIB 3 (350 µl) was then added and the tubes mixed by inversion 6 times followed by incubation on ice for 5 min. The white precipitate formed was pelleted by centrifugation in a F-20/Micro rotor (11,000 rpm/10 min/4°C). The supernatant was removed by pipette and transferred to a second 1.5 ml micro-centrifuge tube. The tubes were centrifuged down again as described above and the supernatant was carefully transferred to a new 1.5 ml micro-centrifuge tube. Isopropanol (565 µl) was added to the cleared supernatant and mixed by gentle inversion. The tube was allowed to stand at room temperature for 30 min and then the DNA was precipitated by centrifugation in a F-20/Micro rotor (11,000 rpm/30 min/4°C). The supernatant was removed and the pellet washed with 750 µl 70% (v/v) aqueous ethanol. The ethanol was aspirated and the pellet air-dried for 10-12 min. The dried pellet was resuspended in 100 µl 10mM Tris buffer (pH 7.5). 1 µl RNase A/T1 mixture (Fermentas, Hanover, MD) was added and the tube mixed followed by incubation at 37°C for 1 hour. The plasmid was then used without further purification.

1.2.1.8 Cyanobacterial DNA Isolation

TE Buffer: 10mM Tris•HCl, pH 8.0, 1mM EDTA, pH 8.0

TES Buffer: 10mM Tris•HCl, pH 8.0, 1mM EDTA, pH 8.0, 40% (w/v) sucrose
Isolation of DNA from cyanobacteria cultures was performed essentially described\textsuperscript{28} but with a few minor modifications. Cyanobacterial cells were pelleted in a SH-3000 rotor (5000 rpm/15 min/RT). The supernatant was removed and the cells were washed three times with 10 volumes (compared to the volume of cells) of TE buffer. The cells were suspended in 50 - 300 µl of TES buffer and kept on ice for 1 hour. Lysozyme (0.1 volumes) (Sigma-Aldrich, 50 mg/ml) was added, the solution mixed thoroughly and incubated at 37°C for 30 - 45 min. 20% SDS (0.1 volumes) and Proteinase K (0.1 volumes) (Fermentas, Hanover, MD, 1 mg/ml) were added to the cells and the mixture stored at 65°C overnight. The resulting brown mixture was diluted to 700 µl with TES buffer and extracted with an equal volume of phenol/chloroform/isoamyl alcohol (25:24:1) with vigorous shaking for 5 min. The resulting emulsion was separated by centrifugation (Eppendorf 5417C centrifuge, 14,000 rpm/10 min). The aqueous layer was carefully removed by pipette and transferred to a new 1.5 ml micro-centrifuge tube. This extraction was repeated until no precipitate was observed at the interface of the two resulting layers. The aqueous layer was washed with an equal volume of chloroform followed by centrifugation to separate the layers (Eppendorf 5417C centrifuge, 14,000 rpm/10 min). The aqueous layer was carefully removed and transferred to a 1.5 ml micro-centrifuge tube where isopropanol (0.7 volumes) was added. The tube was mixed by inversion and stored at room temperature for 30 - 60 min. The DNA was then pelleted by centrifugation in an F20/Micro rotor (11,000 rpm/60 min/4°C). The supernatant was carefully removed and the pellet washed with 1 ml 70% (v/v) aqueous ethanol. The ethanol was removed, the pellet air dried for approximately 10 - 15 min and then suspended in an appropriate volume (100 - 300 µl) of TE buffer. To the mixture was
added 1 μl RNase A/T1 (Fermentas, Hanover, MD) followed by incubation at 37°C for 1 hour. The DNA concentration was determined by using a NanoSpec spectrophotometer (NanoDrop, Wilmington, DE) and then diluted to 100 pmole/μl for use in PCR applications.

1.2.2 Cyanobacterial culture

1.2.2.1 Z+G Media

Solution 1: 0.590 g Ca(NO₃)₂•4 H₂O and 0.476 g NaN₃ in 100 ml dH₂O
Solution 2: 0.410 g K₂HPO₄•3 H₂O in 100 ml dH₂O
Solution 3: 0.250 g MgSO₄•7H₂O in 100 ml dH₂O
Solution 4: 1.680 g NaHCO₃ in 100 ml dH₂O
Solution 5: 1.145 g Na₂EDTA and 0.300 g FeSO₄•7 H₂O in 100 ml dH₂O.

The solution was boiled for 5 minutes in a microwave and the diluted up to 1 L with dH₂O. Solution was then autoclaved for 20 min at 15 lbs/sq. in.

Solution 6: 0.310 g H₃BO₃, 0.169 g MnSO₄, 0.009 g (NH₄)₆Mo₇O₂₄•4 H₂O, 0.029 g ZnSO₄•7 H₂O, 0.015 g Co(NO₃)₂•6 H₂O and 0.013 g CuSO₄•5 H₂O in 100 ml dH₂O

10 ml of solutions 1 – 5 and 80 μl of solution 6 was added to an autoclavable vessel. The mixture was diluted up to 1 L with distilled water and the media autoclaved for 20 min at 15 lbs/sq. in. The media was cooled to room temperature before use.

1.2.2.2 BG-11 media

Solution 1: 15.0 g NaNO₃ in 100 ml dH₂O
Solution 2: 2.0 g K₂HPO₄ in 50 ml dH₂O
Solution 3: 3.75 g MgSO₄•7H₂O in 50 ml dH₂O
Solution 4: 1.80 g CaCl₂•2H₂O in 50 ml dH₂O
Solution 5: 0.30 g Citric acid in 50 ml dH₂O
Solution 6: 0.30 g Ammonium ferric citrate in 50 ml dH₂O
Solution 7: 0.05 g Na₂EDTA in 50 ml dH₂O
Solution 8: 1.00 g Na₂CO₃ in 50ml dH₂O
Solution 9: 2.86 g H₃BO₃, 1.81 g MnCl₂•4H₂O, 0.22 g ZnSO₄•7H₂O, 0.39 g Na₂MoO₄•2H₂O, 0.08 g CuSO₄•5H₂O, 0.05 g Co(NO₃)₂•6H₂O in 1L dH₂O
In an autoclavable vessel 10 ml of solution 1 and 1 ml solutions 2-8 were mixed. The volume was then adjusted to 1 L with dH₂O. The media was autoclaved for 20 min at 15 lbs./sq. in. The media was cooled to room temperature and before use 1 ml solution 9 was added using sterile technique.

1.2.2.3 Cultivation of alga

Cultures of H. hibernicus BZ-3-1 were cultured under conditions described previously with slight modifications. The cyanophyte was cultured in autoclaved 20-L glass carboys containing an inorganic medium (Z+G or BG-11 media). Cultures were continuously illuminated at an incident intensity of 80-100 μmol photons m⁻² s⁻¹ (photosynthetically active radiation) from banks of cool-white fluorescent tubes and sparged with 1% carbon dioxide in air at a rate of 4.5 L/min. The temperature was maintained at 23 ± 1 °C and the culture was harvested after 30 d. The yield was 1.25 g/l dry cell mass.

1.2.3 E. coli culturing

1.2.3.1 Antibiotics

Ampicillin, sodium salt (Fisher Scientific), kanamycin (Fisher Scientific) and chloramphenicol (Fisher Scientific) were prepared according to standard protocols and used at the following concentrations for the plasmids/strains listed below.

<table>
<thead>
<tr>
<th>Plasmid</th>
<th>Antibiotic (conc.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>pDrive</td>
<td>Amp (100 μg/ml)</td>
</tr>
<tr>
<td>pUC19</td>
<td>Amp (100 μg/ml)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>E. coli strain</th>
<th>Antibiotic (conc.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>DH5α</td>
<td>None</td>
</tr>
<tr>
<td>BL21(DE3)</td>
<td>None</td>
</tr>
</tbody>
</table>
1.2.3.2 Agar plate preparation

LB media plus agar was prepared according to standard protocols. After autoclaving and cooling to less than 55°C antibiotics were added to the appropriate concentration. The plates were then poured and allowed to harden and were stored in a sealed bag at 4°C until use. For cases where X-Gal and IPTG were necessary for blue/white screening the plates were dried under a laminar flow hood for 45 min and then 40 μl of both X-Gal (2g/ml in DMF) and IPTG (1M in dH2O) were spread evenly around the plate and allowed to adsorb for 15 min. Then using a glass spreader the cells were applied to the agar plate.

1.2.4 Recombinant DNA procedures

1.2.4.1 Ligation using T4 DNA Ligase

T4 DNA ligase was purchased from Fermentas (Hanover, MD) and stored at –20°C until use. The ligation reaction was performed in a 0.2 ml thin walled PCR tube at 4°C for at least 12 hours. The vector DNA and insert DNA were combined in a ratio of approximately 1:3 as judged by comparison to a known concentration of DNA on an agarose gel. For ligations utilizing DNA cut with restriction enzymes that yield overhanging ends, 1 μl of 10X buffer (supplied by the company) which contained 10mM ATP and 1 μl of T4 DNA ligase (1 Weiss unit) were added to the tube to make the total volume 10 μl. In the case of ligation of blunt end DNA 1μl of 10X buffer, 1 μl PEG 4000 (final concentration of PEG: 5%) and 1 μl of T4 DNA ligase (1 Weiss unit) were added to a final volume of 10 μl.
1.2.4.2 Ligation into pDrive vectors

pDrive TA cloning kits were purchased from Qiagen (Valencia, CA) for use in cloning PCR products. The supplied components, 2X ligation master mix (5 µl) and pDrive cloning vector (1 µl) were aliquoted into 0.2 ml thin walled PCR tube on ice. The aliquots were then stored at \(-20^\circ C\) until use. The aliquots were removed from the freezer and thawed by holding the tube between two fingers. When defrosted the tube was placed on ice and the PCR product (4 µl) was added to the tube and mixed gently by pipetting 2-3 times. The ligation mixtures were then stored at \(4^\circ C\) for at least 12 h.

1.2.4.3 PCR using Taq DNA polymerase from Qiagen

For PCR reactions the following mixture was used unless otherwise stated.

Reagents were added in this order.

\[
\begin{align*}
36.25 \mu l & \text{ dH}_2 \text{O} \\
5 \mu l & 10X \text{ PCR buffer (Qiagen, Valencia, CA)} \\
3 \mu l & \text{MgCl}_2 \text{ solution (25mM)} \\
1.5 \mu l & \text{dNTPs (10 mM each)} \\
1.5 \mu l & \text{Primer 1 (10 pmol/µl)} \\
1.5 \mu l & \text{Primer 2 (10 pmol/µl)} \\
0.25 \mu l & \text{Taq DNA polymerase (1 unit)} \\
1 \mu l & \text{DNA (100 pmol/µl)} \\
\Sigma & 50 \mu l \text{ total volume}
\end{align*}
\]

1.2.4.4 PCR using Advantage 2 PCR kit (BD Biosciences, Mountainview, CA)

For PCR reactions the following mixture was utilized. Reagents were added in this order.

\[
\begin{align*}
40 \mu l & \text{dH}_2 \text{O} \\
5 \mu l & \text{Advantage 2 PCR 10X buffer} \\
1 \mu l & 50X \text{ dNTP solution}
\end{align*}
\]
1 µl Primer 1 (10 pmol/µl)
1 µl Primer 2 (10 pmol/µl)
1 µl DNA polymerase mix
1 µl DNA (100 pmol/µl)
Σ 50 µl total volume

1.2.4.5 Gel electrophoresis for DNA analysis

Agarose (Fisher Bioscience) was dissolved in 1X TAE buffer by use of a
microwave oven. Agarose gels are expressed as percent weight agarose per volume TAE
buffer (i.e. 0.8 g agarose dissolved in 100 ml TAE buffer results in a 0.8% agarose gel).
Prior to use, EtBr (Bio-Rad) was added to a final concentration of 0.4 µg/ml. The gels
were developed in a submersion electrophoresis chamber containing 1X TAE buffer. The
DNA was then visualized using a Transilluminator (Fisher Biotech).

