Title: Identification, heterologous expression, and characterization of the tolypodiol biosynthetic gene cluster through an integrated approach.

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Abstract

Cyanobacteria are tremendous producers of biologically active natural products including the potent anti-inflammatory compound tolypodiol. However, linking biosynthetic gene clusters with compound production in cyanobacteria has lagged behind other bacterial genera. Tolypodiol is a meroterpenoid originally isolated from the cyanobacterium HT-58-2. Here we describe the identification of the tolypodiol biosynthetic gene cluster through heterologous expression in *Anabaena*, and in vitro protein assays of a methyltransferase found in the tolypodiol biosynthetic gene cluster. We have also identified similar biosynthetic gene clusters in cyanobacterial and Actinobacterial genomes suggesting a widespread ecological role for tolypodiol and related compounds. We also report the identification of two new analogs of tolypodiol that we have identified in both the original and heterologous producer. This work further illustrates the usefulness of *Anabaena* as a heterologous expression host for cyanobacterial compounds and how integrated approaches can help link natural product compounds with their producing biosynthetic gene clusters.
Introduction

Cyanobacteria, which are Gram-negative photosynthetic bacteria, are rich sources of specialized secondary metabolites (aka natural products) with a wide range of biological activities including anti-viral, anti-bacterial, and anti-cancer. Peptides (linear and cyclic), depsipeptides, polyketides, polyketide-(depsi)peptide hybrids, meroterpenoids, and terpenes have all been isolated from cyanobacteria. To date, few cyanobacterial compounds have been assessed in pre-clinical trials, but two structural classes have entered clinical trials (cryptophycins and dolastatins), and brentuximab, an antibody drug conjugate containing monomethyl auristatin E (dolastatin analog), has been approved by the FDA.

The cyanobacterial strain HT-58-2, originally classified as a Tolypothrix sp, but subsequently shown to align more closely with Brasilonema sp., is known to produce two unique classes of natural products, the tetrapyrole macrocyclic tolyporphins (e.g., tolyporphin A, 1) and the meroterpenoid tolypodiol (2). More recently, we reported the isolation, structure determination, and biological activity of 6-deoxy-tolypodiol (3) and 11-hydroxytolypodiol (4) from the extract of Brasilonema sp. HT-58-2. Tolypodiol was reported to have potent anti-inflammatory activity in a mouse ear edema assay, with the reported activity similar to hydrocortisone.

Here, we report a biosynthetic gene cluster (BGC) for the production of tolypodiol, successful heterologous expression of the tolypodiol (tyl) BGC in Anabaena sp. PCC 7120, and propose a biosynthetic pathway. We also identified similar BGCs in both cyanobacterial and actinobacterial genomes suggesting that these molecules play an undefined role in the life of the producing organisms. Furthermore, we demonstrate that the methyltransferase, TylH, is responsible for converting a carboxylic acid to a methyl ester, most likely as one of the final steps in the tolypodiol biosynthetic pathway. Lastly, we report two new analogs of tolypodiol, 6-oxo-tolypodiol (5) and 1-oxo-tolypodiol (6) that were isolated from biomass of HT-58-2 after extraction of its lyophilized material as previously reported. The structures of 5 and 6 were determined largely by spectroscopic and spectrometric experiments such as LC-MS and 1D and 2D NMR.

MATERIALS AND METHODS

General. Optical rotation measurements were taken on a JASCO DIP-370 Digital Polarimeter at the sodium line (589 nm) at 22°C. ECD and UV-VIS measurements for 5 and 6 were taken on a Chirascan Circular Dichroism Spectrometer with the sample dissolved in methanol and placed in
a 1 cm quartz cuvette with a solvent subtraction for baseline correction. IR spectra were recorded on a CaF$_2$ salt plate using a Shimadzu IRAffinity-1 Fourier-Transform Infrared Spectrophotometer. All NMR spectra for initial structural characterization of tolypodiol analogs (5 and 6) were acquired on an Agilent Technologies 600 MHz DD2 spectrometer operating at 600 MHz for $^1$H and 150 MHz for $^{13}$C NMR or on a Varian Unity Inova 500 MHz spectrometer operating at 500 MHz for $^1$H or 125 MHz for $^{13}$C. The appropriate residual solvent signal was used as an internal reference. All NMR samples were placed in 3 mm Shigemi tubes for analysis. NMR spectra for tolypodiol and its hydrolysis products were recorded on a Bruker 700 MHz spectrometer with a 5 mm carbon cryo-probe with z-axis gradient. $^1$H-NMR and $^{13}$C-NMR chemical shifts (δ) are expressed in ppm values. CDCl$_3$ was used as the NMR solvent (δ$_H$ = 7.26 ppm, δ$_C$ = 77 ppm). HPLC-HRESIMS was conducted on one of two instruments. System 1, Agilent 6545 LC-MS Q-Tof with ESI ionization was used in the positive mode to acquire high-resolution mass spectrometry (HRMS) data with a C18 Agilent Eclipse Plus (2.1 x 50 mm, 1.8 µm) column (Method A); System 2, an Agilent 6545 Q-ToF equipped with a Jet Stream ion source downstream of an Agilent 1260 Infinity HPLC system consisting of a degasser, quaternary pump, autosampler (held at 12°C), and column compartment (held at 40 °C). Separation was achieved with a Kinetex C18 column (2.1 x 50 mm, 2.6 µm, Phenomenex, Torrance, CA) (Method B). The Q-ToF machines were operated using MassHunter software and data was processed offline using MassHunter Qualitative software. HPLC separations were performed using a Shimadzu system consisting of LC-20AB binary high-pressure gradient solvent delivery unit, SPD-M20A Photodiode Array Detector, and a CBM-20A System Controller. Enzymatic assays were analyzed by HPLC using an Agilent 1100 HPLC instrument consisting of a vacuum degasser, quaternary pump, autosampler (cooled to 4 °C), column thermostat (maintained at 30 °C), and diode array detector.

