Microcystin *mcy*A and *mcy*E Gene Abundances Are Not Appropriate Indicators of Microcystin Concentrations in Lakes

Lucas J. Beversdorf\(^1\)*, Sheena D. Chaston\(^2\)*, Todd R. Miller\(^2\)*, Katherine D. McMahon\(^1,2\)

\(^1\) Department of Civil and Environmental Engineering, University of Wisconsin-Madison, 1415 Engineering Drive, Madison, Wisconsin, 53706, United States of America, \(^2\) Department of Bacteriology, University of Wisconsin-Madison, 1550 Linden Drive, Madison, Wisconsin, 53706, United States of America

These authors contributed equally to this work.

\*lucasb@uwm.edu

Abstract

Cyanobacterial harmful algal blooms (cyanoHABs) are a primary source of water quality degradation in eutrophic lakes. The occurrence of cyanoHABs is ubiquitous and expected to increase with current climate and land use change scenarios. However, it is currently unknown what environmental parameters are important for indicating the presence of cyanobacterial toxins making them difficult to predict or even monitor on time-scales relevant to protecting public health. Using qPCR, we aimed to quantify genes within the microcystin operon (*mcy*) to determine which cyanobacterial taxa, and what percentage of the total cyanobacterial community, were responsible for microcystin production in four eutrophic lakes. We targeted *Microcystis*-16S, *mcy*A, and *Microcystis*, *Planktothrix*, and *Anabaena*-specific *mcy*E genes. We also measured microcystins and several biological, chemical, and physical parameters—such as temperature, lake stability, nutrients, pigments and cyanobacterial community composition (CCC)—to search for possible correlations to gene copy abundance and MC production. All four lakes contained *Microcystis-mcy*E genes and high percentages of toxic *Microcystis*, suggesting *Microcystis* was the dominant microcystin producer. However, all genes were highly variable temporally, and in few cases, correlated with increased temperature and nutrients as the summer progressed. Interestingly, toxin gene abundances (and biomass indicators) were anti-correlated with microcystin in all lakes except the largest lake, Lake Mendota. Similarly, gene abundance and microcystins differentially correlated to CCC in all lakes. Thus, we conclude that the presence of microcystin genes are not a useful tool for eliciting an ecological role for toxins in the environment, nor are microcystin genes (e.g. DNA) a good indicator of toxins in the environment.
**Introduction**

In recent decades, there has been an apparent increase in the frequency and intensity of cyanobacterial harmful algal blooms (cyanoHABs) in freshwater lakes [1]. These cyanoHABs are unsightly resulting in decreased recreational and property value [2]. Upon decay they can deplete dissolved oxygen leading to fish die-offs. Some contain toxins, which impact ecosystem and public health. Microcystin is the most commonly detected and measured cyanobacterial toxin. Although the ecology of cyanobacteria is well described, the relationship between population dynamics and environmental triggers of microcystin production are poorly understood. At least 90 different structural isoforms of microcystin have been identified thus far [3], but little is known about their varying toxicity or how they might differentially affect children and adult development [4]. Microcystins bind to serine/threonine protein phosphatases in animal liver cells [5], eventually leading to tissue hemorrhaging. Acute exposure to microcystin has resulted in fatalities [1], and recent epidemiological studies suggest that chronic exposure to low concentrations of microcystin may lead to liver and colorectal cancer [6,7]. Thus, there is both a need to understand the underlying mechanisms of microcystin production and to develop technologies to better monitor and protect public health.

Microcystin is known to be produced by *Microcystis, Anabaena, Planktothrix* [8] and has been associated with several others [9]. It is synthesized via non-ribosomal peptide synthesis (NRPS) and polyketide synthase (PKS) modules. Ten genes (*mcyA-J*) are involved in microcystin production (55 kbp total) in *Microcystis, Anabaena*, and *Planktothrix* [10,11]. In *Microcystis*, these genes are found in the *mcy* operon and are bidirectionally transcribed from the *mcyA/D* central promoter [12]. In *Anabaena*, these genes are arranged in three operons [13], and in *Planktothrix*, the arrangement is identical to *Microcystis*. Two transcriptional start sites have been identified for *mcyA* and *mcyD*, dependent upon light intensities, with additional transcriptional start sites found in *mcyA, mcyD, mcyE, mcyF, mcyG, mcyH, mcyI*, and *mcyJ* [12,14,15]. However, it is still unknown what environmental conditions trigger microcystin production [16] making them difficult to predict on short time-scales (e.g. daily-weekly).

Cyanobacteria are most often enumerated and identified using microscopic counts and pigment concentrations (e.g. chlorophyll and phycocyanin). This can be an adequate means of monitoring cyanobacterial biomass and coarse population dynamics, but is ineffective at monitoring microcystin-producing populations because only some members of a cyanobacterial genus contain genes capable of encoding for toxin biosynthesis. A toxic (or potentially toxic) cyanobacterium can appear morphologically identical to a non-toxic cyanobacterium (i.e. lacking a complete *mcy* gene cluster) of the same species [17]. Furthermore, toxic and non-toxic cyanobacteria are commonly found in the same cyanobacterial bloom [17]. The percentage of cells containing toxin genes may vary significantly over both space and time [18] and there is little understanding for what environmental factors control changes in the fraction of cells capable of producing toxin. In fact, few studies have examined the relationship between the percentage of *mcy*-containing organisms with actual microcystin concentrations in lakes over both spatial and temporal scales during the cyanobacterial growth season [19,20,21], but none to this extent.

In this study, four lakes in Dane County, Wisconsin, USA were sampled weekly throughout the summer and tested for the presence and abundance of cyanobacteria with and without microcystin genes using quantitative polymerase chain reaction (qPCR). In addition, microcystins were measured using liquid chromatography tandem mass spectrometry (LC-MS/MS). The purpose of this study was to examine the population dynamics of cyanobacteria that have the potential to produce microcystin over the course of a summer using molecular methods, determine if environmental factors can be used to predict the fraction of cells capable of producing microcystins, and establish whether the abundance of such cells relates at all to measured toxin concentrations.
Materials and Methods

Ethics statement

No specific permits were required for the described field study, nor were any specific permissions needed to sample the following locations. None of the lakes in this study were privately owned, and our sampling did not involve endangered or protected species.

Sample Sites

Lakes Mendota, Monona, Wingra, and Kegonsa were sampled weekly from June 1–October 21, 2008. These lakes are located in Dane County, in south central Wisconsin near the city of Madison, and are all connected (either directly or indirectly) via the Yahara River within the Yahara Watershed (S1 Fig). All four lakes are highly eutrophic and suffer from chronic cyanobacterial harmful algal blooms [22], but they vary by size, depth, and surrounding landscape (Table 1) making them a compelling system for comparative studies. We sampled the deepest location (“Deep Hole”) of each lake, which ranged from 4 m in Lake Wingra to 25 m in Lake Mendota. We chose these locations for several reasons. First, previous studies in these lakes have shown that, while some spatial variability exists between deep and very shallow beach locations [23], within lake spatial variability is generally less than vertical heterogeneity [24,25]. Second, the Deep Hole locations of these lakes have been sampled biweekly since 1985 as part of the North Temperate Lakes-Long Term Ecological Research (NTL-LTER) program centered at the University of Wisconsin-Madison Center for Limnology. Third, the Lake Mendota Deep Hole location contains a moored buoy that collects high-resolution biological, physical, and meteorological data. Thus, we selected sample locations that were most representative of the pelagic area and contained historical, as well as high resolution, data.