1.2.5 DNA sequencing

Plasmids were sequenced at the Greenwood molecular biology facility using
standard Sanger sequencing techniques. Sequences were viewed and analyzed using
Chromas v1.45. Plasmids were sequenced using the vector specific primers pDriveSeq+
(5'-ACGACGTTGTAAAACGACGG) and pDriveSeq- (5'-
ACACAGAAACAGCTATGAC)

1.2.6 Sequencing of the mcy cluster

1.2.6.1 Gene amplification of mcyA, -B, -C, and -G

The nucleotide sequences for mcyA, -B, -C, and -G from M. aeruginosa PCC
7806 (accession number AF183408), P. agardhii CYA 126/8 (accession number
*AJ441056* and *A. circinalis* sp. 90 (accession number *AJ536156*) were obtained from Entrez nucleotide (http://www.ncbi.nlm.nih.gov/). The genes were aligned using the MultAlin (http://prodes.toulouse.inra.fr/multalin/multalin.html) algorithm and conserved regions were located. Degenerate primer mixes were then designed for the 5' and 3' end of the genes (Table 2). These mixes had calculated Tm values with ranges of 60-69°C and the degeneracy was kept as low as possible and below 100 except for mcyH5fwd (228-fold degeneracy). The genes were amplified using the Advantage 2 PCR kit (BD Biosciences, Mountainview), using the supplied protocol. The program used was 95°C-1 min then 35 cycles of 95°C-30 sec, 68°C-6 min followed by 68°C-6 min. The reaction mixtures were then applied to a 0.8% agarose gel and visualized using EtBr. The resulting bands were excised from the gel and recovered using the QiaQuick Gel extraction kit (Qiagen, Valencia, CA) according to the manufacturer's supplied protocol. The DNA was then ligated into the pDrive vector (Qiagen, Valencia, CA). The ligation mixture was heated at 65°C for 10 min to inactivate the ligase and the mixture was transformed into *E. coli* DH5α cells. The cells were plated out on LB_Amp agar plates containing X-Gal and IPTG and incubated overnight at 37°C. Positive clones were grown overnight in 3ml LB_Amp media in an incubated shaker and the plasmids were isolated using the High pure plasmid isolation kit (Roche). Plasmids were sequenced with the vector specific primers pDriveseq+/- pDriveseq- to ensure that they contained the desired gene fragment. The gene fragments were then sequenced as described below.
1.2.6.2 Amplification of \textit{mcyD} and \textit{E}

The nucleotide sequences for \textit{mcyD} and \textit{E} from \textit{M. aeruginosa} PCC 7806 (accession number \textit{AF183408}), \textit{P. agardhii} CYA 126/8 (accession number \textit{AJ441056}) and \textit{A. circinalis} sp. 90 (accession number \textit{AJ536156}) were obtained from Entrez nucleotide (http://www.ncbi.nlm.nih.gov/). The genes were aligned using the MultAlin (http://prodes.toulouse.inra.fr/multalin/multalin.html) algorithm\textsuperscript{32} and conserved regions were located. At the 5' and 3' ends of the genes degenerate primer mixes were designed to amplify 1kb fragments of the gene (Table 2). The primer mixes had Tm ranges of 54-59°C and a calculated degeneracy of less than 40. The 1kb fragments were amplified using \textit{Taq} DNA polymerase (Qiagen, Valencia, CA) with the following program: 95°C-3min, 40 cycles of 95°C-30 sec, 55°C-30 sec, 72°C-1 min followed by 72°C-2 min. The products were purified over a 1% agarose gel and visualized with ethidium bromide. The bands were removed from the gel and isolated with the QiaQuick gel elution kit (Qiagen, Valencia, CA) according to the manufacturer's supplied protocol. The DNA was then ligated into the pDrive vector (Qiagen, Valencia, CA). The ligation mixture was heated at 65°C for 10 min and the mixture was transformed into \textit{E. coli} DH5α cells. The cells were plated out on LB\textsubscript{Amp} agar plates containing X-Gal and IPTG and incubated overnight at 37°C. Positive clones were grown overnight in 3 ml LB\textsubscript{Amp} media in an orbital shaker and the plasmids were isolated using the High pure plasmid isolation kit (Roche). Plasmids were sequenced with the vector specific primers pDriveseq+/pDriveseq- to ensure that they contained the desired gene fragment. The sequence was confirmed by homology to the known \textit{mcy} genes by using the blastx search function provided by NCBI (http://www.ncbi.nlm.nih.gov/BLAST/). Once the sequence of the 1kb gene fragments
was known, gene specific primers were designed (Table 2). These primers had a calculated Tm of 66-68°C and overlapped with the known sequence by approximately 125bp. The unknown sequence of the gene was amplified using the Advantage 2 PCR kit (BD Biosciences, Mountainview, CA) with the following program: 95°C-1 min, 35 cycles 95°C-30 sec, 68°C-6 min followed by 68°C-6 min. The bands were purified using a 0.8% agarose gel and isolated with the QiaQuick gel elution kit (Qiagen, Valencia, CA) according to the manufacturer’s supplied protocol. The DNA was then ligated into the pDrive vector (Qiagen, Valencia, CA). The ligation mixture was heated at 65°C for 10 min and the mixture was transformed into E. coli DH5α cells. The cells were plated out on LB_Amp agar plates containing X-Gal and IPTG and incubated overnight at 37°C. Positive clones were grown overnight in 3 ml LB_Amp media in an orbital shaker and the plasmids were isolated using the High pure plasmid isolation kit (Roche). Plasmids were sequenced with the vector specific primers pDriveseq+/pDriveseq- to ensure that they contained the desired gene fragment. The gene fragments were then sequenced as described below.

1.2.6.3 Amplification of mcyH and -J

The nucleotide sequences for mcyH and -J from M. aeruginosa PCC 7806 (accession number AF183408), P. agardhii CYA 126/8 (accession number AJ441056) and A. circinalis sp. 90 (accession number AJ536156) were obtained from Entrez nucleotide (http://www.ncbi.nlm.nih.gov/). The genes were aligned using the MultAlin algorithm and conserved regions were located. Degenerate primer mixes were then designed for the 5’ and 3’ end of the
genes (Table 2). These primer mixtures had calculated Tm values within the range of 65-69°C and the degeneracy was kept as low as possible. The genes were amplified using the Taq DNA polymerase (Qiagen, Valencia, CA). The program used was 95°C-3 min then 40 cycles of 95°C-30 sec, 65°C-30 sec, 72°C-1 min followed by 72°C-2 min. The reaction mixtures were then applied to a 1.0% agarose gel and visualized using EtBr. The resulting bands were excised from the gel and recovered using the QiaQuick Gel extraction kit (Qiagen, Valencia, CA) according to the manufacturer’s supplied protocol. The DNA was then ligated into the pDrive vector (Qiagen, Valencia, CA). The ligation mixture was heated at 65°C for 10 min to inactivate the ligase and the mixture was transformed into E. coli DH5α cells. The cells were plated out on LB Amp agar plates containing X-Gal and IPTG and incubated overnight at 37°C. Positive clones were grown

Table 2: Degenerate primer sequences used in the amplification mcYA,-B,-C,-D,-E,-G,-H, and -J

<table>
<thead>
<tr>
<th>Name</th>
<th>Sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>mcYA5fwd</td>
<td>5'-AGTTTTGATGTTTCGTTGAGGAAATTCTTTTGG</td>
</tr>
<tr>
<td>mcYA3rev</td>
<td>5'-CCTTGATTYAAGTGCGACCKGGAAATC</td>
</tr>
<tr>
<td>mcYB5fwd</td>
<td>5'-CCATGCARSARGGGATTGCTTCATTCATAG</td>
</tr>
<tr>
<td>mcYB3rev</td>
<td>5'-CAAAANSRARGCTAAATACGCYTCGTGATTAAA</td>
</tr>
<tr>
<td>mcYC5fwd</td>
<td>5'-GCATGCGAAATACGTTAGGAAATGGA</td>
</tr>
<tr>
<td>mcYC3rev</td>
<td>5'-GAATCCARAGAAATTAAGTATCCANACTGGT</td>
</tr>
<tr>
<td>mcYDSS5fwd</td>
<td>5'-ATTCTCTAYCTTTTACTCTTAATG</td>
</tr>
<tr>
<td>mcYDSS5rev</td>
<td>5'-GTTCCYGATGCTCCAAAGA</td>
</tr>
<tr>
<td>mcYDSS3fwd</td>
<td>5'-TGCGGATGAGAGCAGTNNATT</td>
</tr>
<tr>
<td>mcYDSS3rev</td>
<td>5'-CTCTAACAATTCTGCTGGCTG</td>
</tr>
<tr>
<td>mcYDLR5fwd</td>
<td>5'-GCGATCGCTAGCTGCTGTTGTTGTTA</td>
</tr>
<tr>
<td>mcYDLR3rev</td>
<td>5'-GGCGTAATTCAAATCTGAAGTGATAGGAGGAAGTTGTTG</td>
</tr>
<tr>
<td>mcYE5F5fwd</td>
<td>5'-ATGGAGGYNATGCAATGCTG</td>
</tr>
<tr>
<td>mcYE5F5rev</td>
<td>5'-CCRAARGAATNACYCCAG</td>
</tr>
<tr>
<td>mcYE3S5fwd</td>
<td>5'-GATAACTTTYTTGAAAATGCGGG</td>
</tr>
<tr>
<td>mcYE3S5rev</td>
<td>5'-GWKMTTCTAAATCRGGATGATTT</td>
</tr>
<tr>
<td>mcYG5fwd</td>
<td>5'-AAGAYTTTTAATCCYGACTNTGCTGTTTT</td>
</tr>
<tr>
<td>mcYG3rev</td>
<td>5'-GARTCAAATCCACATTCTRWNAAATCCTGTTC</td>
</tr>
<tr>
<td>mcYH3fwd</td>
<td>5'-GATARGATTASKCAAGARAGAYTGKAATCTYAAGC</td>
</tr>
<tr>
<td>mcYH3rev</td>
<td>5'-GCCAACTAGARYTTSTRAAGTAGTCYARMACC</td>
</tr>
<tr>
<td>mcYJ5fwd</td>
<td>5'-ATCTTCABGATCYYATDTCCTCCAATGGCT</td>
</tr>
<tr>
<td>mcYJ3rev</td>
<td>5'-AAAAAGATAATCRTCMAHGCCTAATAACCAMCC</td>
</tr>
</tbody>
</table>

R - A or G, M - A or C, S - G or C, W - A or T, K - G or T, Y - C or T, B - C, G or T, D - A, G or T, V - A, C or G, H - A, C or T, N - A, G, C or T
overnight in 3ml LB_Amp media in an orbital shaker and the plasmids were isolated using
the High pure plasmid isolation kit (Roche). The inserts were sequenced with the vector
specific primers pDriveseq+/ pDriveseq-. This section was performed in triplicate to
ensure proper sequencing.