All chemicals and solvents were purchased either from Sigma Aldrich, Alfa Aesar or Tokyo Chemical Industry Co., Ltd. (TCI). SAM was obtained as S-adenosyl-L-methionine disulfate tosylate (Sigma-Aldrich, St. Louis, MO) and used without purification. NdeI, XhoI, T4 DNA ligase, Phusion DNA polymerase, Escherichia coli NEB5α, and Escherichia coli BL21(DE3)-TIR were purchased from New England Biolabs (Ipswich, MA) and used per the manufacturer’s recommendations. Lysogeny broth, Miller (LB) was purchased from EMD Millipore (Billerica, MA). GenCatch Plasmid DNA Mini-Prep Kit and EconoSpin® All-In-One Mini Spin Columns were purchased from Epoch Life Sciences (Sugar Land, TX). PrimeSTAR GXL DNA polymerase was purchased from Takara Bio (Mountain View, CA) and oligonucleotides were purchased from Integrated DNA technologies with standard desalting (Coralville, IA). The UV absorptions were measured on an Eppendorf BioSpectrometer Kinetic and the cell used was Eppendorf µCuvette G1.0 (Eppendorf, New Haven, CT). All microbial procedures were performed using good microbial practices and aseptic technique. Kanamycin (50 µg/mL) and spectinomycin (100 µg/mL) for selection of plasmids in E. coli, while spectinomycin and streptomycin (2.5 µg/mL each) were used for the selection of Anabaena sp. UTEX 2576.

Strain source. Anabaena sp. UTEX 2576 was acquired from the UTEX Culture Collection of Algae, University of Texas at Austin (Austin, TX, 78705). Brasilonema sp. HT-58-2 cultures were revived from cryostorage maintained by the University of Hawaii Cyanobacterial Culture Collection. Escherichia coli strains NEB10β and BL21(DE3) were obtained from New England Biolabs.

Cyanobacterial of Anabaena sp. UTEX 2576. Anabaena sp. UTEX 2576 was cultivated in a Hoffman incubation chamber (Hoffman Manufacturing, Inc, Corvallis, OR, USA) at 28 °C under
24 h constant light illumination. The light was supplied by two GE Lighting 49893 F40/PL/AQ Plant and Aquarium Tube bulbs with a light intensity of 25 microeinsteins/m2/s, and the atmosphere contained 1% CO₂.

_Cultivation and extraction of the cyanobacterium HT-58-2._ Cultures were grown in BG-11(Nitrogen+) media. Over time, they were scaled up to 20 L Pyrex carboys with aeration at a 4-5 L/min flow rate and under continuous fluorescent illumination. The BG-11(Nit+) formulation was composed of the following: NaNO₂ (1.5 g/L), K₂HPO₄·3H₂O (0.04 g/L), MgSO₄·7H₂O (0.075 g/L), Na₂CO₃ (0.02 g/L), CaCl₂·2H₂O (0.036 g/L), Na₂EDTA (0.001 g/L), ferric ammonium citrate (0.006 g/L) and citric acid (0.006 g/L) with trace metals H₂BO₃ (2.86 mg/L), MnCl₂·4H₂O (1.81 mg/L), ZnSO₄·7H₂O (0.22 mg/L), Na₂MoO₄·2H₂O (0.39 mg/L), CuSO₄·5H₂O (0.08 mg/L), Co(NO₃)₂·6H₂O (0.05 mg/L). The phylogenetic characterization and genomic sequencing of this strain has been reported previously.¹³ Cell material was harvested by filtration through the equivalent of a cheese cloth after 45 days of growth in 20 L carboys and lyophilized prior to extraction. The freeze-dried HT-58-2 (43.35 g) cell mass was exhaustedly extracted in 1:1 CH₂Cl₂:i-PrOH to produce the crude extract (2.33 g) after solvent evaporation.

The HT-58-2 strain was originally described in 1992,⁹ but was later subjected to 16S rRNA analysis revealing that it was more aligned with strains of _Brasilonema_,¹³ a genus formally described in 2007,¹⁴ rather than Tolypothrix as originally identified from morphological features. The strain has been maintained in cryostorage at University of Hawaii since the original report,¹¹ periodically removing cryovials to grow the strain.

_Isolation of 6-oxo-tolypodiol (5) and 1-oxo-tolypodiol (6)._ A modified Kupchan partition was used to separate crude into hexanes, dichloromethane, and aqueous methanolic fractions. The dichloromethane fraction yielded the new compounds after C8 silica gel fractionation using a methanol in water gradient (25%, 50%, 75%, 100% MeOH). Along with tolypodiol, the 75% MeOH fraction yielded both 6-oxo-tolypodiol (5) (0.8 mg) and 1-oxo-tolypodiol (6) (0.5 mg) after preparative HPLC (Phenomenex Gemini 5u C18 110A Axia Pac, 250 x 21.20mm) purification using acetonitrile and water with 0.1% formic acid in a linear gradient from 70-100% acetonitrile over 30 min followed by 100% acetonitrile for 15 min. The fractions collected from time 12-14 mins were combined and further purified on a semi-preparative scale (Phenomenex Luna PFP(2) 100A, 250 x 10 mm) with an 80%-100% methanol:water (0.1% formic acid) gradient over 30 min (5, tᵣ 18.5 min; 6, tᵣ 24.0 min) with a purity of 93% and 90%, respectively as estimated by ¹H NMR.

_Assembly and cloning of 12 large meroterpenoid biosynthesis gene clusters (BGCs)._ The meroterpenoid BGC sequence originating from _Brasilonema_ sp. HT-58-2 was codon optimized using the Build OptimizatiOn Software Tools (BOOST)¹⁵ and a _Nostoc_ sp. PCC 7120 codon frequency table to remove all synthesis constraints (refactored sequences are listed in Supplementary Table 1 as Batch202D_p002). Overlapping synthetic DNA fragments were obtained from Twist Bioscience and assembled into the pPJAV550 destination vector (Batch202D_p009 and Batch202D_p010) using yeast Transformation-Associated Recombination (TAR) cloning.¹⁷ Vector sequences are available in Supplementary Table 2. All yeast transformation was performed with an auxotrophic strain CEN.PK2 (Euroscarf, Germany) using the YEAST1 yeast transformation kit (Sigma-Aldrich) and the manufacturer’s protocol. Yeast colonies were then collected from the single-dropout selective plates and plasmid DNA were harvested and transformed into _E. coli_ DH10B cells. The resulting colonies were sequence verified by the PacBio sequencing platform.
During the refactoring of the BGCs, all internal SrfI restriction sites were removed and a unique SrfI site was reintroduced at the 5’ ends of each BGCs. These SrfI sites were then used to clone two promoters by using the NEBuilder HiFi DNA Assembly kit (New England BioLabs) to generate a total of 12 BGC constructs (Batch202D_p007 and p018 in Table S1). Again, all construct sequences were verified by PacBio sequencing.