Sampling Regime

Sampling was performed once or twice weekly from June–October for a total of 17–25 sample dates per lake. Sampling occurred at the deepest site in each lake (Deep Hole herein). First, Secchi disk depth was taken and multiplied by 1.7 [26] to estimate the photic zone depth (PZD). Then, integrated water samples were collected, using a PVC tube, from the surface down to the PZD and stored on ice in a cooler until transported back to the lab for analysis. Dissolved oxygen (DO), pH, and water temperature were measured through the entire water column at discrete 1 m intervals using a 556 MPS YSI meter (YSI, Yellow Springs, OH). Between 100–250 mL of lake water was filtered onto a 0.2 μm SUPOR filter (47 mm diameter; Pall Life Sciences, Ann Arbor, MI).
Ann Arbor, MI) and stored at -80°C in preparation for DNA extractions. For chlorophyll-\textit{a} and phycocyanin quantification, 1 L of water for each was filtered onto a 47 mm glass fiber filter (nominal pore size 0.7 μm; GF/F; #1825–047; Whatman, England) and stored at -20°C until analyses were performed. The GF/F filtrate was collected and frozen at -20°C in preparation for soluble reactive phosphorus (SRP) and nitrate/nitrite analysis. Whole water samples (i.e. unfiltered) were collected and frozen at -20°C in preparation for total phosphorus (TP) and microcystin analysis.

**Analytical measurements**

All analytical procedures used have been previously described [24]. Briefly, SRP was measured by the ascorbic acid-molybdenum blue method 4500 P E [27]. Nitrate and nitrite were measured individually using high-performance liquid chromatography (HPLC) [28] and reported individually, or as combined (N + N). TP was digested as previously described [29] and then determined as SRP. Phycocyanin was extracted in 20 mM sodium acetate buffer (pH 5.5) and measured spectrophotometrically following centrifugation [30]. Chl-\textit{a} was extracted in 90% acetone and measured spectrophotometrically with acid correction [31].

For toxin analysis, whole water samples were lyophilized, resuspended in 5% acetic acid, freeze-thawed three times, separated by solid phase extraction (SPE; Bond Elut C18 column, Varian), and eluted in 100% methanol as previously described [32]. Microcystin (MC) variants of leucine (L) arginine (R), alanine (A), and tyrosine (Y) were detected and quantified at the Wisconsin State Laboratory of Hygiene (SLOH) using electrospray ionization-tandem mass spectrometry (API 3200, MS/MS) after separation by liquid chromatography (LC) [33,34]. All four microcystins were added and reported herein as total microcystin (MC).

**DNA extraction and polymerase chain reaction**

DNA was extracted from filters using a xanthogenate-phenol-chloroform protocol previously described [35,36]. DNA pellets were resuspended in 100 μL of ddH₂O. Resulting DNA concentrations were quantified using Picogreen (Invitrogen, Eugene, OR) and diluted to a concentration of 10 ng μL⁻¹.

All samples were evaluated using end-point PCR for three separate genes using eight previously published primer sets (S1 Table). All PCR reactions were performed using iQ Super mix (Bio-Rad, Hercules, CA) on an Eppendorf thermocycler using previously published primer concentrations and thermocycler parameters [37,38,39,40]. PCR products were run on a 1% agarose gel stained with ethidium bromide to determine the presence or absence of each gene.

All qPCR reactions were performed using iQ SYBR green supermix (Bio-Rad, Hercules, CA) on an iCycler (Bio-Rad, Hercules, CA). Each sample was run in triplicate and each qPCR plate contained a duplicate six-point standard curve with values ranging from 10^3–10^8. Ten ng of sample DNA was added to each reaction. All qPCR was performed using previously published thermocycler parameters (S1 Table). Standards for each plate were generated by cloning PCR product from a microcystin-producing strain (University of Texas-Austin, strain UTEX 2385) into a vector using the TOPO TA cloning kit (Invitrogen, CA), extracting the recombinant plasmid using a QIAGEN plasmid mini kit (QIAGEN, Valencia, CA), cutting it with restriction enzyme \textit{ScaI} (Promega, Madison, WI), purifying using QIAquick PCR purification kit (QIAGEN), and quantifying using a NanoDrop (Thermo Fisher Scientific, Waltham, MA). All plasmids were sequenced to confirm the inserts. All standard curves yielded R^2 > 0.99 and qPCR efficiencies > 90%. At the end of each qPCR reaction, a melt curve was performed, as well as validation via ethidium bromide gel electrophoresis, to check for non-specific amplification. Copy numbers for each sample were calculated using an equation generated from the average of
the standard curves in each plate. Copy numbers were also multiplied by a dilution factor and a water-filtering factor to present data as gene copies mL\(^{-1}\) of filtered lake water. In addition to this, the Microcystis-16S copy numbers were divided by two to account for multiple copies of the 16S gene observed in the Microcystis aeruginosa NIES-843 genome [41].

Cyanobacterial community composition was assessed using automated phycocyanin intergenic spacer analysis (APISA) as previously described [23,24]. Canonical correspondence analyses (CCA) were performed in the R Statistics Environment [42] Vegan Package [43] using a Bray-Curtis similarity matrix that was based on relative abundance data [44]. Prior to analysis, toxin and gene copy data were log transformed. All correlations were performed using the envfit function within the R Vegan Package.

Lake stability calculations

Lake stability characteristics were calculated using the Lake Analyzer program previously described [45]. Briefly, Schmidt Stability (SS), Lake Number (LN), and metalimnion top (metaT) were estimated from water temperature profiles and meteorological data collected at the time of sampling, as well as lake bathymetry. SS represents the resistance to mixing due to the energy stored within the lake (i.e. how stratified the lake is) and LN represents the amount of wind induced mixing within the lake. Very simply, both indices are expected to increase as a lake becomes more thermally stratified as it takes more energy and wind forcing to cause the water column to mix. Lake Analyzer also calculates the thermocline depth, as well as the top and bottom of the metalimnion. The metaT depth to the lake surface, as defined here, is synonymous with the upper mixed layer depth of the epilimnion.