1.2.6.4 Sequencing of mcyA, -B, -C, -D, -E, and -G

To expedite sequencing, the genes were fragmented and sub-cloned into the
pUC19 vector. Plasmids containing the individual genes (10 μg DNA in 100 μl) were
sonicated in a Bachman ultrasonic cleaner in 10 sec intervals until the plasmid was
sheared as observed by gel electrophoresis. The sheared DNA was separated over a 1.2%
agarose gel and the region at approximately 1kb was excised from the gel and purified
using the QiaQuick gel extraction kit. The isolated DNA was then blunted by use of a
Klenow fragment exo- (Fermentas, Hanover, MD) according to the manufacturer’s
protocol. The now blunt ended DNA was ligated into the pUC19 plasmid, which had
been previously restricted by SmaI and dephosphorylated (New England Biolabs), using
T4 DNA ligase. The ligation mixture was transformed into E. coli DH5α cells and plated
on LB_Amp agar plates containing X-Gal and IPTG. Colony PCR was performed on a
variable number of colonies for each gene (number of colonies = kb DNA to be
sequenced x 5). Positive clones were grown overnight in 1.5 ml LB_Amp media in sterile
micro-centrifuge tubes and the plasmids were isolated by hand as described above. The
inserts were sequenced using the vector specific primers pDriveseq+ and pDriveseq-. The
appropriate sequences, as determined by sequence homology with known mcy genes
according to blastx (http://www.ncbi.nlm.nih.gov/BLAST/) were then assembled using
the ContigExpress function of VectorNTI 9.033 with the following parameters, pair wise
assembly algorithm with a minimum overlap of 20bp, minimum identity of 0.85 and minimum cutoff score of 40. Primer walking of the original pDrive plasmid filled in any gaps utilizing sequencing primers that had a calculated Tm of 57°C and overlapped the known sequences by approximately 125bp.

1.2.6.5 Orientation of the mcy gene cluster

Once the 5' and 3' sequences of the genes had been established, gene specific primers that were aimed toward the extremities were designed (Table 3). These primers had a calculated Tm of 57-58°C and overlapped known sequences by approximately 100bp. A variety of combinations based on known cluster orientations were attempted. PCR products were observed on 0.8% gels, (mcyA-G, E-H) and 1.2% gels (all others). The DNA was purified using the QiaQuick Gel extraction kit (Qiagen, Valencia, CA) according to the manufacturer's supplied protocol. The DNA was then ligated into the pDrive vector (Qiagen, Valencia, CA). The ligation mixture was heated at 65°C for 10 min and the mixture was transformed into E. coli DH5α cells. The cells were plated out on LB Amp agar plates containing X-Gal and IPTG and incubated overnight at 37°C.

Table 3: Primers used during the orientation PCR reactions

<table>
<thead>
<tr>
<th>Name</th>
<th>Sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>mcyA5revseq</td>
<td>5'-CTATCATCATTTAAAAAGACAGCAGACAACATAGT</td>
</tr>
<tr>
<td>mcyA5rev</td>
<td>5'-GATAGTAGTAAATTGGTAGCTGTCTCAGTCAGTC</td>
</tr>
<tr>
<td>mcyA3fwd</td>
<td>5'-TTAAAGTTAAGTTGGAGCTACAGTC</td>
</tr>
<tr>
<td>mcyB5rev</td>
<td>5'-ACTTTTTCCTAAGCTGCCTG</td>
</tr>
<tr>
<td>mcyB3fwd</td>
<td>5'-TGCACTTGGTAGATTTCCGTTATTT</td>
</tr>
<tr>
<td>mcyC5rev</td>
<td>5'-ATTGCCGTCATAGGAAAAATCGG</td>
</tr>
<tr>
<td>mcyD3rev</td>
<td>5'-CAAGCATTCAAGCCAGATCGC</td>
</tr>
<tr>
<td>mcyD3fwd</td>
<td>5'-CAAGAAGACTCTACATCTCTAGTAGGAG</td>
</tr>
<tr>
<td>mcyE5rev</td>
<td>5'-GACAGTAAATTCCAAATAAGTTCTCAGGAG</td>
</tr>
<tr>
<td>mcyE3fwd</td>
<td>5'-CATCGAGATTACAAGCGCA</td>
</tr>
<tr>
<td>mcyG3rev</td>
<td>5'-TCTAAATAGTGGACAGACTTGAAG</td>
</tr>
<tr>
<td>mcyG3fwd</td>
<td>5'-GGAGAACGTAGGAAAAATTATTAGG</td>
</tr>
<tr>
<td>mcyH5rev</td>
<td>5'-TTTCGAGAGATTATAATCGCGCGA</td>
</tr>
<tr>
<td>mcyJ5rev</td>
<td>5'-GGGAACATTACGTGTAAGAT</td>
</tr>
<tr>
<td>mcyJ3fwd</td>
<td>5'-TTATGCGCTTTTGACAGATTATCTT</td>
</tr>
</tbody>
</table>

*: all 5rev primers were created as the reverse compliment to the coding strand with the exception of mcyArevseq, which is simply the reverse sequence of the coding strand
Positive clones were grown overnight in 3ml LB<sub>Amp</sub> media in an orbital shaker and the plasmids were isolated using the High pure plasmid isolation kit (Roche). The inserts were sequenced with the vector specific primers pDriveseq+/pDriveseq- and sequencing by primer walking filled in any gaps.

1.2.6.6 Completing mcyH and -C

Because of the high nucleotide similarity between our cluster and the cluster sequenced from <i>A. circinalis</i> sp. 90, the 3' areas of mcyH and -C were amplified using gene specific primers (Table 4) derived from the <i>A. circinalis</i> gene cluster. Taq DNA polymerase (Qiagen, Valencia, CA) was used to amplify the products using the following program, 94°C-3 min, 40 cycles 95°C-30 sec, 57°C-30 sec, 72°C-1 min, then 72°C-2 min. The DNA was purified over a 0.8% agarose gel visualizing with EtBr. The resulting bands were excised from the gel and recovered using the QiaQuick Gel extraction kit (Qiagen, Valencia, CA) according to the manufacturer's supplied protocol, the DNA was then ligated into the pDrive vector (Qiagen, Valencia, CA). The ligation mixture was heated at 65°C for 10 min and the mixture was transformed into <i>E. coli</i> DH5α cells. The cells were plated out on LB<sub>Amp</sub> agar plates containing X-Gal and IPTG and incubated overnight at 37°C. Positive clones were grown overnight in 3ml LB<sub>Amp</sub> media at 37°C at 250 rpm and the plasmids were isolated using the High pure plasmid isolation kit (Roche). The inserts were sequenced with the vector specific primers pDriveseq+/pDriveseq-.

To ensure that the stop codon was indeed true a vector assisted PCR was performed. Cyanobacterial DNA was completely digested with <i>Sau3A</i> (Fermentas,
Hanover, MD) using the buffer supplied by the manufacturer and an appropriate number of units of enzyme. The DNA was purified using of the QiaQuick PCR purification kit (Qiagen, Valencia, CA) according to the supplier’s protocol. The restricted DNA was ligated into a pUC19 vector that had been digested with BamHI and dephosphorylated (New England Biolabs) using T4 DNA ligase (Fermentas, Hanover, MD) under the following conditions.

6.5 µl genomic DNA Sau3AI restricted (150 ng)
0.5 µl pUC19 (250 ng)
1µl 10X T4 DNA ligase buffer (Fermentas)
2 µl T4 DNA ligase (2 Weiss units)

The ligation was stored at 4°C for 18 h and then the DNA was purified by use of a QiaQuick PCR purification kit according to the manufacturer’s protocol.

Gene specific primers were designed as close to the 3’ end of the gene as possible albeit without including the stop codon (Table 4). A standard PCR reaction utilizing Taq DNA polymerase was then performed with one modification, the vector specific primer (pDriveseq+) was used at 100 pmol/µl stock concentration.

The PCR reaction mixture was applied to a 2% agarose gel and visualized using EtBr. The resulting bands were excised from the gel and recovered using the QiaQuick Gel extraction kit (Qiagen, Valencia, CA) according to the manufacturer’s supplied protocol. The DNA was then ligated into the pDrive vector (Qiagen, Valencia, CA). The ligation mixture was heated at 65°C for 10 min and the mixture was transformed into E. coli DH5α cells. The cells were plated out on LB Amp agar plates containing X-Gal and IPTG and incubated overnight at 37°C. Positive clones were grown overnight in 3ml
LB\textsubscript{Amp} media at 37°C at 250 rpm and the plasmids were isolated using the High pure plasmid isolation kit (Roche). The inserts were sequenced with the vector specific primers pDriveseq+/ pDriveseq-.

Table 4: Primers used during the completion of \textit{mcyC} and \textit{mcyH}

<table>
<thead>
<tr>
<th>Name</th>
<th>Sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>\textit{mcyH} AnabaenaRev</td>
<td>5'-CAAAGATTATCGAAATAAAAAATACTTTTTATAAA</td>
</tr>
<tr>
<td>\textit{mcyC} AnabaenaRev</td>
<td>5'-TCAATTCTGACTAAGAATCTCACTTCAATTAATAA</td>
</tr>
<tr>
<td>\textit{mcyH} vectorFwd</td>
<td>5'-AATGGGTTCTGAGATTCACAGGATTC</td>
</tr>
<tr>
<td>\textit{mcyC} vectorFwd</td>
<td>5'-ATGATTGATCCTCCCTATCTCAATCAAAAC</td>
</tr>
</tbody>
</table>
1.3 RESULTS AND DISCUSSION

1.3.1 Amplification/Identification of the \textit{mcy} gene cluster

Since three microcystin gene clusters had been fully characterized at the beginning of this investigation we chose to take a unique approach utilizing PCR to sequence the microcystin gene cluster from \textit{H. hibernicus}. The known nucleotide sequences from \textit{mcyA, -B, -C, -D, -E, -G, -H,} and \textit{-J} were aligned to reveal conserved nucleotide regions which were then used to derive degenerate primer mixes as described in the experimental section. Six of the primer pairs (\textit{mcyA, -B, -C, -H, -G, -J}) gave the expected amplicons, however the primer pairs for the two largest PKS genes (\textit{mcyD, -E}) did not yield a PCR product. One could attribute the lack of PCR products simply to the fact that the desired amplicon was too large (10kb and 11kb respectively). In comparison, the largest fragment that was amplified successfully was \textit{mcyA} (9kb). Another explanation is that the degenerate primer mixes that we used were located in conserved regions of the PKS. This would prevent the desired amplicon from being replicated by scattering the primers throughout the genomic DNA. To circumvent this problem 1kb stretches from the 5' and 3' regions of \textit{mcyD and -E} were amplified and sequenced. Sequence specific primers were then created to amplify the entire gene while overlapping approximately 150 bp with the known sequence. This approach yielded PCR products of the appropriate size that upon sequencing were verified to be the \textit{mcyD} and \textit{mcyE} gene fragments.

