- 1 Routine monitoring of Western Lake Erie to track water quality
- 2 changes associated with cyanobacterial harmful algal blooms

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#### Abstract

Research Laboratory, 2019).

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20 The western basin of Lake Erie has a history of recurrent cyanobacterial harmful algal blooms 21 (HABs) despite decades of efforts by the United States and Canada to limit phosphorus loading, 22 a major driver of the blooms. In response, the National Oceanic and Atmospheric Administration 23 (NOAA) Great Lakes Environmental Research Laboratory (GLERL) and the Cooperative 24 Institute for Great Lakes Research (CIGLR) created an annual sampling program to detect, monitor, assess, and predict HABs in western Lake Erie. Here we describe the data collected 25 26 from this monitoring program from 2012 to 2021. This dataset includes observations on physico-27 chemical properties, major nutrient fractions, phytoplankton pigments, microcystins, and optical properties for western Lake Erie. This dataset is particularly relevant for creating models, 28 29 verifying and calibrating remote sensing algorithms, and informing experimental research to 30 further understand the water quality dynamics that influence. HABs in this internationally 31 significant body of freshwater. The dataset can be freely accessed from NOAA National Centers 32 for Environmental Information (NCEI) at https://doi.org/10.25921/11da-3x54 (Cooperative 33 Institute for Great Lakes Research, University of Michigan; NOAA Great Lakes Environmental

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#### Introduction

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Lake Erie is situated on the international boundary between the United States and Canada and is the smallest by volume of the five Laurentian Great Lakes. It is ecologically, culturally, and economically significant to the approximately 12.5 million people who live in the watershed. Each year Lake Erie supports nearly 14,000 tonnes of commercial and traditional fisheries, over 33,000,000 tonnes of freight, and over \$1.5 million in recreation and tourism business (Sterner et al., 2020). Lake Erie has endured multiple anthropogenic stressors since European settlement in the area, most notably the draining of coastal wetlands for development of agricultural lands in the late 18th century (Allinger and Reavie, 2013). Currently, the ecological state of Lake Erie is considered poor, partially due to excess nutrient input that supports harmful algal blooms (HABs; ECCC and US EPA, 2022). These seasonal HABs are typically dominated by toxin producing cyanobacteria, causing concern for public and ecosystem health (Watson et al., 2016). Humans can be exposed to cyanotoxins through ingestion of contaminated fish and drinking water and through inhalation and dermal exposure during recreational events such as swimming and boating (Carmichael and Boyer, 2016; Buratti et al., 2017). Cyanotoxins can also cause illness and death in aquatic and terrestrial animals (Carmichael and Boyer, 2016). The economic cost of HABs impacts in Lake Erie is estimated to be hundreds of millions of dollars each year (Smith et al., 2019).

To combat the deteriorated state of Lake Erie water quality, bi-national water resource management policies alongside scientific research and water quality monitoring efforts have been underway for decades. The Great Lakes Water Quality Agreement (GLWQA), first signed in 1972, was a commitment between the US and Canada in response to degraded water quality throughout the Great Lakes ecosystem (GLWQA, 2012). Phosphorus was found to be the key nutrient that was promoting excess phytoplankton growth (Charlton et al., 1993), and thus the GLWQA sought to limit total phosphorus input to the lakes in an attempt to reduce

phytoplankton growth and biomass (Steffen et al., 2014). The 1972 Clean Water Act (CWA) was similarly enacted to regulate <a href="mailto:point-source">point-source</a> pollution discharge, including phosphorus, into navigable waters in the United States. After the signing and implementation of the phosphorus load reduction practices outlined in the GLWQA and CWA, the water quality of Lake Erie improved and the lake experienced a period of restoration (Makarewicz and Bertram, 1991). This success was attributed to upgrades to sewage treatment plants and industrial discharges which reduced phosphorus loading from point sources by 50% within ten years of peak levels observed in 1968 (Charlton et al., 1993; Joosse and Baker, 2011; Steffen et al., 2014).

While the water quality of Lake Erie rebounded in the 1980s and early 1990s, by the mid 1990s and early 2000s annual HAB events were occurring in Lake Erie again, particularly in the warm, shallow western basin (Allinger and Reavie, 2013; Kane et al., 2015; Watson et al., 2016). Total phosphorus loading has been relatively stable in Lake Erie from the 1980s onward (Dolan and Chapra, 2012; Watson et al., 2016), and although point-source phosphorus loading controls had been a successful mitigation measure at one point, several anthropogenic stressors within the watershed were exacerbating the issue of poor water quality. An increase in agricultural sources of biologically available soluble nutrients, legacy phosphorus in the Lake Erie watershed, altered nutrient cycling by invasive dreissenid mussels, and climate change are thought to be primarily responsible for the HABs resurgence (Vanderploeg et al., 2001; Conroy et al., 2005; Bridoux et al., 2010; Michalak et al., 2013; Matisoff et al., 2016; Huisman et al., 2018; Van Meter et al., 2021).

The post-recovery period HABs have predominantly been composed of the cyanobacteria species *Microcystis aeruginosa* along with genera *Anabaena, Aphanizomenon, Dolichospermum,* and *Planktothrix* (Steffen et al., 2014; Watson et al., 2016). These cyanobacteria can produce an array of several types of phycotoxins, with the most common being a suite of hepatotoxins known as microcystins (MCs). Microcystins primarily affect the liver but can also cause adverse health effects on the kidneys, brain, and reproductive organs

(Carmichael and Boyer, 2016). Phycotoxins are commonly present during Lake Erie HABs, and in August 2014 the city of Toledo, OH drinking water supply was contaminated with MCs, leaving >400,000 without clean drinking water (Steffen et al., 2017).

To understand HAB events in US waterways, Congress authorized the Harmful Algal Bloom and Hypoxia Research and Control Act in 1998 (HABHRCA; Public Law 115-423) which mandated the National Oceanic and Atmospheric Administration (NOAA) to "advance the scientific understanding and ability to detect, monitor, assess, and predict HAB and hypoxia events". Under HABHRCA, the NOAA Great Lakes Environmental Research Lab (GLERL), NOAA National Centers for Coastal Ocean Science (NCCOS), and the Cooperative Institute for Great Lakes Research (CIGLR; formerly CILER - Cooperative Institute for Limnology and Ecosystems Research) developed an ecological forecast to predict HAB events in Lake Erie. Starting in 2008, researchers at these institutes began using remote sensing to monitor seasonal HABs, created a seasonal forecast system based on spring P loads, and developed models to predict short-term bloom changes to alert stakeholders and the public (Rowe et al., 2016). Products from these efforts, known as Lake Erie Harmful Algal Bloom Forecasts, are freely available during the bloom season at <a href="https://coastalscience.noaa.gov/research/stressor-impacts-mitigation/hab-forecasts/lake-erie/">https://coastalscience.noaa.gov/research/stressor-impacts-mitigation/hab-forecasts/lake-erie/</a>.

In-situ sampling of the bloom was necessary to calibrate and validate the remote sensing images and models as well as measure microcystin concentration. Sampling events were led by personnel at GLERL and CIGLR starting in 2008 and were designed to collect discrete samples within the extent of the bloom area. At first, samples were taken opportunistically within the bloom and sampling locations and analytical parameters were inconsistent. In 2009, regular sampling stations were identified based on spatial patterns of the bloom. From 2009 to 2011, in addition to opportunistic samples, nine main stations in the western basin of Lake Erie were sampled intermittently from June through October (Bertani et al., 2017; Rowland et al., 2020). While these sampling efforts initially began to complement

existing research products, the experimental nature of the 2008 to 2011 sampling cruises also provided insight into creating a regular monitoring program that would support critical research and product development related to western Lake Erie HABs.

In 2012, researchers at GLERL and CIGLR, with support from the Great Lakes

Restoration Initiative (GLRI), formalized a sampling regimen to monitor the spatial and temporal variability of seasonal HAB events in western Lake Erie (WLE). The establishment of this monitoring program corresponded with increased federal emphasis on evaluating trends and drivers of WLE HABs and water quality. Four monitoring stations were identified and regular surface samples were collected from May to September and analyzed for nutrient, pigment, and particulate microcystin concentrations (Figs. 1 & 2). In following years, the monitoring program evolved and expanded. New stations were added to better characterize the bloom and complement other observing systems. Sampling parameters were adjusted and added based on the needs of current research (Table 1). Results of these sampling cruises were compiled and distributed informally upon request until 2019 when the data were organized and archived on the NOAA National Centers for Environmental Information (NCEI) open-access data repository (<a href="https://www.ncei.noaa.gov/">https://www.ncei.noaa.gov/</a>).

Long term monitoring of WLE is fundamental to the continual assessment of water quality changes in response to both stressors and water quality management efforts (Hartig et al., 2009, 2021). The GLERL/CIGLR monitoring data has been used by numerous researchers to develop and assess models (Rowe et al., 2016; Weiskerger et al., 2018; Fang et al., 2019; Liu et al., 2020; Qian et al., 2021; Wang and Boegman, 2021; Hellweger et al., 2022; Maguire et al., 2022), to calibrate remote sensing algorithms (Sayers et al., 2016, 2019; Avouris and Ortiz, 2019; Bosse et al., 2019; Vander Woude et al., 2019; Pirasteh et al., 2020; Xu et al., 2022), and to elucidate ecological mechanisms and complement experimental data (Cory et al., 2016; Reavie et al., 2016; Berry et al., 2017; Steffen et al., 2017; Kharbush et al., 2019, 2023; Newell

et al., 2019; Den Uyl et al., 2021; Smith et al., 2021, 2022; Hoffman et al., 2022; Marino et al., 2022; Yancey et al., 2022a, b).

The objective of this paper is to inform users of the dataset "Physical, chemical, and biological water quality monitoring data to support detection of Harmful Algal Blooms (HABs) in western Lake Erie, collected by the Great Lakes Environmental Research Laboratory and the Cooperative Institute for Great Lakes Research since 2012" by describing the data generated from this monitoring program and detailing how samples were collected and analyzed. This paper contextualizes this long-term data set so that it can continue to be used to benefit our collective ecological knowledge of western Lake Erie.

Table 1. Description of stations sampled in western Lake Erie from 2012 to 2021. Latitude and longitude (decimal degree) coordinates for each station are target locations as the boat was allowed to drift at each site during *in-situ* sampling.