**Heterologous expression of tolypodiol in Anabaena sp. UTEX 2576.** The plasmids pMero_HT583 and pMero_HT583_PpetE were introduced into Anabaena sp. UTEX 2576 through triparental mating with *E. coli* TOP10 cells containing either pMero_HT583 and pMero_HT583_PpetE and *E. coli* JCM113 according to our previously published protocol. The exoconjugates were selected on BG11(N+) media supplemented with spectinomycin and streptomycin (2.5 µg/mL each) for 4 weeks at 28°C with 24-hour illumination in a 1% CO2 atmosphere. Single colonies were restreaked onto BG11(N+) media supplemented with spectinomycin and streptomycin. The colonies were then cultivated for an additional 5 weeks followed by collection of the cells by scraping them from the plate. The cells were added to a tared 20 mL scintillation vial, lyophilized, and then extracted with 20 mL ethyl acetate overnight with stirring provided by a magnetic stir bar. The cellular debris were removed via filtration through a Buchner funnel and the extract was concentrated to near dryness in vacuo using a 30 °C water bath. The extract was transferred to a 11 mL glass vial and concentrated to dryness in a miVac Quattro Concentrator (ATS Life Sciences Scientific Products, Warminster, PA). The extract was redissolved in HPLC grade methanol at a concentration of 1 mg/mL and then diluted to a concentration of 0.2 mg/mL for LCMS analysis.

**LCMS analysis of tolypodiol.** **Method A.** An Agilent 6545 LC-MS Q-Tof with ESI ionization was used in the positive mode to acquire high-resolution mass spectrometry (HRMS) data with a C18 Agilent Eclipse Plus (2.1 x 50 mm, 1.8 µm) column. A flow rate of 0.5 mL/min with a water (A)-acetonitrile (B) mobile phase system containing 0.1% (v/v) formic acid was used. The column was pre-equilibrated with 95% A/5% B. Upon injection the mobile phase composition was maintained for 0.75 minutes followed by changing the mobile phase to 0% A/100% B over 9.25 minutes using a linear gradient and 100% B was maintained for an additional 7 min.

The following parameters were used with positive polarity; Gas Temp, 350 °C; Drying gas, 10 L/min; Nebulizer, 55 psi; Sheath gas temp, 400 °C; Sheath gas flow, 10 L/min; VCap, 3500 V; Nozzle voltage (Expt), 1000 V; Fragmentor, 175 V; Skimmer, 65 V; Oct 1 RF Vpp, 750 V; Mass range, 100-3000 m/z; Acquisition rate, 10 spectra/s; Time, 100 ms/spectrum. The AutoMS/MS settings were as follows: Mass range, 100-3000 m/z; 1 spectra/s; 1000 ms/spectrum; Transients/spectrum, 5889.

**Method B.** High-resolution mass spectrometry was obtained using an Agilent 1260 HPLC upstream of an Agilent 6545 Q-ToF downstream of an Agilent 1260 Infinity HPLC system. Separation was achieved using Kinetex C18 column (50 x 2.1 mm, 2.6 µm, Phenomenex, Torrance, CA) at a flow rate of 0.4 mL/min and the following gradient. Line A was water with 0.1% (v/v) formic acid and line B was acetonitrile with 0.1% (v/v) formic acid. The column was pre-equilibrated with 90% A/10% B. Upon injection the mobile phase composition was maintained for 0.75 minutes followed by changing the mobile phase to 10% A/90% B over 7.5 minutes using a linear gradient and then switching the mobile phase to 0% A/100%B over 0.1 minutes. The mobile phase is then maintained at 0%A / 100%B for 3.65 minutes.

The Agilent Q-ToF mass spectrometer was equipped with an Agilent JetSpray source operated with the following parameters: Auto MS/MS mode, Positive polarity; Gas Temp, 325 °C; Drying gas, 7 L/min; Nebulizer, 20 psi; Sheath gas temp, 270 °C; Sheath gas flow, 12 L/min; VCap, 4000
V; Nozzle voltage (Expt), 600 V; Fragmentor, 175 V; Skimmer, 65 V; Oct 1 RF Vpp, 750 V; Mass range, 100-3000 m/z; Acquisition rate, 10 spectra/s; Time, 100 ms/spectrum. The AutoMS/MS settings were as follows: Mass range, 50-3000 m/z; 10 spectra/s; 100 ms/spectrum; Transients/spectrum, 429; Isolation Width, Medium (~4 m/z); The Precursor selection parameters were as follows: 5 Max Precursor Per Cycle; Abs. Threshold 1000 counts, Rel. Threshold (%) 0.01%, Excluded after 4 spectra; Released after 0.01 min.

Computational docking and MD simulations of TylF. The AlphaFold predicted structure of TylF was downloaded from the AlphaFold Protein Structure Database (Locus BZZ01_00205). Next, the protein’s orientation in the membrane was calculated using the PPM 2.0 web server. Ligands were docked into the protein using AutoDock Vina in UCSF Chimera. Ligand binding was restricted to within the inside of the protein near the hypothetical active site, and conformations were picked which placed the conserved residues in proximity to the predicted site of initialization of cyclization in the ligand.