The previously published gene clusters contained three other genes of which two, \textit{mcyF}, and \textit{-I}, have been found in two of the three clusters,\textsuperscript{16,18} while \textit{mcyT} was discovered only in the \textit{Planktothrix} cluster.\textsuperscript{17} It was therefore decided that these genes
would be sequenced during the PCR reactions used for joining the gene fragments or discovered during the sequencing of the cluster's flanking regions.

1.3.2 Organization of the microcystin gene cluster

After the genes were amplified and sequenced, gene specific primers were designed to amplify regions between the known sequences. PCR reactions were tried in the order and orientation known from the other clusters. The previously sequenced gene clusters all contained \textit{mcyB} flanked by \textit{mcyA} at the 5' end and \textit{mcyC} at the 3' end. Since this was observed to be the only commonality in gene orientation between the known clusters, it was assumed to be present in the \textit{H. hibernicus} cluster and therefore was the first set of PCR reactions attempted. The next step was to establish the orientation of the other genes. The next three PCR reactions that were attempted were between the primer pairs \textit{mcyASrevseq/mcyD5rev}, \textit{mcyASrevseq/mcyG5rev} and \textit{mcyA5rev/mcyH3fwd} which mimic the arrangement of the genes in \textit{Microcystis}, \textit{Anabaena} and \textit{Planktothrix}, respectively. Only the primer pair \textit{mcyA5revseq/mcyG5rev} gave an amplicon of the appropriate size that upon sequencing was confirmed to be the desired product. To limit the amount of PCR reactions that were performed the next set was modeled after the arrangement in the \textit{Anabaena} cluster. All the reactions afforded predominant bands of the expected size. Given the similarities between the \textit{H. hibernicus} and \textit{A. circinalis} clusters it was anticipated that \textit{mcyF} and \textit{mcyI} would be located between \textit{mcyE} and \textit{mcyH} and this was discovered to be the case during sequencing. The PCR reactions established the order and orientation of the genes as seen in Figure 5. The resulting sequences were assembled using the ContigExpress platform in the VectorNTI software. Each of the original gene
fragments amplified from the degenerate primer mixtures were assembled into contigs first. These contigs along with the sequences from the second round of PCR were then assembled into the gene cluster.

The gene cluster is 55.5kb in length and was discovered to contain two putative operons. The first operon \((mcyABC)\) is transcribed in the opposite direction, from a putative bi-directional promoter, as the second operon \((mcyGDJEFIH)\). The promoter region is 891 bp in length and is proposed to be bi-directional in nature. Use of BPROM, a web based program for sigma70 promoter recognition, yielded possible \(-10\) and \(-35\) boxes in the analyzed region for each direction.\(^{34}\)

\[\text{Figure 5: Schematic diagram of the mcy cluster sequenced from } H. \text{hibernicus as well as the known clusters from } M. \text{aeruginosa, P. agardhii, A. circinalis and the nodularin gene cluster from } N. \text{spumigena. Genes not drawn to scale.} \]
1.3.3 Characterization of the microcystin genes

Upon assembly of the gene cluster the nucleotide sequences of the individual genes were translated and compared to the corresponding proteins of the known microcystin gene clusters and the nodularin gene cluster using the Basic Local Alignment Search Tool (BLAST) available from the NCBI (Table 5). All genes held significant similarity on both the amino acid and nucleotide level. The percent similarity on the amino acid level ranges from 53% (NdaA-N. spumigena) to 92% (McyJ-A. circinalis). The lowest similarity to a mcy gene was 67% (McyB-P. agardhii/McyH-M. aeruginosa). The similarity on the nucleotide level ranges from 79% (mcyD-M. aeruginosa) to 89% (ndaH-N. spumigena/mcyJ-A. circinalis).

Table 5: Percent similarity of genes from H. hibernicus to known mcy genes. % homology to amino acid (% homology to nucleotide)

<table>
<thead>
<tr>
<th>Gene</th>
<th>mcyA</th>
<th>mcyB</th>
<th>mcyC</th>
<th>mcyD</th>
<th>mcyE</th>
<th>mcyF</th>
<th>mcyG</th>
<th>mcyH</th>
<th>mcyJ</th>
</tr>
</thead>
<tbody>
<tr>
<td>A. circinalis</td>
<td>77(85)</td>
<td>82(86)</td>
<td>80(85)</td>
<td>78(86)</td>
<td>78(84)</td>
<td>72(80)</td>
<td>79(86)</td>
<td>75(85)</td>
<td>78(85)</td>
</tr>
<tr>
<td>N. spumigena</td>
<td>53(82)</td>
<td>N/A</td>
<td>77(84)</td>
<td>74(85)</td>
<td>79(85)</td>
<td>79(81)</td>
<td>77(85)</td>
<td>75(84)</td>
<td>83(89)</td>
</tr>
<tr>
<td>P. agardhii</td>
<td>62(84)</td>
<td>67(80)</td>
<td>75(82)</td>
<td>71(82)</td>
<td>75(84)</td>
<td>N/A</td>
<td>73(80)</td>
<td>70(80)</td>
<td>N/A</td>
</tr>
<tr>
<td>M. aeruginosa</td>
<td>64(83)</td>
<td>68(81)</td>
<td>70(85)</td>
<td>67(79)</td>
<td>73(81)</td>
<td>69(79)</td>
<td>69(81)</td>
<td>67(80)</td>
<td>69(79)</td>
</tr>
</tbody>
</table>

The low homology to NdaA (53%) can be explained by the proposed evolution of the nda cluster. Nodularin is missing two amino acid residues when compared to microcystin, D-Ala and the L-X residue. These amino acids are activated by mcyA-A2 and mcyB-A1 respectively. Moffitt and Neilan proposed that the nodularin cluster arose from a deletion event involving excision of the second CAT from mcyA and the first CAT from mcyB, which would result in ndaA.\(^{35}\) When McyA and McyB are compared to NdaA, similarity values of 37% for both are obtained. When NdaA-A1T1 and NdaA-
C$_2$A$_2$T$_2$ are compared to the corresponding region from McyA and McyB the similarity is 54% and 73%, respectively. The lower similarity of NdaA-A$_1$T$_1$ can be attributed the fact that nodularin contains Mdhb (derived from Thr) and microcystin contains Mdha (derived from Ser).

The first operon (mcyABC) contains 3 ORFs. The proposed ORFs for mcyA and mcyC begin with an ATG start codon while the ORF for mcyB is proposed to begin with a GTG codon. A section of 20bp separates the termination codon of mcyA for the start codon of mcyB while 2bp separates the TAA of mcyB from the ATG of mcyC. The lengths of mcyA, -B, -C are 8,388bp, 6,390bp and 3,858bp and encode apo proteins of a predicted mass of 314,605 Da, 241,944 Da and 147,174 Da respectively.

The second operon contains the PKS genes (mcyG, -D, -E), the tailoring enzymes (mcyJ, -F, -I) and the putative ABC transporter (mcyH). There are 71bp between mcyG and mcyD, 78 bp separating mcyD and mcyJ, 35 bp between mcyJ and mcyE, 47 bp separating mcyE and mcyF, 35 bp separating mcyF and mcyI and a 21 bp spacer between mcyI and mcyH. The lengths of mcyG, -D, -J, -E, -F, -I, -H are 7,923 bp, 11,626 bp, 924 bp, 10,392 bp, 766 bp, 1,026 bp and 1,780 bp, respectively, and, correspondingly, encode for apo proteins of 294,142 Da, 43,127 Da, 36,193 Da, 387,621 Da, 28,723 Da, 37,189 Da, 67,498 Da in size. McyG, -D, and -J were most similar to A. circinalis with similarities ranging from 69 to 79%, 67 to 78%, 81 to 92%, respectively, while McyE, -F, -I held the highest similarity to N. spumigena and had ranges of 73 to 79%, 75 to 79%, and 69 to 83%. McyH was equally similar (75%) to both A. circinalis and N. spumigena.

The second operon (mcyGDJEFIH) from H. hibernicus is most similar in arrangement to that of the nodularin gene cluster. In Anabaena there is a putative
transcriptional start point for mcyH in the 295 bp spacing between mcyI and mcyH. In Hapalosiphon, as in Nodularia, that spacing is less than 30bp (21 bp and 23 bp respectively) and no promoter region can be identified. Not surprisingly the homology for McyF, -I and -H is highest when compared to the corresponding nda genes (Ndg, -H and -I).

1.3.4 NRPS adenylation domain specificities

The substrate specificity conferring amino acid residues of the adenylation domains from M. aeruginosa, P. agardhii, A. circinalis and H. hibernicus were analyzed by the method of Stachelhaus et al.36 (Table 6). The signature sequences of mcyA-A1, -A2, -BA1, -B-A2 and -C-A from H. hibernicus are very tightly conserved with respect to the known microcystin signature sequences. Oddly, the signature sequences from H. hibernicus and A. circinalis for mcyB-A1 and mcyC-A are identical although they supposedly activate different amino acids. A. circinalis has been shown to produce MC-LR and MC-RR while H. hibernicus has previously been shown to produce MC-LA. However, further analysis indicates the presence of MC-AR (R. Kurmayer, G. Christiansen, personal communication). The large differences in amino acid side chain size and polarity raise questions as to the discrimination ability/process. Further biochemical studies are needed to address this issue and are currently in progress.

