

Original Article

The Lake Erie HABs Grab: A binational collaboration to characterize the western basin cyanobacterial harmful algal blooms at an unprecedented high-resolution spatial scale

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ABSTRACT

Monitoring of cyanobacterial bloom biomass in large lakes at high resolution is made possible by remote sensing. However, monitoring cyanobacterial toxins is only feasible with grab samples, which, with only sporadic sampling, results in uncertainties in the spatial distribution of toxins. To address this issue, we conducted two intensive “HABs Grabs” of microcystin (MC)-producing *Microcystis* blooms in the western basin of Lake Erie. These were one-day sampling events during August of 2018 and 2019 in which 100 and 172 grab samples were collected, respectively, within a six-hour window covering up to 2,270 km² and analyzed using consistent methods to estimate the total mass of MC. The samples were analyzed for 57 parameters, including toxins, nutrients, chlorophyll, and genomics. There were an estimated 11,513 kg and 30,691 kg of MCs in the western basin during the 2018 and 2019 HABs Grabs, respectively. The bloom boundary poses substantial issues for

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spatial assessments because MC concentration varied by nearly two orders of magnitude over very short distances. The MC to chlorophyll ratio (MC:chl) varied by a factor up to 5.3 throughout the basin, which creates challenges for using MC:chl to predict MC concentrations. Many of the biomass metrics strongly correlated ($r > 0.70$) with each other except chlorophyll fluorescence and phycocyanin concentration. While MC and chlorophyll correlated well with total phosphorus and nitrogen concentrations, MC:chl correlated with dissolved inorganic nitrogen. More frequent MC data collection can overcome these issues, and models need to account for the MC:chl spatial heterogeneity when forecasting MCs.

1. Introduction

Harmful algal blooms (HABs) are a global issue and are a growing concern due to their documented expansion driven by increased anthropogenic nutrient pollution and climate change (O'Neil et al., 2012; Paerl et al., 2016). Many human health, ecological, and economic problems are associated with HABs, but a significant risk is the toxin contamination of recreational and drinking waters by freshwater cyanobacterial HABs (cyanoHABs) (Carmichael, 1992; He et al., 2016; Qin et al., 2010; Sitoki et al., 2012; Steffen et al., 2017). Among the many types of toxins produced by cyanoHABs, microcystins (MCs) are the most commonly documented and often found in the highest concentrations (Carmichael and Boyer, 2016; Graham et al., 2020; Harke et al., 2016; Loftin et al., 2016). During bloom conditions, MC concentrations can exceed the World Health Organization (WHO) and local government guidelines for drinking water and recreational uses by several orders of magnitude (Davis et al., 2019; Krausfeldt et al., 2019). Several major cities worldwide have recently issued 'do not drink advisories' due to MCs in treated tap water (Qin et al., 2010; Sitoki et al., 2012; Steffen et al., 2017). Annual summertime cyanoHABs dominated by *Microcystis*, a globally distributed MC-producer (Harke et al., 2016), have plagued Lake Erie's western basin for the past two decades (Bridgeman et al., 2013; Steffen et al., 2014; Stumpf et al., 2012). In August 2014, the City of Toledo issued a three-day do not drink advisory due to MCs in tap water, which affected nearly 500,000 residents (Qian et al., 2015; Steffen et al., 2017). Although the 2014 Lake Erie cyanoHAB was not particularly expansive compared to previous years (Davis et al., 2019), the accumulation of MC-producing cyanoHAB biomass at Toledo's drinking water intake posed a threat to human health. Therefore, the development of forecasting capabilities for cyanoHAB toxins is paramount to protect human health and mitigate the multiple negative impacts of blooms.

Unlike monitoring for MCs, monitoring the biomass of annual cyanoHABs is made possible over large areas and over time by remote sensing. Airborne or satellite remote sensing allows for surface cyanoHAB biomass in lakes to be quantified with high spatial resolution and frequent revisits (Wynne and Stumpf, 2015). Regular (daily to weekly) satellite images of a lake allow for an annual assessment of cyanoHAB biomass, and multiple years of images document interannual trends and variability. Throughout the Lake Erie bloom season, the U.S. National Oceanic and Atmospheric Administration (NOAA) disseminates semi-weekly bulletins that display bloom location and biomass quantified by remote sensing. The bulletins also forecast the bloom's location over the next several days using wind predictions and resulting water currents (Wynne et al., 2013). Additionally, Environment and Climate Change Canada's (ECCC) EOLakeWatch delivers a suite of satellite-derived products for Lake Erie, mapping total chlorophyll *a* and providing quantitative indices describing the bloom spatial extent, intensity, and duration (Binding et al., 2021). Coupling the extensive satellite-derived biomass data with an examination of environmental drivers has led to the development of regression models and mechanistic models driven by springtime Maumee River cumulative discharge and phosphorus loading (GLWQA, 2015; Sayers et al., 2016; Stumpf et al., 2016b; Verhamme et al., 2016). These models largely explain the interannual variability of cyanoHAB biomass and allow for seasonal predictions months in advance of the actual bloom (Stumpf et al., 2012).

Unfortunately, neither ECCC nor NOAA remote sensing products and bulletins include annual MC assessments or MC forecasts because cyanoHAB toxins cannot be directly detected through remote sensing (Stumpf et al., 2016a). Neither may cyanoHAB biomass be used as a proxy for MC concentration because blooms comprise both MC-producing and non-MC-producing strains, and there is no consistent correlation between cyanoHAB biomass and MC concentration (Liu et al., 2020; Stumpf et al., 2016a). Therefore, in order to estimate the mass of MCs throughout a lake or basin and develop annual assessments of toxicity and predictive models, in the same manner that remote sensing has allowed for cyanoHAB biomass, extensive MC data acquired by vessel-based water grab samples are needed.

Collecting water samples at sufficiently high spatial resolution in Lake Erie so that MC data might be reasonably compared with satellite cyanoHAB biomass data is a formidable challenge. Lake Erie is the 11th largest lake in the world by surface area (Herdendorf, 1982), and the western basin, where cyanoHABs are most prevalent, has an area of approximately 3000 km². Two U.S. states (Ohio and Michigan) and a Canadian province (Ontario) have jurisdiction over the western basin's waters, thus making the cyanoHABs an international problem. While the management of environmental problems that span international boundaries can be challenging (Perz et al., 2010), the U.S. and Canada have a long history of collaboration on the Great Lakes (e.g., International Joint Commission, Great Lakes Water Quality Agreement, Great Lakes Fisheries Commission; McKindles et al., 2020). While many researchers and agencies are studying Lake Erie's cyanoHABs, the basin's large size and multiple jurisdictions lead to discrepancies in grab sample collection and analysis methods for routine monitoring, making data amalgamation difficult (Golnick et al., 2016). Some researchers (Fang et al., 2019) have pooled cyanoHAB biomass data from the various institutions to make annual assessments, but correction factors need to be introduced and caveats acknowledged to make the combined dataset useful. Thousands of grab samples from Lake Erie have been analyzed for MCs in recent years, but before those data can be combined to make spatial assessments of MCs throughout the lake, a tool is needed to quantify the uncertainty associated with pooling multiple sources of MCs data. As part of a more extensive multi-institution study oriented toward developing forecasting of conditions favorable for MC production by Lake Erie cyanoHABs, investigators determined that the study would benefit from the collection of one or more MCs datasets from a high spatial resolution survey with coordinated methods collected across the lake basin. Such a dataset was expected to allow for calculating the total mass and the average concentration of MCs in the basin at a single point in time, which would be analogous to a satellite image displaying cyanoHAB biomass.

This study's primary objective was to coordinate two six-hour, high-resolution sampling events on Lake Erie's western basin to estimate the total mass of MCs during the peaks of the annual cyanoHAB on 9 August 2018 and 7 August 2019. We termed these one-day sampling events "HABs Grabs." The 2018 HABs Grab had a smaller number of participating institutions and was limited to the U.S. waters. Due to the attention of the initial HABs Grab, Canadian institutions also participated in 2019. Additionally, because HABs Grab partners have a wide range of expertise on cyanoHABs (molecular scale to ecosystem modeling), the HABs Grab represented an opportunity to collect data to answer many additional scientific questions. All HABs Grab samples

were collected and analyzed for 57 different parameters by consistent methods, including toxins, pigments, nutrients, and cyanobacterial DNA. In this manuscript, we present: 1) the coordination elements of the HABs Grab events, 2) how the HABs Grabs aligned with the seasonal blooms, 3) the total mass of MC in the western basin of Lake Erie, 4) the environmental parameters (i.e., nutrients, temperature) during the HABs Grabs, 5) comparison among biomass metrics, and 6) correlations between MCs, biomass, the MC-to-biomass ratio, and environmental parameters collected during the HABs Grab.

2. Materials and methods

2.1. Sample collection and handling methods

The field and laboratory crews met at the University of Toledo Lake Erie Center several days before the HABs Grab to calibrate water quality sondes (EXO2 YSI Inc., Yellow Springs, OH, USA), distribute sample equipment and bottles, and demonstrate sample collection and handling methods. This ensured that all samples were collected with the same methods. The laboratory was set up to facilitate sample processing (filtering and dispensing aliquots).

During six hours on 9 August 2018, four institutions (Ohio State University [OSU] Stone Laboratory, University of Toledo Lake Erie Center [UT-LEC], Bowling Green State University [BGSU], and Limno-Tech) collected a total of 100 samples in the U.S. waters of the western basin of Lake Erie. The NOAA Lake Erie Harmful Algal Bloom Bulletin was used to determine sample locations in advance of the field operation. On 7 August 2019, eight institutions and agencies (the aforementioned, NOAA's Great Lakes Environmental Research Laboratory [GLERL], Environmental and Climate Change Canada [ECCC], Fisheries and Oceans Canada, and University of Windsor's Great Lakes Institute for Environmental Research [GLIER]) collected a total of 172 samples across the entire western basin, with sample locations pre-determined in a grid-style pattern. Four research vessels were used in 2018, and eight were used in 2019. The vessels sampled areas of the lake in proximity to their marina. In both years, samples were collected between 8:00 AM and 2:00 PM local time to minimize variability caused by the bloom movement (by advection and vertical migration) during sampling.

The same grab sample collection and handling methods were used both years and among all groups. The vessels did not anchor at each site thus facilitating rapid sampling in order to collect the high number of sampling stations per vessel targeted during the six-hour sampling window. Upon arriving at the location, time, GPS, and water depth were recorded. Sample equipment and bottles were first rinsed with lake surface water. A two-meter-long tube sampler was used to collect an integrated water sample from the surface to 2 m depth, and the water was deposited into a clean and pre-rinsed 20-L bucket (Golnick et al., 2016). Lake water from the bucket was poured into transparent 2.4-L polyethylene terephthalate glycol (PETG) bottles and stored in a dark cooler while being transported back to the laboratory. No on-vessel processing of the water took place. Water was poured into the calibration cup of an EXO2 sonde (YSI Inc., Yellow Springs, OH, USA) immediately after sampling to record field parameters (water temperature, pH, specific conductivity, turbidity, and chlorophyll and phycocyanin fluorescence as relative fluorescence units (RFU)). After the water was collected, the vessel moved to the next location, usually within 5 to 10 min after arriving.

All samples collected in the US waters were transported to the UT-LEC, whereas all Canadian samples were transported to GLIER for processing. Both laboratories used the same methods and supplies. Upon arriving at the processing laboratory, the sample bottles were vigorously inverted several times, and aliquots were distributed for different parameters, including filtration before sample splitting for some parameters.

2.2. Laboratory analytical methods

2.2.1. Microcystins

Every sample was analyzed for microcystins (MCs) with two analytical methods, and the sample water for both methods came from the same aliquot. For total MCs, 25 mL was poured into 60-mL amber glass vials and frozen at -20°C . All MCs samples were transported to OSU for three freeze/thaw cycles to lyse cells, and following the third thaw, water was filtered into two separate glass vials with glass micro-fiber (GMF, $0.45\ \mu\text{m}$) syringe filter to remove cellular debris. One vial was delivered to the City of Toledo Collins Park Water Treatment Plant for analysis of MCs by enzyme-linked immunosorbent assay (ELISA) and the second to the Lumigen Instrument Center at Wayne State University for analysis of 12 MC congeners and Nodularin by liquid chromatography with tandem mass spectrometry (LC-MS/MS). Extracellular MCs were analyzed similarly, albeit using water filtered the day of collection without freeze/thaw lysis.