Molecular dynamics (MD) simulations of TylF embedded in a lipid bilayer were carried out with the GROMACS simulation package v2022 using the CHARMM36 force-field (FF) for lipids and the CHARMM36m FF for proteins. Water was simulated using CHARMM TIP3 parameters. The glutamate residue in the predicted active site (E51) was modified to be protonated, and ligands with a carboxylic acid were specified to be deprotonated. The membranes were composed of POPG, DGDG, MGDG, SQDG to roughly match the experimental values measured for cyanobacteria. The upper and lower membrane leaflets were symmetrical and contained 4 POPG lipids, 10 DGDG lipids, 18 MGDG lipids, and 10 SQDG lipids. Potassium and chloride ions were added to a concentration of 150 mM KCl. The combined protein-membrane system was set up using the CHARMM-GUI webserver. Two different simulations were setup with the 3-geranylgeranyl-4-hydroxybenzoic acid and compound 10 bound in the active site. FF parameters for the two substrates were generated using the CHARMM-GUI interface and CGenFF. Systems were simulated for 100 ns after initial energy minimization and equilibration using position restraints. Equations of motion were integrated with a leap-frog algorithm using a 2-fs time step. Van der Waals interactions were computed using a force-switched Lennard-Jones potential between 1.0 and 1.2 nm. Electrostatic interactions were computed using the particle-mesh Ewald method with a real space cutoff 1.2 nm and a Fourier grid spacing of 0.12 nm. Temperature was held constant using a velocity-rescaling algorithm with a time constant of 1 ps, and pressure was held constant with a stochastic cell-rescaling algorithm using a time constant of 5 ps. Particle positions were saved in 5 ps intervals for trajectory analysis.

Synthesis of desmethyltolypodiol (12). In order to hydrolyze the methyl ester of tolypodiol, a protocol using lithium hydroxide in aqueous methanol was adapted. In brief, approximately 1 mg of tolypodiol was dissolved in LCMS grade MeOH (0.7 mL) and H2O (0.2 mL) in a 5 mL round bottom flask to which a few drops of 0.1 N LiOH dissolved in LCMS grade water was added. The reaction was placed in a 40 °C water bath and stirred overnight while connected to a reflux condenser. The reaction was quenched by acidification with a few drops of 0.1 N HCl. This reaction was done twice to secure sufficient material. The reaction mixtures were combined, dried down under a gentle stream of air, reconstituted in a minimal amount of LCMS methanol/water (8:2), and then purified by HPLC (Phenomenex Luna 5u C18(2) 150 x 4.6 mm; A: H2O, B: MeOH (0.1% formic acid in both); gradient of 70% B (0-2 min), 70-100% B (2-17 min), 100% B (17-22 min); flow rate = 0.6 mL/min) to yield 0.7 mg of the hydrolyzed product and 0.6 mg of recovered
tolypodiol. For NMR data, see Supplementary Figures 1-2 and Supplementary Table 3. HRESIMS m/z 443.2791 [M+H]+ (calc. for C_{27}H_{39}O_{5}^+, 443.2792, 0.2 ppm error)

*Creation of expression vector for overproduction of TylH.* The DNA fragment corresponding to TylH (protein accession number ARV57258) was amplified from *Brasilomena* sp. strain HT-58-3 genomic DNA by PCR using PrimeSTAR GXL DNA polymerase and the oligonucleotides TylH_Fwd-Ndel (5'-ATCTCTAAGCATATGTCTAAATGGAATGCACAGATTTACC-3') and TylH_Rev-XhoI (5'-ATATAGACTCGAGAGACCCGACGTATGAGTGAA-3') with an annealing temperature of 58 °C using an extension time of 2:00 min and 35 cycles. The restriction sites of Ndel and XhoI are indicated by the underlined bases while the italicized bases indicate random bases added to increase restriction enzyme efficiency. The PCR ampiclon was purified using the EconoSpin® All-In-One Mini Spin Column followed by digestion with Ndel and XhoI. The restricted fragment was purified using the EconoSpin® All-In-One Mini Spin Column followed by ligation into pET28a, which had been previously digested with Ndel and XhoI and purified using the EconoSpin® All-In-One Mini Spin Column using T4 DNA ligase. The ligation mixture was then introduced in chemically competent *E. coli* NEB3α cells. Plasmid containing cells were then selected for on LB agar plates supplemented with kanamycin after overnight incubation at 37 °C. Random colonies were selected and the plasmids isolated with the GenCatch Plasmid DNA Mini-Prep Kit. The DNA inserts were sequenced with Sanger sequencing at the Center for Qualitative Life Sciences at Oregon State University to ensure insert sequence fidelity. The resulting sequence verified plasmid was named pET28-TylH-1.

*Overexpression, and purification of TylH.* Chemically competent *E. coli* strain BL21(DE3) cells were transformed with pET28-TylH-1 and the transformation mixture was plated on LB agar plates supplemented with kanamycin. A single colony of *E. coli* BL21(DE3) cells containing pET28-TylH-1 was picked and grown overnight at 37 °C in LB media containing kanamycin (25 mL) and cultured in a shaker at 200 rpm at 37 °C overnight. A portion of the seed culture (5 mL) was transferred into LB medium (2 x 500 mL) in 2 x 2 L Erlenmeyer flasks containing kanamycin and grown at 37 °C until the OD_{600} reached 0.6 (approximately 3.5 hr). Then, the temperature was reduced to 15 °C and after 1 h adaptation, Isopropyl β-D-1-thiogalactopyranoside (IPTG, 0.1 mM final concentration) was added to each flask. After further growth overnight at 200 rpm (approximately 16 hours), the cells were harvested by centrifugation (3,200 x g, 20 min, 4 °C), the media was decanted, and the cell pellet was stored at -20 °C until use.