Table 6: Amino acid specificity conferring sequences of adenylation domains as Stachelhaus et al.36

<table>
<thead>
<tr>
<th></th>
<th>McyA-A1</th>
<th>McyA-A2</th>
<th>McyB-A1</th>
<th>McyB-A2</th>
<th>McyC-A</th>
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<tr>
<td>H. hibernicus</td>
<td>DVWHISLIV DLFNNALTS DVFGLYVA</td>
<td>DARHIGIFX DVFGLYVA</td>
<td>DARHIGIFX DVFGLYVA</td>
<td>DARHIGIFX DVFGLYVA</td>
<td></td>
</tr>
<tr>
<td>P. agardhii</td>
<td>DVWHISLIV DLFNNALTS DLFGLYVA</td>
<td>DARHIGIFX DPFGLYVA</td>
<td>DARHIGIFX DPFGLYVA</td>
<td>DARHIGIFX DPFGLYVA</td>
<td></td>
</tr>
<tr>
<td>M. aeruginosa</td>
<td>DVWHISLIV DLFNNALTS DAWFLGNTV</td>
<td>DARHIGIFK DVFGLYVA</td>
<td>DARHIGIFK DVFGLYVA</td>
<td>DARHIGIFK DVFGLYVA</td>
<td></td>
</tr>
<tr>
<td>A. circinalis</td>
<td>DVWHISLIV DLFNNALTS DVFGLYVA</td>
<td>DARHIGIFK DVFGLYVA</td>
<td>DARHIGIFK DVFGLYVA</td>
<td>DARHIGIFK DVFGLYVA</td>
<td></td>
</tr>
</tbody>
</table>
1.3.5 Phylogenetic analysis of adenylation domains

Using the nucleotide sequence of the adenylation domains from the conserved motif A1 \((L(TS)YxEL)\) up to and including the amino acid just prior to the conserved motif of the thiolation domain \((DxFFxxLGG(HD)S(LI))\), a phylogenetic analysis was performed, the results of which can be seen in Figure 6. Most of the adenylation domains cluster together as one would expect due to the proposed linear evolution of the microcystin synthetase gene cluster (i.e. all \(mcyA\)-A domains show higher similarity to each other than to other A domains). However an abnormality can be observed when analyzing \(mcyB\)-A1 and \(mcyC\)-A of both \(H. hibernicus\) and \(A. circinalis\). This clustering seems to be the result of a duplication of the \(mcyC\)-A domain followed by recombination as suggested by the closer relation to \(P. agardhii mcyC\)-A (Figure 6a). However, we find no evidence that the condensation domains have undergone a similar evolution (Figure 6b).

The microcystin gene cluster is purported to be a site of frequent recombination and mutation, which may help explain the wide variety of microcystin variants seen in nature. Also evidence of recombination of small stretches\(^{38}\) and entire domains\(^{39}\) has been reported. When aligning the adenylation domain amino acid sequences obtained from \(mcyB\)-A1 and \(mcyC\)-A, one can observe only eleven amino acid substitutions in the entire adenylation domain. And more importantly only one amino acid substitution between the conserved motifs A4 and A5, the purported site for amino acid selectivity.\(^{40}\)
Figure 6: (a) Evolutionary tree diagram of the adenylation domains found in the microcystin clusters and the nodularin cluster using GrsA-A1 as an out-group. (b) Evolutionary tree diagram of the condensation domains found in the microcystin clusters as well as the nodularin cluster. Generated using Vector NTI version 9.0
1.4 CONCLUSION

A 55.5 kb gene cluster responsible for the biosynthesis of microcystin in *H. hibernicus* was sequenced using a unique degenerate PCR method. It is envisioned that this procedure could be used on any strain that produces microcystins and could be a valuable tool in genetic analysis of the *mcy* clusters. The time necessary for sequencing the *mcy* gene cluster was significantly reduced (approximately 5 weeks compared to a minimum of 1 year for the standard approach using hybridization of genomic libraries). As an additional benefit, the problems associated with the creation of a genomic library were avoided. This method is quick, inexpensive and relies on materials/kits already contained in all molecular biology labs (PCR reagents, cloning vectors, and DNA sequencing equipment). It is envisioned that this technique will be helpful in studying the *mcy* gene cluster in any isolated strain.
2. Genetic knockout of mcyT in *P. agardhii* 126/8

2.1 INTRODUCTION

2.1.1 McyI

The biosynthesis of microcystin requires the formation of Mdha, which is postulated to arise from dehydration of a serine residue. The gene clusters from *Anabaena, Microcystis, Hapalosiphon* and the nodularin gene cluster from *Nodularia* contain a *D*-3-phosphoglycerate dehydrogenase homolog, *mcyI* (*ndaH*), while it was absent from the sequenced gene cluster found in *Planktothrix*. It has been hypothesized that McyI is responsible for the formation of the Mdha residue. However, as of this date no *in vitro* characterizations of this protein have been reported.

*D*-3-Phosphoglycerate dehydrogenases are known to contain three domains, the catalytic domain, the NAD binding domain and the ACT domain. The ACT domain is where the allosteric inhibitor serine binds to induce a conformational change thereby inhibiting the formation of *3*-phosphopyruvate.\(^4\) During the comparison of the amino acid sequence of McyI to known *D*-3-phosphoglycerate dehydrogenases it was observed that the ACT domain was missing in McyI (Figure 7). This suggested that McyI might not be involved in serine biosynthesis but perhaps was the protein that catalyzed Mdha formation instead. With McyI having such high homology to known *D*-3-phosphoglycerate dehydrogenases (42% identity, *Rubrobacter xylanophilus* DSM 9941) it was postulated that the reactive intermediate in dehydration is phosphoserine, which would reduce the energy of activation necessary for dehydration and produce inorganic phosphate as a by-product as seen in the lantibiotics.
Figure 7: Multalign of 3-D-phosphoglycerate dehydrogenases from *E. coli* W3110, *E. coli* K-12, *Tersina pestis*, *Mycobacterium tuberculosis*, *Synechocystis* sp. PCC 6803, *Homo sapiens* and Mcyf from *H. hibernicus* BZ-3-1. The ACT domain is underlined in red.
The lantibiotics are a group of ribosomally produced peptide antibiotics that are characterized by the presence of lanthionine (Lan) and/or methylanthionine (MeLan). These residues are formed by post-translational dehydration of either serine or threonine, followed by Michael addition of the thiol group of a cysteine. The initial step in the dehydration involves the transfer of the gamma phosphate group of ATP onto the serine/threonine hydroxyl group resulting in phosphoserine and ADP. With the hydroxyl group activated, the dehydration can occur with a lower activation barrier. The dehydration and cyclization can be catalyzed by a pair of proteins as in the case of nisin, where NisB and NisC catalyze the dehydration and cyclization respectively, or by a single enzyme as in the case of lacticin 481, where LctM catalyzes both reactions. Interestingly, the C terminus of LctM showed some homology to LanC but the N terminus showed no homology to LanB making in silico prediction of the dehydratase activity impossible.

2.1.2 MctT

Since mcyI is absent from the Planktothrix gene cluster efforts were undertaken to locate it at a different locus in the genome. These attempts, which utilized PCR and degenerate primers, were unsuccessful (results not shown). This observation left the question of how the Planktothrix genus synthesizes the Mdha residue unanswered. While lacking mcyI, Planktothrix does contain mcyT, a type II thioesterase homolog (68% identity, Nostoc punctiforme PCC 73102).

Type II thioesterases are known to have two functions in vitro. They are known to cleave either misprimed amino acids or acetyl groups tethered to the 4'-phosphopantetheinyl (4'-PP) arm of thiolation domains. In both cases the thioesterases
are responsible for binding and interacting with thioesters, which they cleave hydrolytically to generate the free thiol and acid. Therefore it is conceivable that the *Planktothrix* genus evolved to form Mdha from O-acetyl-serine convergently.

To probe the function of McyT in the biosynthesis of MC-RR insertional mutagenesis was decided to be the best course of action. Christiansen *et al.* had previously used this technique to inactivate *mcyJ* in CYA 126/8. Analyzing the mutant strain with MALDI-TOF revealed that (D-Asp$^3$)-MC-RR was no longer present, but a new variant reduced in mass by 14 units was observed. This variant was identified as (DMAAdda, D-Asp$^3$)-MC-RR that contains a desmethoxy-Adda side chain (11). It was envisioned that insertional mutagenesis of *mcyT* might provide insight into the function of McyT by yielding a new structural microcystin variant.
2.2 EXPERIMENTAL SECTION

2.2.1 Insertional mutagenesis of mcyT

2.2.1.1 Construction of the homologous recombination plasmid

The mcyT mutant amplicon, consisting of mcyT, the bi-directional promoter and the 5’-end of mcyD, was amplified using the Advantage 2 PCR kit® (BD Biosciences) with the primers mcyTfwd (5’-CGACACTAGTGGGATACGGTAGGCGATG) and mcyTre5 (5’-ACTCAGAAAAATTTCCAGCCCTTCTGC). As expected an amplicon that was 3kb in size was obtained. It was purified over a 0.8% agarose gel and isolated using the QiaQuick Gel extraction Kit according to the manufacturer’s protocol. The amplicon was then cloned into the pDrive vector (Qiagen) and transformed into E. coli DH5α cells. The cells were grown overnight at 37°C on an LBAAMP plate containing X-Gal and IPTG. Positive colonies were then grown overnight in LBAMP media in an orbital shaker and the plasmid was isolated using the High Pure Plasmid Isolation Kit (Roche Applied Sciences). The plasmid was designated pDT-1.

2.2.1.2 Creating the chloramphenicol disrupted mcyT gene

After the amplicon was verified via sequencing, the restriction with PfoI was performed. This procedure was modified from Parker et al.45. In a 0.2 ml thin walled PCR tube (Corning) 5 µl of pDT-1 in TE buffer, pH 8.0 (500 ng) was combined with 3.8 µl dH2O, 1 µl Buffer Tango (Fermentas), 0.2 µl EtBr (1 mg/ml) and 2 units of PfoI (Fermentas). The reaction was performed as 16 replicate samples to provide suitable amounts of DNA for the next step. Incubation with the enzyme was performed at 37°C for 20 hours at which time the reactions were run on a 0.8% agarose gel (20 cm, 100V, 45 min) to ensure separation of the singly digested form from the fully restricted form.
The correct band was identified by comparison against pDT-1 linearized with SacI and fully restricted with PfoI according to the supplier's protocols. The linearized form from all 16 reactions was combined and isolated from the gel using the QiaQuick Gel extraction kit (Qiagen) according to the supplier's protocol. The 5'-overhangs generated by PfoI were blunted by exposure to the Klenow exo-- fragment (Fermentas) according to the manufacturer's protocol. The linearized mcyT-pDrive construct was ligated to the chloramphenicol resistance cassette of pACYC184 (NEB). The ligation was accomplished using T4 DNA ligase (Fermentas) as described above. The chloramphenicol resistance cassette was isolated by restriction of pACYC184 using sufficient units of BsaAI (New England Biolabs). The 1.2kb fragment was isolated from a 0.8% agarose gel using the QiaQuick Gel extraction Kit (Qiagen). The ligation mixture was transformed into electro-competent E. coli DH5α cells. The cells were grown overnight on LBA plates containing 100 µg/ml ampicillin and 50 µg/ml chloramphenicol. Colonies were then checked for the appropriate insert location by colony PCR using the primers mcyT5seq (5'-GTTAGGTGGGGTAATATGGC) and mcyT3seq (5'-CGACTATTCAGTCCTCATTATGC). The positive colony obtained was grown overnight in LB media plus 100 µg/ml ampicillin and 50 µg/ml chloramphenicol and the plasmid isolated with the High Pure Plasmid Isolation Kit (Roche Applied Sciences) to give plasmid pDTCm-1.