MCs by ELISA were quantified with Abraxis kits (#520011; Eurofins Abraxis, Warminster, PA, USA) on an Abraxis Cyanotoxin Automated Assay System at the Toledo Water Treatment Plant. Samples that exceeded $4\ \mu\text{g/L}$ were diluted and reanalyzed, and samples with analytical duplicates that differed by more than 10% were reanalyzed (Ohio Environmental Protection Agency, 2018).

Quantitative analysis of MCs by LC-MS/MS was conducted using an online concentration method (Birbeck et al., 2019). Briefly, using a Thermo Scientific TSQ Altis™ triple quadrupole mass spectrometer (Thermo Scientific, Waltham, MA, USA) with an EQuan MAX Plus™ system, 1 mL of sample was injected onto a loading column (Thermo Scientific Hypersil GOLD aQ $2.1 \times 20\ \text{mm}$, $12\ \mu\text{m}$ particle size) using an HTC PAL autosampler (CTC Analytics, Zwingen, Switzerland). The analytical column used was a Thermo Accucore aQ, $50 \times 2.1\ \text{mm}$, $2.6\ \mu\text{m}$ particle size column, kept at a stable temperature of 35°C for gradient analysis. Mass spectrometry analysis was performed using positive electrospray ionization source (ESI) mode. Quantitation data results were accomplished using TraceFinder™ EFS 4.1. The congeners analyzed for included MC-LR, RR, YR, WR, HtyR, HilR, [D-Asp³]-RR, [D-Asp³]-LR, LA, LF, LY, and LW, as well as nodularin. The MCs were purchased from Enzo Life Sciences, Inc. (Farmingdale, NY, USA). The surrogate C₂D₅ MC-LR was purchased from Cambridge Isotope Laboratories, Inc. (Tewksbury, MA, USA). It is noteworthy that [d-Asp³] MC-RR has been reported misidentified and has been a putative identification as [d-Asp³, Dhb] MC-RR (Birbeck et al., 2019).

2.2.2. Nutrients

A 100-mL aliquot was filtered to measure dissolved nutrients (nitrate, nitrite, ammonium, dissolved reactive phosphorus, and dissolved reactive silicate) directly upon return to the lab. First, two $\sim 10\text{-mL}$ sub-aliquots were filtered through a $0.45\text{-}\mu\text{m}$ membrane filter to rinse the equipment, with the remainder filtered and poured into a 60-mL PETG bottle, which was frozen until analyzed. A sub-sample was taken for extracellular MCs. Unfiltered aliquots for total phosphorus (TP) and total Kjeldahl nitrogen (TKN) were poured directly from the 2.4-L bottle into separate 250 mL bottles and frozen at -20°C until analysis. Total N concentration was calculated by the sum of nitrate, nitrite, and TKN. All nutrient samples were transported to OSU for analysis following standard procedures (Chaffin et al., 2019).

2.2.3. Chlorophyll and phycocyanin

For chlorophyll (chl) *a* concentration, 50 to 100 mL, depending on phytoplankton biomass, was filtered onto a 25 mm glass fiber filter (GF/F; $07\ \mu\text{m}$ nominal pore size), noting the volume. The filters were stored on silica gel in mini-Petri dishes at -80°C . Chl *a* was extracted from the filters using dimethylformamide (DMF) and quantified by fluorometry (Golnick et al., 2016). Additionally, samples were analyzed on the day of collection by a FluoroProbe (bbe Moldaenke, GmbH) with a benchtop cuvette reader for chl *a* associated with four phytoplankton groups: 1)

green algae, 2) phycocyanin-rich cyanobacteria, 3) diatoms, chryso-phytes and dinoflagellates, 4) cryptophytes and phycoerythrin-rich cyanobacteria (Beutler et al., 2002; Chaffin et al., 2013). The ratio between the FluoroProbe total chl *a* concentrations and the DMF chl *a* concentrations was used to adjust the chl *a* concentrations attributed to each algal group (Bridgeman et al., 2012). The Canadian samples from the 2019 event ($n = 60$) were not analyzed with a FluoroProbe because the instrument was available at the time of the event.

Because the Canadian samples were not analyzed with a FluoroProbe, cyanobacteria-chl *a* concentration was estimated using step-wise regression with the U.S. data. Input parameters were DMF-chl *a*, total ELISA MCs, *mcyE*, and phycocyanin concentration. There was a very tight relationship between DMF-chl *a* and cyanobacteria-chl *a* ($P < 0.001$, $R^2 = 0.974$), and no other variables were included ($P > 0.05$). FluoroProbe cyanobacteria-chl *a* concentration for the Canadian samples was therefore estimated using the equation:

$$\text{Eq. (1) } \text{FLP}_{\text{cyano}} = (0.5926 * \text{DMF}_{\text{chla}}) + 0.0935$$

Where $\text{FLP}_{\text{cyano}}$ is the FluoroProbe cyanobacteria-chl *a* concentration ($\mu\text{g/L}$) and DMF_{chla} is the chl *a* concentration ($\mu\text{g/L}$) measured from the DMF method.

For phycocyanin and phycoerythrin concentration, 400 mL was filtered onto a 47 mm GF/C (1.2 μm nominal pore size) filter and immediately frozen at -80°C until extraction and analysis. Extraction of the two pigments was done using 3 mL of 50 mM potassium phosphate buffer (pH 6.7) at -4°C for 24 h and subsequently placed at $+4^\circ\text{C}$ for another 24 h. The extract was centrifuged (20 min, $+5^\circ\text{C}$, $3210 \times g$, Beckman GS-6R rotor, Beckman Coulter Life Sciences, Indianapolis, IN, USA) to remove the filter and cell debris. Analyses of phycocyanin and phycoerythrin pigments were carried out according to Sarada et al. (1999) and Thoisen et al. (2017), respectively. Briefly, the supernatant's absorbance was measured at 455, 564, 592, and 750 nm on a USB2000 spectrophotometer (Ocean Optics, Dunedin, FL, USA) using potassium phosphate buffer as a blank. The absorbance values were scatter-corrected by subtracting the absorbance at 750 nm. The entire process was conducted under red light to avoid pigment degradation due to exposure to ambient light.

2.2.4. Molecular samples

DNA was extracted from 25 mm 1.2 μm pore size filters (Pall Versapor® acrylic copolymer; p/n 66,393; Port Washington, NY, USA) using a Qiagen DNeasy® Blood and Tissue Kit (Qiagen, Germantown, MD, USA). Briefly, samples were incubated with 100 μL Qiagen ATL tissue lysis buffer, 300 μL Qiagen AL lysis buffer, and 30 μL proteinase K at 56°C for 1 h with agitation, followed by vortexing at maximum speed for 10 min. Lysates were homogenized with a QiaShredder™ spin-column before purification, according to the DNeasy® protocol. DNA quantity and quality were assessed using a NanoDrop spectrophotometer (Thermo Fisher Scientific, NanoDrop Products, Wilmington, DE, USA). Genes indicative of the genetic potential to produce microcystins/nodularin (*mcyE*), saxitoxins (*sxtA*) and cylindrospermopsins (*cyrA*) were enumerated using a commercially available multiplex qPCR kit (Phytoxigene CyanoDTec™ Toxin Genes Test; Diagnostic Technology, Sydney, Australia) modeled after the multiplex qPCR assay described elsewhere (Al-Tebrineh et al., 2012). The primers were general to all cyanobacteria and not targeted to a specific genus. Briefly, molecular grade water (80 μL) was added to each tube of a CyanoDTec™ cyanotoxin detection kit and processed following kit directions. A synthetic standard of known toxin gene copy (Diagnostic Technology) was assayed in serial dilutions to generate a standard curve spanning five orders of magnitude (100–1000,000 copies) for each target toxin gene. Amplification was conducted on a Quantabio Q Real-Time PCR system (Quantabio, Beverly, MA, USA) in a total volume of 25 μL . Each sample was run in duplicate. Gene copies in each reaction were calculated using the Quantabio Q software and back-calculated to copies/L. Only the *mcyE* gene abundances will be reported in this study.

Random shotgun sequencing of the whole community was

performed on 25 samples from the 9 August 2018 HABs Grab to estimate the proportion of the *Microcystis* population capable of producing MCs. These 25 sampling sites ranged from 14 km to 52 km away from the Maumee River's mouth. Metagenomic shotgun reads were subjected to the Geomicrobiology Quality Check protocol as part of the Metagenomics Pipeline, which can be found at <https://github.com/Geo-omic/s/scripts>. Briefly, reads were trimmed for quality and removal of Illumina adapters and spike-ins using BBDuk. Quality checked reads were then de-replicated and forward and reverse reads were pooled. The relative abundance of *Microcystis* was quantified with a Basic Local Alignment Search Tool (BLAST) v.2.8.1 search of reads against the v4 region of all *Microcystis* 16S rRNA genes in the SILVA database v. 138.1. To avoid counting non-specific matches (i.e., from other cyanobacteria), the 16S v4 regions from all cyanobacteria deposited in the SILVA database v. 138.1 were included in the BLAST database as competitors for reads (Supplemental File 2). The relative abundance of *Microcystis mcyD* and *mcyE* genes was quantified with a BLAST search to all *Microcystis mcy* genes *D* and *E* genes publicly available in the IMG/MER database (<https://img.jgi.doe.gov/>). To avoid non-specific mapping, *mcy* genes from taxa other than *Microcystis* were also included in the BLAST database as competitors for reads (Supplemental File 3). Aligned reads were counted as positive matches if they had 80% query coverage and 95% sequence identity. These cutoff parameters, database metrics, and mapping tools were tested exhaustively to ensure specificity and sensitivity by cross-checking matches to the full SILVA database and a custom universal database (https://github.com/TealFurnholm/Meta-NGS_Reference_Database). Filtered reads per gene were then quantified and normalized by the length of the gene and sequence library size. Counts for ambiguous assignments (alignment to multiple subjects with identical bitscores) were divided by the total number of subjects. Normalized read counts for the *mcy* gene were then divided by normalized read counts for the *Microcystis* 16S rRNA gene (assuming one *mcy* and 16S rRNA gene copy per cell) to give an estimate of the proportion of the *Microcystis* population within each sample that contained each *mcy* gene within their genome.

2.3. Data analysis

2.3.1. Water currents (FVCOM)

The hydrodynamic conditions were simulated using the Lake Erie operational version of the Finite Volume Community Ocean Model (FVCOM; Chen et al., 2003) maintained by NOAA. FVCOM is a three-dimensional (3D), free-surface, primitive-equation model that solves the integral form of the governing equations on an unstructured, sigma-coordinate mesh. The advantage of an unstructured grid mesh for shoreline fitting and local mesh refinement makes the model particularly attractive in applications to coastal waters. FVCOM has been applied in many coastal systems characterized by geometric complexities and highly variable flow patterns, including various applications to the Great Lakes (for example: Anderson et al., 2015; Rowe et al., 2016; Xue et al., 2017).

The Lake Erie (LE)-FVCOM was configured with horizontal resolution ranging from 200 to 2500 m and 21 vertical terrain-following sigma layers. The inflow and outflow to the lake include Detroit, Maumee, and Niagara Rivers. The LE-FVCOM was run in a hindcast mode for the year 2018 and 2019 and driven by hourly meteorological surface forcing from the High-Resolution Rapid Refresh (HRRR), a cloud-resolving and convection-allowing weather forecast and data assimilation system running real-time at a 3-km grid resolution. The LE-FVCOM utilized the Mellor and Yamada level 2.5 (MY-2.5) and Smagorinsky turbulent closure schemes for vertical and horizontal mixing, respectively (Mellor and Yamada, 1982; Smagorinsky, 1963).

2.3.2. Total microcystin mass

Total MC mass was determined for each of the HABs Grab sampling events by applying similar geospatial methods. Geolocated

concentrations determined for the homogenized samples from the upper 2 m of the water column at each sampling station by ELISA were used to construct a geodatabase for each sampling event, along with existing bathymetric data rather than field-reported water depths. Sampling station data were bounded by a set of points that was developed by making conservative assumptions. Zero-concentration open lake boundary points were created along the August 2018 bloom edge from the 9 August forecast bloom position reported in the 6 August NOAA HABs Bulletin based on a 5 August Sentinel-3 satellite image. The entire basin was sampled in the 9 August 2019 event, so the creation of a set of open-water boundary points was not necessary. Shoreline point values were projected as constants from the nearest offshore sampling station in both years, including for island shorelines as appropriate.