Station	Latitude	Longitude	Avg. Depth (m)	Years Monitored
WE02	41.762	-83.330	5.4	2012-2021
WE04	41.827	-83.193	8.4	2012-2021
WE06	41.705	-83.385	2.9	2012-2021
WE08	41.834	-83.364	4.8	2012-2021
WE09	41.718	-83.424	2.7	2016-2021
WE12	41.703	-83.254	6.6	2014-2021
WE13	41.741	-83.136	8.9	2014-2021
WE14	41.720	-83.010	9.3	2015
WE15	41.617	-83.009	4.5	2015-2017
WE16	41.660	-83.143	6.2	2018-2021

### Methods

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## Study Site

Based on the lake's bathymetry, Lake Erie can be divided into the eastern, central, and western basins which in turn influence physical and biological processes (Allinger and Reavie, 2013). The data presented in this paper were collected from the western basin, which encompasses the western part of the lake to Point Pelee, ON, Canada and Cedar Point, OH, USA (Fig. 1). The well-mixed western basin is the shallowest (maximum average depth of 11 m), warmest, and most productive of the three basins. Although it's typical for temperate WLE to have ice cover in the winter (Jan to Mar), summer (Jul to Sep) surface water temperatures often reach or exceed 25 °C. The western basin receives 95% of its hydraulic inflow from the Detroit River, which connects Lake Erie hydrologically to Lake Huron via the St. Clair River and Lake St. Clair (Cousino et al., 2015). Among the other tributaries to WLE (including River Raisin, Portage River, Ottawa River, Stony Creek, Swan Creek, and Sandusky River), the Maumee River discharges into the western basin near the city of Toledo, Ohio and contributes a significant amount of sediments and nutrients to the entire Lake Erie basin (Baker et al., 2014a, b; Rowland et al. 2020; see NCWQR 2022 for Maumee River water quality data). Nutrient and sediment loads from the Maumee River can vary with precipitation, where stormwater runoff can provide a pulse of nutrients into the basin, potentially altering cyanobacteria dynamics (Baker et al., 2014a; King et al., 2022). Land use in the Lake Erie watershed is 75% agricultural and 11% urban, both of which contribute to the large amounts of soluble reactive phosphorus into the basin (Mohamed et al., 2019; Myers et al., 2000). This dataset includes water quality data from ten monitoring stations on the United

States side of WLE that were sampled from 2012 to 2021 (Figs. 1 & 2, Tables 1 & 2). The

average depth of monitoring stations ranged from 2.7 m at WE9 to 9.3 m at WE14. These sites were chosen to reflect the various nutrient and hydrologic inputs and gradients into WLE, as well as represent areas of the basin that are prone to HABs. The Maumee River inflow was a major consideration in determining these sites. The initial 4 stations sampled in this program (WE02, WE04, WE06, and WE08) were selected because they were consistently within the WLE blooms occurring at the time. Additional sites were later added to better represent the spatial extent of HABs and to augment existing data provided by moored buoy continuous monitoring systems, advanced monitoring technologies, such as Environmental Sample Processors (Den Uyl et al., 2022), and other monitoring programs in WLE.

#### Field Sampling

Western Lake Erie discrete field sampling was accomplished using NOAA GLERL research vessels. Sampling took place during ice-free months and aimed to quantify the environmental conditions prior to, during, and at the end of the bloom (Fig. 2). Sampling stations represent approximate locations (Table 1; Fig. 1); *in situ* measurements and sampling were collected once the boat reached the targeted location and then proceeded to drift during sampling. The frequency and timing of those cruises varied over the first few years but has been consistent since 2017 (Fig. 2). Sampling was disrupted in 2020 due to the global COVID-19 pandemic and resulting public health restrictions. In 2020, sampling was initiated in mid-June at a reduced number of sites for select water quality parameters. In July, sampling stations and parameters were expanded and all stations and parameters were sampled and measured by August 2020. The prior standard sampling schedule resumed in April 2021.

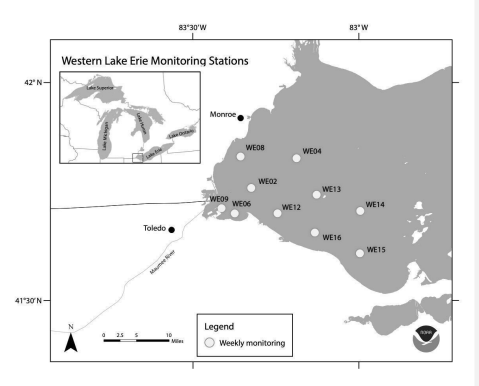


Figure 1. Location of western Lake Erie water quality monitoring stations. This map was provided by NOAA for use in this publication.

	WE2	WE4	WE6	WE8	WF9	
2021			• • • • • • • • • • • • • • • • • • • •	• • • • • • • • • • • • • • • • • • • •		
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2014 -		0 0 0000 0000 000 000 000	0 0 000000000000000000	0 0 00000000000000000000000000000000000		
2013	0 000 00 00 00000 0000 000	0 000 00 000 0000 000 00	0 000 000000000 000 00	0 000 00 000000000000000000000000000000		
_ 2012		0 0 0 0 0 000000000	0 0 0 000000000			
Year	WE12	WE13	WE14	WE15	WE16	
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2012						
	Apr May Jun Jul Aug Sep Oct	Apr May Jun Jul Aug Sep Oct	Apr May Jun Jul Aug Sep Oct	Apr May Jun Jul Aug Sep Oct	Apr May Jun Jul Aug Sep Oct	
	_	_	Date	_	_	

Figure 2. Sampling frequency for each monitoring station for years sampled between 2012 to 2021.

In-situ measurements for conductivity, temperature, dissolved oxygen (DO), beam attenuation, transmission, and photosynthetically active radiation (PAR) were taken with a Sea-Bird 19plus V2 conductivity, temperature, and depth (CTD) profiler attached to a hydraulic crane. Data were collected on the downcast and were reported as the mean of recorded values within ± 0.5 m of the discrete sample depth. In 2012, sample temperature was taken on the boat with a Vee Gee Scientific IP67-rated digital thermometer. Sky conditions were recorded at the discretion of the field technician at each station during the sampling cruise. A Secchi disk was lowered into the water on the shaded side of the boat at each station and the depth at which the Secchi disk was no longer visible was recorded (Wetzel and Likens, 2000).

Water column samples were collected using a 5 L vertical Niskin bottle (General Oceanics model 1010). Niskin casts were evenly distributed between one or more high-density

polyethylene bottles that were rinsed with site water and stored in a cooler. Three to four Niskin casts were used to fill the bottles, such that each bottle is a composite sample of the water collected. Surface samples were taken 0.75 m below the water's surface, mid-column samples were taken at approximately 4.25 m below surface, and benthic or bottom samples were taken at approximately 0.5 m above the lake bottom at each station. Surface samples were taken at all stations while mid-column and benthic sample collection varied between sites and years. Scum samples of dense cyanobacterial accumulation on the surface of the water were collected opportunistically using a 2 L modified Van Dorn water sampler. Sampling times were reported as Eastern Daylight Time (UT -4:00). Upon arrival at the laboratory, raw water samples were immediately subsampled and preserved until analysis.

Wind speed and wave height data were obtained from moored buoy continuous monitoring systems in proximity to sampling stations for a timestamp that corresponded to the time samples were collected at that station. Wave height data for all stations were obtained from the Toledo Intake Buoy (owned and maintained by Limnotech Inc.). Wind speed data for stations WE02, WE06, WE09, WE12, WE14, WE15, and WE16 were also collected from this buoy. Data for this buoy is available through the Great Lakes Observing System (GLOS; platform ID 45165, <a href="https://seagull.glos.org/data-console/71">https://seagull.glos.org/data-console/71</a>). Wind speed data for stations WE04, WE08, and WE13 were obtained from the Toledo Harbor Light no. 2 buoy (Station THLO1, owned and maintained by GLERL). Data for this buoy is available through NOAA's National Data Buoy Center (<a href="https://www.ndbc.noaa.gov/station\_realtime.php?station=THLO1">https://www.ndbc.noaa.gov/station\_realtime.php?station=THLO1</a>).

#### Laboratory analysis of samples

Water collected from WLE was subsampled to make a range of analytical measurements in the laboratory (Table 2).

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collected from moored buoy continuous monitoring systems which provide the data in Imperial

units.

Parameter	Years monitored	Method
Surface samples (n=1296)	2012-2021	n/a
Mid-column samples (n=19)	2015	n/a
Benthic samples (n=512)	2015-2021	n/a
Station depth_(m)	2012-2021	Sea-Bird 19plus V2 CTD profiler
Time of sampling (Eastern Daylight Time UTC -4:00)	2012-2021	n/a
Latitude (decimal degree)	2012-2021	n/a
Longitude (decimal degree)	2012-2021	n/a
Wind speed (knots)	2015-2021	Moored buoy continuous monitoring systems
Wave height (ft)	2012-2021	Moored buoy continuous monitoring systems
Cloud cover (sky)	2012-2021	Qualitative description,
Secchi depth (m)	2012-2021	Wetzel and Likens (2000)
Sample temperature (°C)	2012	Vee Gee Scientific digital thermometer
CTD temperature (°C)	2013-2021	Sea-Bird 19plus V2 CTD profiler
CTD specific conductivity (µS cm <sup>-1</sup> )	2013-2021	Sea-Bird 19plus V2 CTD profiler
CTD beam attenuation (m <sup>-1</sup> )	2013-2021	Sea-Bird 19plus V2 CTD profiler
CTD transmission (%)	2013-2021	Sea-Bird 19plus V2 CTD profiler
CTD dissolved oxygen (DO; mg L <sup>-1</sup> )	2013-2021	Sea-Bird 19plus V2 CTD profiler
CTD photosynthetically active radiation (PAR; µE m <sup>-2</sup> s <sup>-1</sup> )	2013-2021	Sea-Bird 19plus V2 CTD profiler