The frozen cells from 1 L of culture were thawed on ice and then suspended in 30 mL lysis buffer [Tris (50 mM, pH 8), NaCl (300 mM), imidazole (10 mM)]. The suspension was sonicated using a Qsonica sonicator (9 mm probe, 30% amplitude, 45 sec, five times) on ice, then cell debris were removed by centrifugation at (13,751 x g, 4 °C for 30 min). The supernatant was applied to a Ni-NTA resin (750 μL, BioRad, Hercules, CA) by batch binding at 4 °C for 1 h followed by pelleting the resin by centrifugation at 3,200 x g for 30 sec. The supernatant was discarded, and the resin was washed with lysis buffer (2.5 mL) followed by washing two times with 1 mL Wash buffer [Tris (50 mM, pH 8), NaCl (300 mM), imidazole (25 mM), β-mercaptoethanol (5 mM)]. The resin was then washed twice with 1.5 mL Elution buffer [Tris-HCl (50 mM, pH 8), NaCl (300 mM), imidazole (250 mM), β-mercaptoethanol (5 mM)] followed by another wash with elution buffer (0.5 mL). The fractions from the washing buffer contained the highest concentration of TylH (determined by Bradford stain and SDS-PAGE, Supplementary Figure 3). The protein solution was desalted using an Econo-Pac 10DG Desalting column (BioRad) equilibrated with desalting buffer [Tris (50 mM, pH 8), glycerol (10% w/v), NaCl (50 mM), β-mercaptoethanol (5
The protein concentration of the fractions collected from the washing buffer was determined using an Eppendorf BioSpectrometer at (280 nm, 1 mm cell) using a calculated extinction coefficient of 29,910 cm$^{-1}$ M$^{-1}$.

**In vitro enzymatic activity assays.** Reaction mixtures containing 100 mM Tris buffer (pH 7.5), 5 mM SAM dissolved in DI water (stock: 10 mg/ml), 1 mM substrate dissolved in MeOH (stock: 10 mg/mL), 10 µM TylH (stock: 0.7 mg/mL in desalting buffer), and 1mM β-mercaptoethanol in a final volume of 100 µL were mixed gently by pipetting and incubated at 28 °C overnight. The protein was removed by the addition of acetonitrile (50 µL) followed by centrifugation (15,871 x g) for 3 min. All negative control reactions were performed in the same manner by replacing the omitted reagent with the respective solvent. A portion of the quenched reaction (15 µL) was then analyzed by HPLC as described below.

**HPLC analysis of in vitro TylH reaction mixtures.** All enzymatic reactions were analyzed on an Agilent 1100 HPLC instrument consisting of a vacuum degasser, quaternary pump, autosampler (cooled to 4 °C), column thermostat (maintained at 30 °C), and diode array detector using a Luna C18 column (15 cm x 4.6 mm, 5 µm, Phenomenex, Torrance, CA) where line C was water + 0.1 % (v/v) formic acid and line D was acetonitrile with 0.1 % (v/v) formic acid at a flow rate of 1 mL/min. The column was pre-equilibrated with 90% C / 10% D and held at this composition for 1 min. The composition was changed to 100% D over the next 20 min using a linear gradient. The composition was then held at 100% D and this composition was held for 5 min. The mobile phase was then changed to 90% C/10% D over 8 min and the composition was held for 8 min prior to the next injection.

**Results and Discussion:**

**Identification of the tolypodiol biosynthetic gene cluster (BGC) and similar BGCs in bacteria.**

A retrobiosynthetic analysis of tolypodiol predicted it to be biosynthesized from p-hydroxybenzoic acid and geranylgeranyl diphosphate as key building blocks. Given the structural similarities between merosterol A and tolypodiol, we first attempted to search the published genome of HT-58-2 by BLAST analysis using MstE as the probe, given its role as a terpene cyclase in the biosynthesis of the merosterol A. While a similar protein was identified, BZZ01_14670, it did not appear to be in proximity to other encoded proteins that were predicted to be involved in the biosynthesis of tolypodiol. In bacteria, p-hydroxybenzoic acid is known to be synthesized by UbiC, therefore, we used All0938 (UbiC from *Anabaena* sp. PCC 7120) as the BLAST probe sequence and identified two proteins with significant similarity in the genome of HT-58-2, BZZ01_27380 (87% identity/92% similarity) and BZZ01_00185 (73% identity/81% similarity). Examination of the genomic region near BZZ01_00185 showed the presence of encoded proteins that have the predicted activities necessary for the biosynthesis of tolypodiol (Figure 1, Supplementary Table 4). This same genomic region was recently postulated to be involved in the production of the tolypodiolides by the Lindsey group. We subsequently renamed BZZ01_00185 as tylB and it is predicted to encode a chorismate lyase, which is responsible for converting chorismate to p-hydroxybenzoic acid. TylC, TylD, TylII, TylK, and TylM have similarity to enzymes involved in the formation of the isoprenoid precursors dimethylallyl diphosphate (DMAP) and isopentenyl diphosphate (IPP). TylE belongs to the polyprenyl synthase family and is presumably involved in the biosynthesis of geranylgeranyl diphosphate (GGPP). TylA is predicted to be 4-hydroxybenzoate solanesyltransferase, which is likely responsible for the transfer
of the geranylgeranyl unit to the 3-position of 4-hydroxybenzoate. TylG has similarity to aromatic ring hydroxylases and is likely responsible for introduction of one of the hydroxyl groups while TylJ (annotated as a cytochrome P450) is likely responsible for the installation of the second hydroxyl group. TylH is predicted to be a SAM dependent methyltransferase, which is envisioned to methylate a carboxylic acid precursor to form the methyl ester seen in the tolypodiols. TylL is annotated as a member of the AAA ATPase protein family\(^{41}\) and could be a regulator of the tolypodiol BGC expression or responsible for the transport of tolypodiol through membranes. No other transport proteins were identified in the vicinity of the tyl BGC. TylF is annotated as a hypothetical protein but is presumed to be the terpene cyclase (see subsequent section).

**Figure 1.** A) Predicted biosynthetic gene cluster for the biosynthesis of tolypodiols identified in the genome of Brasilonema sp. HT-58-2. B) Predicted biosynthetic pathway for the formation of tolypodiol in HT-58-2.