2.2.1.3 Preparation of the transformation construct

Plasmid pDTCm-1 was retransformed into E. coli DH5α cells and grown overnight at 37°C on an LB agar plate containing 100 µg/ml ampicillin and 50 µg/ml
chloramphenicol. Four individual colonies were grown overnight in 5ml LB media plus 100 μg/ml ampicillin and 50 μg/ml chloramphenicol in an orbital shaker. The plasmids were isolated by hand preparation followed by digestion with sufficient units of SacI. Two restriction digests were combined and the DNA purified using the QiaQuick PCR purification kit according to the manufacturer’s protocol with one exception. The DNA was eluted using two 50 μl aliquots of Buffer EB and stored on ice until use.

2.2.1.4 Transformation of P. agardhii CYA 126/8

P. agardhii CYA 126/8 was grown until mid log phase growth (approximately 3 weeks) at 22°C and a light intensity of 80-100 μmol photons m⁻² s⁻¹ in Z+G media. 50 ml of cell culture was centrifuged down in a 50 ml Falcon tube using the SLA-1500 rotor (11,000 rpm, 15 min, RT). 25 ml of the media was added to a 125 ml Erlenmeyer flask containing 25 ml freshly prepared Z+G media, this was stored for later use. The cells were washed twice with 20 ml 1 mM HEPES (filter sterile) followed by centrifugation (SLA-1500, 11,000 rpm, 15 min, RT). The cells were then resuspended in 1 ml 1 mM HEPES and transferred to a 1.5 ml eppendorf tube and pelleted again, this time using a F-20/Micro rotor (11,000 rpm, 15 min, RT). The supernatant was drained via aspiration and the cells were placed on ice until use.

The linearized DNA was heated at 100°C for 15 min in a heat block followed by cooling on ice. After 5 min on ice the tube was centrifuged for 10 sec to recover the condensed water and then the DNA was stored on ice until use. The cells were then resuspended in the denatured DNA and the slurry was transferred to a 0.2 cm Gene Pulser cuvette (BioRad) (pre-cooled to -20°C). The electroporation was performed on a Gene Pulser II
As quickly as possible after electroporation the cells were removed from the electroporation cuvette and transferred to the 125 ml Erlenmeyer flask containing Z+G media prepared above by washing the cuvette with 1 mL media. The flask was covered with a sheet of white paper and kept under the normal culture conditions. After 24 h the paper was removed and the culture was allowed to grow for an additional 24 h. At this time chloramphenicol was added to a final concentration of 1 μl/ml and the culture left alone. Within one week the culture became completely bleached of all color and after five weeks a faint green tint appeared in the flask. The mutant was then sub-cultured two times in Z+G media containing 1 μg/ml chloramphenicol to ensure that all the wild type strain had been killed. The mutant strain was named *P. agardhii* CYA 126/8 ΔmcyT.

### 2.2.1.5 Confirmation of mutant identity

DNA was isolated from CYA 126/8 ΔmcyT as described above in section 1.2.1.8 and a PCR reaction was performed using the primers mcyT5seq and mcyT3seq using *Taq* DNA polymerase. The resulting amplicon was compared to wild type DNA and construct DNA amplicons on a 0.8% agarose gel.

### 2.2.2 Peptide analysis

#### 2.2.2.1 Peptide extraction of *P. agardhii* CYA 126/8 ΔmcyT

CYA 126/8 ΔmcyT was cultured in 20 L of Z+G media containing 1 μg/ml chloramphenicol with air bubbled through at a rate of 3 L/min and a light intensity of 80-100 μmol photons m⁻² s⁻¹. When the cells reached saturation they were concentrated to a
pellet (8.9 g wet weight). The cells were then extracted twice with 50% aqueous methanol (20 ml) by sonicating for 15 min in a Branson 1510 ultrasonic cleaner followed by shaking at 250 rpm at room temperature for 15 min. The debris was removed by centrifugation in an International clinical centrifuge model CL on setting 6 for 10 min. The methanol was removed from the green supernatant at reduced pressure. The residue was then applied to a 4 g ODS-A column (YMC-GEL, 70 mesh) that had been pre-equilibrated with water. The residue was loaded onto the column and then batch eluted using 30 ml each of 25% aqueous methanol, 50% aqueous methanol, 75% aqueous methanol and 100% methanol. The fractions were dried using a SPD Speed Vac (Thermo Savant) without heating. The 50% methanolic fraction yielded 14.1 mg, the 75% methanolic fraction yielded 2.1 mg and the 100% methanol fraction yielded 2.2 mg. Due to the low mass in the 75% methanolic fraction it was combined with the 50% aqueous methanol fraction. The samples were then ready for analysis by HPLC and LC-MS.

Extracts of the wild type were prepared in an analogous fashion.

### 2.2.2.2 HPLC analysis of CYA 126/8 ΔmcyT peptide content

The combined 50% and 75% aqueous methanol fractions were dissolved in methanol at a concentration of 10 mg/ml and filtered through a 0.22 μm filter. The extracts were then analyzed using a Shimadzu LC-10AS HPLC equipped with a HiBar® RT 250-4 column (RP-18, 5 μm 5 mm X 22 cm). The sample was run at 1 ml/min flow rate from 30% aqueous acetonitrile plus 0.5% TFA to 100% acetonitrile plus 0.5% TFA over 45 min and observed at 210 and 240 nm. The (D-Asp³)-MC-RR was a large peak with a retention time of 14.27 min in the wildtype extract.
2.2.2.3 LC-MS analysis of CYA 126/8 ΔamcyT peptide content

The combined 50% and 75% aqueous methanol fractions (conc.: 10 mg/ml) for CYA 126/8 ΔamcyT and wild type were analyzed using a Agilent 1100 LC-MS equipped with a HiBar® RT 250-4 column (RP-18, 5 μm 5mm X 22cm). The sample was run at 1 ml/min flow rate from 30% aqueous acetonitrile plus 0.5% trifluoroacetic acid to 90% aqueous acetonitrile plus 0.5% trifluoroacetic acid over 30 min. The ions were observed using an Agilent TOF-MSD in positive ESI mode and the data was analyzed using Analyst QS software (ver 1.1, Applied Biosystems).

2.2.3 Anabaenopeptin I

2.2.3.1 Isolation of Anabaenopeptin I (14)

The combined 50% and 75% aqueous methanol fractions were repeatedly injected on a Shimadzu LC-10AS equipped with a LUNA 10 μ C-18 column (250 X 10 mm, Phenomenex). The sample was run at 3 ml/min using the gradient outlined in Table 7. Pure anabaenopeptin I was isolated at a retention time of 21.4 min. The structure was elucidated through the use of NMR and MS techniques.

Table 7: HPLC gradient used for the isolation of Anabaenopeptin I. Mobile phase contained 0.05% trifluoroacetic acid.

<table>
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<tr>
<th>Time (min)</th>
<th>AcCN (%)</th>
<th>H₂O (%)</th>
</tr>
</thead>
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<tr>
<td>0</td>
<td>30:70</td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>30:70</td>
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<td>25</td>
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<td>30</td>
<td>50:50</td>
<td></td>
</tr>
<tr>
<td>32</td>
<td>100:0</td>
<td></td>
</tr>
</tbody>
</table>
2.2.3.2 Anabaenopeptin I (14) was isolated as a white amorphous powder: \([\alpha]_{D}^{25} -21.43^\circ (c 0.35 \text{ mg ml}^{-1}, \text{MeOH})\); UV (MeOH) \(\lambda_{\text{max}} (\log e)\) 280 (2937); IR (film) \(\nu_{\text{max}}\)3426, 3960, 2922, 1639 cm\(^{-1}\); HR-ESI TOF \(m/z\) 916.4825 (calcd for \(\text{C}_{4840}\text{N}_{57}\text{O}_{11}\) 916.4821, 0.4 ppm error)

2.2.4 N-Me homotyrosine synthesis

2.2.4.1 DL-\(N\)-Acetyl-homotyrosine

\(DL\)-Homotyrosine (250 mg/1.28 mmol) was added to glacial acetic acid (13 mL) to create a suspension. Acetic anhydride (133 \(\mu\)L/1.40 mmol) was added and then the reaction was kept at 110\(^\circ\)C overnight. The acetic acid was removed and the solid product was suspended in a minimal amount of 1N HCl. The aqueous layer was extracted 3 times with EtOAc. The organic layers were combined and dried over MgSO\(_4\), filtered and then the solvent was removed yielding \(N\)-Acetyl-homotyrosine. Yield: 290 mg (95%) \(^1\)H NMR (300 MHz, MeOH-\(d_4\)) \(\delta\) 1.99 (s, 3H), 2.05 (m, 2H), 2.59 (m, 2H), 4.32 (d, \(J = 3.6\) Hz, 1H), 6.69 (d, \(J = 8.4\) Hz, 2H), 7.00 (d, \(J = 8.4\) Hz, 2H); \(^{13}\)C NMR (75 MHz, MeOH-\(d_4\)) \(\delta\) 175.6, 173.4, 156.7, 132.9, 130.4, 116.2, 34.7, 32.2, 22.3

2.2.4.2 L and D-Homotyrosine

\(DL\)-\(N\)-Acetyl-homotyrosine (290mg/ 1.28mmol) was dissolved in 0.1M K\(_2\)HPO\(_4\) buffer (pH 7.0) (25 mL). The pH was then adjusted to 7 by addition of NaOH. Acylase I from Aspergillus sp. [Sigma-Aldrich] (50 mg/2.35 units) was dissolved in 0.1M K\(_2\)HPO\(_4\) buffer (pH 7.0) (0.5 mL). The acylase was added to the \(DL\)-\(N\)-Acetyl-homotyrosine solution and allowed to sit at RT for 36h. 75 mL AcCN was added and the resulting solid
filtered through Celite. The AcCN was removed and the remaining aqueous layer was titrated to pH 1 using HCl. The aqueous layer was extracted 4 times with EtOAc.

The organic layers were combined and dried over MgSO₄, filtered and then the solvent removed too yield a yellow oil. This oil was suspended in 3N HCl (10ml) and heated at 65°C overnight. The aqueous HCl was removed yielding D-Homotyrosine.

The extracted aqueous layer from above was titrated to pH 5.5 with solid NaHCO₃ the concentrated to dryness to yield L-Homotyrosine. ¹H NMR (300 MHz, MeOH-d₄) δ 2.01 (m, 2H), 2.52 (dd, J = 7.5, 7.8 Hz, 2H), 3.88 (t, J = 5.4 Hz, 1H), 6.67 (d, J = 7.8 Hz, 2H), 7.00 (d, J = 7.8 Hz, 2H)

2.2.4.3 Homotyrosine methyl ester

Resolved homotyrosine (125mg/0.64 mmol) was dissolved in dry MeOH (10mL) and TMS-Cl (813μL/ 6.4 mmol) was added. The reaction was allowed to stir overnight at RT. The reaction mixture was concentrated to dryness and then dissolved in a minimal amount of dry MeOH (2-3mL). The dissolving and resuspending was repeated 3 times. The crude product was used without further purification. ¹H NMR (300 MHz, MeOH-d₄) δ 2.121 (m, 2H), 2.67 (m, 2H), 3.82 (s, 3H), 4.03 (t, J = 5.4 Hz, 1H), 6.72 (d, J = 7.8 Hz, 2H), 7.06 (d, J = 7.8 Hz, 2H); ¹³C NMR (75 MHz, MeOH-d₄) δ 169.9, 156.1, 130.8, 129.9, 115.7, 54.3, 53.2, 33.2, 30.6; IR (film) 3619, 1741, 1646, 1244; HR-ESI

C₁₁H₁₆NO₃ calcd. 210.1131, obs.210.1136, 2.4 ppm error

2.2.4.4 N,N-Bn,Me homotyrosine methyl ester
Homotyrosine methyl ester was dissolved in MeOH (−0.1M solution) NaOAc (1.1eq) was added, followed by addition of benzaldehyde (1.05eq). The mixture was stirred for 1h at RT, at which time NaBH₃(CN) (3eq) was added and the mixture stirred for 24h. Powdered paraformaldehyde (1.05eq) was added, the solution was allowed to stir for 4-5h at RT, then NaBH₃(CN) (3eq) was added and the reaction was stirred for an additional 24h.