The entire MC dataset of sampling stations and constructed bounding values was interpolated using inverse distance weighting (IDW) methods (Philip and Watson, 1982; Watson and Philip, 1985) within the ArcMap 10.7.1 Spatial Analyst extension software package (Environmental Systems Research Institute, Redlands, CA). The interpolated MC concentrations were converted to 100-m by 100-m grid cells and multiplied by the average water depth in each grid cell to determine a total mass of MC per grid cell for the whole water column. Masses were then summed for all column grid cells to calculate a total MC mass for each sampling event. Recent studies of the vertical distribution of biomass and MC in western Lake Erie have revealed that significant cyanobacteria and toxin remain at depth, even on days when cyanobacterial colonies are observed at the surface (Kramer, 2018). Surface and bottom MC concentrations were also similar in weekly samples collected by NOAA GLERL during summer 2019 (Supplemental Fig. 1). Furthermore, the highest MC concentrations usually occur in the shallow nearshore waters where the upper 2 m make up a larger proportion of the water column, whereas MC concentrations are an order of magnitude less at deeper sites (Palagama et al., 2020). The calculation assumed a uniform water column concentration for each grid cell. This assumption may yield an overestimate of total MC mass if concentrations declined significantly between the upper two meters of the water column and the deeper waters. However, other available data collected around the HABs Grab dates do not indicate that the assumption of consistent top to bottom MC concentrations is unreasonable, and it would be overly conservative to assume all MCs were within the upper 2 m.

2.3.3. Spatial interpolations

Spatial interpolations of the HABs Grab data were conducted in ArcGIS v10.3 with the Kriging function. All default settings were used. The Kriging outputs were clipped to the area sampled.

2.3.4. Correlation among parameters

A Pearson correlation coefficient (r) matrix was generated among metrics used to estimate cyanobacterial biomass during the 9 August 2018 and 7 August 2019 HABs Grab. For our analysis, we considered r values less than 0.30 to have a negligible correlation, 0.31 to 0.50 to be weakly correlated, 0.51 to 0.70 to be moderately correlated, 0.71 to 0.90 to be strongly correlated, and 0.91 to 1.0 to be very strongly correlated. The parameters included those measured aboard the research vessels with the YSI handheld units (chl and PC relative fluorescence units), measured in the laboratory with the FluoroProbe (cyanobacteria-specific chl a and total chl a concentrations), and filtered extracted chl a , PC, and cyanobacteria-specific 16S gene copies. Total MCs and extracellular MCs measured by ELISA were also included in the matrix to determine how toxin concentration correlated with the biomass metrics.

Scatter plots of MCs, total chl a , and the ratio of MCs to chl a against potential environmental explanatory variables (for example, temperature, pH, nutrient concentrations) for samples with MCs concentrations greater than 1 $\mu\text{g/L}$ were generated to display correlations and patterns visually. Additionally, residuals from the MC-ELISA vs. chl a regression equation were calculated and plotted against environmental

explanatory variables. IBM SPSS Statistics v23 were used for all data analysis.

2.4. Seasonal progression of the Lake Erie bloom

Routine monitoring (every ~ 10 days) data collected by the UT-LEC was used to put the 2018 and 2019 HABs Grab into the context of the seasonal bloom progression. Data from five sites representative of the western basin, extending 14–31 km from the Maumee River's mouth (Supplemental Fig. 2), were averaged for each sample date. The five sites selected have been routinely monitored by UT-LEC since 2002, and *Microcystis* biovolume in this dataset aligns well with remote sensing data (Bridgeman et al., 2012, 2013). Water samples were collected throughout the water column and analyzed by a benchtop FluoroProbe, as described above, to determine the relative contribution of green algae, cyanobacteria, diatoms, and cryptophytes to total chl a concentration. ELISA was used to measure total MC concentrations.

Seasonal progression of the bloom spatial extent was documented by ECCO's EOLakeWatch, using Sentinel-3 OLCI satellite imagery. In order to minimize cloud-related uncertainties in image products, 14-day rolling average chl a maps were produced (Binding et al., 2021). Bloom extent was delineated using the threshold of satellite-derived chl $a > 10 \mu\text{g/L}$.

3. Results

3.1. Progression of the blooms

Cyanobacteria-specific chl a concentrations increased from lows in June to peaks in late July and early August and then decreased throughout August and September in both years (Fig. 1A & 1B). The August chl a peak of 2018 ($7.9 \pm 2.1 \mu\text{g/L}$) was lower than that of 2019 ($37.8 \pm 9.7 \mu\text{g/L}$). Total MCs followed a similar pattern as cyanobacteria chl a and peak MCs concentrations in 2018 ($2.2 \pm 0.8 \mu\text{g/L}$) were less than that of 2019 ($5.5 \pm 1.4 \mu\text{g/L}$). The peak bloom spatial extent in 2018 lasted from mid-August through mid-September, covering an area of approximately 400 km^2 (Fig. 1C). In 2019, the bloom extent reached 1171 km^2 in mid-August, declined slightly in late August, and peaked again during mid-September at 1310 km^2 (Fig. 1D).

Total chl a (determined by the DMF method) increased throughout summer during 2018, peaking at $19.1 \pm 3.4 \mu\text{g/L}$ in early September. Total chl a in 2019 was greater than 30 $\mu\text{g/L}$ from late July throughout early September, with the highest concentration on 1 August of $45.7 \pm 11.5 \mu\text{g/L}$ (Fig. 1E and 1F). Pre-bloom of both years, diatoms accounted for more than 60% of total chl a , and cyanobacteria made up a relatively low percentage of total chl a ($< 25\%$). Cyanobacteria accounted for 52% of the chl a during the peak bloom in 2018, but in the larger bloom year of 2019, cyanobacteria accounted for 81%. Diatoms returned to dominance following the cyanobacterial bloom peak in both years. These data indicate that both HABs Grab events were conducted during the peak bloom conditions for biomass, bloom extent, MC concentrations, and cyanobacterial dominance of the phytoplankton community.

3.2. Spatial data of biomass and microcystins

During the 9 August 2018 HABs Grab, the highest concentrations of chl a (measured with the DMF method) occurred within several kilometers from the Ohio and Michigan shorelines, and lower concentrations were measured in the center of the basin (Fig. 2). All 2018 samples had a filter-extracted chl a concentration less than 21 $\mu\text{g/L}$, except the sample closest to the Maumee River, which had 54.3 $\mu\text{g/L}$ and the FluoroProbe determined 61% of the chl a was from green algae and diatoms. Much higher chl a concentrations, nearly an order of magnitude greater, were measured on 7 August 2019 (Fig. 2). The highest concentrations occurred in Maumee Bay near the mouth of the Maumee River ($> 150 \mu\text{g/L}$), and concentrations decreased with increasing

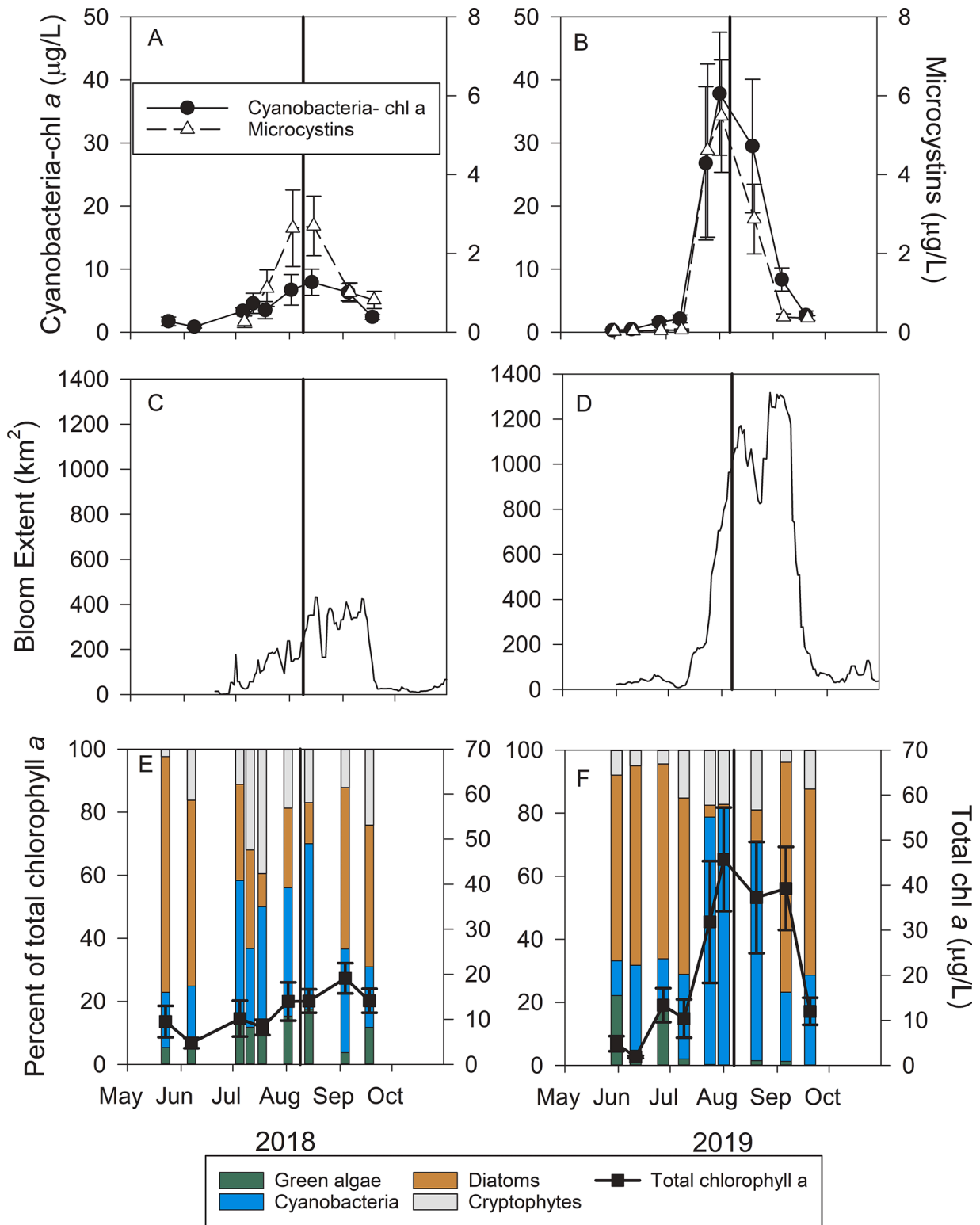


Fig. 1. Time series of cyanobacterial biomass and microcystins (A & B), 14 day rolling average bloom spatial extent (C & D), and total phytoplankton biomass and the relative abundance of green algae, cyanobacteria, diatoms, and cryptophytes (E & F) in western Lake Erie on 9 August 2018 and 7 August 2019. The values in A, B, E, and F are the mean of five sites (± 1 standard error) monitored by the University of Toledo’s Lake Erie Center. The bold vertical line in each indicates the dates of the HABs Grab event and shows the event occurred during peak bloom conditions each year.

distance from Maumee Bay. There was a ‘finger’ of higher chl a concentrations that extended northward in the center of the basin into Ontario waters. Cyanobacteria-specific chl a measured by the FluoroP-probe showed nearly identical spatial pattern as the DMF method (Supplemental Fig. 3).