*TyIF is the founding member of a new family of bacterial terpene cyclases with similarity to fungal terpene cyclases.*

Using TylF as a query sequence for a BLAST analysis against the Swiss-Prot/UniProt database retrieves terpene cyclase proteins from the BGCs involved in the biosynthesis of a variety of fungal natural products (Supplementary Table 5). The top hit was DpmpB, which is involved in the biosynthesis of decalin-containing diterpenoid pyrones in *Macrophomina phaseolina* MS6\(^{42}\) (28%
amino acid similarity/50% amino acid identity). Just outside of the top 100 sequences returned from the BLAST query was Trt1 (25% amino acid similarity/47% amino acid identity), which has been characterized as a terpene cyclase in the terretonin biosynthetic pathway. Trt1 was previously shown to cyclize 5-farnesyl-3,5-dimethylorsellinic acid methyl ester and not 5-farnesyl-3,5-dimethylorsellinic acid.43

TyIF does not contain an aspartate rich motif found in Class I (DDxDD) or Class II (DxDD) terpene cyclases.44 Multiple sequence alignment of the proteins identified by BLAST identified only one conserved motif Lxx(D/N)xxWExx(Y/F) found in the aligned proteins (Supplementary Figure 4), and we propose this motif to be the involved in cyclization. When Trt1 is included in the multiple sequence alignment (MSA) the conserved motif was identified as (L/I)xx(D/N)xxWExx(Y/F) (Supplementary Figure 4). Downloading the predicted protein structure of TyIF from the AlphaFold Protein Structure Database (entry A0A1Y0RDB4)19 allowed the prediction of the location of the conserved motif (Supplementary Figure 5). The L44SANIAWEFLF54 sequence was identified in a region that had very high prediction confidence (pLDDT > 90, Supplementary Figure 6, green residues) and the conserved Trp and Glu residues are located inside a cavity that is presumed to be the active site (see below). The AlphaFill database entry (AF-A0A1Y0RDB4-F1_model_v4, accessed Jan 27, 2023) did not have any ligands, metal ions or substrates predicted to be bound to the protein. We have also been unable to produce soluble protein in E. coli, most likely due to the fact that TyIF has seven predicted transmembrane sections (Supplementary Figure 7), and therefore we have been unable to directly test the role of TyIF as a terpene cyclase and the role of each individual residue in the L44SANIAWEFLF54 motif.

We turned to MD simulations to interrogate substrate binding to TyIF as Trt1 was previously shown to accept methyl 5-farnesyl-3,5-dimethylorsellinate43 while MstE was shown to cyclize 3-geranylgeranyl-4,5-dihydroxybenzoate.38-39 Therefore the identity of the cyclization substrate was unclear as TyIF is a cyanobacterial protein (as is MstE) but it is membrane bound (as is Trt1). We examined the predicted binding of 3-geranylgeranyl-4-hydroxybenzoic acid to determine if it is a substrate for cyclization. Docking of 3-geranylgeranyl-4-hydroxybenzoic acid followed by 100 ns of MD simulations resulted in a model with stable rmsd values (Supplementary Figure 8). The 3-geranylgeranyl-4-hydroxybenzoate fit within a cavity that was bound at the bottom by three Trp residues (Trp20, Trp50, Trp72) and the geranylgeranyl side chain was located in proximity to the L44SANIAWEFLF54 motif (Figure 2A). The top of the cavity was bordered by Arg62 and Arg126. The guanidino groups of the Arg residues fluctuated between 2.95 and 11.87 Å (Arg62) and between 3.17 and 11.11 Å (Arg126) from the carboxylate of 9 during the course of the MD simulation. This suggests that a hydrogen bond can be formed between the substrate carboxylate and Arg residues and that 9 is the correct substrate. We then modeled 10, the putative cyclized product (Figure 2B, Supplementary Figure 9) and found that the carboxylate of 10 was also anchored by Arg62 and Arg126 and the E ring is positioned in proximity of the conserved glutamate residue (Glu51), supporting the role of Glu51 as the catalytic acid for cyclization.

MstE utilizes a similar but distinct strategy for binding its substrate, geranylgeranyl-3,4-dihydroxybenzoate.39 Four aromatic side chains interact with the geranylgeranyl side chain (Phe49, Y100, Trp59, and Trp210) while Asp109 catalyzes the cyclization, similar to the situation envisioned for TyIF. In contrast the carboxylate of geranylgeranyl-3,4-dihydroxybenzoate is anchored through a hydrogen bond with Tyr157, while the 3’-OH group is bound by Glu339. While a structure of Trt1 is not available, the MSA shows that Trt1 contains an Ala and a Leu at the equivalent positions, which is consistent with binding a non-charged methyl ester.
Meroterpenoid BGCs encoding TylF-like proteins are widespread in bacterial genomes.

We next investigated the prevalence of BGCs encoding TylF-like proteins in sequenced bacterial genomes. Using TylF as a BLAST probe for an EFI-EST search (e-value cutoff, 5; Maximum number of retrieved sequences, 1,000) and creation of the clustered network with a cutoff score of 50 resulted in the identification of 30 clusters and 33 singletons (Supplementary Figure 10). Proteins from cyanobacteria were identified in clusters 1 and 5 (hexagons, Supplementary Figure 10). The TylF-like proteins in cluster 1 included cyanobacterial proteins from Nostoc sp. (Peltigera membranacea cyanobiont), but the majority of these proteins were from other bacterial genera including Lysinibacillus (most prevalent), Rufibacter, etc. Examination of the genes upstream and downstream of the TylF-like protein encoding gene revealed that they were typically in small or non-existent BGCs and are likely not responsible for the biosynthesis of natural products. Cluster 8 (Lavender, Supplementary Figure 10) contained fungal sequences including the known terpene cyclases NvfL from Aspergillus novofumigatus (strain IBT 16806), AtmB from Byssoschlamys spectabilis, and DpmpB from Macrophomina phaseolina (strain MS6). Cluster 5 contained proteins encoded in the genomes of cyanobacteria (Brasilonema, Tolypothrix, Scytonema, and Nostoc) and Actinobacteria (Streptomycyes, Actinoplanes, Saccharopolyspora, and Frankia). Closer inspection of these genome neighborhoods revealed that these loci encoded Rieske 2Fe-2S proteins, prenyl synthase, terpene precursor biosynthetic proteins, prenyl transferases, and cytochrome P450 proteins, and transporter proteins (e.g., ABC transporters) (Figure 4 and Supplementary Figure 11). This suggests that these BGCs have the capacity to biosynthesize meroterpenoids and that TylF-like proteins are likely terpene cyclases. These data suggest that TylF is the first described member of a class/family of previously unrecognized
bacterial terpene cyclases that are distributed across bacterial genera. The current products that these BGCs produce are under investigation in our labs.