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Turbidity (NTU)	2013-2021	EPA Method 180.1
Particulate microcystins (μg L <sup>-1</sup> )	2012-2021	Wilson et al. (2008)
Dissolved microcystins (µg L <sup>-1</sup> )	2014-2021	Wilson et al. (2008)
Phycocyanin (µg L <sup>-1</sup> )	2012-2021	Horvath et al. (2013)
Chlorophyll a (µg L <sup>-1</sup> )	2012-2021	Speziale et al. (1984)
Total phosphorus (TP; μg L <sup>-1</sup> )	2012-2021	Standard Method 4500-P
Total dissolved phosphorus (TDP; μg L <sup>-1</sup> )	2012-2021	Standard Method 4500-P
Soluble reactive phosphorus (SRP; μg L <sup>-</sup> <sup>1</sup> )	2012-2021	Standard Method 4500-P
<u>Ammonium-N</u> (μg L <sup>-1</sup> )	2012-2021	Standard Method 4500-nh3-nitrogen (Ammonium)
Nitrate-N + Nitrite-N (mg L <sup>-1</sup> )	2012-2021	Standard Method 4500-no3-nitrogen (nitrate)
Urea <u>-N</u> (μg L <sup>-1</sup> )	2016-2017	Milvenna and Savidge (1992), Goeyens et al. (1998), Chaffin and Bridgeman (2014)
Particulate organic carbon (POC; mg L <sup>-1</sup> )	2012-2021	Hedges and Stern (1984)
Particulate organic nitrogen (PON; mg L <sup>-1</sup> )	2012-2021	Hedges and Stern (1984)
Colored dissolved organic material (CDOM; m <sup>-1</sup> )	2014-2021	Binding et al. (2008), Mitchell et al. (2003)
Dissolved organic carbon (DOC; mg L <sup>-1</sup> )	2012-2017	APHA Standard Method 5310 B
Total suspended solids (TSS; mg L <sup>-1</sup> )	2012-2021	APHA Standard Method 2540
Volatile suspended solids (VSS; mg L <sup>-1</sup> )	2012-2021	APHA Standard Method 2540

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Turbidity was measured on raw samples using a Hach 2100AN Turbidimeter following US EPA method 180.1 (1993). Colored dissolved organic material (CDOM, also defined as chromophoric dissolved organic matter) was determined by filtering lake water through an acid

rinsed 0.2 µm nuclepore polycarbonate filter into acid-washed and combusted borosilicate vials. Optical density of the filtered samples was then measured using a Perkin Elmer UV/VIS Lambda 35 spectrophotometer at wavelengths from 300-800 nm. CDOM absorption was calculated at 400 nm (Mitchell et al., 2003; Binding et al., 2008).

Dissolved organic carbon (DOC) concentrations were determined following American Public Health Association (APHA) Standard Method 5310 B. Briefly, lake water was filtered through 0.45 µm polyvinylidene difluoride membrane filters into combusted borosilicate glass vials and frozen at -20°C until analysis. The filtrate was acidified with HCl and sparged with air for 6 min before being analyzed on a Shimadzu total organic carbon analyzer.

Duplicate samples for particulate organic carbon (POC) and particulate organic nitrogen (PON) were collected onto pre-combusted glass fiber filters and analyzed following Hedges and Stern (1984) Samples were stored at -20 °C until analysis. The filters were then acidified by fumigation with 10% HCl and dried at 70°C for 24 h before being quantified on a Perkin Elmer 2400 or a Carlo-Erba 1110 CHN elemental analyzer.

Total suspended solids (TSS) and volatile suspended solids (VSS) were determined via gravimetric analysis following APHA Standard Method 2540. A known volume of lake water was filtered through a pre-combusted, pre-weighed Whatman GF/F glass fiber filter. The filters were then dried at 60° C for at least 24 h and reweighed. The difference in mass between the pre-weighed and processed filter was reported as TSS. Volatile suspended solids concentrations were quantified by combusting the filters used for TSS analysis at 450 °C for 4 h, weighing the combusted filters, and calculating the mass lost.

#### **Nutrient fractions**

Total phosphorus (TP) and total dissolved phosphorus (TDP) samples were collected in duplicate by subsampling 50 mL (2012 to 2019) or 20 mL (2020 to 2021) of lake water into acid washed glass tubes and by filtering 20 mL of lake water through a 0.2 µm membrane filter and

collecting the filtrate, respectively. Samples for TP and TDP were refrigerated until samples were digested with potassium persulfate solution and autoclaved at 121°C for 30 min, modified from APHA Standard Method 4500-P. Digested TP and TDP samples were stored at room temperature until concentrations were measured on a Seal QuAAtro continuous segmented flow analyzer (SEAL Analytical Inc.) from 2012 to 2019 and a Seal AA3 from 2020 to 2021 using the ascorbic acid molybdenum method as detailed by the instrument manual and APHA Standard Method 4500-P. Analytical detection limits for the analyses were taken from the instrument manufacturer's documentation.

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Soluble reactive phosphorus (SRP), ammonium, nitrate + nitrite, and urea were each determined by filtering 12 mL of lake water through a 0.2 µm membrane filter into 15 mL centrifuge tubes during field sampling. Sample filtrates were stored at -20 °C upon receipt at the laboratory. Soluble reactive phosphorus, ammonium, and nitrate + nitrite concentrations were determined simultaneously on a Seal AA3 continuous segmented flow analyzer. Soluble reactive phosphorus concentrations, like TP and TDP concentrations, were measured using the ascorbic acid molybdenum method as detailed by the instrument manual and APHA Standard Method 4500-P. Ammonium concentrations were measured using Bertholet reactions according to the instrument manual and APHA Standard Method 4500-nh3-nitrogen. Nitrate + nitrite concentrations were measured using copper-cadmium reduction methods according to the instrument manual and APHA Standard Method 4500-no3-nitrogen. Analytical detection limits for these inorganic nutrient analyses were taken from the instrument manufacturer's documentation. Urea samples were measured by adding diacetyl monoxime and thiosemicarbazide to the filtrate and briefly vortexing to mix, followed by adding sulfuric acid and ferric chloride to the solution and briefly vortexing to mix. Samples were then incubated in the dark for 72 h at room temperature before absorbance at 520 nm was read on a Perkin Elmer UV/VIS Lambda 35 spectrophotometer. Urea concentrations were then quantified using a

standard curve (Mulvenna and Savidge, 1992; Goeyens et al., 1998; Chaffin and Bridgeman,

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2014). The detection limit was calculated using the standard deviation of repeated measurements.

#### Photopigments and microcystins

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Particulate phycocyanin and chlorophyll a concentrations were determined by filtering a known volume of lake water under low vacuum (<200 mm Hg) onto 47 mm Whatman GF/F glass fiber filters (Cytiva Life Sciences). Particulate phycocyanin sample filters were stored in 15 mL conical polypropylene centrifuge tubes and chlorophyll a sample filters were stored in amber glass vials at -20 °C until analysis. Analysis methods for particulate phycocyanin were derived from Horváth et al. (2013) where 9 mL of phosphate buffer was added to sample tubes and samples were agitated using a shaker at 5 °C for 15 min at 100 rpm then vortexed for 10 s each. To encourage cell lysis, samples were subjected to three freeze/thaw cycles at -20 °C followed by sonication for 20 min using a Fisher FS110 H sonicator. Fluorescence of the extracted samples was measured using an Aquafluor 8000-010 fluorometer (Turner Designs) with excitation from 400-600 nm and emission filter of >595 nm. Particulate phycocyanin was calibrated annually against C-Phycocyanin material from Sigma-Aldrich. Analysis methods for chlorophyll a were derived from Speziale et al. (1984) where chlorophyll a was extracted from samples using dimethylformamide and placed into a 65 °C water bath for 15 min. Samples were then cooled to room temperature and vortexed for 15-20 s before being quantified using a 10 AU fluorometer (Turner Designs) with excitation filter of 436 nm and emission at 680 nm. Phycocyanin and chlorophyll a procedures were performed under low or green light to reduce pigment degradation within the cell.

Dissolved and particulate microcystins were quantified using a procedure adapted from Wilson et al. (2008). Dissolved microcystins (dMC) were determined through duplicate samples of  $\sim 2$  mL filtrate that was passed through a 0.2  $\mu$ m membrane filter and stored in glass vials at -20 °C until analysis. Particulate microcystins (pMC) were collected by filtering a known volume of lake water onto a Whatman GF/F glass fiber filter (2012 to 2015) or a 3  $\mu$ m pore size

polycarbonate membrane filter (2016 to 2021). Particulate MC was then extracted from the filters. In sampling years 2012 to 2015, glass fiber filters were submerged in a glass vial containing a 75:25 methanol:water solution (MeOH/H<sub>2</sub>O) and sonicated in an ice bath for 2 min. The samples were centrifuged for 15 min and the supernatant was transferred to a clean glass vial. An additional 5 mL of MeOH/H<sub>2</sub>O was added to the filter/precipitate and the sample was incubated at -20 °C for 5 h. The sample was then sonicated for 2 min, centrifuged, and the supernatant was removed and added to the first extract vial. The composite supernatant was then centrifuged under a vacuum until dry. The dried extract was then stored at -20 °C until analysis. Particulate MC concentrations were then determined by adding 1 mL of MIlliQ water to the sample and using sonication to dissolve the dried extract. For sampling years 2016 to 2021, filters were stored in 2 mL sterile microcentrifuge tubes at -20 °C until analysis. During analysis, pMC were extracted from the membrane filters by adding 1 mL of MilliQ water and subjecting samples to three freeze/thaw cycles at -20 °C followed by addition of Abraxis QuickLyse reagents according to the manufacturer (Eurofins/Abraxis). Particulate MC samples for all sampling years were analyzed immediately after extraction. For all sampling years, dMC and pMC concentrations were determined using a congener-independent enzyme-linked immunosorbent assay (ELISA) kit designed to detect and quantity microcystins and nodularins using the ADDA moiety (Envirologix brand used from 2012 to 2015; Eurofins/Abraxis microcystins/nodularins (ADDA) (EPA ETV) (EPA method 546), ELISA, 96-test kit used from 2016 to 2021). Analytical detection limits for the analyses were taken from the manufacturer's documentation.

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## Results and Discussion

This dataset demonstrates the temporal and spatial variability in water quality parameters in western Lake Erie from 2012 to 2021. Overall, sites closest to the Maumee River inflow (i.e., WE06 and WE09) had the highest median concentrations of nutrients, sediments, pigments, and microcystins compared to sites further out in the basin (i.e., WE02, WE04, and WE13; Table 3). Stations WE06 and WE04 were sampled since the initiation of the monitoring program and consistently represented the high and low extremes of water quality observations during a given time point, respectively, (Table 3) and select parameters for these two sites are represented in figs. 3 and 4. Supplemental figs. 1-16 display the same parameters as figs. 3 and 4 for the remaining stations.