Within- and between-lake statistics

Pearson correlations were performed for in-lake comparisons to look for relationships between the abundance of the Microcystis-16S rDNA (Mic-16S), mcyA, and Microcystis-mcyE (Mic-mcyE) genes with each other and with all available environmental data. In addition, the gene copy number for Mic-mcyE was divided by the number of adjusted gene copies for the Mic-16S to estimate the proportion of Microcystis cells containing toxin genes in each lake for each sampling event. Mic-mcyE was also divided by the general mcyA to determine what proportion of the potentially toxic cyanobacterial population was Microcystis. Gene copy numbers and environmental parameters were log-transformed for statistical analysis. All comparisons were performed in MatLab [46] with calculated R and p values reported (including with or without Bonferroni Correction). We chose to use a Bonferroni Correction to reduce potential false positives that can result from multiple comparisons, as well as to focus on the strongest relationships that may exist. All between-lake comparisons were performed in MatLab using a one-way analysis of variance (ANOVA). Boxplots generated display the median and 25th and 75th percentiles. Boxes that have non-overlapping notches are considered significantly different (p values reported). Whiskers represent the minima and maxima, which are not considered outliers (± 2.7σ), and the symbols are the outliers.

Results

Within lake comparisons

Pair-wise comparisons of the three target genes in Lake Mendota yielded strong (≥ 0.92) Pearson correlation coefficients between mcyA, Mic-16S, and Mic-mcyE (S2 Table). Lake Monona yielded correlation coefficient values of a similar strength with (≥ 0.89) between mcyA, Mic-16S, and Mic-mcyE (S3 Table). Lake Kegonsa also yielded strong correlation coefficient values.
Pair-wise correlation values for Lake Wingra were weaker, yielding 0.58 between mcyA and Mic-16S, 0.73 between Mic-16S and Mic-mcyE, and 0.82 between Mic-mcyE and mcyA (S4 Table). In Lakes Mendota, Monona, and Kegonsa, all mcyA, Mic-mcyE, and Mic-16S pairwise comparisons were significant at \( p < 0.0007 \) and Wingra at \( p < 0.05 \). Since mcyA was highly correlated to Mic-mcyE and Mic-16S, Mic-mcyE was highly correlated to Mic-16S, and there was a lack of Anabaena and Planktothrix detects (S6 Table), it is likely that MC production in these four eutrophic lakes originated from Microcystis only. We then proceeded to see which environmental parameters, if any, correlated with these genes.

In most cases, each of the four lakes exhibited characteristically strong seasonal gradients (i.e. increasing temperature and thermal stratification) with nutrient depletion throughout the sampling period (Figs 1–3, Table 2). Thus, we expected mcyA, Mic-mcyE, and Mic-16S to positively correlate to temperature, lake stability, increased biomass and nutrient depletion within a lake. In Lake Mendota, Mic-16S, Mic-mcyE, and mcyA were all significantly negatively correlated to SRP and positively correlated to metaT (\( p < 0.0007 \), S2 Table). Though not significant, all genes were strongly negatively correlated to Secchi and PZD as well. Weak negative correlations were detected between nitrate, nitrite, and TP and weak positive correlations were found with phycocyanin and MC. In Lake Monona, mcyA, Mic-mcyE, and Mic-16S were not significantly correlated to any environmental parameters (\( p < 0.0007 \); S3 Table). However, mcyA and Mic-mcyE trended with strong negative correlation to nitrate, nitrite + nitrate, TP, and PZD and positively with water temperature. Lake Kegonsa was the only lake in which TP and metaT increased during the summer (Figs 3 and 1, respectively). As such, Mic-mcyE and mcyA were significantly positively correlated to TP (\( p < 0.0007 \); S4 Table) and trended negatively (but not significantly) with metaT. Mic-mcyE, mcyA and Mic-16S also trended negatively, non-significantly, with Secchi and PZD and positively with water temperature. In Lake Wingra, mcyA, Mic-16S, and Mic-mcyE were not significantly correlated to any environmental parameters at a \( p \)-value of 0.0007 (S5 Table). Gene copies were the lowest and most variable in Lake Wingra as well (Table 3). However, Mic-16S trended positively with water temperature and SS and negatively with DO. Mic-mcyE also trended positively with DO and water temperature, but was only weakly correlated to all other environmental parameters measured.

Contrary to gene copy abundance, MC concentrations did not significantly correlate with any genes or environmental parameters in any of the lakes (\( p < 0.0007 \); S2–S5 Tables). Again, this underscores how toxin concentrations, and the conditions that constrain toxin production, vary
on daily to weekly time-scales. MC concentrations were highest in Lakes Monona and Kegonsa, and MC trended with measured parameters in only those lakes. In Lake Monona, MC concentrations were negatively correlated to $mcyA$, $Mic-mcyE$, and $Mic$-16S at a $p$-value of 0.05 (i.e. without Bonferroni correction). Also, MC concentrations trended negatively with water temperature, TP, SRP in Lake Kegonsa and positively with nitrate, N + N, and metaT (all $p < 0.05$).

**Between lake comparisons**

All 4 lakes in this study were eutrophic and similar biologically and chemically (Table 2). Physically, the lakes were quite different due to their varying size, shape, and maximum depth. All lakes were significantly different ($p << 0.01$) in terms of thermal stability (i.e. SS; Fig 1) with the deeper lakes exhibiting stronger thermal stratification as expected. However, Lakes Monona and Mendota were not significantly different from each other ($p > 0.9$) by Lake Number ($L_N$), which incorporates wind shear, nor were Lakes Kegonsa and Wingra significantly different.
Table 2. Average (range) environmental characteristics of the 4 eutrophic lakes sampled in this study from May-October. BDL = below detection limit.

| Lake characteristics | Mendota | Monona | Kegonsa | Wingra* |
|----------------------|---------|--------|---------|---------|
| Water temperature (°C) | 22.4    | 22.8   | 22.8    | 22.5    |
| (14.0–28.7)          | (14.6–29.7) | (15.6–30.2) | (12.0–29.9) |
| Dissolved oxygen (mg L⁻¹) | 10.8    | 10.3   | 11.4    | 10.1    |
| (6.3–18.3)           | (6.4–14.3) | (8.7 15.4) | (7.5–12.2) |
| Photic zone depth (m) | 3.7     | 3.1    | 2.4     | 2.0     |
| (1.7–16.1)           | (1.7–10.2) | (1.7–8.5) | (1.7–3.4) |
| Total P (μg L⁻¹)     | 50      | 40     | 100     | 20      |
| (BDL—190)            | (BDL—100) | (15–200) | (BDL—50) |
| Soluble reactive P (μg L⁻¹) | 20      | 20     | 40      | 10      |
| (BDL—100)           | (BDL—80) | (BDL—150) | (BDL—50) |
| Nitrate (μg L⁻¹)     | 180     | 70     | 60      | BDL     |
| (65–450)             | (10–550) | (7–220) | BDL     |
| Nitrite (μg L⁻¹)     | 60      | 180    | 40      | BDL     |
| (BDL—230)           | (5–920) | (BDL—420) | BDL     |
| Nitrate + Nitrite (μg L⁻¹) | 240    | 260    | 105     | BDL     |
| (80–675)             | (20–940) | (20–615) | BDL     |
| Chlorophyll-a (μg L⁻¹) | 54      | 107    | 43      | 128     |
| (BDL—210)           | (BDL—250) | (BDL—240) | (BDL—230) |
| Phycocyanin (μg L⁻¹) | 104     | 107    | 112     | 112     |
| (4–560)             | (6–290) | (2–370) | (BDL—290) |
| Total microcystins (μg L⁻¹) | 7.1    | 10.7   | 13.2    | 3.4     |
| (BDL—66)            | (BDL—30) | (BDL—70) | (BDL—8.5) |
| Schmidt Stability (J m⁻²) | 367    | 134    | 10.1    | 1.1     |
| (1.8–685)           | (0.1–287) | (0.0–32) | (0–2.9)  |
| Lake number (unitless) | 6.7     | 6.7    | 0.5     | 0.2     |
| (0.0–27.3)          | (0.0–25.3) | (0.0–2.9) | (0.0–0.9) |
| Metalimnion top (m) | 9.8     | 9.9    | 6.3     | 2.8     |
| (2.5–17.1)          | (2.5–16.4) | (4.0–8.5) | (2.4–3.5) |