Microcystins (measured by ELISA) during 9 August 2018 were highest along the Ohio shoreline and around the Bass Islands on the eastern edge of the western basin ranging from 2 to 5 µg/L (Fig. 2), and MCs concentrations were lower in the center of the basin. In the 7 August 2019 HABs Grab the highest MCs concentrations, 15 to 50 µg/L, were

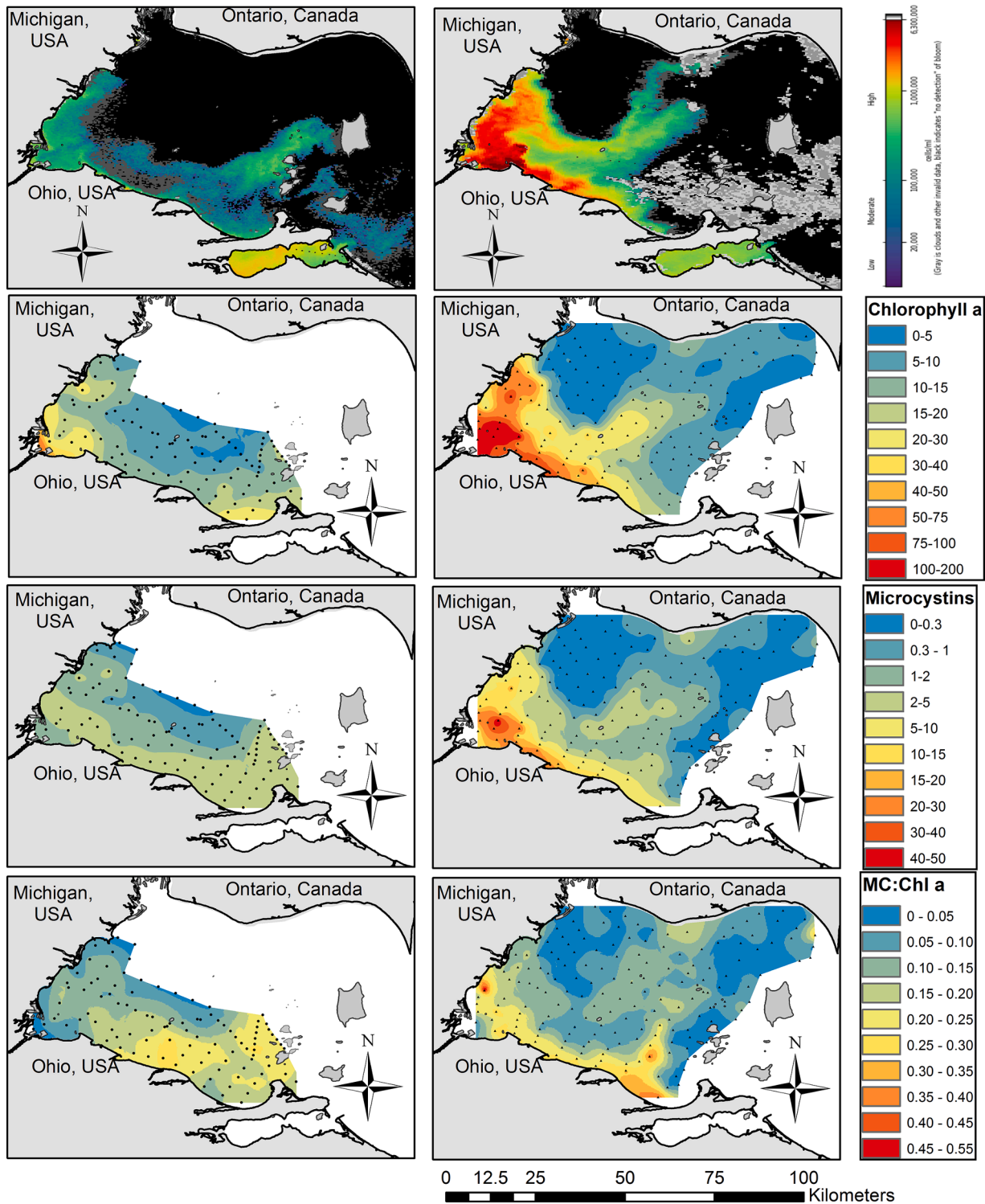


Fig. 2. Cyanobacterial biomass and microcystins during the 9 August 2018 (left column) and 7 August 2019 (right column) HABs Grab. Top row: NOAA Cyanobacterial Index taken on 5 August 2018 and 7 August 2019. The heat map ranges from 20,000 cells/mL in dark blue to 6300,000 cells/mL in dark red. Black is cells not detected and gray is cloud cover. Second row: Chlorophyll *a* ($\mu\text{g/L}$, top). Third row: total microcystins measured by ELISA ($\mu\text{g/L}$, middle). Bottom row: the ratio of microcystins to chlorophyll *a*. Small dots on the maps represent the sample collection locations.

measured in Maumee Bay (western corner of the basin) and decreased with distance from Maumee Bay. Microcystins were below detection ($<0.3 \mu\text{g/L}$) in the northwest portion of the basin where the Detroit River outflows into Lake Erie and below detection around the Bass Islands. Likewise, there was a ‘finger’ of MCs ranging from 1 to 5 $\mu\text{g/L}$ in the center of the basin, reaching into Ontario waters.

The ratio of total MC to chlorophyll concentrations (MC:chl *a*) was not consistent across the basin, and the spatial pattern was not the same between years. On 9 August 2018 HABs Grab there were ‘hot spots’ with MC:chl *a* ratios greater than 0.20 in the south-center portion of the basin and around the islands (Fig. 2). Lower ratios were recorded in Maumee Bay and the northern half of the basin. On the 7 August 2019 HABs Grab,

the majority of the western basin had MC:chl *a* ratios less than 0.15, and hot spots greater than 0.20 were confined to within ~10 km from the Ohio shoreline and within the ‘finger’ of higher biomass extended northward in the center of the basin (Fig. 2).

Percent of the *Microcystis* population capable of producing MCs ranged from 15.8% to 34.0%, and the overall average was 21.9% (Fig. 3). Both MC genes, *mcyD* and *mcyE*, gave very similar results (Fig. 3). Nineteen of the 25 samples were less than 25%. The percent toxigenic *Microcystis* did not correlate with distance from the Maumee River or with any environmental parameter measured. Overall, this indicated that less than one-quarter of the *Microcystis* cells during the 9 August 2018 HABs Grab were capable of producing MCs.

3.3. Water currents

Simulations show that daily average circulations in the surface mixed layer on both HABs Grab days were characterized by a general pattern of west-to-east flow exiting the basin (Michalak et al., 2013; Schwab et al., 2009). Detailed flow patterns reveal how the spatial variability of the observed blooms was influenced by hydrodynamic transport. Stronger flow occurred during the 9 August 2018 HABs Grab (Fig. 4A). The water mass associated with the Detroit River flowed southward and could reach the central part (41.7 °N – 41.8 °N) of the basin before turning eastward and exiting through the north and middle channels divided by Pelee Island. Accordingly, the water mass and cyanobacteria associated with the Maumee River were compressed around the west coast out of Maumee Bay and moved along the southern part of the basin.

During the 7 August 2019 HABs Grab, the circulation was weaker, which led to less eastward hydrodynamic transport of the cyanobacteria and allowed for longer residence time in the basin (Fig. 4B). The majority of Detroit River water was more constrained to the northern part of the basin with an anti-clockwise turn to the northeast to exit through the north channel. Correspondingly, the weakened southward intrusion allowed the water mass and cyanobacteria associated with the Maumee River to extend further north, compared to the bloom pattern during the 9 August 2018 HABs Grab. The bloom was distributed mostly in the southern and central parts of the basin. A portion of the bloom was mixed into the anti-clockwise circulation in the northern part of the basin described above, leaving a ‘finger’ of bloom biomass pointing

towards Canada and Pigeon Bay.

3.4. Nutrient concentrations

Except for dissolved inorganic N (DIN), higher concentrations of nutrients were observed on 7 August 2019 (Fig. 5). DIN concentrations on 9 August 2018 ranged from 40 to 60 $\mu\text{mol/L}$ in Maumee Bay, and concentrations near the Michigan and Ohio shorelines (20–40 $\mu\text{mol/L}$) were greater than the center of the basin (15–20 $\mu\text{mol/L}$). On 7 August 2019, DIN concentrations were highest near Little Cedar Point (20–40 $\mu\text{mol/L}$) and were less than 20 $\mu\text{mol/L}$ in the majority of the western basin. The lowest DIN concentrations (5–10 $\mu\text{mol/L}$) were recorded in the center of the basin within the ‘finger’ of biomass extending into Ontario waters. Nitrate made up the majority of the measured DIN. Ammonium concentrations on 9 August 2018 were less than 2 $\mu\text{mol/L}$ nearshore and less than 1 $\mu\text{mol/L}$ in the center of the basin. On 7 August 2019, ammonium concentrations between 6 and 8 $\mu\text{mol/L}$ were measured in Maumee Bay, and concentrations decreased with increasing distance from Maumee Bay.

Total N concentrations (the sum of nitrate, nitrite, and TKN) in 9 August 2018 followed the same spatial pattern as DIN concentrations. Total N in 7 August 2019 was greatest in Maumee Bay (150–275 $\mu\text{mol/L}$) where the highest levels of cyanobacteria biomass occurred, and TN concentrations were lower (25–50 $\mu\text{mol/L}$) in the rest of the basin.

In both years, total P concentrations were greatest in Maumee Bay and decreased with increasing distance from the bay. Total P concentrations were greater during 7 August 2019 in Maumee Bay and the rest of the basin. Dissolved reactive P concentrations were less than 0.1 $\mu\text{mol/L}$ in most samples in both years and often below detectable levels (<0.03 $\mu\text{mol/L}$).

The ratio of total N to total P concentrations (TN:TP) exceeded 30 (by moles) during both HABs Grabs. On the 9 August 2018 HABs Grab, TN:TP was between 75 and 100 for most of the area sampled. On 7 August 2019, most of the southern half of the basin had TN:TP between 50 and 125, whereas the northern half of the basin had much higher TN:TP of 150 to 520. The higher TN:TP in the northern half of the basin was due to lower TP concentrations.

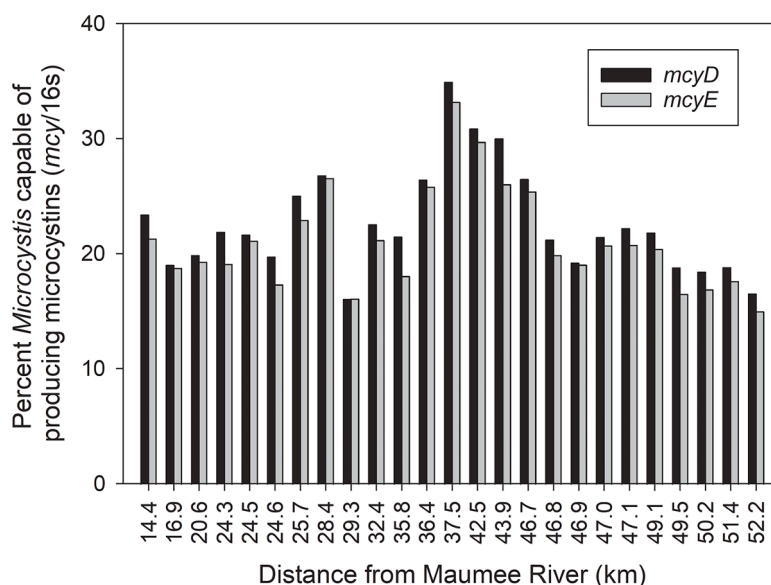


Fig. 3. The percentage of the *Microcystis* population capable of producing microcystins (relative abundance of *mcy* genes counts normalized to the gene counts of the *Microcystis*-specific 16S derived from shotgun metagenome read-mapping) during the 9 August 2018 HABs Grab as a function of distance from the mouth of the Maumee River.

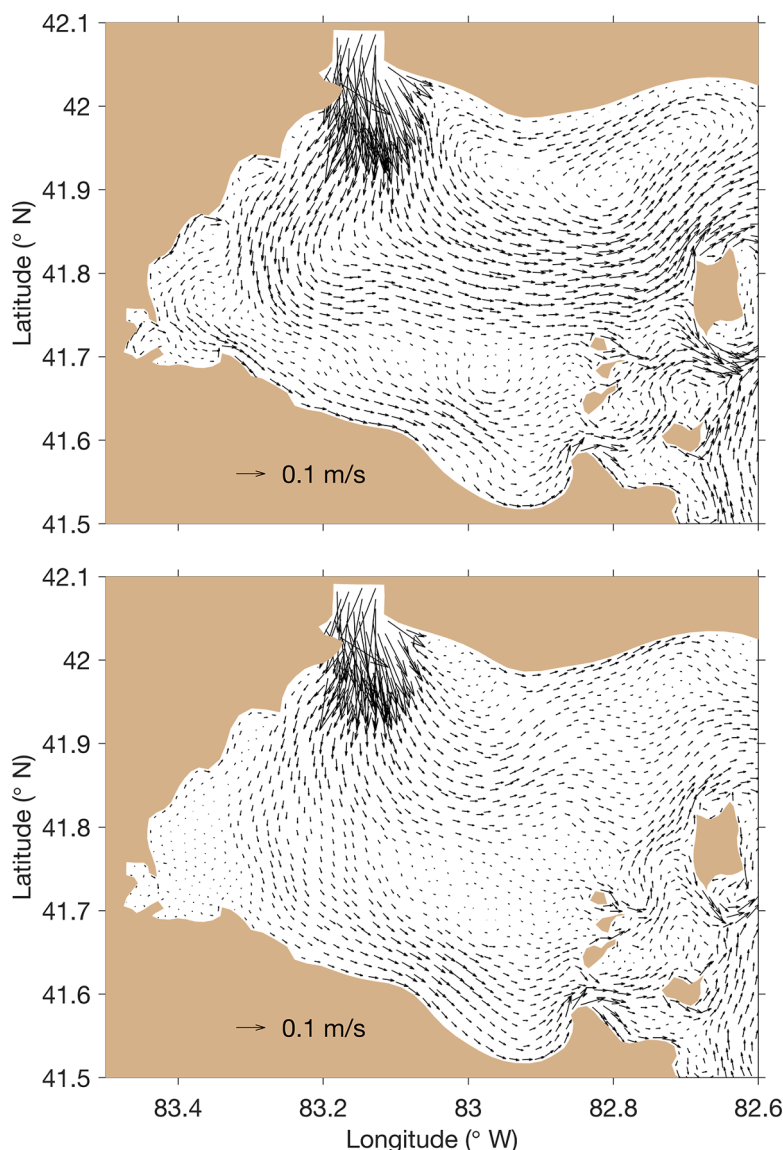


Fig. 4. The hydrodynamic conditions of the western basin during the 9 August 2018 (top) and 7 August 2019 HABs Grab as simulated with the Finite Volume Community Ocean Model (FVCOM).