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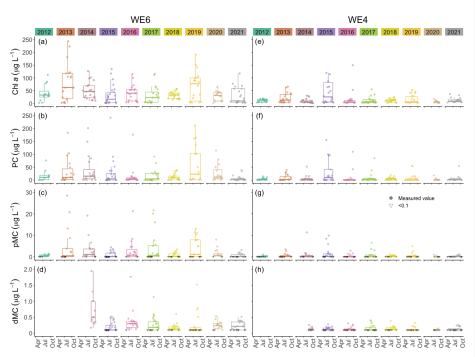


Fig 3. Comparison of chlorophyll *a* (Chl *a*), phycocyanin (PC), particulate microcystins (pMC), and dissolved microcystins (dMC) between stations WE04 and WE06 from 2012 to 2021. Boxplots represent the median and 25% and 75% quartiles with whiskers extending to the highest or lowest point within 1.5x the interquartile range. A scatterplot is overlaid on the boxplots.

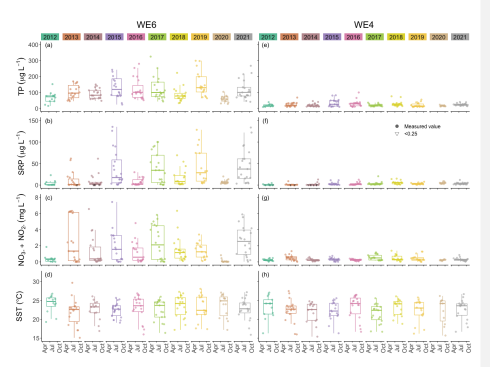


Fig 4. Comparison of total phosphorus (TP), soluble reactive phosphorus (SRP), nitrate plus nitrite ( $NO_3^- + NO_2^-$ ), and sea surface temperature (SST) between stations WE04 and WE06 from 2012 to 2021. Boxplots represent the median and 25% and 75% quartiles with whiskers extending to the highest or lowest point within 1.5x the interquartile range. A scatterplot is overlaid on the boxplots.

Table 3. Median values of each parameter at each monitoring station for all surface samples collected between 2012 to 2021.

1			i	Γ			ľ	1	ı
0.99	0.34	2.0	1.1	2.4	0.81	0.38	09.0	0.54	0.71
0.23	0.10	0.38	0.25	0.42	0.20	0.14	0.27	0.18	0.16
1.4	0.63	2.4	1.5	2.5	1.2	0.78	1.7	1.1	1.0
0.44	0.27	0.83	0.45	1.4	0.31	0.25	0.17	0.27	0.30
12.6	12.9	11.8	13.8	43.1	8.4	10.2	2.9	23.9	10.6
5.7	2.2	8.7	5.8	29.5	5.4	2.7	1.5	2.0	4.0
12.8	4.5	18.7	12.3	8.44	10.1	9.0	4.7	5.5	7.2
53.3	19.2	90.1	6.03	133	47.6	22.3	31.0	34.8	30.2
17.5	7.7	33.0	19.5	32.6	15.1	9.8	40.0	12.7	12.3
4.8	1.2	8.0	5.7	5.2	2.9	2.6	17.0	2.7	3.4
0.20	0.17	0.28	0.22	0.26	0.16	0.15	0.16	0.19	0.18
0.78	0.46	1.5	0.88	96.0	19.0	99.0	08.0	98.0	0.91
9.9	3.0	14.8	9.0	23.2	11.0	4.3	7.2	6.3	6.3
28.2	58.4	20.5	34.4	4.3	25.9	52.4	40.2	43.0	40.8
5.1	2.2	6.4	4.3	12.6	5.4	2.7	3.7	3.4	3.6
264	377	173	166	127	266	456	962	391	297
7.7	9.7	9.7	7.7	7.1	7.7	7.8	8.1	7.7	7.4
287	244	346	299	395	276	244	238	261	269
23.1	22.9	23.0	23.3	23.9	23.1	22.9	23.2	23.0	24.1
8.0	2.0	9.0	1.0	0.3	8.0	1.5	1.4	1.0	13
WE02	WE04	WE06	WE08	WE09	WE12	WE13	WE14	WE15	WE16
	0.8 23.1 287 7.7 264 5.1 28.2 9.9 0.78 0.20 4.8 17.5 53.3 12.8 5.7 12.6 0.44 1.4 0.23	0.8 23.1 287 7.7 264 5.1 28.2 9.9 0.78 0.20 4.8 17.5 53.3 12.8 5.7 12.6 0.44 1.4 0.23 2.9 2.44 7.6 3.77 2.2 58.4 3.0 0.46 0.17 1.2 7.7 19.2 4.5 2.2 12.9 0.27 0.83 0.10	3.8         23.1         28.7         7.7         28.4         5.1         28.2         9.9         0.78         0.28         4.8         17.5         53.3         12.8         6.7         12.6         6.4         14.4         14.9         0.23           2.0         2.2.9         2.4         7.6         0.46         0.17         1.2         7.7         19.2         4.5         2.2         12.9         0.53         0.10           0.5         2.3         3.4         7.6         1.7         1.2         7.7         19.2         4.5         1.2         0.53         0.10           0.5         2.3         3.4         7.6         1.7         1.2         0.2         1.8         0.2         1.8         0.3         0.01         1.8         7         1.8         0.8         0.0	3. 2. 3. 2. 3. 2. 3. 2. 3. 3. 2. 3. 3. 3. 3. 3. 3. 3. 3. 3. 3. 3. 3. 3.	3         3         4         5         4         6         4         6         7         7         6         4         7         6         7	3. 2. 2. 2. 2. 2. 2. 2. 2. 2. 2. 2. 2. 2.	3         3         4         5         4         4         6         4         6         6         4         6         6         4         6         7         1         6         7         1         6         7         1         6         7         1         6         7         1         6         7         1         7         1         6         7         1         1         7         1         2         2         2         2         1         1         2         1         1         2         1         1         2         1         2         2         2         2         2         2         2         2         2         2         2         2         2	4.8         2.3.1         2.8.4         7.7         2.8.4         5.1         2.8.4         6.7.5         6.3.4         6.7.5         6.3.3         1.7.5         6.3.4         6.7         6.2.4         6.3.7         6.2.4         6.3.7         6.2.4         6.3.7         6.2.4         6.3.7         6.2.4         6.2.7         6.2.4         6.2.7         6.2.4         6.2.7         6.2.4         6.2.7         6.2.4         6.2.7         6.2.4         6.2.7         6.2.4         6.2.7         6.2.4         6.2.7         6.2.4         6.2.7         6.2.4         6.2.7         6.2.4         6.2.7         6.2.4         6.2.7         6.2.4         6.2.7         6.2.4         6.2.7         6.2.4         6.2.7 </td <td>4.8         2.3.1         2.8.4         7.7         2.8.4         5.1         2.8.4         1.7.5         5.8.4         9.0         0.7.8         0.4.8         1.7.5         5.3.3         1.2.8         5.1         1.2.5         5.3.4         1.7.5         1.8.4         1.8.6         0.1.7         1.2.5         1.8.7</td>	4.8         2.3.1         2.8.4         7.7         2.8.4         5.1         2.8.4         1.7.5         5.8.4         9.0         0.7.8         0.4.8         1.7.5         5.3.3         1.2.8         5.1         1.2.5         5.3.4         1.7.5         1.8.4         1.8.6         0.1.7         1.2.5         1.8.7

#### Physicochemical properties

Median surface temperatures for all samples across all years ranged from 22.9 to 24.1 °C and median benthic temperatures ranged from 22.8 to 23.2 °C (Table 3, Fig. 4), indicating that WLE was thermally well mixed throughout the sampling period. A summary of the dataset indicates that 23.8% of surface temperatures were ≥25 °C, and these higher temperatures all occurred from mid-June through the end of September. Bloom forming cyanobacteria species in Lake Erie, including *Microcystis spp.*, often reach maximum growth rates at warmer temperatures (≥ 25 °C) than eukaryotic phytoplankton (Steffen et al., 2014; Huisman et al., 2018). Despite having warmer temperatures that promote recurring HABs, there was only one recorded instance of hypoxia (DO <2 mg L⁻¹) in the dataset and it occurred at WE13 on 08 July 2019. Median DO was 7.62 mg L⁻¹ in all surface samples and 7.02 mg L⁻¹ in all benthic samples from 2012 to 2021 (Table 3), again indicating minimal stratification in WLE during sampling. Median conductivity from 2012 to 2021 was highest at sites WE06 and WE09, which are closest to the Maumee River input, and lowest at sites WE04 and WE13 near the middle of the basin (Table 3). WE06 and WE09 were the only sites to have median conductivity values above 300 μS cm⁻¹.

#### Optical properties

Biotic and abiotic particulate concentrations and movement patterns in WLE are prone to spatial and seasonal variations and are heavily influenced by loading from the Maumee River (Prater et al., 2017; Maguire et al., 2022). Secchi depth, turbidity, and PAR measurements have been correlated with distance from Maumee Bay, where light penetration was lowest near the Maumee River (Chaffin et al., 2011). Variability in optical property measurements in WLE is also dependent on Maumee River inputs, and changes in optical properties can potentially be used

in remote sensing algorithms to detect changes in water quality (Sayers et al., 2019). Median Secchi disk depth over the entire dataset was highest at WE04 and lowest at WE06 and WE09, which are closest to the Maumee River (Table 3). Other optical properties, such as PAR, beam attenuation, and transmittance also followed this spatial pattern. In a summary of all samples, median PAR measured at 0.5 m below surface was highest at WE13 and WE14 and lowest at WE09; median transmittance was highest at WE04 and lowest at WE09; and median beam attenuation and turbidity were highest at WE09 and lowest at WE04 (Table 3). Median turbidity values at each site over the 2012 to 2021 period were within the range of previously reported values in the WLE basin (Barbiero and Tuchman, 2004). Median CDOM absorbance and DOC, TSS, and VSS concentrations were again highest at WE09 and lowest at WE04 (Table 3). CDOM gradients in WLE are likewise affected by loading from the Maumee River (Cory et al., 2016) and DOC and CDOM values from this dataset have been used as predictor variables in models estimating PAR attenuation variation in WLE (Weiskerger et al., 2018).