*Lake Wingra is primarily fed by ground water and urban run-off

doi:10.1371/journal.pone.0125353.t002

Table 3. Summary qPCR statistics for each of the four lakes.

| Mendota | Monona | Kegonsa | Wingra |
|---------|--------|---------|--------|
| Microcystis-16S | 100 | 100 | 330 | 20 |
| (BLQ—202) | (BLQ—256) | (BLQ—521) | (BLQ—31) |
| McyA | 65 | 45 | 120 | 5 |
| (BDL—240) | (BLQ—180) | (BLQ—550) | (BDL—10) |
| Mic-mcyE | 50 | 30 | 95 | 5 |
| (BDL—180) | (BLQ—120) | (BLQ—350) | (BDL—10) |
| Mic-mcyE/Mic-16S* | 97% | 74% | 52% | 21% |
| (11–187%) | (15–142%) | (13–99%) | (0–52%) |
| Mic-mcyE/mcyA** | 78% | 82% | 74% | 4400% |
| (16–138%) | (52–127%) | (37–95%) | (54–4400%) |

Values are expressed as the number of gene copies x 10³ mL⁻¹ of lake water filtered. BDL = below detection limit. BLQ = below level of quantification.

*Assuming two 16S gene copies cell⁻¹ (Kaneko et al., 2007), Mic-mcyE/Mic-16S represents the percentage of the Microcystis community that is toxic.

**Mic-mcyE/mcyA represents the percentage of the toxic community that is Microcystis

doi:10.1371/journal.pone.0125353.t003
from each other \( (p > 0.1) \). By this measure, Mendota and Monona were significantly more stable than Kegonsa and Wingra \( (p < 0.01) \). Additionally, the low SS and LN values for Lakes Kegonsa and Wingra suggest that they were polymeric in 2008. MetaT in all lakes was significantly different \( (p < 0.01) \) from one another, except in comparison of Lakes Mendota and Monona \( (p > 0.9) \); Fig 1). As stated earlier, metaT increased throughout the summer in polymeric Lake Kegonsa, whereas in Lakes Mendota and Monona it decreased throughout the summer as expected with warming upper-mixed-layer waters and a deepening of the thermocline.

Despite the varying thermal stability conditions, water temperature, DO, and PZD were not significantly different by a one-way ANOVA between the lakes \( (p > 0.01) \); Fig 2 inset), PZD variability significantly decreases and Lakes Mendota \( (\text{mean} \sim 3.4 \text{ m}) \) and Monona PZD \( (\text{mean} \sim 2.7 \text{ m}) \) both become significantly greater \( (p < 0.01) \) than Lakes Kegonsa \( (\text{mean} \sim 2.5 \text{ m}) \) and Wingra \( (\text{mean} \sim 1.0 \text{ m}) \), albeit by \( \sim 2 \text{ m} \). Additionally, the photic zone did extend through the entire water column in Lake Wingra in some cases. Nutrients were lowest in Lake Wingra, and both nitrate and nitrite were below detection for the entire sampling period \( (\text{Fig } 3) \). This may be a result of being fed by urban storm water run-off and groundwater, and/or the large number of macrophytes in Lake Wingra, whereas Lakes Mendota, Monona, and Kegonsa are larger and receive heavy nutrient loading from agricultural run-off from the Yahara Watershed. On average, TP and SRP were higher in Lake Kegonsa \( (\text{TP significantly}; p < 0.05) \), whereas nitrate was highest in Lake Mendota.

Interestingly, average phycocyanin concentrations were greatest in Lake Wingra, despite having the lowest nutrients, though phycocyanin concentrations were not significantly different between the four lakes throughout the sampling period \( (p > 0.99); \text{S2 Fig}) \). Chl-a was also highest in Lake Wingra, and was significantly greater than all other lakes \( (p < 0.01) \). Lake Monona chl-a was also significantly greater than that of Lakes Mendota and Kegonsa \( (p < 0.05) \). Conversely, chl-a was lowest in Lake Kegonsa despite having the highest TP and SRP concentrations.

There were common trends among the different target genes within the various lakes, with the exception of Lake Wingra, in that they were low in spring, peaked in mid-summer, and then declined into autumn \( (\text{S3 Fig}; \text{Table 3}) \). However, gene copy numbers were highly variable throughout the sampling period, demonstrating the importance of making multiple measurements through time. On average, Lake Kegonsa had the highest number of mcyA, Mic-mcyE, and Mic-16S gene copy numbers across the sampling period, though mcyA and Mic-mcyE were not significantly different between Lakes Kegonsa and Mendota \( (p > 0.05) \). In all other cases, Lake Kegonsa gene copy numbers were significantly greater than the other lakes \( (p < 0.05) \). Lake Wingra had the lowest number of all genes, which were significantly lower than all other lakes \( (p < 0.05) \). Lake Mendota and Lake Monona did not have significantly different copy numbers of any of the measured genes \( (p > 0.5) \).