The solvent was removed and the oily product was resuspended in 1N HCl. The aqueous layer was extracted 2 times with Et₂O. The aqueous layer was then adjusted to pH 9 with solid K₂CO₃ at 0°C followed by extraction with EtOAc (4X). The organic layers were combined and dried over MgSO₄, filtered and the solvent removed to yield a N,N-Bn,Me homotyrosine methyl ester. ¹H NMR (300 MHz, MeOH-d₄) δ 1.96 (m, 2H), 2.24 (s, 3H), 2.56 (dd J = 7.6, 7.7 Hz, 2H), 3.54 (dd, J = 9.81 Hz, 1H), 3.68 (s, 3H), 6.66 (d, J = 8.4 Hz, 2H), 6.94 (d, J = 8.4 Hz, 2H), 7.27 (m, 5H). L: [α]₂₅D −21.3 (c 0.79, MeOH), D: [α]₂₅D 14.4 (c 0.99, MeOH); ¹³C NMR (75 MHz, MeOH-d₄) δ 174.4, 156.4, 140.3, 133.5, 130.4, 129.9, 129.2, 128.0, 116.1, 65.6, 59.6, 51.5, 38.0, 32.6, 32.3; IR (film) 3425, 2951, 1728, 1643, 1514, 1452, 1247; HR-ESI C₁₉H₂₄NO₃ calcd. 314.1757, obs. 314.1759, 0.6 ppm error

2.2.4.5 N-Me homotyrosine methyl ester

N,N-Bn,Me homotyrosine methyl ester was suspended in EtOH acidified with HCl (0.06M solution). Pd(OH)₂ was added and the reaction was stored under hydrogen at atmospheric pressure for 18h. The reaction was filtered through Celite using EtOH acidified with HCl. The ethanol was removed and the solid was resuspended in dH₂O.
Solid $K_2CO_3$ was added to adjust the pH to 9 and then the aqueous layer was extracted 4 times with EtOAc. The organic layers were combined and dried over $MgSO_4$, filtered and the solvent removed to yield a $N$-Me homotyrosine methyl ester. $^1H$ NMR (300 MHz, MeOH-$d_4$) $\delta$ 1.90 (m, 2H), 2.54 (m, 2H), 3.17 (t, $J = 6.5$ Hz, 1H), 3.70 (s, 3H), 6.68 (d, $J = 8.6$ Hz, 2H), 6.99 (d, $J = 8.6$ Hz, 2H). $L$: $[\alpha]^{25}_D$ 21.8 (c 0.38, MeOH), $D$: $[\alpha]^{25}_D$ –18.4 (c 0.46, MeOH); $^{13}$C NMR (75 MHz, MeOH-$d_4$) $\delta$ 175.9, 156.7, 133.1, 130.4, 116.2, 63.4, 52.3, 35.8, 34.3, 31.9; IR (film) 3403, 1643, 1454, 1248; HR-ESI $C_{12}H_{18}NO_3$ calcd. 224.1287, obs. 224.1264, 10.2 ppm error

2.2.4.6 $N$-Me homotyrosine

$N$-Me homotyrosine methyl ester was dissolved in dry EtOAc and LiI (5 eq) was added. The reaction was refluxed for 20h in darkness. The reaction mixture was concentrated to dryness, resuspended in EtOAc and the solid was collected and washed with EtOAc. The solid was dissolved in $H_2O$, filtered and then the water was evaporated to yield $N$-Me homotyrosine. $^1H$ NMR (300 MHz, MeOH-$d_4$) $\delta$ 1.90 (m, 2H), 2.43 (m, 2H), 2.45 (s, 3H), 3.31 (t, $J = 6.1$ Hz), 6.68 (d, $J = 8.5$ Hz, 2H), 7.01 (d, $J = 8.5$, 2H). $L$: $[\alpha]^{25}_D$ 13.4 (c 0.36, 1N HCl), $D$: $[\alpha]^{25}_D$ –12.43 (c 0.17, 1N HCl); $^{13}$C NMR (75 MHz, MeOH-$d_4$) $\delta$ 170.9, 157.0, 131.8, 130.4, 116.4, 61.5, 32.6, 32.5, 30.9; IR (film) 3566, 1646; HR-ESI $C_{11}H_{16}NO_3$ calcd. 210.1130, obs. 210.1127, 1.4 ppm error
2.3 RESULTS AND DISCUSSION

2.3.1 Creation of CYA 126/8 ΔmcyT

McyT is a putative type II thioesterase. Enzymes of this class have been shown to be responsible for editing and proofreading during natural product biosynthesis. Walsh and co-workers have demonstrated that these Te domains are responsible for cleaning off the 4'-PP arm of thiolation domains by either removing wrongly activated amino acids or acetyl groups (incorporated by attack on acetyl-coenzyme A instead of co-enzyme A during phosphopantetheinylation).\textsuperscript{44} The lack of \textit{mcyI} in the genome of CYA 126/8 led us to hypothesize that \textit{mcyT} might be involved in the formation of Mdha with O-Ac-Ser as the activated intermediate. Given that previous experiments with \textit{mcyJ} knockouts produced (DMAdda, D-Asp\texttextsuperscript{3})-MC-RR, we hoped that insertional inactivation of \textit{mcyT} would produce (O-AcSer\texttextsuperscript{1}-D-Asp\texttextsuperscript{3})-MC-RR (12) or (L-Ser\texttextsuperscript{1}-D-Asp\texttextsuperscript{3})-MC-RR (13).

The construction of the recombination plasmid required special restriction conditions to only cleave at one of the \textit{Pfol} sites located in the \textit{mcyT} sequence. Previously attempted homologous recombination experiments with \textit{P. agardhii} revealed the need to have approximately 1.5kb regions flanking the chloramphenicol resistance cassette (G.}

\textsuperscript{12} R = Ac
\textsuperscript{13} R = H
Christiansen, personal communication). To ensure the appropriate size of the flanking regions we needed to insert the chloramphenicol resistance cassette at the 5'-end of the \textit{mcyT}. Upon \textit{in silico} analysis of the putative amplicon, we observed that the only site suitable for disruption of \textit{mcyT} corresponded to the recognition sequence of \textit{PfoI} (T↓CCNGGA). However another \textit{PfoI} site was located 750bp downstream of the desired insertion site. This created a problem because we could not cleave at both sites as this would reduce the flanking region and result in very low recombination efficiencies if any recombinants were created. An extensive literature search provided a solution to the problem, ethidium bromide. Ethidium bromide is a known DNA intercalation agent and studies by Parker \textit{et al} demonstrated that ethidium bromide could be used to stop restriction after linearization at one site. Ethidium bromide intercalates at a much higher rate into linearized (a double strand cut) and nicked (a single strand cut) plasmids than into the closed circular form that plasmids are usually found in.\textsuperscript{45} By optimizing the amount of ethidium bromide present in the reaction mixture (0.02 mg/ml), the amount of singly restricted plasmid could be maximized. After isolation of the linearized plasmid, the 5'-overhanging ends were blunted by use of a Klenow fragment and then ligated to the chloramphenicol (Cm) resistance cassette that had been restricted from the pACYC184 plasmid using \textit{BsaI} (YAC↓GTR). In order to ensure that the plasmids that were isolated had the Cm cassette in the correct \textit{PfoI} site, a colony PCR was performed. Out of the 20 individual colonies that were tested, only 1 clone contained the chloramphenicol cassette at the desired location.

The DNA was then introduced into the cyanobacterium by electroporation as linear single stranded DNA (See experimental section). After adding the chloramphenicol
to a final concentration of 1 μg/ml the culture became mostly bleached within 2 days and completely bleached within 1 week. After 5 weeks of growing under low light conditions, a faint green tint could be seen. The mutant was then grown to sufficient mass for DNA extraction and peptide analysis. In order to ensure that the desired mutant had been grown and not a random chloramphenicol resistant mutant, we checked the isolated mutant DNA by PCR. The mutant gave an amplicon the same size as the construct indicating that the desired homologous recombination had occurred (Figure 8).

2.3.2 Peptide analysis

Upon comparison of the peptide content of CYA 126/8 ΔmcyT to CYA 126/8 wild type, several differences could be observed. The wild type (2.1 g wet weight) yielded 40 mg of material from a C18 flash column eluted with 50% aqueous methanol and 40 mg in the fraction eluted with 75% methanol. The latter fraction was shown to contain the microcystins by NMR analysis. CYA 126/8 ΔmcyT (1.2 g wet weight), in contrast yielded 10 mg in the 50% aqueous methanol fraction but only 2 mg in the 75% aqueous methanol fraction. The characteristic peaks of the Adda side chain were absent from the crude NMR spectra of the 75% aqueous MeOH fraction (data not shown). This was corroborated by HPLC and LC-MS analysis as the extremely large peak due to (D-Asp$^3$)-
MC-RR had been severely reduced (See appendix), indicating that \textit{mcyT} is crucial for microcystin biosynthesis.

2.3.3 Anabaenopeptin I

![Anabaenopeptin I structure image]

Some cyanobacterial strains have been shown to upregulate or produce previous unknown peptides when a gene cluster is knocked out. Therefore the combined 50% and 75% aqueous methanol fractions were then analyzed to see if the same held true for CYA 126/8 \textit{}Δ\textit{mcyT}. In the absence of the huge (\textit{D-Asp}^3)-MC-RR peak characteristic of the wildtype, one uncharacterized peak stood out and was isolated pure after one HPLC column with an increased yield when compared to the wild type (0.59 mg per g dry mass vs. 0.33 mg per g dry mass). It should be noted that the increase in mass of anabaenopeptin I cannot be solely attributed the creation of the mutant and reduction of (\textit{D-Asp}^3)-MC-RR. It is uncertain whether this increase in mass is due to collection bias or represents a real increase in production by \textit{P. agardhii}. The first thing noticed in the \textit{1H}-NMR data (methanol-\textit{}d}_4) was the presence of 3 \textit{para} substituted benzene rings indicating
homotyrosine or tyrosine residues. Using $^1$H-NMR (methanol-$d_3$), HMQC and 1D TOCSY experiments, the compound was found to contain two homotyrosine, tyrosine, valine and an isoleucine residue with one of the residues containing an N-methyl group. It also contained either a lysine or an ornithine residue but the identity could not be determined using the $^1$H-NMR data alone. The presence of a lysine was confirmed by FABMS ($m/z$ $M+H^+ 916.4$) and upon review of the $^1$H NMR data it was seen that the resonances of two of the methylenes overlapped. HRESI-TOF gave a mass of 916.4727 ($M+H^+$, 0.17 ppm error), which corresponds to a molecular formula of $C_{48}H_{65}N_{7}O_{11}$.