#### **Nutrient fractions**

The Maumee River is a major contributor of nutrients to Lake Erie (Steffen et al., 2014; Kast et al., 2021). Median TP concentrations in WLE from 2012 to 2021 were lowest at WE04 and highest at WE09 (Table 3, Fig. 4). Median concentrations at each station from 2012 to 2021 were above the GLWQA Annex 4 goals for TP concentration in open waters, which is 15 μg P L<sup>-1</sup> for WLE. This goal was met in 92 of 1275 (7.2%) samples and these target values were primarily recorded from stations WE04 and WE13. Sites closer to the mouth of the Maumee River had higher median TP values. While TP loading from the Maumee River tributary declined between 1982 to 2018 (Rowland et al., 2020) the proportion of dissolved P has increased (Joosse and Baker, 2011; Stow et al., 2015). Median TDP values in the WLE dataset were lowest at WE04 and highest at WE09 (Table 3) with a highest recorded value of 274 μg P L<sup>-1</sup> at

WE08 in 2015. Median SRP concentrations for each station in this dataset were lowest at WE14 and WE15 and were highest at WE09 (Table 3). The maximum recorded SRP concentration was 135.4 μg P L<sup>-1</sup> at WE06 in 2015 (Fig. 4). Using this dataset, Newell et al. (2019) found that the Maumee River N loading has become more chemically reduced over time where ammonium and PON have increased. Median ammonium concentrations in WLE from 2012 to 2019 were lowest at WE12 and WE14 and highest at WE09 (Table 3) with a recorded maximum concentration of 2109 μg N L<sup>-1</sup> at WE12 in 2017. Median nitrate + nitrite was lowest at WE13 and WE14 and highest at WE09 (Table 3), with a maximum recorded value of 9.5 mg N L<sup>-1</sup> at WE09 in 2016. See Fig. 4 for a comparison of nitrate + nitrite concentrations between WE04 and WE06. Median PON concentrations were lowest at WE04 and highest at WE09 (Table 3) with a recorded max of 40.93 mg N L<sup>-1</sup> at WE08 in 2015.

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### Photopigments and microcystins

Median extracted chlorophyll *a* concentrations in surface waters from 2012 to 2021 were lowest at WE04 and highest at WE06 (Table 3, Fig. 3). The highest recorded surface concentration of chlorophyll *a* was 6784 μg L<sup>-1</sup> on 10 August 2015 at WE08 during the most severe bloom year in this dataset, according to the CI Index (Wynne et al., 2013; Lunetta et al., 2015). The highest measured levels of particulate phycocyanin, pMC, and TP were also recorded at WE06 on 10 August 2015. Other notably high chlorophyll *a* concentrations were measured during severe bloom years in 2017 (532 μg L<sup>-1</sup> at WE09 on 04 August) and 2019 (593 μg L<sup>-1</sup> at WE09 on 05 August). Similarly, median surface particulate phycocyanin concentration for 2012 to 2021 was highest at WE06 and lowest at WE04 (Table 3, Fig. 4). The highest recorded phycocyanin value was from WE08 on 10 August 2015 (8228 μg L<sup>-1</sup>), followed by 3315 μg L<sup>-1</sup> at WE06 in 2013 during another severe bloom year.

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Particulate MC concentrations had highest median concentrations at WE06 and were lowest at WE04 (Table 3, Fig. 4), similar to particulate chlorophyll *a* and phycocyanin observations. The highest recorded particulate MC concentration in this dataset was from 10 August 2015 at WE08 during a severe bloom year (297 μg L<sup>-1</sup>), followed by 289 μg L<sup>-1</sup> at WE06 in 2017 during another severe bloom year according to the CI Index (Wynne et al., 2013; Lunetta et al., 2015). Median dMC concentrations were highest at WE06 and lowest at WE13 (Table 3). The maximum dissolved MC in the dataset was 8.19 μg L<sup>-1</sup> at WE09 on 05 August 2019, which correlates with high chlorophyll *a* concentrations.

Although the United States does not federally enforce water quality criteria or regulations for cyanotoxins in drinking water, the US EPA has a recommended health advisory of 1.6 µg L<sup>-1</sup> microcystins in drinking water for school-age children through adults (US EPA, 2015) while the WHO and the Ohio EPA use 1 µg L<sup>-1</sup> microcystins as a guideline (WHO, 2020). From 2012 to 2021, 44.4% of pMC samples in this dataset exceeded the WHO guidelines and 34.1% exceeded the US EPA health advisory. Monitoring MC concentrations in western Lake Erie has become especially pertinent since August 2014 when the Toledo, OH drinking water treatment plant was contaminated with microcystins in excess of 1 µg L<sup>-1</sup> and customers were alerted to not drink their tap water until toxin levels were decreased (Steffen et al., 2017). The pMC concentrations at our WLE monitoring stations varied from 1.2-10.1 µg L<sup>-1</sup> on 04 August 2014 during this crisis.

# Data Availability

475	The entire dataset detailed in this manuscript can be freely accessed through the NOAA
476	National Centers for Environmental Information (NCEI) data repository at
477	https://www.ncei.noaa.gov/. The data collection is titled "Physical, chemical, and biological water
478	quality monitoring data to support detection of Harmful Algal Blooms (HABs) in western Lake
479	Erie, collected by the Great Lakes Environmental Research Laboratory and the Cooperative
480	Institute for Great Lakes Research since 2012". The digital object identifier is
481	https://doi.org/10.25921/11da-3x54. The data presented in this manuscript are available in three
482	separate accession files within this collection including: 2012 to 2018 data is available under
483	NCEI Accession 0187718 v2.2 at <a href="https://www.ncei.noaa.gov/archive/accession/0187718">https://www.ncei.noaa.gov/archive/accession/0187718</a> ; 2019
484	data is available under NCEI Accession 0209116 v1.1 at
485	https://www.ncei.noaa.gov/archive/accession/0209116; 2020 to 2021 data is available under
486	NCEI Accession 0254720 v1.1 at <a href="https://www.ncei.noaa.gov/archive/accession/0254720">https://www.ncei.noaa.gov/archive/accession/0254720</a>
487	(Cooperative Institute for Great Lakes Research, University of Michigan; NOAA Great Lakes
488	Environmental Research Laboratory, 2019). Future data will be added to this collection as it
489	becomes available.

### Conclusions

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The western Lake Erie data collected and compiled by NOAA GLERL and CIGLR represent ten years of routine water quality monitoring to detect, track, and predict cyanobacterial HAB events in an area of the Great Lakes that has experienced significant environmental degradation. While this monitoring initiative started in conjunction with remote sensing efforts, it eventually became a standalone program. This ongoing program provides a service to the region and contributes data for investigating the nuanced dynamics of potentially toxic HABs fueled by excess nutrient loading into the WLE basin. For instance, this dataset has assisted in assessing progress toward binational nutrient loading reduction efforts on lake basin concentrations of phosphorus. Long-term monitoring programs like this one provide consistent data which is useful for identifying patterns and variations within the ecosystem and in determining the root cause of those changes. As the sites and parameters of this monitoring program have already changed to adapt to the needs of research, this program will continue to evolve as we consider adding parameters that encompass other aspects of bloom dynamics. For example, lake samples can be analyzed for genomic data that will provide insights on the ability of the current phytoplankton community to produce microcystins. This decadal history has already been an invaluable resource for the research community, and it will continue to enrich our collective scientific knowledge of water quality dynamics in western Lake Erie.

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### **Author Contributions**

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Anna G Boegehold prepared the manuscript. Ashley M. Burtner performed field sampling, laboratory processing, data processing, QA/QC and data management, manuscript revision, data curation. Andrew Camilleri performed field sampling, laboratory processing, manuscript revision. Glenn Carter performed field sampling, laboratory processing, data processing, methodology. Paul DenUyl performed field sampling, laboratory processing, manuscript revision. David Fanslow performed field sampling, laboratory processing. Deanna Fyffe Semenyuk performed field sampling, laboratory processing, manuscript revision. Casey Godwin was responsible for project administration, supervision, visualization, manuscript revision, methodology, field sampling, sample processing. Duane Gossiaux performed field sampling, laboratory processing, manuscript revision, methodology. Tom Johengen was responsible for project administration, supervision, field sampling, methodology. Holly Kelchner performed field sampling, laboratory processing, manuscript revision. Christine Kitchens performed field sampling, laboratory processing, data processing, manuscript revision. Lacey A. Mason was responsible for data curation, manuscript revision. Kelly McCabe performed field sampling, laboratory processing, manuscript revision, methodology. Danna Palladino performed field sampling, laboratory processing, data processing, manuscript revision. Dack Stuart performed field sampling, data processing. Henry Vanderploeg was responsible for project administration, supervision. Reagan Errera was responsible for project administration, supervision, Visualization, manuscript revision, methodology.