<table>
<thead>
<tr>
<th>Bacterial Species</th>
<th>GenBank Accession Numbers</th>
</tr>
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<tbody>
<tr>
<td>Brasilonema sp. HT-58-2</td>
<td></td>
</tr>
<tr>
<td>Tolyphothrix campylomenoides</td>
<td>VBS11288 (SD81_035220 - SD81_035280)</td>
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<tr>
<td>Scytomonema sp. NIES-4073</td>
<td>(NIES4073_28450 - NIES4073_28670)</td>
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<td>Nostoc flagelliforme</td>
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<td>Nostoc sp. ATCC 43529</td>
<td>(A6S26_33945 - A6S26_34035)</td>
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<tr>
<td>Nostoc sphaeroides</td>
<td>CCNC1 (GXM_09181 - GXM_09200)</td>
</tr>
</tbody>
</table>

Scale: 3 kbp

**Figure 3.** BGCs identified using TylF as the BLAST probe for an ESI-EFT search in cyanobacterial genomes.

Heterologous expression of the tolypodiol (tyl) BGC in Anabaena sp.

As genetic disruption of the tolypodiol producing *Brasilonema* sp. HT-58-2 has not been reported and the culture is unialgal and not axenic, we elected to use heterologous expression to prove that tylA-tylM was responsible for the biosynthesis of tolypodiol. We and others have previously used *Anabaena* sp. PCC 7120 (herein *Anabaena* 7120) to successfully heterologously express cyanobacterial BGCs, and elected to express the tyl BGC in this cyanobacterial host. The tyl BGC was cloned into pPJAV550 after codon optimization for expression in Anabaena 7120. We cloned two versions of the tyl BGC, one with the native promoter and one under the control of the PpetE promoter (see Experimental section for full description) to yield pHT583_Mero-I and pHT583_Mero_PpetE-1 respectively.

Introduction of pHT583_Mero-I and pHT583_Mero_PpetE-1 to *Anabaena* sp. UTEX 2576 (=*Anabaena* sp. PCC 7120) through triparental mating with *Escherichia coli* JCM113 and NEB10β containing the plasmid of interest, resulted in exoconjugates after selection on BG-11(Nit⁺)SpSm agar plates (BG-11 media containing sodium nitrate as the nitrogen source, supplemented with spectinomycin and streptomycin (2.5 µg/mL each)). Colonies were selected, restreaked, and cultivated for an additional 5 weeks on BG-11(Nit⁺)SpSm agar plates. The cells were then collected, lyophilized, extracted with ethyl acetate, and analyzed by LC-HRMS. We were able to detect tolypodiol in the extract of pHT583_Mero_PpetE-1/*Anabaena* 7120 but not pHT583_Mero-I/*Anabaena* 7120 (Figure 4 and Supplementary Figure 12). The produced compound had a protonated ion consistent with the molecular formula C₂₈H₄₁O₅⁺ (M+H obs. 457.2940, calc. 457.2949, 2.0 ppm error) as well as an identical retention time (8.1 min), and MS/MS fragmentation pattern as the tolypodiol standard (Supplementary Figure 13-14). We used molecular networking to identify other analogs present in the extract (Figure 4B,
Supplementary Figure 15) and were able to observe 6-deoxotolypodiol, 11-hydroxytolypodiol, 1-oxotolypodiol, 6-oxotolypodiol, and a compound with the same mass and MS/MS fragmentation pattern as tolypodiol with a different chromatographic retention time (Supplementary Figure 12-14) in the extracts of the Anabaena strains containing pH583_Mero_PpetE-1, thereby linking these metabolites to the tolypodiol BGC.

Characterizing the role of TylH in tolypodiol biosynthesis.

To better understand the biosynthesis and gather further evidence to link the tolypodiol BGC to compound production, we characterized the enzymatic activity of TylH. The encoding gene was expressed in E. coli and TylH was purified using Ni-NTA chromatography. Enzymatic assays were then performed by incubating the purified TylH with hydrolyzed tolypodiol (2), SAM, and β-mercaptoethanol at room temperature overnight. Analysis of the enzymatic assay, by HPLC with standards, revealed almost complete conversion of hydrolyzed tolypodiol into tolypodiol (Figure 5 and Supplementary Figure 16). This conversion was not observed in any of the control reactions lacking TylH, SAM, or 2 (Supplementary Figure 16).

TylH substrate specificity.

The activity of TylH was then probed to determine if TylH has stringent specificity towards desmethyltolypodiol (12) or if it can be used for the O-methylation of a variety of carboxylic acids. To test this, an enzymatic assay was performed the same as with 12 but using chroman-6-carboxylic acid (13) as the substrate. Chroman-6-carboxylic acid was chosen as the substrate as it is structurally identical to a portion of hydrolyzed tolypodiol containing the carboxylic acid that undergoes O-methylation. Analysis of the enzymatic assay by HPLC revealed that the substrate did not undergo conversion (Supplementary Figure 17) into the corresponding methyl ester. In complimentary experiments, TylH did not convert 4-hydroxybenzoic acid to the corresponding methyl ester (data not shown) thus showing that TylH has high specificity towards substrates with pentacyclic structure and that the C, D, or E rings play a role in substrate binding, although this needs to be confirmed with future experiments. The failure to convert chroman-6-carboxylic acid and 4-hydroxybenzoic acid into the corresponding methyl ester also suggests that

![Figure 4. Production of tolypodiol and related compounds by expression of the tyl BGC in Anabaena 7120. (A) Extracted ion chromatogram for tolypodiol (457.2940-457.2960 m/z) in Anabaena 7120 containing pPJAV361 (empty vector), pMero_HT583_PpetE (tyl BGC), and purified tolypodiol. (B) Molecular network of metabolites found in Anabaena 7120 containing pMero_HT583_PpetE. The full GNPS network can be seen in Supplementary Figure 14.](image-url)
the O-methylation of 7 occurs later in the biosynthetic pathway. We examined unpurified and fractionated extract of Brasilonema sp. HT-58-2 but could not observe a signal with a m/z value consistent with 12. This suggests that the conversion of the carboxylic acid to the methyl ester is not the final step in the biosynthetic pathway suggesting the pathway seen in Figure 1, however due to the low ionization efficiency of the tolypodiols this statement is made noting that caveat.