We defined \( \text{Mic-mcyE divided by mcyA (Mic-mcyE/mcyA)} \) as the proportion of potential microcystin producers that were Microcystis, since the mcyA primers we used should encompass Microcystis, Planktothrix, and Anabaena, while the Mic-mcyE primers target only Microcystis. In Lakes Mendota, Monona, and Kegonsa, the average Mic-mcyE/mcyA percentages were 78\%, 82\%, and 74\%, respectively, and were not significantly different across lakes \( (\text{Table 3}; \text{Fig 4}) \). In Lake Wingra, the average Mic-mcyE/mcyA percentage was 4400\% and was likely due to gene copy numbers that were near the detection limit. In any case, all of the existing data suggest that the majority of microcystin producers in these lakes were Microcystis. Furthermore, no Anabaena-mcyE was detected in 2008, and Planktothrix-mcyE was detected by only end-point PCR later in the season \( (\text{S5 Table}) \). We then defined the proportion of the Microcystis community that was potentially toxic by dividing Mic-mcyE/Mic-16S \( (\text{Table 3}; \text{Fig 4}) \). In rank order, lakes with higher potentially toxic Microcystis were Mendota (97\%), Monona (74\%), Kegonsa (52\%) and Wingra (21\%), with Wingra being significantly lower than all other lakes.
Lakes Mendota and Monona were also significantly greater than Lake Kegonsa ($p < 0.01$ and 0.05, respectively). Lakes Wingra had the lowest average MC concentration (3.4 $\mu$g L$^{-1}$), which was significantly different from Lakes Monona (10.7 $\mu$g L$^{-1}$) and Kegonsa ($p < 0.05$). However, Lake Wingra also had the highest percentage of samples above 1 $\mu$g L$^{-1}$ (78%; Fig 4), the World Health Organization level for safe drinking water. The low MC concentrations in Lake Wingra are consistent with the low Microcystis, Microcystin and Microcystin Genes across 4 Lakes

Variability in toxin genes and total microcystin with cyanobacterial community composition

We asked whether variability in toxin gene abundance or MC concentration was correlated with variability in cyanobacterial community composition (CCC). In Lake Mendota, mcyA, Mic-mcyE, Mic-16S, and MC ($R^2 = 0.62, 0.61, 0.55,$ and 0.32, respectively) were all significantly correlated to changes in CCC ($p < 0.05$; Fig 5). However, MC was mostly correlated to changes in CCC that occurred earlier in the summer, whereas toxin gene abundance correlated to CCC much later in season when gene copy numbers peaked. In Lake Monona, only mcyA was significantly correlated to CCC ($R^2 = 0.47, p < 0.01$), and all toxin gene abundances were anti-correlated to MC, which was most correlated with spring/early summer samples. Similarly, all toxin gene abundances were anti-correlated to MC in Lake Kegonsa. Though none of the variables were significantly correlated to CCC, an opposite trend was observed in Lake Kegonsa (compared to Lakes Monona and Mendota) where MC was most correlated to...
late summer/fall CCC and toxin gene abundances correlated to early summer CCC. Toxin gene abundances and MC were lowest in Lake Wingra and no significant relationships were observed with CCC ($p > 0.25$). In all lakes, toxin gene abundance did not follow the same trend with CCC, as did MC. Thus while toxin gene abundance is clearly related to the abundance of Microcystis, the influence of CCC on MC production is not simply due to the dominance of Microcystis in the population.

**Discussion**

**Microcystis as the dominant microcystin producer**

One of the initial goals of this study was to determine which cyanobacterial genera were capable of producing microcystin in four eutrophic lakes. These lakes were chosen because of their similar landscape and connectivity, they all suffer from noxious cyanobacterial blooms, and they vary in shape and size. The $mcyA$ primer set published by Hisbergues *et al.* [37] was designed to target all three of the major microcystin-producing genera: *Microcystis*, *Anabaena*, and *Planktothrix*. We used this primer set to gain an overall view of the cyanobacterial community members with the potential to produce microcystins. In addition, three primer sets were used to target the $mcyE$ gene [39,47] of each specific microcystin-producing genus in order to better differentiate among the genera potentially involved in toxin production. Mic-$mcyE$ and $mcyA$ were detected via end-point PCR in all samples collected between June-October 2008 (S1 Table). *Planktothrix-mcyE* was detected via end-point PCR analysis in only two of the four lakes, Monona and Kegonsa, late in the season (September and October), and no *Anabaena-mcyE* was detected throughout the sampling period (S6 Table). Thus, taken together, the strong correlation between Mic-$mcyE$ and $mcyA$ (0.82, 1.00, 0.98, and 0.97 in Wingra, Kegonsa, Monona, and Mendota, respectively), the high Mic-$mcyE$/$mcyA$ percentages, and the small number of times *Planktothrix* was detected, suggests that *Microcystis* was indeed the dominant microcystin producer in the four lakes studied.

At least 90 different cyanobacterial taxa were detected in these lakes forming a diverse community potentially influencing toxin levels. Variation in CCC was significantly correlated with
toxin gene abundance and MC concentration in some lakes, but these two variables did not correlate with CCC in the same direction. Toxin gene abundance was correlated with the abundance of *Microcystis*, which explains the significant correlation between toxin gene abundance and CCC. Yet, MC levels correlated with CCC in an opposite direction in a canonical correspondence analysis. This indicates that MC concentration is not simply related to the abundance of the potentially dominant toxin producer.

**Microcystin correlations with the mcy genes**

*Microcystis aeruginosa* appears to be the most common microcystin producer of this genus and sequencing efforts have revealed that *Microcystis aeruginosa* may contain two copies of the 16S rRNA gene [41]. A study revealed that among *Microcystis* isolates from a lake, 73% of *Microcystis aeruginosa* contained the mcyB gene, whereas only 17% of *Microcystis ichthyoblabe* and no *Microcystis wesenbergii* contained the mcyB gene [48]. In addition, *Microcystis aeruginosa* likely carries only one copy of the mcy genes [12]. None of the genes monitored—Mic-16S, Mic-mcyE, and mcyA—were significantly correlated to MC concentrations (following Bonferroni Correction; S2–S5 Tables). The same observation has been made in other systems [49,50], while at the same time some studies have shown positive correlations with these variables. We conclude that at least within these lakes, MC concentration varies with time and environmental conditions to such a degree that simple measures of gene copy numbers cannot explain the variation, particularly at a fixed sampling point. Additionally, there were some sampling dates in Lakes Mendota, Monona, and Wingra when the number of Mic-mcyE gene copies exceeded the corrected number of Mic-16S. Although it is theoretically impossible for the number of toxigenic *Microcystis* to exceed the number of total *Microcystis*, this has been observed in similar studies [51,52] and was viewed as an indication of extreme dominance of *Microcystis* carrying toxin genes in relation to the entire *Microcystis* community. It also reflects uncertainty about the actual gene copy number per genome in native populations, as compared to sequenced genomes. Finally, differential primer specificity and limitations to quantification using qPCR may also explain these unreasonably high ratios.

**Microcystin correlations with environmental parameters**

MC concentrations did not significantly correlate (following Bonferroni Correction) to any environmental parameter in the four study lakes. Prior to Bonferroni Correction, only Lake Kegonsa had significant correlations with MC, which was strongly correlated to nitrate, nitrate + nitrite, SRP and TP, water temperature, and metaT. We note that Lake Kegonsa was an exception in this study in that TP increased during the summer and metaT decreased, which could have been caused by particle resuspension due to lake mixing (supported by the low Schmidt Stability and Lake Number; Fig 1). However, water temperature, which was nearly identical in the four lakes, was significantly negatively correlated to MC in Lake Kegonsa. Water temperature in Lake Kegonsa was above 15°C throughout the study period, which is generally considered the minimum growth temperature for *Microcystis*. However, on days when the temperature was above 25°C, microcystin concentrations were below the limit of detection, which is not consistent with prior studies citing increased microcystins with increasing temperature [53]. Furthermore, while the photic zone water temperature in all four lakes was essentially the same throughout this study (Fig 2), MC was very different. Thus, despite the multiple effects temperature can have on the increase in cyanobacterial dominance [54], temperature cannot be the sole controlling factor in cyanobacterial growth or toxin production in eutrophic lakes. Therefore, while climate projections may suggest an increase in cyanobacterial
dominance due to rising temperatures, this does not necessarily translate to an increase in toxin production.