3.5. Total MC mass and average concentration

The total mass of MCs (measured by ELISA) in the western basin on 7 August 2018 was estimated as 11,513 kg, and the total mass on 9 August 2019 was estimated as 30,691 kg (Table 1). The MC mass estimated assumed the MC concentration measured in the 0–2 m sample was representative of the unsampled water deeper than 2 m (see methods). The average MC concentration in samples with detectable MCs ($> 0.30 \mu\text{g/L}$) was $1.94 \mu\text{g/L}$ and $5.02 \mu\text{g/L}$ on the 2018 and 2019 HABs Grabs, respectively. The approximate area of the western basin is 2780 km^2 and the approximate mean depth is 8 m, which gives a volume of $2.224 \times 10^{10} \text{ m}^3$. The basin-wide average MC concentration would have been $0.52 \mu\text{g/L}$ and $1.38 \mu\text{g/L}$ on 9 August 2018 and 7 August 2019, respectively, if the cyanoHAB was evenly spread out throughout the basin.

3.6. Metrics comparison and correlations to microcystins

One sample on 9 August 2018 that was collected near the mouth of the Maumee River was removed from the correlation matrix because it had chl values two to three times greater than the second-highest sample

(depending on metric) due to high levels of green algae and diatoms. Nearly all cyanobacterial biomass metrics significantly ($p < 0.05$) correlated with each other, but more importantly, r values (strength of correlation) ranged from 0.43 (very weak correlation) to 0.99 (very strongly correlated), and correlation coefficients often differed between years within a parameter (Table 2). For this section, only the r values greater than 0.70 are highlighted (strongly correlated). The handheld YSI chl RFU strongly correlated with 2018 PC RFU and with 2018 FluoroProbe total chl a , but no correlations in 2019 had a r value greater than 0.70. It is important to note that 2018 had greater chl RFU values, but lower filter-extracted chl a concentrations than 2019. Phycocyanin RFU strongly correlated with 2019 FluoroProbe cyanobacteria-chl a , FluoroProbe total chl a in both years, and with 2019 filter-extracted chl a . Of special note, chl RFU and PC RFU did not strongly correlate with their respective traditional laboratory methods. The FluoroProbe-estimated cyanobacteria-specific chl a very strongly correlated with FluoroProbe total chl a and filter-extracted chl a and, in 2019 only, with cyanobacteria 16S gene copies, PC RFU, and *mcyE* gene copies. The FluoroProbe-estimated total chl a concentrations were strongly correlated with filter-extracted chl a ($r = 0.90$ and 0.98), and with cyanobacteria-specific, PC RFU, and 2019 chl RFU. Filter-extracted-chl a

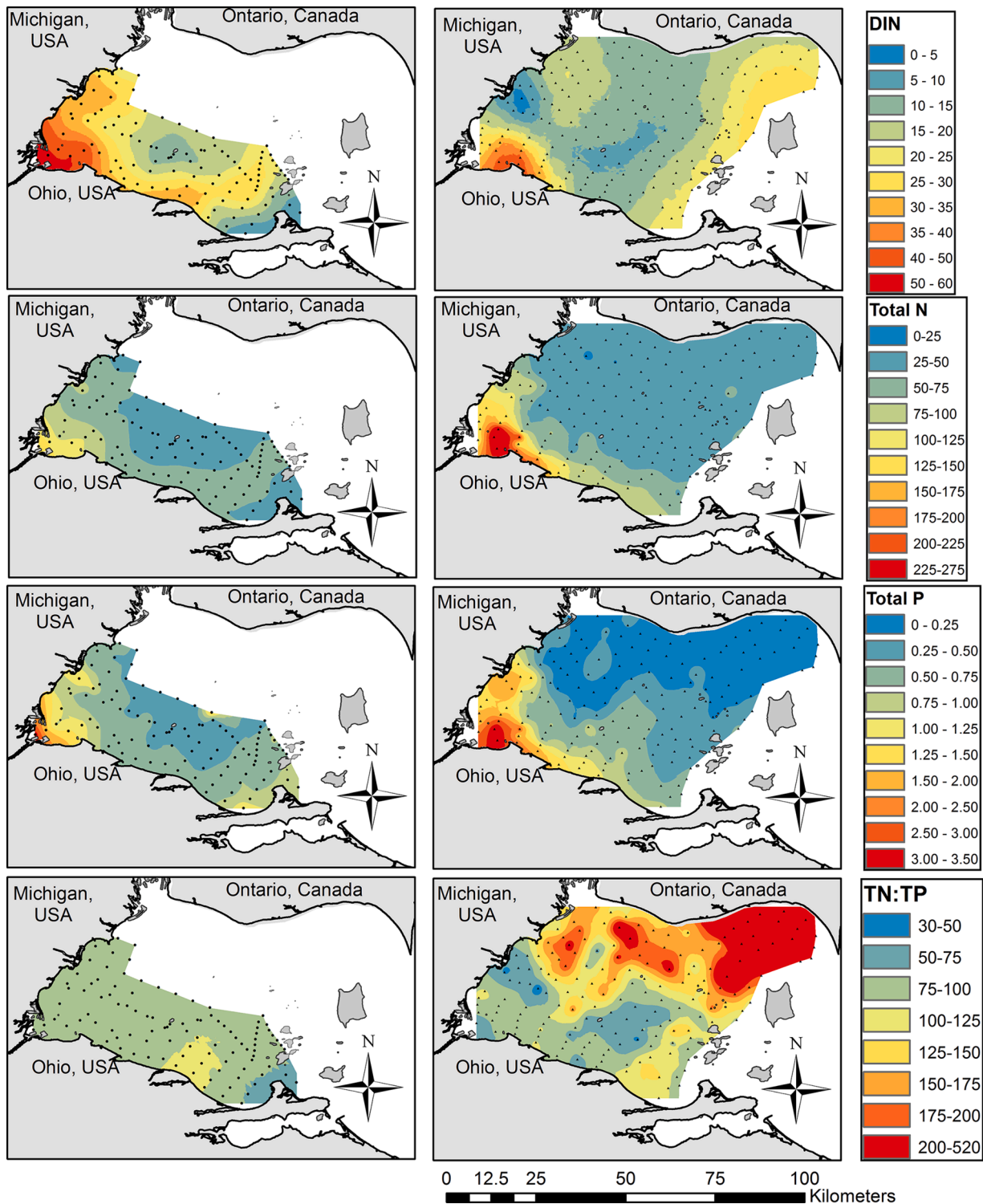


Fig. 5. Nutrient concentrations ($\mu\text{mol/L}$) measured during the HABs Grabs. Dissolved inorganic nitrogen (sum of nitrate, nitrite, and ammonium; top), total nitrogen (second row), total phosphorus (third row), and the ratio of total N to total P concentration (bottom) of the 9 August 2018 (left) and 7 August 2019 (right) HABs Grabs. Small dots on the maps represent the sample collection locations.

concentrations were very strongly correlated with both FluoroProbe parameters and cyanobacterial 16S gene copies, and strongly correlated with 2019 PC RFU and 2019 *mcyE* gene copies. Filter-extracted-PC concentrations did not strongly correlate with any parameter. Cyanobacterial 16S gene copies very strongly correlated with both FluoroP-probe parameters, filter-extracted chl *a*, and *mcyE* gene copies. Gene copies of *mcyE* measured during 2019 strongly correlated with both

FluoroProbe parameters and filtered-extracted chl *a*, but *mcyE* did not correlate with any parameter in 2018.

On 9 August 2018 total MCs concentrations measured with the ELISA method correlated with cyanobacteria-chl *a* and chl *a* extracted from a filter. On 7 August 2019 MCs strongly correlated with every biomass metric to some degree ($r > 0.71$) except for filter-extracted PC, and 2019 MCs were very strongly correlated with cyanobacteria-specific chl *a*,

Table 1

Total microcystins mass (kg) in the western basin of Lake Erie during the two HABs Grabs events, the maximum and average (standard deviation in parenthesis) MC concentrations (µg/L) measured in water samples collected during the HABs Grab on 9 August 2018 and 7 August 2019, and theoretical basin-wide average concentration. The microcystin data were from the ELISA method.

	Total MCs (kg)	Max MC measured	Average MC in samples		Theoretical basin-wide
			All samples	In samples with detectable MC	
2018	11,513	6.38	1.81 (1.21)	1.94 (1.16)	0.52
2019	30,691	46.56	3.50 (6.77)	5.02 (7.70)	1.38

FluoroProbe total chl *a*, and filter-extracted chl *a* ($r = 0.90 - 0.92$). Extracellular MCs concentrations correlated only with cyanobacterial 16S gene copies.

Total MCs measured by ELISA ranged from less than detectable levels (< 0.15) to 46.56 µg/L, whereas the LC-MS/MS sum of 12 congeners ranged from 0.01 to 26.53 µg/L (Fig. 6). In both years, the four MC congeners found in the highest concentrations were MC-LR (44.4%, 35.9% in 2018, 2019, respectively), RR (23.1%, 33.6%), LA (16.7%, 13.5%), and YR (13.6%, 15.7%), and the other congeners analyzed for were collectively usually less than 3% (Supplemental Fig. 4). Despite the differences in concentrations, the two methods were highly correlated. Linear regressions conducted separately for each year gave similar slopes (0.5125 for 2018 and 0.5462 for 2019), and the R² was 0.79 for 2018 and 0.95 for 2019. This indicates that MCs concentrations measured by ELISA were approximately twice as high as measured by LC-MS/MS.

3.7. HAB biomass and MC correlations with environmental parameters

Filter-extracted chl *a* concentrations were selected for comparison with the environmental parameters due to the high degree of correlation with the other cyanobacterial biomass metrics (Table 2). Cyanobacteria-specific chl *a* measured with the FluoroProbe could have been displayed

Table 2

Pearson correlation coefficient (*r*) matrix among metrics used to estimate cyanobacterial biomass during the 9 August 2018 and 7 August 2019 HABs Grab and the range of values observed. Strong ($r = 0.71$ to 0.90) and very strong ($r > 0.90$) correlations between two parameters are bolded. “NS” = not significant correlation ($P > 0.05$). ND = no data was recorded. *One 2018 outlier sample was removed from the correlation that had chl values approximately 2 to 3 times greater than the highest sample reported in the last column.