New tolypodiol structures.

The molecular formula of 5 was determined by HRESIMS to be C\textsubscript{28}H\textsubscript{38}O\textsubscript{5} based on the detection of a protonated molecule at m/z 455.2790 [M+H]\textsuperscript{+} (calc for C\textsubscript{28}H\textsubscript{39}O\textsubscript{5}\textsuperscript{+}, 455.2792; -0.4 ppm). It became clear early in the structure elucidation that 5 bore many similarities with tolypodiol, which is a well-characterized compound isolated from HT-58-2.\textsuperscript{11} Specifically, the \textsuperscript{1}H NMR spectrum revealed the 1,2,4-substituted benzene ring, methoxy functional group and five methyl signals. After negotiating through the remainder of the \textsuperscript{1}H and \textsuperscript{13}C NMR spectra (Supplementary Figure 18-19), it became apparent that the major difference was an oxidation at the C-6 position to a carbonyl (\(\delta_C = 210.7\)) with regard to tolypodiol's hydroxy (\(\delta_C = 68.8\)),\textsuperscript{11} which would account for the observed mass difference of 2 amu. In addition, the diastereotopic protons at C-7 are further deshielded (\(\delta_H 5\): 2.32, 2.20; tolypodiol: 2.04, 1.23) and appear as doublets, while H-5 exhibits a change in multiplicity from a doublet in tolypodiol to a singlet in 5. The remainder of the planar structure was confirmed with a set of 2D NMR experiments (Supplementary Figure 20-22). 2D-NOESY (Supplementary Figure 23-24) along with relevant coupling constants were utilized to decipher the relative stereochemistry, which was determined to be the same as tolypodiol, except, of course, at the C-6 position where a carbonyl replaced the secondary alcohol.

HRESIMS was utilized to determine the molecular formula of 6 to be C\textsubscript{28}H\textsubscript{38}O\textsubscript{5} based on the identification of a protonated molecule at m/z 455.2788 [M+H]\textsuperscript{+} (calc for C\textsubscript{28}H\textsubscript{39}O\textsubscript{5}\textsuperscript{+}, 455.2792; -0.9 ppm). As in the case of 5, the molecular formula was 2 amu less than tolypodiol and the planar structure of 6 was recognized to be very similar to that of tolypodiol based on the 1D and 2D NMR spectra (Supplementary Figure 25-29). It became evident that the major difference was an oxidation at the C-1 position from the hydroxyl of tolypodiol to a carbonyl based on the lack of a proton resonance around 3.5 ppm that corresponded to the carbinol proton at C-1 of tolypodiol. An HMBC correlation indicated the presence of a carbonyl (\(\delta_C = 214.7\)) at C-1 and the multiplicity of

![Figure 5. Conversion of Tolypodiol-COOH (12) to tolypodiol (2) by TylH. HPLC chromatograms observing at 270 nm of (i) Full reaction containing 12, SAM, and TylH; (ii) Reaction containing 12, and SAM; (iii) standard of tolypodiol; (iv) standard of tolypodiol-COOH. Full chromatograms are shown in Supplementary Figure 13.](image-url)
H-2a/b indicated coupling only to their diastereomeric partner and the adjacent diastereomeric protons, H-3a/b. Additionally, H2a/b were further deshielded (δH 5: 3.15, 2.12) than those of tolypodiol (δH 6: 1.68, 1.65) providing further evidence C-1 was oxidized. Analysis of the JH-H coupling (H-5: 2.4 Hz; H-6: 2.4, 6.0 Hz) revealed the relative stereochemistry at C-6 was the same as tolypodiol.

For the determination of absolute configuration, ECD spectra of 5 and 6 were collected and compared to that of tolypodiol (Supplementary Figure 30). The negative Cotton effect at around 265 nm suggests 5 and 6 have the same absolute configuration as tolypodiol, which is supported by the fact that the terpene cyclization is predicted to happen early in the biosynthetic pathway.

Conclusion

Cyanobacteria are a source of unique chemical structures, bioactivities, and enzymology in which the tolypodiols provide all three. Here we describe the identification and successful heterologous expression of tolypodiol (2) and related compounds 3, 4, 5, and 6. Located in the biosynthetic gene cluster is a previously undescribed terpene cyclase, TylF, that belongs to a new family of membrane-bound bacterial terpene cyclases. We were also able to identify similar terpene cyclases in other bacterial genomes including cyanobacteria and actinobacteria. We identified a SAM-dependent methyltransferase, TylH, encoded in the biosynthetic gene cluster and characterization by in vitro assays demonstrated that it was capable of methylating 12 to form 2, but was not able to methylate chroman-6-carboxylic acid or 4-hydroxybenzoic acid suggesting that the C, D or E rings are necessary for binding. While we were unable to detect desmethyltolypodiol (12) in either Anabaena sp. UTEX 2576 containing the tyl BGC or in the native producer Brasilonema sp. HT-58-2, the substrate specificity of TylH towards the cyclized substrate suggests that methylation occurs later in the biosynthetic pathway. The identification of similar BGCs in other bacterial genomes suggests that these meroterpenoid compounds play an important role in inter-organismal interactions and our successful heterologous expression opens the potential to investigate the role of the tolypodiol family in the structure of cyanobacterial based communities and having a stable production host allows the investigation of potential therapeutic applications.

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