To our knowledge, no studies have measured both MC concentrations and toxin genes, both temporally and spatially, in multiple lakes at weekly time scales. Rinta-Kanto et al. [51] did an extensive 3 year study in the Lake Erie Western Basin and observed that microcystin, total phosphorus, mcyD, and Microcystis-16S were all positively correlated on monthly time scales. In our study in Lake Mendota, MC significantly positively correlated to mcyA, Mic-mcyE, and Mic-16S but significantly negatively correlated to Chl-a (R² = 0.07, 0.74, 0.87, and 0.84, respectively; p < 0.05) on average monthly time-scales. However, on monthly time-scales in Lake Kegonsa, MC significantly negatively correlated to mcyA, Mic-mcyE, Mic-16S, water temperature, SS, and positively correlated to metaT (R² = 0.82, 0.94, 0.96, 0.89, 0.89, and 0.73, respectively; p < 0.05). And, in Lake Wingra, only phycocyanin correlated to MC, strongly negatively (R² = 0.94; p < 0.05). No parameters were correlated to MC on monthly scales in Lake Monona, and no nutrients (e.g. N and P or N: P ratios) correlated with MC in any lake. Thus, depending on the lake and time-scale for which the samples were collected, different parameters could be strongly correlated, weakly or negatively, or not correlated at all. Therefore, any studies aimed at using gene abundance as a predictor of toxin production in near real time, or studies aimed at modeling future scenarios, might misrepresent the extent of toxin production based solely on correlations.

An unresolved issue in cyanotoxin production is the mechanisms for which cyanotoxins (including microcystin and other secondary metabolites) are triggered in the environment. In culture-based studies, gene abundance and MC production have been related to N and P [55,56,57], iron (Fe)[15,58], sulfur (S)[59], light [8], growth rates [60,61], and nitrate uptake rates [62], as well as N stress [63] and C and N metabolism [64]. In the environment however, direct relationships are confounded by uncontrolled conditions and complex community interactions, but have been linked positively to P [65,66] and light [66,67] and negatively to N [24,68]. In our four study lakes, lake size played an interesting role. In lakes that were thermally stable (Mendota and Monona), gene copy numbers and biomass increased with temperature and nutrient depletion, as one might expect. In Lakes Kegonsa and Wingra, which are likely polymeric, that trend was less pronounced. Lake Mendota, the largest of the four lakes, had the highest percentage of toxic Microcystis (mean of 97%) but the lowest detects of MC concentrations above 1 μg L⁻¹. Conversely, the smallest lake, Lake Wingra, had the lowest percentage of toxic Microcystis but the highest number of detects above 1 μg L⁻¹. In Lake Wingra, nitrate and nitrite were both below detection for the entire sampling period. In a previous study, we suggested the N stress might stimulate toxin production in eutrophic lakes [24]. In Lake Wingra, MC production may be constitutively expressed due to the low amount of combined N available. However, because of the low biomass, as indicated by Mic-16S abundance, extremely high concentrations of MC couldn’t be obtained (Fig 4). In Lake Mendota, nitrate was the highest but MC concentrations were compared to Lakes Monona and Kegonsa, which had less nitrate. Thus, nitrogen may play an important, albeit complex, role in these lakes in both stimulating toxin production (deplete N) and for growth (replete N).

Toxin genes, biomass indicators, and toxin concentrations measured in this study were highly variable temporally. Additionally, there are several limitations to using gene quantification when compared to actual toxin measurements. It is possible for cyanobacteria to contain the target gene of interest, and therefore yield a positive result, but not produce toxin. This could be due to the absence of one or more of the required mcy genes via gene deletion, recombination, or transformation or due to gene disruption and inactivation by transposons or phage [69,70]. Second, qPCR of DNA provides little information about the levels at which that gene is being expressed and whether, or how much of, the toxin is being produced. Based on
this study, we conclude that quantification of toxin gene abundance is unlikely to provide data leading to a mechanistic or predictable ecological model of MC production in lakes. We suggest that future studies focus on measuring the expression of toxin genes, along with toxin concentrations, at higher resolution and include lake physics to elucidate the role of toxins in the freshwater systems.

Supporting Information

S1 Fig. Map of the Yahara Lakes chain, which includes Lakes Mendota, Monona, and Kegonsa and Lake Wingra. Lakes Mendota, Monona, and Kegonsa are all connected via the Yahara River. Lake Wingra is fed by ground water and urban run-off and feeds into Lake Monona.

S2 Fig. Time-series comparison of chlorophyll-a (Chl-a) and phycocyanin, as well as boxplot results of one-way analysis of variance of variables (ANOVA) between the lakes. Lakes are in order from largest to smallest; left to right: ME = Mendota, MO = Monona, KE = Kegonsa, and WI = Wingra.

S3 Fig. Log transformed time-series comparison of Microcystis-16S, Microcystis-mcyE, and mcyA, as well as boxplot results of one-way analysis of variance of variables (ANOVA) between the lakes. Lakes are in order from largest to smallest; left to right: ME = Mendota, MO = Monona, KE = Kegonsa, and WI = Wingra.

S1 Table. Primer sets used for PCR and qPCR analyses.

S2 Table. Pearson correlation coefficients for Lake Mendota (R values). Bold values indicate significant correlations (p < 0.0007) after Bonferroni Correction and italics indicate significant correlations (p < 0.05) before Bonferroni correction.

S3 Table. Pearson correlation coefficients for Lake Monona (R values). Bold values indicate significant correlations (p < 0.0007) after Bonferroni Correction and italics indicate significant correlations (p < 0.05) before Bonferroni Correction.

S4 Table. Pearson correlation coefficients for Lake Kegonsa (R values). Bold values indicate significant correlations (p < 0.0007) after Bonferroni Correction and italics indicate significant correlations (p < 0.05) before Bonferroni Correction.

S5 Table. Pearson correlation coefficients for Lake Wingra (R values). Bold values indicate significant correlations (p < 0.0007) after Bonferroni Correction and italics indicate significant correlations (p < 0.05) before Bonferroni Correction. No correlations were calculated for nitrate, nitrate and N + N because no concentrations were above the limit of detection.

S6 Table. Presence/absence end-point PCR for detection of the general mcyA primers and mcyE primers specific for Microcystis and Planktothrix. ND = no data.
Acknowledgments
We would like to thank Curtis Hedman of the Wisconsin State Laboratory of Hygiene for his expertise in toxin analysis. In addition, we wish to thank Dave Harring of the Center for Limnology for maintenance of boats, vehicles, trailers, and other sampling equipment necessary for this project.