	Year	Chl RFU	PC RFU	Cyano. Chl <i>a</i>	Total chl <i>a</i>	Chl <i>a</i>	PC	16S	<i>mcyE</i>	Total MCs	Exc. MCs	Range of values*
Chlorophyll RFU - YSI	2018	1	.85	.51	.75	.54	ND	ND	NS	NS	-0.21	0.30 - 3.67
	2019	1	.66	.65	.68	.70	.43	.66	.63	.71	.43	0.05 - 3.85
Phycocyanin RFU - YSI	2018	.85	1	.57	.76	.61	ND	ND	NS	NS	-0.35	-0.06 - 2.90
	2019	.66	1	.77	.78	.74	.55	.62	.65	.78	.48	0.00 - 21.13
Cyanobacteria chlorophyll <i>a</i> µg/L - FluoroProbe	2018	.51	.57	1	.86	.94	ND	ND	NS	.73	NS	0.28 - 11.15
	2019	.65	.77	1	.99	.98	.54	.92	.80	.90	.60	0.43 - 110.40
Total chlorophyll <i>a</i> µg/L - FluoroProbe	2018	.75	.76	.86	1	.90	ND	ND	NS	.48	NS	2.82 - 20.99
	2019	.68	.78	.99	1	.98	.53	.92	.79	.90	.58	3.08 - 132.63
Chlorophyll <i>a</i> µg/L - filter extracted	2018	.54	.61	.94	.90	1	ND	ND	NS	.72	NS	3.50 - 31.76
	2019	.70	.74	.98	.98	1	.51	.95	.83	.92	.60	1.88 - 196.16
Phycocyanin µg/L - filter extracted	2018	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	No data
	2019	.43	.55	.54	.53	.51	1	.49	.55	.52	.42	0 - 18.22
Cyanobacteria 16S gene copies/L - qPCR	2018	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	No data
	2019	.66	.62	.92	.92	.95	.49	1	.97	.89	.89	$3.3 \times 10^7 - 7.5 \times 10^8$
Cyanobacteria <i>mcyE</i> gene copies/L - qPCR	2018	NS	NS	NS	NS	NS	ND	ND	1	.40	.50	$0 - 4.7 \times 10^8$
	2019	.63	.65	.80	.79	.83	.55	.97	1	.83	.55	$3.1 \times 10^4 - 2.8 \times 10^7$
Total Microcystins µg/L - ELISA	2018	NS	NS	.73	.48	.72	ND	ND	.40	1	.24	<0.1 - 6.38
	2019	.71	.78	.90	.90	.92	.52	.89	.83	1	.62	<0.1 - 46.56
Extracellular Microcystins µg/L - ELISA	2018	-0.21	-0.35	NS	NS	NS	ND	ND	.50	.24	1	<0.1 - 0.75
	2019	.43	.48	.60	.58	.60	.42	.89	.55	.62	1	<0.1 - 1.38

here, which gave very similar patterns as total chl *a*, but we chose total chl *a* owing to the greater number of samples, and FluoroProbe data are less common than filter-extracted chl *a*. Cyanobacteria-specific chl *a* metrics are displayed in the supplemental document.

MCs increased with filter-extracted chl *a* concentration on 7 August 2019 but not 9 August 2018 (Fig. 7A); however, in 2018 MCs did increase with cyanobacteria-chl *a* (Supplemental Fig. 5). The ratio of MCs to chl *a* (MC:Chl *a*) ranged from near 0 (MCs lower than detection) to 0.31 for all samples but three (Fig. 7b). On 9 August 2018, there was no relationship between MC:Chl *a* and cyanoHAB biomass (as chl *a* concentration). On 7 August 2019, the minimum MC:Chl *a* increased with increasing chl *a* concentration. For example, MC:Chl *a* ranged from 0 to 0.3 for chl *a* concentration less than 10 µg/L, but MC:Chl *a* ranged from 0.15 to 0.3 for chl *a* concentrations greater than 50 µg/L. Regarding the MCs to cyanobacteria-specific chl *a* ratio (MC:cyanobacteria-chl *a*), 7 August 2019 was very similar to MC:Chl *a* because cyanobacteria were the majority of the chlorophyll (Supplemental Fig. 5). However, on 9 August 2018, most samples had a MC:cyanobacteria-chl *a* between 0.1 and 0.6, but MC:Chl *a* ranged from near 0 to 0.3. Green algae and diatoms were in relatively greater concentrations during the 2018 HABs Grab, and their contribution to total chl *a* decreased MC:Chl *a*.

The correlations between MCs and chl *a* concentration with environmental parameters showed similar patterns (Figs. 8 and 9), which was due to the high correlation between MCs and chl *a* (Table 2). The highest MCs and chl *a* concentrations were recorded in lake water temperatures between 25 °C and 27 °C (Fig. 8). Higher pH values were observed on 7 August 2019 than 9 August 2018, and on 7 August 2019 highest MCs and chl *a* concentrations occurred at pH greater than 9.0, but low concentrations of both MCs and chl *a* were also observed at high pH values. MCs and chl *a* increased with turbidity, but this pattern was expected because algal biomass is captured in turbidity measurements. On 7 August 2019, the highest concentrations of MCs and chl *a* concentrations occurred at specific conductivity between 270 and 320 µS/cm, which reflects water closest to the Maumee River’s mouth. The toxin-to-biomass ratio, MC:Chl *a*, did not correlate with temperature, turbidity, pH, or specific conductivity. The residuals between MC vs. chl

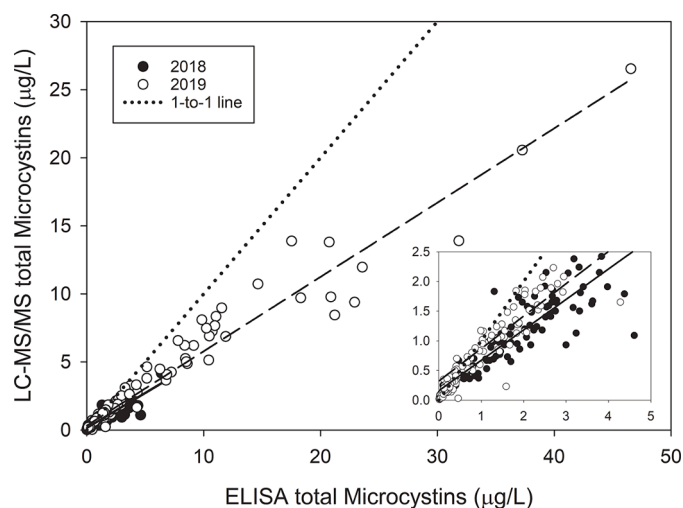


Fig. 6. The relationship between total microcystins measured by ELISA and the calculated total microcystins from 13 different congeners during the 9 August 2018 (filled circles, solid regression line) and 7 August 2019 (open circles, dashed regression line) HABs Grab. The dotted line is a 1-to-1 line. The inset panel shows a zoomed in view of the lower range where most of the 2018 samples occurred. 2018 regression equation: $LC-MS/MS = (0.5125 * ELISA) + 0.1596$, $R^2 = 0.79$. 2019 regression equation: $LC-MS/MS = (0.5462 * ELISA) + 0.3229$, $R^2 = 0.95$.

a were centered around 0 throughout the range of the observed physical parameter, indicating no relationship.

Regarding nutrients, MCs and chl *a* concentrations increased linearly with TKN, TN, and TP concentrations (Fig. 9). MCs measured on 7 August 2019 also increased with DIN, but that pattern was not observed on 9 August 2018. The highest concentrations of MCs and chl *a* concentrations occurred at TN:TP concentration ratios between 50 and 100 (molar). The MC:chl *a* did not correlate with any N or P parameter during 9 August 2018. On 7 August 2019, the majority of lower toxin-to-biomass ratios (MC:chl *a* < 0.2) occurred at low DIN (< 20 µmol/L) and low TP (< 1 µmol/L) concentrations, and higher concentrations had greater MC:chl *a*. For TKN concentrations less than 75 µmol/L, MC:chl *a* ranged from 0.05 to 0.4, but MC:chl *a* increased with increasing TKN concentrations greater than 75 µmol/L. The residuals between MC vs. chl *a* were centered around 0 across the range of TKN, TN, TP, and TN:TP. The residuals increased with DIN concentration on 7 August 2019 ($R^2 = 0.32$). The regression line crossed the DIN axis at 17.9 µmol/L, which indicates samples collected from waters with DIN concentrations less than 17.9 µmol/L had less MC per chl *a* compared to samples collected from higher DIN concentrations.

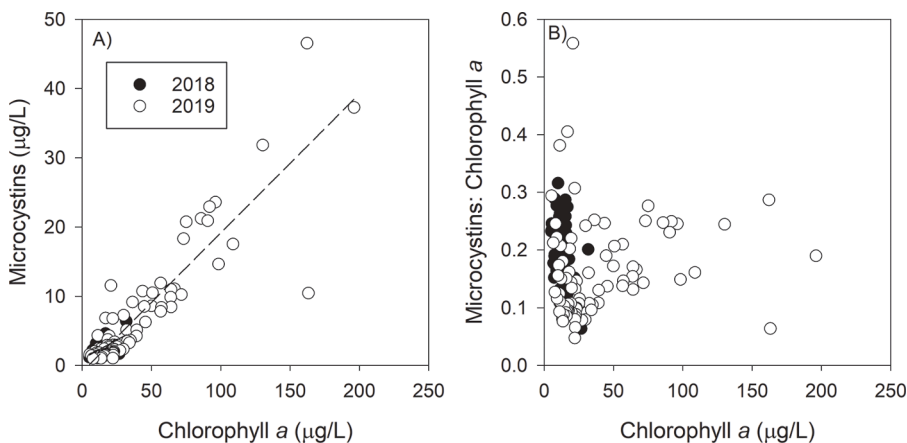


Fig. 7. Total microcystins (measured by ELISA) concentration (A) and the microcystins-to-chlorophyll ratio (B) as a function of chlorophyll *a* concentration for each year. 9 August 2018 regression equation: $MC = (0.109 * Chl) + 0.955$, $R^2 = 0.32$. 7 August 2019 regression equation: $MC = (0.201 * Chl) - 0.927$, $R^2 = 0.80$. During the large bloom year of 2019, higher chlorophyll *a* concentrations corresponded to high toxin-to-biomass ratios, but high toxin-to-biomass ratios were also observed at low chlorophyll *a* concentrations.

4. Discussion

4.1. Comparison to other years

The tremendous inter-annual variation of cyanoHAB biomass in Lake Erie is primarily driven by springtime (March-July) P load from the Maumee River (Stumpf et al., 2016b), and cyanoHAB biomass in 2018 and 2019 followed the expected pattern. Much higher biomass was recorded during 2019 than in 2018 (Fig. 1), which corresponded to heavy springtime rainfall and nutrient loading. Despite the difference in annual cyanoHAB biomass, temporal patterns are usually consistent from year to year, with cyanobacteria first appearing in Maumee Bay during July, peaking between mid-August to early September, and decreasing throughout the autumn (Binding et al., 2021; Bridgeman et al., 2013; Stumpf et al., 2016b), although the very large 2011 bloom peaked in mid-October (Binding et al., 2012; Stumpf et al., 2012). Overall, the cyanoHABs during 2018 and 2019 were typical blooms and followed the expected patterns based on previous years. Furthermore, the 2018 and 2019 HABs Grabs occurred during the peak biomass of each year.

Analysis of random shotgun sequencing data and quantification of the microcystin-producing (*mcy*) gene and *Microcystis* 16S rRNA genes showed that, on average, only 22.6% and 21.1% of the *Microcystis* cells (based on *mcyD* and *mcyE*, respectively) present during the 9 August 2018 HABs Grab had the potential to produce MCs (Fig. 3). There was close agreement between data from *mcyD* and *mcyE*, supporting the precision of the method. The percentage of potential MC-producers was not related to distance from the Maumee River, which agrees with previous studies that the *Microcystis* populations in the river are separate from those in the lake (Kutovaya et al., 2012). The percentage of MC-producers could have looked different if we conducted the HABs Grab earlier or later during the bloom's progression. Potential MC-producers inoculate the water column sooner than non-MC-producing strains (Kitchens et al., 2018), which suggests the percentage of potential MC-producers could have been higher if we sampled earlier in the bloom. However, the percentage of potential MC-producers could have been lower later in the bloom when N and light become limiting (Chaffin et al., 2013), and the competitive balance between MC-producing and non-MC-producing strains is shifted towards the latter strains (Davis et al., 2010; Kardinaal et al., 2007). Furthermore, N-limitation and the shift to non-MC-producing strains manifests in lower MC to biomass ratios in late summer and fall (Gobler et al., 2016; Horst et al., 2014).