## 538 Competing Interests

539 The authors declare that they have no conflict of interest

#### References

- 541 Allinger, L. E. and Reavie, E. D.: The ecological history of Lake Erie as recorded by the
- 542 phytoplankton community, J. Gt. Lakes Res., 39, 365–382,
- 543 https://doi.org/10.1016/j.jglr.2013.06.014, 2013.
- 544 Avouris, D. M. and Ortiz, J. D.: Validation of 2015 Lake Erie MODIS image spectral
- 545 decomposition using visible derivative spectroscopy and field campaign data, J. Gt. Lakes Res.,
- 546 45, 466–479, https://doi.org/10.1016/j.jqlr.2019.02.005, 2019.
- 547 Baker, D. B., Ewing, D. E., Johnson, L. T., Kramer, J. W., Merryfield, B. J., Confesor, R. B.,
- 548 Peter Richards, R., and Roerdink, A. A.: Lagrangian analysis of the transport and processing of
- 549 agricultural runoff in the lower Maumee River and Maumee Bay, J. Gt. Lakes Res., 40, 479-
- 550 495, https://doi.org/10.1016/j.jglr.2014.06.001, 2014a.
- 551 Baker, D. B., Confesor, R., Ewing, D. E., Johnson, L. T., Kramer, J. W., and Merryfield, B. J.:
- 552 Phosphorus loading to Lake Erie from the Maumee, Sandusky and Cuyahoga rivers: The
- importance of bioavailability, J. Gt. Lakes Res., 40, 502–517,
- 554 https://doi.org/10.1016/j.jglr.2014.05.001, 2014b.
- 555 Barbiero, R. P. and Tuchman, M. L.: Long-term Dreissenid Impacts on Water Clarity in Lake
- 556 Erie, J. Gt. Lakes Res., 30, 557–565, https://doi.org/10.1016/S0380-1330(04)70371-8, 2004.
- 557 Berry, M. A., Davis, T. W., Cory, R. M., Duhaime, M. B., Johengen, T. H., Kling, G. W., Marino,
- 558 J. A., Den Uyl, P. A., Gossiaux, D., Dick, G. J., and Denef, V. J.: Cyanobacterial harmful algal
- 559 blooms are a biological disturbance to Western Lake Erie bacterial communities, Environ.
- 560 Microbiol., 19, 1149–1162, https://doi.org/10.1111/1462-2920.13640, 2017.
- 561 Bertani, I., Steger, C. E., Obenour, D. R., Fahnenstiel, G. L., Bridgeman, T. B., Johengen, T. H.,
- 562 Sayers, M. J., Shuchman, R. A., and Scavia, D.: Tracking cyanobacteria blooms: Do different
- monitoring approaches tell the same story?, Sci. Total Environ., 575, 294–308,
- 564 https://doi.org/10.1016/j.scitotenv.2016.10.023, 2017.
- 565 Binding, C. E., Jerome, J. H., Bukata, R. P., and Booty, W. G.: Spectral absorption properties of
- dissolved and particulate matter in Lake Erie, Remote Sens. Environ., 112, 1702–1711,
- 567 https://doi.org/10.1016/j.rse.2007.08.017, 2008.
- 568 Bosse, K. R., Sayers, M. J., Shuchman, R. A., Fahnenstiel, G. L., Ruberg, S. A., Fanslow, D. L.,
- 569 Stuart, D. G., Johengen, T. H., and Burtner, A. M.: Spatial-temporal variability of in situ
- 570 cyanobacteria vertical structure in Western Lake Erie: Implications for remote sensing
- observations, J. Gt. Lakes Res., 45, 480–489, https://doi.org/10.1016/j.jglr.2019.02.003, 2019.
- 572 Bridoux, M., Sobiechowska, M., Perez-Fuentetaja, A., and Alben, K. T.: Algal pigments in Lake
- 573 Erie dreissenids, pseudofeces and sediments, as tracers of diet, selective feeding and
- 574 bioaccumulation, J. Gt. Lakes Res., 36, 437–447, https://doi.org/10.1016/j.jglr.2010.06.005,
- 575 2010.
- 576 Buratti, F. M., Manganelli, M., Vichi, S., Stefanelli, M., Scardala, S., Testai, E., and Funari, E.:
- 577 Cyanotoxins: producing organisms, occurrence, toxicity, mechanism of action and human health

- 578 toxicological risk evaluation, Arch. Toxicol., 91, 1049–1130, https://doi.org/10.1007/s00204-016-
- 579 1913-6, 2017.
- 580 Carmichael, W. W. and Boyer, G. L.: Health impacts from cyanobacteria harmful algae blooms:
- Implications for the North American Great Lakes, Harmful Algae, 54, 194–212,
- 582 https://doi.org/10.1016/j.hal.2016.02.002, 2016.
- 583 Chaffin, J. D. and Bridgeman, T. B.: Organic and inorganic nitrogen utilization by nitrogen-
- stressed cyanobacteria during bloom conditions, J. Appl. Phycol., 26, 299–309,
- 585 https://doi.org/10.1007/s10811-013-0118-0, 2014.
- 586 Chaffin, J. D., Bridgeman, T. B., Heckathorn, S. A., and Mishra, S.: Assessment of Microcystis
- 587 growth rate potential and nutrient status across a trophic gradient in western Lake Erie, J. Gt.
- 588 Lakes Res., 37, 92–100, https://doi.org/10.1016/j.jglr.2010.11.016, 2011.
- 589 Charlton, M. N., Milne, J. E., Booth, W. G., and Chiocchio, F.: Lake Erie Offshore in 1990:
- Restoration and Resilience in the Central Basin, J. Gt. Lakes Res., 19, 291–309,
- 591 https://doi.org/10.1016/S0380-1330(93)71218-6, 1993.
- 592 Conroy, J. D., Kane, D. D., Dolan, D. M., Edwards, W. J., Charlton, M. N., and Culver, D. A.:
- 593 Temporal Trends in Lake Erie Plankton Biomass: Roles of External Phosphorus Loading and
- 594 Dreissenid Mussels, J. Gt. Lakes Res., 31, 89–110, https://doi.org/10.1016/S0380-
- 595 1330(05)70307-5, 2005.
- 596 Cooperative Institute for Great Lakes Research, University of Michigan; NOAA Great Lakes
- 597 Environmental Research Laboratory: Physical, chemical, and biological water quality monitoring
- 598 data to support detection of Harmful Algal Blooms (HABs) in western Lake Erie, collected by the
- 599 Great Lakes Environmental Research Laboratory and the Cooperative Institute for Great Lakes
- Research since 2012, NOAA National Centers for Environmental Information [data set],
- 601 https://doi.org/10.25921/11da-3x54, 2019.
- 602 Cory, R. M., Davis, T. W., Dick, G. J., Johengen, T., Denef, V. J., Berry, M. A., Page, S. E.,
- Watson, S. B., Yuhas, K., and Kling, G. W.: Seasonal Dynamics in Dissolved Organic Matter,
- Hydrogen Peroxide, and Cyanobacterial Blooms in Lake Erie, Front. Mar. Sci., 3, 2016.
- 605 Cousino, L. K., Becker, R. H., and Zmijewski, K. A.: Modeling the effects of climate change on
- water, sediment, and nutrient yields from the Maumee River watershed, J. Hydrol. Reg. Stud., 4,
- 607 762–775, https://doi.org/10.1016/j.ejrh.2015.06.017, 2015.
- 608 Den Uyl, P. A., Harrison, S. B., Godwin, C. M., Rowe, M. D., Strickler, J. R., and Vanderploeg,
- 609 H. A.: Comparative analysis of Microcystis buoyancy in western Lake Erie and Saginaw Bay of
- 610 Lake Huron, Harmful Algae, 108, 102102, https://doi.org/10.1016/j.hal.2021.102102, 2021.
- 611 Den Uyl, P. A., Thompson, L. R., Errera, R. M., Birch, J. M., Preston, C. M., Ussler, W. I.,
- 612 Yancey, C. E., Chaganti, S. R., Ruberg, S. A., Doucette, G. J., Dick, G. J., Scholin, C. A., and
- 613 Goodwin, K. D.: Lake Erie field trials to advance autonomous monitoring of cyanobacterial
- 614 harmful algal blooms, Front. Mar. Sci., 9, https://doi.org/10.3389/fmars.2022.1021952, 2022.
- 615 Dolan, D. M. and Chapra, S. C.: Great Lakes total phosphorus revisited: 1. Loading analysis
- and update (1994–2008), J. Gt. Lakes Res., 38, 730–740,
- 617 https://doi.org/10.1016/j.jglr.2012.10.001, 2012.

- 618 Environment and Climate Change Canada and the U.S. Environmental Protection Agency.
- 619 2022. State of the Great Lakes 2022 Technical Report. Cat No. En161-3/1E-PDF. EPA 905-
- 620 R22-004. Available at binational.net, 2022.
- 621 Fang, S., Del Giudice, D., Scavia, D., Binding, C. E., Bridgeman, T. B., Chaffin, J. D., Evans, M.
- 622 A., Guinness, J., Johengen, T. H., and Obenour, D. R.: A space-time geostatistical model for
- 623 probabilistic estimation of harmful algal bloom biomass and areal extent, Sci. Total Environ.,
- 624 695, 133776, https://doi.org/10.1016/j.scitotenv.2019.133776, 2019.
- 625 Goeyens, L., Kindermans, N., Abu Yusuf, M., and Elskens, M.: A Room Temperature Procedure
- for the Manual Determination of Urea in Seawater, Estuar. Coast. Shelf Sci., 47, 415–418,
- 627 https://doi.org/10.1006/ecss.1998.0357, 1998.
- 628 Hartig, J. H., Zarull, M. A., Ciborowski, J. J. H., Gannon, J. E., Wilke, E., Norwood, G., and
- 629 Vincent, A. N.: Long-term ecosystem monitoring and assessment of the Detroit River and
- 630 Western Lake Erie, Environ. Monit. Assess., 158, 87–104, https://doi.org/10.1007/s10661-008-
- 631 0567-0, 2009.
- 632 GLWQA: Great Lakes Water Quality Agreement; Protocol Amending the Agreement Between
- 633 Canada and the United States of America on Great Lakes Water Quality, 1978, as Amended on
- 634 October 16, 1983 and on November 18, 1987, https://binational. net/2012/09/05/2012-glwqa-
- 635 aqegl/ (last access: November 2022), 2012.
- 636 637 Hartig, J. H., Francoeur, S. N., Ciborowski, J. J. H., Gannon, J. E., Sanders, C. E., Galvao-
- 638 Ferreira, P., Knauss, C. R., Gell, G., and Berk, K.: An ecosystem health assessment of the
- 639 Detroit River and western Lake Erie, J. Gt. Lakes Res., 47, 1241–1256,
- 640 https://doi.org/10.1016/j.jglr.2021.05.008, 2021.
- 641 Hedges, J. I. and Stern, J. H.: Carbon and nitrogen determinations of carbonate-containing
- 642 solids1, Limnol. Oceanogr., 29, 657–663, https://doi.org/10.4319/lo.1984.29.3.0657, 1984.
- Hellweger, F. L., Martin, R. M., Eigemann, F., Smith, D. J., Dick, G. J., and Wilhelm, S. W.:
- 644 Models predict planned phosphorus load reduction will make Lake Erie more toxic, Science,
- 376, 1001–1005, https://doi.org/10.1126/science.abm6791, 2022.
- Hoffman, D. K., McCarthy, M. J., Boedecker, A. R., Myers, J. A., and Newell, S. E.: The role of
- 647 internal nitrogen loading in supporting non-N-fixing harmful cyanobacterial blooms in the water
- column of a large eutrophic lake, Limnol. Oceanogr., n/a, https://doi.org/10.1002/lno.12185,
- 649 2022.
- 650 Horváth, H., Kovács, A. W., Riddick, C., and Présing, M.: Extraction methods for phycocyanin
- 651 determination in freshwater filamentous cyanobacteria and their application in a shallow lake,
- 652 Eur. J. Phycol., 48, 278–286, https://doi.org/10.1080/09670262.2013.821525, 2013.
- Huisman, J., Codd, G. A., Paerl, H. W., Ibelings, B. W., Verspagen, J. M. H., and Visser, P. M.:
- 654 Cyanobacterial blooms | Nature Reviews Microbiology, Nat. Rev. Microbiol., 16, 471–483,
- 655 https://doi.org/10.1038/s41579-018-0040-1, 2018.
- 656 Joosse, P. J. and Baker, D. B.: Context for re-evaluating agricultural source phosphorus
- 657 loadings to the Great Lakes, Can. J. Soil Sci., 91, 317–327, https://doi.org/10.4141/cjss10005,
- 658 2011.