Author Contributions
Conceived and designed the experiments: TRM SDC LJB. Performed the experiments: TRM SDC LJB. Analyzed the data: SDC LJB. Contributed reagents/materials/analysis tools: TRM SDC LJB KDM. Wrote the paper: LJB SDC TRM KDM.

References
1. Carmichael WW. Health effects of toxin-producing cyanobacteria: “The CyanoHABs”. Human and Ecological Risk Assessment. 2001; 7: 1393–1407.
2. Dodds WK, Bouska WW, Eitzmann JL, Pilger TJ, Pitts KL, et al. Eutrophication of US freshwaters: analysis of potential economic damages. Environ Sci Technol. 2009; 43: 12–19. PMID:19209578
3. Meriluoto J, Spoon L. Cyanotoxins: sampling, sample processing and toxin uptake. In: Hudnell HK, editor. Cyanobacterial harmful algal blooms: state of the science and research needs. Springer New York. 2008; pp. 483–499. PMID:18561363
4. Weirich CA, Miller TR. Freshwater harmful algal blooms: toxins and children’s health. Curr Probl Pediatr Adolesc Health Care. 2014; 44: 2–24. doi:10.1016/j.cppeds.2013.10.007 PMID: 24439026
5. MacKintosh C, Beattie KA, Klumpp S, Cohen P, Codd GA. Cyanobacterial microcystin-LR is a potent and specific inhibitor of protein phosphatases 1 and 2A from both mammals and higher plants. FEBS Lett. 1990; 264: 187–192. PMID:2162782
6. Li Y, Chen JA, Zhao Q, Pu C, Qiu Z, et al. A cross-sectional investigation of chronic exposure to microcystin in relationship to childhood liver damage in the Three Gorges Reservoir Region, China. Environ Health Perspect. 2011; 119: 1483–1488. doi:10.1289/ehp.1002412 PMID: 21561830
7. Zhou L, Yu H, Chen K. Relationship between microcystin in drinking water and colorectal cancer. Biomed Environ Sci. 2002; 15: 166–171. PMID: 12244757
8. Sivonen K, Jones G. Cyanobacterial toxins. In: Chorus I, Bartram J, editors. Toxic cyanobacteria in water: a guide to their public health consequences, monitoring, and management. London: E & FN Spon. 1999; pp. 41–112.
9. Dittmann E, Fewer DP, Neillan BA. Cyanobacterial toxins: biosynthetic routes and evolutionary roots. FEMS Microbiol Rev. 2013; 37: 23–43. doi: 10.1111/j.1574-6976.2012.12000.x PMID: 23051004
10. Tillett D, Dittmann E, Erhard M, von Döhren H, Börner T, et al. Structural organization of microcystin biosynthesis in Microcystis aeruginosa PCC7806: an integrated peptide—polyketide synthetase system. Chem Biol. 2000; 7: 753–764. PMID: 11053079
11. Nishizawa T, Ueda A, Asayama M, Fujii K, Harada K, et al. Polyketide synthase gene coupled to the peptide synthetase module involved in the biosynthesis of the cyclic heptapeptide microcystin. J Biochem (Tokyo). 2000; 127: 779–789. PMID: 10788786
12. Kaebernick M, Dittmann E, Börner T, Neillan BA. Multiple alternate transcripts direct the biosynthesis of microcystin, a cyanobacterial nonribosomal peptide. Appl Environ Microbiol. 2002; 68: 449–455. PMID: 11823177
13. Rouhiainen L, Vakkilainen T, Siemer BL, Buikema W, Haselkorn R, et al. Genes coding for hepatotoxic heptapeptides (microcystins) in the cyanobacterium Anabaena strain 90. Appl Environ Microbiol. 2004; 70: 686–692. PMID: 14766543
14. Sevilla E, Martin-Luna B, Vela L, Teresa Bes M, Luisa Peleato M, et al. Microcystin-LR synthesis as response to nutrition: transcriptional analysis of the mcyD gene in Microcystis aeruginosa PCC7806. Ecotoxicology. 2010; 19: 1167–1173. doi: 10.1007/s10646-010-0500-5 PMID: 20326219
15. Sevilla E, Martin-Luna B, Vela L, Bes MT, Fillat MF, et al. Iron availability affects mcyD expression and microcystin-LR synthesis in Microcystis aeruginosa PCC7806. Environ Microbiol. 2008; 10: 2476–2483. doi:10.1111/j.1462-2920.2008.01663.x PMID: 18647335
16. Neillan BA, Pearson LA, Muenchhoff J, Moffitt MC, Dittmann E. Environmental conditions that influence toxin biosynthesis in cyanobacteria. Environ Microbiol. 2012; 15: 1239–1253. doi: 10.1111/j.1462-2920.2012.02729.x PMID: 22429476
17. Sivonen K, Jones GJ. Cyanobacterial toxins. In: Chorus I, Bartram J, editors. Toxic cyanobacteria in water: a guide to their public health consequences. London: Spon. 1999; pp. 41–111.

18. Briand E, Escoffier N, Straub C, Sabart M, Quiblier C, et al. Spatiotemporal changes in the genetic diversity of a bloom-forming Microcystis aeruginosa (cyanobacteria) population. ISME J. 2009; 3: 419–429. doi: 10.1038/ismej.2008.121 PMID: 19902863

19. Sabart M, Pobel D, Briand E, Combouérie B, Salençon MJ, et al. Spatiotemporal variations in microcystin concentrations and in the proportions of microcystin-producing cells in several Microcystis aeruginosa populations. Appl Environ Microbiol. 2010; 76: 4750–4759. doi: 10.1128/AEM.02531-09 PMID: 20511422

20. Conradie KR, Barnard S. The dynamics of toxic Microcystis strains and microcystin production in two hypertrophic South African reservoirs. Harmful Algae. 2012; 20: 1–10.

21. Fortin N, Aranda-Rodriguez R, Jing H, Pick F, Bird D, et al. Detection of microcystin-producing cyanobacteria in Missisquoi Bay, Quebec, Canada, using quantitative PCR. Appl Environ Microbiol. 2010; 76: 5105–5112. doi: 10.1128/AEM.00183-10 PMID: 20562282

22. Lathrop RC. Perspectives on the eutrophication of the Yahara lakes. Lake Reservoir Manag. 2007; 23: 345–365.

23. Miller TR, Beversdorf L, Chaston SD, McMahon KD. Spatiotemporal molecular analysis of cyanobacteria blooms reveals Microcystis-Aphanizomenon interactions. PLoS One. 2013; 8: e74933. doi: 10.1371/journal.pone.0074933 PMID: 24086400

24. Beversdorf LJ, Miller TR, McMahon KD. The role of nitrogen fixation in cyanobacterial bloom toxicity in a temperate, eutrophic lake. PLoS One. 2013; 8: e56103. doi: 10.1371/journal.pone.0056103 PMID: 23405255

25. Brock TD A eutrophic lake—Lake Mendota, Wisconsin. New York: Springer-Verlag. 1985:308 p.

26. Poole HH, Atkins WRG. Photo-electric measurements of submarine illumination throughout the year. Journal of the Marine Biological Association of the United Kingdom (New Series). 1929; 16: 297–324.