4.2. Microcystin mass notes

The HABs Grabs surveys allowed us to estimate, with reasonably high confidence, 11,513 kg and 30,691 kg of MCs in the western basin

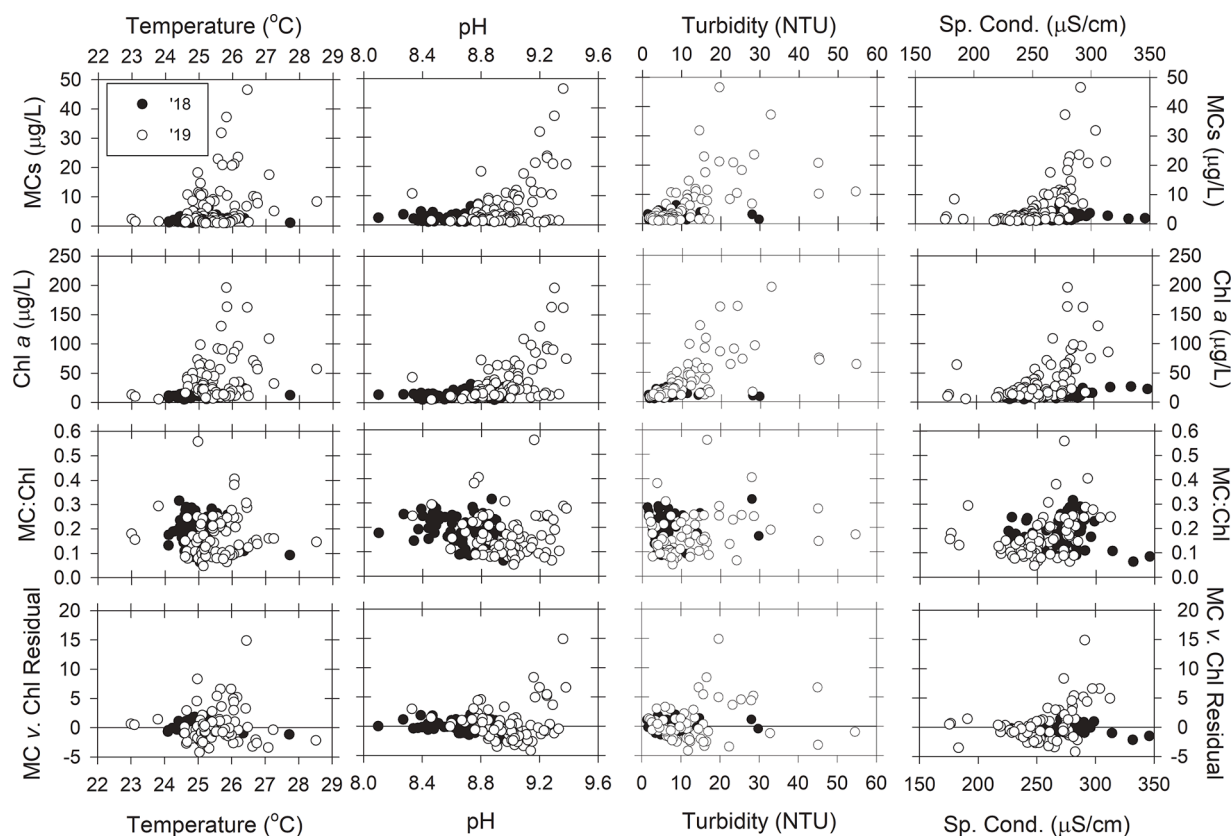


Fig. 8. The relationships between microcystins (ELISA), chlorophyll *a*, the ratio of microcystins to chlorophyll *a* concentration, and residuals from the microcystins *v.* chlorophyll regression (Fig. 7A) with water temperature, pH, turbidity, and specific conductivity during the 9 August 2018 (black circles) and 7 August 2019 (white circles) HABs Grab for samples with greater than 1 µg/L of total microcystins.

on a single day during peak cyanoHAB conditions on 9 August 2018 and 7 August 2019, respectively. The uncertainty associated with these estimates would be higher with fewer samples due to the spatial heterogeneity of biomass and MC concentrations. The cyanoHAB bloom boundary is particularly problematic for these estimates. For example, in the northwest portion of the 7 August 2019 cyanoHAB (Michigan shore), only 5 km separated high biomass (chl *a* > 50 µg/L) and high MCs (> 5 µg/L) from low biomass (chl *a* < 5 µg/L) and MC below detection (< 0.3 µg/L). A rapid gradient was observed in Maumee Bay between the highest biomasses and MC concentrations and intermediate levels over a similarly short distance. Without the high number of samples with high spatial resolution, we would have missed these cyanoHAB edge gradients and would have had lower-confidence in MC total mass estimates. The strong spatial gradients in cyanoHAB biomass and MCs and the potential for missing fine-scale variability with discrete sampling further demonstrate the value of synoptic high-resolution satellite observations in capturing these dynamic events.

The revelation of tons of MCs in the western basin from the HABs Grab events can be alarming, especially to managers and the general public, without the proper context. The high mass of MCs not only reflects bloom expanse but also reflects the large area (~2780 km²), and hence large volume, of the western basin. Total mass estimates are useful for researchers studying cyanoHAB toxin dynamics and the relationships to environmental triggers on an ecosystem scale. However, local MC concentrations are more important for managers and the public because concentrations are associated with human health risks. For example, if the cyanobacterial biomass and MCs were spread throughout the basin by stronger currents, the basin-wide MC concentration would have been 0.52 µg/L or 1.38 µg/L, on 9 August 2018 and 7 August 2019, respectively. It is critically essential to message these results in the proper context.

4.3. Microcystin to biomass ratio

The HABs Grab dataset is the largest single-day characterization of the Lake Erie cyanobacteria-dominated blooms. While there have been previous studies in Lake Erie that presented relationships between MC and cyanobacterial biomass with environmental parameters, those studies grouped data collected from a few sites and throughout the growing season (Horst et al., 2014; Millie et al., 2009; Rinta-Kanto et al., 2009; Wang et al., 2009). Studies that group data from many dates overlook how the relationship between two parameters may change throughout the seasons, which may skew trends and result in larger uncertainty. The relationships presented here from the one-day surveys omit the temporal impacts among parameters; however, these relationships would likely be different if the HABs Grabs were conducted later in the year when P and N become co-limiting to cyanobacterial growth and MC production, and there is a shift towards non-MC-producing strains (Gobler et al., 2016).

Recently, Liu et al. (2020) used the MC:chl *a* ratio to hindcast MC concentrations from satellite-derived chl *a* concentrations for Lake Erie 2009 – 2016. Liu et al. (2020) stated that MC:chl *a* varied more temporally than spatially; however, our HABs Grab surveys showed that MC:chl *a* was not consistent spatially. MC:chl *a* ranged from 0.06 to 0.32 across the entire western basin during both HABs Grabs when MC were detected at concentrations greater than 1.0 µg/L, a range that is comparable to the seasonal variability of MC:Chl *a* documented in Liu et al. (2020). Furthermore, waters within a few kilometers of the Ohio shoreline, which serve as sources of drinking water for several municipalities, had the highest MC:chl *a* ratios. Thus, forecasting MC concentrations based on chl *a* concentrations and an assumed constant MC:chl *a* ratio could result in modeled MC concentrations that are off by a factor of up to 5.3 during the early stages of the peak bloom.

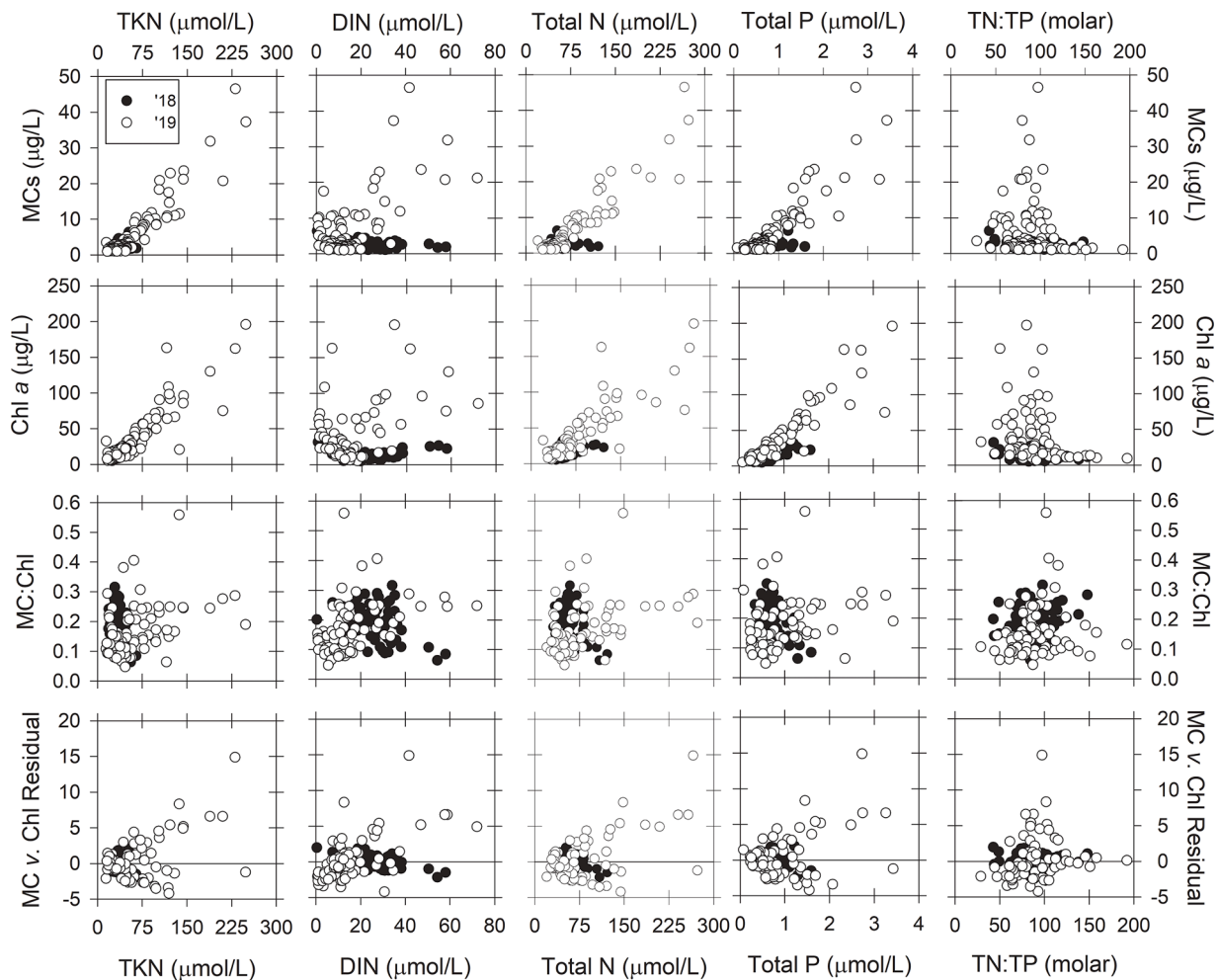


Fig. 9. The relationships between microcystins (ELISA), chlorophyll *a*, the ratio of microcystins to chlorophyll *a* concentration, and residuals from the microcystins *v.* chlorophyll regression (Fig. 7A) with total Kjeldhal nitrogen, dissolved inorganic nitrogen, total nitrogen, total phosphorus, and the total nitrogen-to-total phosphorus ratio during the 9 August 2018 (black circles) and 7 August 2019 (white circles) HABs Grab for samples with greater than 1 $\mu\text{g/L}$ of total microcystins.

During the large bloom year of 2019, the MC:chl *a* ratio did appear to have some dependency on cyanobacteria biomass (as measured by chl *a*). As biomass increased, so did the minimum MC:chl *a* per chl *a* concentration. For example, at chl *a* concentrations of 25 $\mu\text{g/L}$, MC:chl *a* ranged from 0.05 to 0.30, but at chl *a* concentrations from 50 to 200 $\mu\text{g/L}$, MC:chl *a* ranged from 0.15 to 0.30. Thus, high cyanobacterial biomasses were more likely to be associated with higher MC concentrations that would warrant a recreational contact advisory. Nevertheless, lower biomasses may also warrant the advisories if the MC:chl *a* is on the high side of the range. For example, the range of predicted MC concentrations for chl *a* concentrations of 25 $\mu\text{g/L}$ and 50 $\mu\text{g/L}$ would be 1.25 to 7.5 $\mu\text{g/L}$ and 7.5 to 15 $\mu\text{g/L}$, respectively. In the state of Ohio, recreational advisories are posted for MC concentrations exceeding 6 $\mu\text{g/L}$. The uncertainty associated with MC:chl *a* at chl *a* concentration of 25 $\mu\text{g/L}$ would give predicted MC concentrations both above and below Ohio's recreational guideline, but a higher chl *a* concentration of 50 $\mu\text{g/L}$ would exceed Ohio's recreation guideline under the entire MC:chl *a* range observed in this study. In terms of drinking water treatment, the large uncertainty in MC:chl *a* makes it difficult for water treatment plant operators to estimate MC concentrations for the optimization of treatment. If managers estimate MC exposure risk from models based on MC:chl and chl *a* concentrations, it would be wise to take a more conservative approach and assume the high range of projected MC concentrations. The difficulty in estimating MC concentration from chl *a* highlights the need for rapid MC tests and a better understanding of the factors that regulate MC production.