When totaling the molecular formula for the known residues all but CHO$_2$ was represented. The $^{13}$C NMR contained a signal at 159.83, which hinted at the presence of an ureido bond leaving an OH unaccounted for. These pieces of information led us to believe that we had isolated a member of the anabaenopeptin family. Using 1D TOCSY and HMBC experiments, the 2D structure was established (Table 8). As in the case of all the known anabaenopeptins the lysine 2-NH and the alpha carbon (C2) show a correlation with the ureido carbonyl (C39). Similarly the NH of the tyrosine residue also showed a large correlation with C39. With the Lys and Tyr residues bound to the ureido carbonyl and due to the lack of HMBC correlation to the carbonyl of the Tyr it was determined that C48 was not involved in an amide bond and therefore must be a free carboxylic acid. This last piece of evidence accounted for the OH that had still to be accounted for in the molecular formula. Interestingly the $\delta_C$ of the carbonyl carbon of the tyrosine subunit could only be determined through an HMBC correlation from
Table 8: NMR data for Anabaenopeptin I (14)

<table>
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<tr>
<th>Unit</th>
<th>C/H number</th>
<th>δH (J in Hz)(^a)</th>
<th>δC (^b)</th>
<th>HMBC (^c,d)</th>
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<td>2</td>
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<tr>
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<td>3</td>
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<td>5</td>
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<td>3.54, m</td>
<td>40.38</td>
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<tr>
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<td>7</td>
<td>6-NH 7.31, dd (7.03, 3.02)</td>
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<td>8,13.</td>
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<td></td>
<td>11</td>
<td>1.10, m</td>
<td></td>
<td></td>
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<td>12</td>
<td>0.79, t (7.38)</td>
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<td>10</td>
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<td>N-Me-Hty</td>
<td>13</td>
<td>171.69</td>
<td></td>
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<td>14</td>
<td>4.52, t (6.82)</td>
<td>61.42</td>
<td>13,23,24</td>
<td></td>
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<td>2.12, m</td>
<td>34.85</td>
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<td></td>
<td>16</td>
<td>1.80, m</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>17</td>
<td>2.26, dt (-12.44, 5.02)</td>
<td>33.03</td>
<td>17</td>
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<td>2.32, dt (-12.48, 4.63)</td>
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<td>116.33</td>
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<td>31.94</td>
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<tr>
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<td>116.48</td>
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<td>31</td>
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<td>61.17</td>
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<td>20.07</td>
<td>35,36</td>
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<td>38</td>
<td>0.98, d (6.60)</td>
<td>19.76</td>
<td>35,36</td>
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<tr>
<td>Tyr</td>
<td>39</td>
<td>159.83</td>
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<td>40</td>
<td>4.45, m</td>
<td>56.6*</td>
<td>48</td>
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<td>6.06, d (8.03)</td>
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<td>42,44/46,45</td>
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<td>177.0**</td>
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<tr>
<td></td>
<td>48</td>
<td>177.0**</td>
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</table>

\(^a\) Recorded at 500 MHz \(^b\) Recorded at 125 MHz \(^c\) Protons showing long range correlation with indicated carbon \(^d\) Correlations were observed for \(J_{CH} = 7\) Hz \(\delta_C\) of this carbon could only be determined through HMQC \(\delta_C\) of this carbon could only be determined through HMBC
H40. The rest of the residues were placed in their respective positions using the HMBC correlations of the NH and alpha carbons to the carbonyl signals.

2.3.4 \textit{N-Me} homotyrosine synthesis

After determining that \textit{N-Me-Hty} was present in the molecule, a synthesis had to be undertaken to create the enantiomerically pure standards for configurational analysis (Figure 9). The first step was to resolve the \textit{DL}-homotyrosine mixture. This was accomplished by exposing N-Ac-Hty to Acylase I isolated from \textit{A. mellus} to remove the acetamide unit from the \textit{L} enantiomer. The resulting \textit{D} enhanced mixture was deacetylated by exposure to 1N HCl. The synthesis of each enantiomer was then done in parallel using standard methodology. The benzylation and methylation were performed in one pot using an adaptation of a recently developed protocol.\textsuperscript{46} Instead of exposing the Hty methyl ester•HCl salt to Amberlite IRA-410 resin to make the free base form, 1.1 eq of NaOAc was added to the Hty methyl ester•HCl in methanol prior to benzaldehyde addition to act as a proton scavenger. Creation of the free base was essential for the benzylation reaction to proceed as planned. Failure to do this resulted in incomplete benzylation, which in turn resulted in multiple products being isolated from the reaction. Powdered formaldehyde was used as suggested, however owing to the suspended NaOAc from the neutralization step, the dissolution of the formaldehyde could not be observed visually as described.\textsuperscript{46} Simultaneously in the lab the synthesis of \textit{D-N-Me-Phe} was being attempted. It was observed that exposure of \textit{D-N-Me-Phe-OMe} to LiOH in MeOH resulted in racemization. It was assumed that the same problem would be encountered with \textit{N-Me-Hty-OMe}. A literature search revealed that a few research groups had
encountered similar problems with protected amino acids. Therefore the removal of the methyl ester was performed using LiI in refluxing EtOAc. The standards were then used for stereochemical analysis without further purification.

![Synthesis of enantiomerically enriched N-Me homotyrosine](image)

**Figure 9:** Synthesis of enantiomerically enriched N-Me homotyrosine

(i) Ac₂O, AcOH, 110°C, 16h; (ii) Acylase 1, 0.1M K₂HPO₄ buffer (pH 7.0); (iii) 1N HCl, 100°C; (iv) TMS-Cl, MeOH; (v) (a) NaOAc (1.1 eq), C₂H₅CHO (1.05 eq), MeOH, 1h then NaBH₄(CN) (3 eq), 24h (b) CH₂O (1.05 eq) 4.5h then NaBH₄(CN) (3 eq), 24h; (vi) H₂, Pd(OH)₄, EtOH + HCl, O/N; (vii) LiI, EtOAc reflux, O/N

### 2.3.5 Stereochemistry of Anabaenopeptin I (14)

The stereochemistry of the amino acid residues was determined by advanced Marfey analysis using FDAA as the derivatization reagent and observed with an LC-MS equipped with an ESI source either in positive or negative mode. This setup greatly reduced the work required in identification because it removed the need for co-injection of standards. After hydrolysis of 14 with 6N HCl, D-Lys, L-Val were determined to be present as well as either L-Ile or L-\textit{allo}-Ile. The Ile species could not be resolved using a non-chiral column. Use of a chiral column later revealed that L-Ile was present in the molecule (Cyril Portman, personal communication). The analysis of the stereochemistry
of the Tyr, Hty and N-Me-Hty residues was extremely problematic, as analysis of the hydrolysate did not show any traces of the phenolic residues. Itou et al resorted to use of anhydrous hydrazine in order to determine the stereochemistry of the Tyr side chain in anabaenopeptin H. However making the required reagent is time consuming and the by-products toxic, which makes the approach unappealing. Therefore it was decided to scan less harsh hydrolysis conditions to see if the Tyr and Hty residues could be observed. Degradation with 2N HCl at 55°C for 2h surprisingly was harsh enough to liberate the Tyr residue from the ureido bond. Detection of the Marfey derivatized residue using negative ESI-TOF allowed the Tyr to be assigned the L configuration. However, the Hty and N-Me-Hty residues were still absent from the spectrum. The configurations of the Hty residues is proposed on the basis of the similarity of the NMR spectra of anabaenopeptin I with those of another anabaenopeptin (15) isolated from wild type CYA 126/8, which differed from 14 simply by replacement of the Tyr residue with an Arg residue. Anabaenopeptin J (0.1mg) was degraded in 6N HCl for 12 at 110°C, followed by derivatization with 3eq of Marfey's reagent. The products were then observed using LC-MS in negative ESI mode. Both the Hty and the N-Me-Hty residues coeluted with the L enantiomer standards (Cyril Portman, personal communication)
2.4 CONCLUSION

Insertional mutagenesis was utilized to create a knockout mutant of mcyT from *P. agardhii* CYA 126/8. This was done to investigate the role of *mcyT* in microcystin biosynthesis. The generated mutant did not produce a structural variant of MC-RR as expected and microcystin biosynthesis reduced to a level far below that of the wild type (<5%) was observed instead. This indicates that *mcyT* is essential for microcystin biosynthesis, even if its precise function remains a mystery. If McyT did function as a general type II thioesterase, one would expect that upon disruption all non-ribosomal peptide production would be reduced. However this is not observed. One explanation for why only microcystin biosynthesis is affected is that each NRPS cluster may have its own dedicated type II thioesterase. Therefore, a knockout of *mcyT* would not necessarily lead to disruption of peptide biosynthesis by other clusters. In this connection it is interesting to note that recent studies have elucidated the fact that a population of *P. agardhii* contains individuals carrying defective *mcy* genes, which result from insertions or deletions of gene fragments. All of the resulting mutants are deficient in microcystin production. It is noteworthy, however, that to this date no strain has yet been found containing a defective variant of *mcyT* as ascertained by PCR (G. Christiansen, personal communication). This may suggest that several closely related type II thioesterases are present which could result in false positives in the PCR screen. Unfortunately, at this time too little genomic information exists about *P. agardhii*, or other cyanobacteria with a well-studied spectrum of secondary metabolites, to accurately assess the role and specificity McyT and type II thioesterases in general.
Upon peptide analysis of *P. agardhii* CYA 126/8 ΔmcyT an uncharacterized peak was isolated and the structure determined. The 2D structure of anabaenopeptin I (14) was elucidated using NMR and HR-FABMS techniques. The stereochemistry of the residues was determined by advanced Marfey analysis. The derivatized residues were observed using LC-MS, which greatly reduced the workload associated with identifying the enantiomer. The phenolic residues (Hty, Tyr) proved problematic to identify and were therefore identified through $^{1}$H NMR comparison to a closely related compound, anabaenopeptin J (15). The similarities between the proton resonances of the residues in the macrolactam ring residues and the fact that the Lys, Ile and Val residues were of identical configuration in each compound led us to believe that the stereochemistry of the Hty and Tyr residues would also be identical.
HPLC trace of (a) *P. agardhii* CYA 126/8 wild type and (b) *P. agardhii* CYA 126/8 ΔmcyT. HPLC was performed as described in section 2.2.2.2 and observed at 240 nm.
125 MHz 13C NMR spectrum of Ambabenopeptin I (14) in MeOD-4
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media was autoclaved for 20 min at 15 lbs./sq. in.


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