27. Greenberg AE, Clesceri LS, Eaton AD. Standard methods for the examination of water and wastewater. Washington, EUA: American Public Health Association. 1992; pp.

28. Flowers JJ, He S, Yilmaz S, Noguera DR, McMahon KD. Denitrification capabilities of two biological phosphorus removal sludges dominated by different 'Candidatus Accumulibacter' clades. Environmental Microbiology Reports. 2009; 1: 583–589. PMID: 20898723

29. White AE, Karl DM, Björkman K, Beversdorf LJ, Letelier RM. Production of organic matter by Trichodesmium IMS101 as a function of phosphorus source. Limnol Oceanogr. 2010; 55: 1755–1767.

30. Demarsac NT, Houmard J. Complementary chromatic adaptation—physiological conditions and action spectra. Methods Enzymol. 1988; 167: 318–328.

31. Tett P, Kelly MG, Hornberger GM. Method for spectrophotometric measurement of chlorophyll-a and pheophytin-a in benthic microalgae. Limnol Oceanogr. 1975; 20: 887–896.

32. Harada K-I, Matsuura K, Suzuki M, Oka H, Watanabe MF, et al. Analysis and purification of toxic peptides from cyanobacteria by reversed-phase high-performance liquid chromatography. J Chromatogr A. 1988; 448: 275–283. PMID: 3147286

33. Eaglesham GK, Norris RL, Shaw GR, Smith MJ, Chiswell RK, et al. Use of HPLC-MS/MS to monitor cylindrospermopin, a blue-green algal toxin, for public health purposes. Environ Toxicol. 1999; 14: 151–154.

34. Hedman CJ, Krick WR, Karner Perkins DA, Harrah EA, Sonzogni WC. New measurements of cyanobacterial toxins in natural waters using high performance liquid chromatography coupled to tandem mass spectrometry. J Environ Qual. 2008; 37: 1817–1824. doi: 10.2134/jeq2007.0368 PMID: 18689743

35. Tillett D, Neilan BA. Xanthogenate nucleic acid isolation from cultured and environmental cyanobacteria. J Phycol. 2000; 36: 251–258.

36. Miller TR, McMahon KD. Genetic diversity of cyanobacteria in four eutrophic lakes. FEMS Microbiol Ecol. 2011; 78: 336–348. doi: 10.1111/j.1574-6941.2011.01162.x PMID: 21707672

37. Hisbergues M, Christiansen G, Rouhiainen L, Sivonen K, Börner T. PCR-based identification of microcystin-producing genotypes of different cyanobacterial genera. Arch Microbiol. 2003; 180: 402–410. PMID: 14551674

38. Neilan BA, Jacobs D, Therese DD, Blackall LL, Hawkins PR, et al. rRNA sequences and evolutionary relationships among toxic and nontoxic cyanobacteria of the genus Microcystis. Int J Syst Bacteriol. 1997; 47: 693–697. PMID: 9226902
Vaitomaa J, Rantala A, Halinen K, Rouhiainen L, Tallberg P, et al. Quantitative real-time PCR for determination of microcystin synthetase E copy numbers for *Microcystis* and *Anabaena* in lakes. Appl Environ Microbiol. 2003; 69: 7289–7297. PMID: 14660378

Rantala A, Fewer DP, Hisbergues M, Rouhiainen L, Vaitomaa J, et al. Phylogenetic evidence for the early evolution of microcystin synthesis. Proc Natl Acad Sci U S A. 2004; 101: 568–573. PMID: 14701903

Kaneko T, Nakajima N, Okamoto S, Suzuki I, Tanabe Y, et al. Complete genomic structure of the bloom-forming toxic cyanobacterium *Microcystis aeruginosa* NIES-843. DNA Res. 2007; 14: 247–256. doi: 10.1093/dnares/dsm026 PMID: 18192279

R Core Team. R: A language and environment for statistical computing. Vienna, Austria: R Foundation for Statistical Computing. 2014. doi: 10.1016/j.jneumeth.2014.06.019 PMID: 24970579

Oksanen J, Blanchet FG, Kindt R, Legendre P, Minchin PR, et al. vegan: Community Ecology Package. Vol. 20). Amsterdam: Elsevier. 1998. 853 p.

Read JS, Hamilton DP, Jones ID, Muraoka K, Winslow LA, et al. Derivation of lake mixing and stratification indices from high-resolution lake buoy data. Environ Model Softw. 2011; 26: 1325–1336.

MATLAB. Natick, Massachusetts: The MathWorks Inc. 2012.

Watanabe MF, Oishi S. Effects of environmental factors on toxicity of a cyanobacterium (*Microcystis aeruginosa*) under culture conditions. 1985; 49: 1342–1348. PMID: 3923932

Pattanaik B, Wulff A, Roleda MY, Garde K, Mohlin M. Production of the cyanotoxin nodularin-A multifactorial approach. Harmful Algae. 2010; 9: 342–350. PMID: 20453121
64. Zilliges Y, Kehr JC, Meissner S, Ishida K, Mikkat S, et al. The cyanobacterial hepatotoxin microcystin binds to proteins and increases the fitness of *Microcystis* under oxidative stress conditions. PLoS One. 2011; 6: e17615. doi: 10.1371/journal.pone.0017615 PMID: 21445264

65. Oh HM, Lee SJ, Kim JH, Kim HS, Yoon BD. Seasonal variation and indirect monitoring of microcystin concentrations in Daechung Reservoir, Korea. Appl Environ Microbiol. 2001; 67: 1484–1489. PMID: 11282594

66. Kotak BG, Zurawell RW, Prepas EE, Holmes CF. Microcystin-LR concentration in aquatic food web compartments from lakes of varying trophic status. Can J Fish Aquat Sci. 1996; 53: 1974–1985.

67. Wicks RJ, Thiel PG. Environmental factors affecting the production of peptide toxins in floating scums of the cyanobacterium *Microcystis aeruginosa* in a hypertrophic African reservoir. Environ Sci Technol. 1990; 24: 1413–1418.

68. Kotak BG, Lam AKY, Prepas EE, Kenefick SL, Hrudey SE. Variability of the hepatotoxin microcystin-LR in hypereutrophic drinking water lakes. J Phycol. 1995; 31: 248–263.

69. Kurmayer R, Christiansen G, Fastner J, Börner T. Abundance of active and inactive microcystin genotypes in populations of the toxic cyanobacterium *Planktothrix* spp. Environ Microbiol. 2004; 6: 831–841. PMID: 15250885

70. Christiansen G, Molitor C, Philmus B, Kurmayer R. Nontoxic strains of cyanobacteria are the result of major gene deletion events induced by a transposable element. Mol Biol Evol. 2008; 25: 1695–1704. doi: 10.1093/molbev/msn120 PMID: 18502770