Several environmental parameters were investigated as explanatory variables for the relationships between MC and chl *a* observed during our HABs Grab. The data collected on 9 August 2018, the low HAB biomass year, did not show any patterns or correlations, likely due to the high range of MC:chl *a* observed at low biomasses (as discussed above). On 7 August 2019, MC:chl *a* showed a negligible relationship with total P ($R^2 = 0.10$), which the tautologous relationship between chl *a* and total P could explain this trend. MC:chl *a* and the residuals of MC *v.* chl *a* showed better relationships with DIN ($R^2 = 0.18, 0.32$, respectively). The lowest within-bloom MC:chl *a* ratios occurred in the center of the basin (Fig. 2) where the DIN concentrations were less than 10 $\mu\text{mol/L}$ (Fig. 5), which is known to constrain MC production in *Microcystis* dominated blooms (Chaffin et al., 2018a; Gobler et al., 2016). Moreover, the residuals from the MC *v.* chl *a* plot showed less than expected MCs at similar DIN concentrations, further supporting N-limitation of MC synthesis (Fig. 9). So even at these larger spatial scales, these results were consistent with the prediction by the theory of ecological stoichiometry of phytoplankton toxins, which links the production of the N-rich MCs by cyanobacteria to N availability (Brandenburg et al., 2020; Van de Waal et al., 2014; Wagner et al., 2019). Furthermore, MC:chl *a* has been shown to decrease throughout the season (Chaffin et al., 2018b; Horst et al., 2014; Liu et al., 2020), and the decrease of MC:chl *a* paralleled drawdown of DIN concentrations (Chaffin et al., 2018b; Horst et al., 2014). Nitrate concentration data, or knowledge of how nitrate concentration changes throughout the season, could improve MC concentration predictions based on HAB biomass and the MC:chl *a* ratio.

Ammonium is often in low concentrations during cyanobacterial blooms (< 1 $\mu\text{mol/L}$) because uptake rates exceed regeneration rates (Blomqvist et al., 1994; Hampel et al., 2019), and there can be an inverse relationship between ammonium and MC concentrations (Kelly et al., 2019).

The MC:chl *a* and MC:cyanobacteria-chl *a* gave similar results for the 7 August 2019 HABs Grab because cyanobacteria dominated the phytoplankton community and green algae and diatoms were in low concentration. On the 9 August 2018 HABs Grab, MC:Chl *a* ranged up to 0.3, whereas MC:cyanobacteria-chl *a* was between 0.1 and 0.6. Green algae and diatoms were in relatively greater concentrations during the 2018 HABs Grab, and their contribution to total chl *a* decreased MC:Chl *a*. MC:cyanobacteria-chl *a* on 9 August 2018 was on average about double that of 7 August 2019 HABs Grab, which suggests the cyanobacteria on 9 August 2018 were more toxic than on 7 August 2019. However, year-to-year comparisons are difficult with one-day snapshots. Bloom progression, nitrogen availability, light climate histories, and water temperature (Gobler et al., 2016; Kardinaal et al., 2007; Kitchens et al., 2018; Martin et al., 2020) would have interacted to result in the MC to chl *a* ratio observed on the HABs Grab, and we would have observed different ratios if the HABs Grab were conducted earlier or later in the bloom.

4.4. CyanoHAB metric comparison

Finding that many cyanoHAB biomass metrics correlated was not surprising and should have raised alarms if they did not correlate. However, correlation coefficients differed between the two years, which is problematic for comparing data across multiple metrics and years. The two parameters that had the lowest correlations with other parameters were chl RFU from the handheld sonde and PC concentration extracted from filters. It is also important to note the chl RFU did not correlate with chl *a* concentrations and PC RFU did not correlate with PC concentrations, and previous studies have highlighted the lack of correlation between handheld sensors and traditional methods (Cotterill et al., 2019; Hodges et al., 2017). While handheld fluorometers and filter extractions quantify the same pigments, fundamental differences will affect the data. The physiological state of the cells will impact the amount of fluorescence per cell, increasing it under stressful conditions such as low nutrients and photo-inhibition (Campbell et al., 1998). The morphology of the cyanobacterial colonies (single cell, small colonies, large colonies) affects the fluorescence signals, especially when large colonies drift in front of the sensor (Hodges et al., 2017). Finally, the amount of PC and chl *a* per cell is variable, with PC being much more variable than chl *a*, and is affected by light climate history (Chaffin et al., 2012), nutrient status (Beardall et al., 2001; Kirk, 1994), and growth phase (Chang et al., 2012). That variability likely explains the result that chl *a* has stronger relationships with MC, 16S rRNA, and *mcyE* concentrations than does PC. Given how PC and chl *a* fluorescence signals and cellular content are affected by environmental parameters, care must be taken when comparing dataset with different pigment-based metrics.

Both ADDA-ELISA and targeted LC-MS/MS have been used to quantify concentration of MCs in natural waters with each platform having its own sets of strengths and weaknesses. The ADDA-hapten is a conserved amino acid in MCs and is responsible for the primary mode of toxicity by inhibiting protein phosphatase 1 and 2a. To make an ELISA have cross-reactivity with all MCs, the ADDA-hapten was used to create an ADDA antibody (Fischer et al., 2001). The disadvantage with the ADDA-ELISA is that the cross-reactivity with several prevalent MCs produces artificially high concentrations (Guo et al., 2017; Thees et al., 2019). The target LC-MS/MS platform provides the concentrations for twelve MCs with standards. The primary weakness is that there are hundreds of MC congeners (Spoof and Catherine, 2017), and the potential of not quantifying all the prevalent MCs is real. As in this study, several other studies report ADDA-ELISA to have a higher total MC concentration than the LC-MS/MS platform (Birbeck et al., 2019; Foss

and Aubel, 2015; Guo et al., 2017; Thees et al., 2019). Three explanations for this scenario are: 1) the standards are not available for MCs in the bloom and not quantitated by LC-MS/MS (Foss and Aubel, 2015), 2) degradation products such as ADDA tetrapeptide, ADDA, and linear chain are present and cross-react with ADDA-ELISA (Thees et al., 2019), and 3) several of the prevalent MCs with standards have cross-reactivities are higher than 100% (Fischer et al., 2001; Guo et al., 2017). The MC congeners detected during the HABs Grabs (Supplemental Fig. 4) agree with recent 2016–2017 Lake Erie MC congener studies, in which the four major MC congeners were MC-RR, MC-LR, MC-YR and MC-LA (Matson et al., 2020; Palagama et al., 2020). Over 21 MCs were present in 2016 and 2017 blooms (Palagama et al., 2020) and unknown MC congeners were reported in the 2015 bloom (Foss and Aubel, 2015), thus these data support that the ADDA-ELISA is detecting MC congeners that do not have commercially available standards. Since the MC congeners have a wide range of toxicity and have differing fate and transport because of the wide range of hydrophobicity, it is important to identify the prevalent MCs in Lake Erie and make standards available.

4.5. Lessons learned from the HABs Grab

The HABs Grab highlighted limitations of the current cyanoHABs monitoring network, but the collection and laboratory processing of 100 (or 172) water samples in one day is not feasible to conduct on a routine basis. The majority of water samples collected in Lake Erie for long-term monitoring of water quality are from fixed locations that are sampled at a variable frequency ranging from weekly to seasonally. While this type of sampling scheme produces useful temporal data, it lacks spatial resolution and could miss essential areas of the bloom, like the ‘finger’ of biomass and MCs extending into Canadian waters observed in 2019 (Fig. 2). There are several potential options to overcome the spatial resolution limitation. Obviously, remote sensing can be used as a proxy for cyanobacterial biomass data over large spatial areas, but it cannot measure MCs directly (Stumpf et al., 2016a) and the spatial heterogeneity of MC:Chl *a* documented here suggests there would be significant uncertainty introduced when using a fixed ratio to extrapolate MCs from Chl *a* distributions. Gliders and autonomous underwater vehicles (AUV), such as the third generation of the environmental sample processor (ESP), can collect data and samples in areas of the lake not routinely sampled and transmit the data wirelessly to researchers and managers (Anderson et al., 2019; Scholin et al., 2017). Volunteers and citizen scientists can be trained to collect and process samples. For example, NOAA’s phytoplankton monitoring network and OSU Stone Lab’s charter boat water sampling program are citizen science projects; however, these volunteers need their own boats to collect offshore samples. Finally, the advancement of drones may facilitate the collection of water samples from bloom hot spots or areas not monitored by scientists, but drone technology is still in its infancy (Lally et al., 2019). While these options to overcome limitations of spatial resolution can augment routine monitoring programs and remote sensing, they do not replace existing methods. Therefore, it is paramount to develop and validate models that can forecast MCs over a large spatial area.

Our HABs Grab events occurred in an aquatic environment with the primary goal of quantifying the total mass of MCs, but the methods we used can be applied to other environments. Future efforts could expand this event to encompass connecting channels impacted by cyanobacterial blooms, including the Maumee River and the Detroit River-Lake St. Clair-Thames River continuum (Davis et al., 2014; Matson et al., 2020; McKay et al., 2020), priority systems for the United States and Canadian governments, respectively. Environment and Climate Change Canada’s already strong presence in the latter could be leveraged in these future efforts and benefit from the models suggested above. Likewise, HABs Grab-like methods can be used in the watershed to identify hot spots of nutrient loss from fields and other high contributors of nutrients.

Coordination of the HABs Grab in 2019 presented a few challenges.

For the HABs Grab to occur on a particular date, we needed: 1) crewed vessels, 2) sufficient lab personnel and equipment to process all samples the day of collection, 3) favorable weather for the smaller vessels to safely and quickly transit their assigned sectors, and 4) a cyanoHAB present in the lake. Added value to the HABs Grab dataset was presented by 5) cloud-free conditions to validate satellite-derived bloom products. Early during planning, we targeted the first two weeks of August for the HABs Grab because there are cyanobacteria present in the western basin, albeit with much interannual variation in biomass in early August (Bridgeman et al., 2013), and university researchers are available prior to the return to classes in late-August. Then we surveyed interested groups for their availability during the first two weeks of August and picked several dates to conduct the HABs Grab when most groups were available. We also came to an understanding that not all groups might be available on the date selected. On the first Monday of our two-week window, we met at the Lake Erie Center for a logistics meeting, distributed sample equipment, and set up sample filtering stations to facilitate efficient sample processing. We monitored weather forecasts and sampled on the first decent weather day even though all groups might not have been available. We took this approach because we did not want to risk passing up the opportunity to collect 175 samples on a decent day for potentially collecting 200 samples, for example, on a later date that could get canceled due to storms. The decision to sample was made on the day prior to the event. The short notice decision was challenging for the researchers involved and even extended to the media wishing to cover the binational event. Overall, future HABs Grab-like events can be successful if logistics are thoroughly planned, and there is an understanding that not all involved may be able to participate, given the constraints of the particular project. These exercises also represent a valuable way to involve media on both sides of the border, thereby increasing public awareness and understanding of cyanoHAB and MC issues in Lake Erie and reducing the likelihood of an 'out-of-sight, out-of-mind' mentality among members of the public.

4.6. Conclusions

There was an estimated 11,513 kg and 30,691 kg of MCs in the western basin of Lake Erie on a single day during peak cyanoHAB conditions on 9 August 2018 and 7 August 2019, respectively. These estimates were made possible with a high spatial resolution dataset collected throughout the entire basin using consistent methods. The fact that cyanoHABs can produce tons of toxins can be alarming; therefore, these numbers must be put into context when messaging to managers and the public. For example, the high MC mass estimates are a function of bloom expanse and the large area of the basin (~2780 km²) that is prone to cyanoHABs. The basin-wide average estimated concentration was 0.52 µg/L and 1.38 µg/L, respectively, if the bloom was spread throughout the basin. The bloom boundary poses substantial issues for spatial interpolations because MC concentration can vary by nearly two orders of magnitude over very short distances. The dataset also showed that the MC:chl *a* ratio varied by a factor up to 5.3 throughout the basin, creating challenges for using the ratio to predict MC concentrations. These issues can only be overcome with more frequent data collection. Models designed to forecast MC concentrations must account for the spatial heterogeneity in MC:chl *a* and the rapid gradients at the bloom boundary. Furthermore, the HABs Grabs dataset will continue to be utilized in upcoming studies ranging from molecular characterization of the microbial communities, more detailed toxin analysis, ecosystem modeling, water mass and cyanoHAB transport, and remote sensing validation.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Supplementary materials

Supplementary material associated with this article can be found, in the online version, at [doi:10.1016/j.hal.2021.102080](https://doi.org/10.1016/j.hal.2021.102080).

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