- 659 Kane, D. D., Ludsin, S. A., Briland, R. D., Culver, D. A., and Munawar, M.: Ten+years gone:
- 660 Continued degradation of offshore planktonic communities in U.S. waters of Lake Erie's western
- and central basins (2003–2013), J. Gt. Lakes Res., 41, 930–933,
- 662 https://doi.org/10.1016/j.jglr.2015.06.002, 2015.
- Kast, J. B., Apostel, A. M., Kalcic, M. M., Muenich, R. L., Dagnew, A., Long, C. M., Evenson, G.,
- and Martin, J. F.: Source contribution to phosphorus loads from the Maumee River watershed to
- 665 Lake Erie, J. Environ. Manage., 279, 111803, https://doi.org/10.1016/j.jenvman.2020.111803,
- 666 2021.
- 667 Kharbush, J. J., Smith, D. J., Powers, M., Vanderploeg, H. A., Fanslow, D., Robinson, R. S.,
- 668 Dick, G. J., and Pearson, A.: Chlorophyll nitrogen isotope values track shifts between
- 669 cyanobacteria and eukaryotic algae in a natural phytoplankton community in Lake Erie, Org.
- 670 Geochem., 128, 71–77, https://doi.org/10.1016/j.orggeochem.2018.12.006, 2019.
- 671 Kharbush, J. J., Robinson, R. S., and Carter, S. J.: Patterns in sources and forms of nitrogen in
- a large eutrophic lake during a cyanobacterial harmful algal bloom, Limnol. Oceanogr., n/a,
- 673 https://doi.org/10.1002/lno.12311, 2023.
- 674 King, W. M., Curless, S. E., and Hood, J. M.: River phosphorus cycling during high flow may
- 675 constrain Lake Erie cyanobacteria blooms, Water Res., 222, 118845,
- 676 https://doi.org/10.1016/j.watres.2022.118845, 2022.
- 677 Liu, Q., Rowe, M. D., Anderson, E. J., Stow, C. A., Stumpf, R. P., and Johengen, T. H.:
- 678 Probabilistic forecast of microcystin toxin using satellite remote sensing, in situ observations and
- numerical modeling, Environ. Model. Softw., 128, 104705,
- 680 https://doi.org/10.1016/j.envsoft.2020.104705, 2020.
- Lunetta, R. S., Schaeffer, B. A., Stumpf, R. P., Keith, D., Jacobs, S. A., and Murphy, M. S.:
- 682 Evaluation of cyanobacteria cell count detection derived from MERIS imagery across the
- 683 eastern USA, Remote Sens. Environ., 157, 24–34, https://doi.org/10.1016/j.rse.2014.06.008,
- 684 2015.
- 685 Maguire, T. J., Stow, C. A., and Godwin, C. M.: Spatially referenced Bayesian state-space
- 686 model of total phosphorus in western Lake Erie, Hydrol. Earth Syst. Sci., 26, 1993–2017.
- 687 https://doi.org/10.5194/hess-26-1993-2022, 2022.
- Makarewicz, J. C. and Bertram, P.: Evidence for the Restoration of the Lake Erie Ecosystem:
- Water quality, oxygen levels, and pelagic function appear to be improving, BioScience, 41, 216–
- 690 223, https://doi.org/10.2307/1311411, 1991.
- 691 Marino, J. A., Denef, V. J., Dick, G. J., Duhaime, M. B., and James, T. Y.: Fungal community
- 692 dynamics associated with harmful cyanobacterial blooms in two Great Lakes, J. Gt. Lakes Res.,
- 693 48, 1021–1031, https://doi.org/10.1016/j.jglr.2022.05.007, 2022.
- 694 Matisoff, G., Kaltenberg, E. M., Steely, R. L., Hummel, S. K., Seo, J., Gibbons, K. J.,
- 695 Bridgeman, T. B., Seo, Y., Behbahani, M., James, W. F., Johnson, L. T., Doan, P., Dittrich, M.,
- 696 Evans, M. A., and Chaffin, J. D.: Internal loading of phosphorus in western Lake Erie, J. Gt.
- 697 Lakes Res., 42, 775–788, https://doi.org/10.1016/j.jglr.2016.04.004, 2016.

- 698 Michalak, A. M., Anderson, E. J., Beletsky, D., Boland, S., Bosch, N. S., Bridgeman, T. B.,
- Chaffin, J. D., Cho, K., Confesor, R., Daloğlu, I., DePinto, J. V., Evans, M. A., Fahnenstiel, G. L.,
- He, L., Ho, J. C., Jenkins, L., Johengen, T. H., Kuo, K. C., LaPorte, E., Liu, X., McWilliams, M.
- 701 R., Moore, M. R., Posselt, D. J., Richards, R. P., Scavia, D., Steiner, A. L., Verhamme, E.,
- 702 Wright, D. M., and Zagorski, M. A.: Record-setting algal bloom in Lake Erie caused by
- agricultural and meteorological trends consistent with expected future conditions, Proc. Natl.
- 704 Acad. Sci., 110, 6448–6452, https://doi.org/10.1073/pnas.1216006110, 2013.
- 705 Mitchell, B.G., Kahru, M., Wieland, J., and Stramska, M.: Determination of spectral absorption
- 706 coefficients of particles, dissolved material and phytoplankton for discrete water samples, In:
- 707 Mueller, J.L., G.S. Fargion, and C.R. McClain [Eds.] Ocean Optics Protocols for Satellite Ocean
- 708 Color Sensor Validation, Revision 4, Volume IV: Inherent Optical Properties: Instruments,
- 709 Characterizations, Field Measurements and Data Analysis Protocols. NASA/TM- 2003-211621,
- NASA Goddard Space Flight Center, Greenbelt, MD, Chapter 4, pp 39-64, 2003.
- 711 Mohamed, M. N., Wellen, C., Parsons, C. T., Taylor, W. D., Arhonditsis, G., Chomicki, K. M.,
- 712 Boyd, D., Weidman, P., Mundle, S. O. C., Cappellen, P. V., Sharpley, A. N., and Haffner, D. G.:
- 713 Understanding and managing the re-eutrophication of Lake Erie: Knowledge gaps and research
- 714 priorities, Freshw. Sci., 38, 675–691, https://doi.org/10.1086/705915, 2019.
- 715 Mulvenna, P. F. and Savidge, G.: A modified manual method for the determination of urea in
- seawater using diacetylmonoxime reagent, Estuar. Coast. Shelf Sci., 34, 429–438,
- 717 https://doi.org/10.1016/S0272-7714(05)80115-5, 1992.
- 718 Myers, D.N., Thomas, M.A., Frey, J.W., Rheaume, S.J., and Button, D.T.: Water Quality in the
- 719 Lake Erie-Lake Saint Clair Drainages Michigan, Ohio, Indiana, New York, and Pennsylvania,
- 720 1996–98: U.S. Geological Survey Circular 1203, 35 p., https://pubs.water.usgs.gov/circ1203/,
- 721 2000.
- 722
- 723 NCWQR: Heidelberg Tributary Loading Program (HTLP) Dataset. Zenodo.
- 724 https://doi.org/10.5281/zenodo.6606949, 2022.
- Newell, S. E., Davis, T. W., Johengen, T. H., Gossiaux, D., Burtner, A., Palladino, D., and
- 726 McCarthy, M. J.: Reduced forms of nitrogen are a driver of non-nitrogen-fixing harmful
- 727 cyanobacterial blooms and toxicity in Lake Erie, Harmful Algae, 81, 86–93,
- 728 https://doi.org/10.1016/j.hal.2018.11.003, 2019.
- 729 Pirasteh, S., Mollaee, S., Fatholahi, S. N., and Li, J.: Estimation of Phytoplankton Chlorophyll-a
- 730 Concentrations in the Western Basin of Lake Erie Using Sentinel-2 and Sentinel-3 Data, Can. J.
- 731 Remote Sens., 46, 585–602, https://doi.org/10.1080/07038992.2020.1823825, 2020.
- 732 Prater, C., Frost, P. C., Howell, E. T., Watson, S. B., Zastepa, A., King, S. S. E., Vogt, R. J., and
- 733 Xenopoulos, M. A.: Variation in particulate C: N: P stoichiometry across the Lake Erie
- vatershed from tributaries to its outflow, Limnol. Oceanogr., 62, S194–S206,
- 735 https://doi.org/10.1002/lno.10628, 2017.
- 736 Qian, S. S., Stow, C. A., Rowland, F. E., Liu, Q., Rowe, M. D., Anderson, E. J., Stumpf, R. P.,
- 737 and Johengen, T. H.: Chlorophyll a as an indicator of microcystin: Short-term forecasting and
- 738 risk assessment in Lake Erie, Ecol. Indic., 130, 108055,
- 739 https://doi.org/10.1016/j.ecolind.2021.108055, 2021.

- 740 Reavie, E. D., Cai, M., Twiss, M. R., Carrick, H. J., Davis, T. W., Johengen, T. H., Gossiaux, D.,
- Smith, D. E., Palladino, D., Burtner, A., and Sgro, G. V.: Winter-spring diatom production in
- 742 Lake Erie is an important driver of summer hypoxia, J. Gt. Lakes Res., 42, 608–618,
- 743 https://doi.org/10.1016/j.jglr.2016.02.013, 2016.
- Rowe, M. D., Anderson, E. J., Wynne, T. T., Stumpf, R. P., Fanslow, D. L., Kijanka, K.,
- 745 Vanderploeg, H. A., Strickler, J. R., and Davis, T. W.: Vertical distribution of buoyant Microcystis
- blooms in a Lagrangian particle tracking model for short-term forecasts in Lake Erie, J.
- 747 Geophys. Res. Oceans, 121, 5296-5314, https://doi.org/10.1002/2016JC011720, 2016.
- 748 Rowland, F. E., Stow, C. A., Johengen, T. H., Burtner, A. M., Palladino, D., Gossiaux, D. C.,
- 749 Davis, T. W., Johnson, L. T., and Ruberg, S.: Recent Patterns in Lake Erie Phosphorus and
- 750 Chlorophyll a Concentrations in Response to Changing Loads, Environ. Sci. Technol., 54, 835–
- 751 841, https://doi.org/10.1021/acs.est.9b05326, 2020.
- 752 Sayers, M., Fahnenstiel, G. L., Shuchman, R. A., and Whitley, M.: Cyanobacteria blooms in
- 753 three eutrophic basins of the Great Lakes: a comparative analysis using satellite remote
- 754 sensing, Int. J. Remote Sens., 37, 4148–4171, https://doi.org/10.1080/01431161.2016.1207265,
- 755 2016.
- 756 Sayers, M. J., Bosse, K. R., Shuchman, R. A., Ruberg, S. A., Fahnenstiel, G. L., Leshkevich, G.
- 757 A., Stuart, D. G., Johengen, T. H., Burtner, A. M., and Palladino, D.: Spatial and temporal
- 758 variability of inherent and apparent optical properties in western Lake Erie: Implications for
- 759 water quality remote sensing, J. Gt. Lakes Res., 45, 490–507,
- 760 https://doi.org/10.1016/j.jglr.2019.03.011, 2019.
- 761 Smith, D. J., Tan, J. Y., Powers, M. A., Lin, X. N., Davis, T. W., and Dick, G. J.: Individual
- 762 Microcystis colonies harbour distinct bacterial communities that differ by Microcystis oligotype
- 763 and with time, Environ. Microbiol., 23, 3020–3036, https://doi.org/10.1111/1462-2920.15514,
- 764 2021.
- 765 Smith, D. J., Berry, M. A., Cory, R. M., Johengen, T. H., Kling, G. W., Davis, T. W., and Dick, G.
- 766 J.: Heterotrophic Bacteria Dominate Catalase Expression during Microcystis Blooms, Appl.
- 767 Environ. Microbiol., 88, e02544-21, https://doi.org/10.1128/aem.02544-21, 2022.
- Smith, R. B., Bass, B., Sawyer, D., Depew, D., and Watson, S. B.: Estimating the economic
- costs of algal blooms in the Canadian Lake Erie Basin, Harmful Algae, 87, 101624,
- 770 https://doi.org/10.1016/j.hal.2019.101624, 2019.
- 771 Speziale, B. J., Schreiner, S. P., Giammatteo, P. A., and Schindler, J. E.: Comparison of N,N-
- 772 Dimethylformamide, Dimethyl Sulfoxide, and Acetone for Extraction of Phytoplankton
- 773 Chlorophyll, Can. J. Fish. Aquat. Sci., 41, 1519–1522, https://doi.org/10.1139/f84-187, 1984.
- 774 Standard Methods Committee of the American Public Health Association, American Water
- 775 Works Association, and Water Environment Federation: Standard Methods For the Examination
- of Water and Wastewater, 23<sup>rd</sup> edition, Sections 2540 Solids, 4500-P Phosphorus, 4500-nh3-
- 777 nitrogen (ammonia), 4500-no3-nitrogen (nitrate), 5310 Total Organic Carbon, edited by: Lipps
- WC, Baxter TE, Braun-Howland E, APHA Press, Washington, DC, ISBN 1625762402, 2017.

- 779 Steffen, M. M., Belisle, B. S., Watson, S. B., Boyer, G. L., and Wilhelm, S. W.: Status, causes
- and controls of cyanobacterial blooms in Lake Erie, J. Gt. Lakes Res., 40, 215–225,
- 781 https://doi.org/10.1016/j.jglr.2013.12.012, 2014.
- 782 Steffen, M. M., Davis, T. W., McKay, R. M. L., Bullerjahn, G. S., Krausfeldt, L. E., Stough, J. M.
- A., Neitzey, M. L., Gilbert, N. E., Boyer, G. L., Johengen, T. H., Gossiaux, D. C., Burtner, A. M.,
- 784 Palladino, D., Rowe, M. D., Dick, G. J., Meyer, K. A., Levy, S., Boone, B. E., Stumpf, R. P.,
- 785 Wynne, T. T., Zimba, P. V., Gutierrez, D., and Wilhelm, S. W.: Ecophysiological Examination of
- 786 the Lake Erie Microcystis Bloom in 2014: Linkages between Biology and the Water Supply
- 787 Shutdown of Toledo, OH, Environ. Sci. Technol., 51, 6745–6755,
- 788 https://doi.org/10.1021/acs.est.7b00856, 2017.
- 789 Sterner, R. W., Keeler, B., Polasky, S., Poudel, R., Rhude, K., and Rogers, M.: Ecosystem
- 790 services of Earth's largest freshwater lakes, Ecosyst. Serv., 41, 101046,
- 791 https://doi.org/10.1016/j.ecoser.2019.101046, 2020.
- 792 Stow, C. A., Cha, Y., Johnson, L. T., Confesor, R., and Richards, R. P.: Long-Term and
- 793 Seasonal Trend Decomposition of Maumee River Nutrient Inputs to Western Lake Erie, Environ.
- 794 Sci. Technol., 49, 3392–3400, https://doi.org/10.1021/es5062648, 2015.
- 795 US EPA United States Environmental Protection Agency: Method 180.1: Determination of
- 796 Turbidity by Nephelometry, Revision 2.0, Edited by: O'Dell, J.W., 1993.
- 797 US EPA United States Environmental Protection Agency: Drinking Water Health Advisory for
- 798 the Cyanobacterial Microcystin Toxins, EPA Document Number 820R15100, 2015.
- 799 Van Meter, K. J., McLeod, M. M., Liu, J., Tenkouano, G. T., Hall, R. I., Van Cappellen, P., and
- 801 Basu, N. B.: Beyond the Mass Balance: Watershed Phosphorus Legacies and the Evolution of
- the Current Water Quality Policy Challenge, Water Resour. Res., 57, e2020WR029316,
- 803 https://doi.org/10.1029/2020WR029316, 2021.
- Vander Woude, A., Ruberg, S., Johengen, T., Miller, R., and Stuart, D.: Spatial and temporal
- 805 scales of variability of cyanobacteria harmful algal blooms from NOAA GLERL airborne
- hyperspectral imagery, J. Gt. Lakes Res., 45, 536–546,
- 807 https://doi.org/10.1016/j.jglr.2019.02.006, 2019.
- 808 Vanderploeg, H. A., Liebig, J. R., Carmichael, W. W., Agy, M. A., Johengen, T. H., Fahnenstiel,
- 809 G. L., and Nalepa, T. F.: Zebra mussel (Dreissena polymorpha) selective filtration promoted
- 810 toxic Microcystis blooms in Saginaw Bay (Lake Huron) and Lake Erie, Can. J. Fish. Aquat. Sci.,
- 811 58, 1208–1221, https://doi.org/10.1139/f01-066, 2001.
- 812 Wang, Q. and Boegman, L.: Multi-Year Simulation of Western Lake Erie Hydrodynamics and
- 813 Biogeochemistry to Evaluate Nutrient Management Scenarios, Sustainability, 13, 7516.
- 814 https://doi.org/10.3390/su13147516, 2021.
- 815 Watson, S. B., Miller, C., Arhonditsis, G., Boyer, G. L., Carmichael, W., Charlton, M. N.,
- 816 Confesor, R., Depew, D. C., Höök, T. O., Ludsin, S. A., Matisoff, G., McElmurry, S. P., Murray,
- 817 M. W., Peter Richards, R., Rao, Y. R., Steffen, M. M., and Wilhelm, S. W.: The re-eutrophication
- of Lake Erie: Harmful algal blooms and hypoxia, Harmful Algae, 56, 44–66,
- 819 https://doi.org/10.1016/j.hal.2016.04.010, 2016.

- 820 Weiskerger, C. J., Rowe, M. D., Stow, C. A., Stuart, D., and Johengen, T.: Application of the
- 821 Beer-Lambert Model to Attenuation of Photosynthetically Active Radiation in a Shallow,
- 822 Eutrophic Lake, Water Resour. Res., 54, 8952–8962, https://doi.org/10.1029/2018WR023024,
- 823 2018.
- Wetzel, R.G., and Likens G.E.: Limnological Analyses, 3<sup>rd</sup> edition, Springer New York, NY,
- 825 https://doi.org/10.1007/978-1-4757-3250-4, 2000.
- 826 WHO World Health Organization: Cyanobacterial toxins: microcystins. Background document
- 827 for development of WHO Guidelines for drinking-water quality and Guidelines for safe
- 828 recreational water environments, WHO/HEP/ECH/WSH/2020.6, 2020.
- 829
- 830 Wilson, A. E., Gossiaux, D. C., Höök, T. O., Berry, J. P., Landrum, P. F., Dyble, J., and
- 831 Guildford, S. J.: Evaluation of the human health threat associated with the hepatotoxin
- 832 microcystin in the muscle and liver tissues of yellow perch (Perca flavescens), Can. J. Fish.
- 833 Aquat. Sci., 65, 1487–1497, https://doi.org/10.1139/F08-067, 2008.
- 834 Wynne, T. T., Stumpf, R. P., Tomlinson, M. C., Fahnenstiel, G. L., Dyble, J., Schwab, D. J., and
- Joshi, S. J.: Evolution of a cyanobacterial bloom forecast system in western Lake Erie:
- 836 Development and initial evaluation, J. Gt. Lakes Res., 39, 90–99,
- 837 https://doi.org/10.1016/j.jglr.2012.10.003, 2013.
- 838 Xu, J., Liu, H., Lin, J., Lyu, H., Dong, X., Li, Y., Guo, H., and Wang, H.: Long-term monitoring
- 839 particulate composition change in the Great Lakes using MODIS data, Water Res., 222,
- 840 118932, https://doi.org/10.1016/j.watres.2022.118932, 2022.
- Yancey, C.E., Mathiesen, O., and Dick, G.J.; Transcriptionally active nitrogen fixation and
- 842 biosynthesis of diverse secondary metabolites by Dolichospermum and Aphanizominom-like
- 843 Cyanobacteria in western Lake Érie Microcystis blooms, bioRxiv [preprint],
- 844 https://doi.org/10.1101/2022.09.30.510322 01 October 2022a.
- 845 Yancey, C. E., Smith, D. J., Den Uyl, P. A., Mohamed, O. G., Yu, F., Ruberg, S. A., Chaffin, J.
- 846 D., Goodwin, K. D., Tripathi, A., Sherman, D. H., and Dick, G. J.: Metagenomic and
- 847 Metatranscriptomic Insights into Population Diversity of Microcystis Blooms: Spatial and
- 848 Temporal Dynamics of mcy Genotypes, Including a Partial Operon That Can Be Abundant and
- 849 Expressed, Appl. Environ. Microbiol., 88, e02464-21, https://doi.org/10.1128/aem.02464-21,
- 850 2022b.