Spatial distributions of external and internal phosphorus loads in Lake Erie and their impacts on phytoplankton and water quality

Hongyan Zhang¹,*, Leon Boegman², Donald Scavia³, and David A. Culver⁴

1. Cooperative Institute of Limnology and Ecosystems Research (CILER), School of Natural Resources and Environments, University of Michigan, Ann Arbor, Michigan 48108 USA
2. Department of Civil Engineering, Queen’s University, Kingston, Ontario K7L 3N6 Canada
3. Water Center, Graham Sustainability Institute, University of Michigan, 625 East Liberty Road, Ann Arbor, MI 48193, USA
4. Department of Evolution, Ecology, and Organismal Biology, The Ohio State University, Columbus, Ohio 43210 USA

*Corresponding author. Address: 4840 South State Road, Ann Arbor, MI 48108, USA. Telephone: (734) 741-2354; Fax: (734) 741-2205.
E-mail: zhanghy@umich.edu
Abstract

Re-eutrophication in Lake Erie has led to new programs to reduce external phosphorus loads, and it is important to understand the interrelated dynamics of external and internal phosphorus loads. In addition to developing phosphorus load response curves for algal biomass in the western basin and hypoxia in the central basin, we used a two-dimensional (vertical-longitudinal) hydrodynamic and ecological model to show that both external and internal phosphorus loads were distributed homogeneously in the water column in Lake Erie’s western basin. In the stratified central and eastern basins phosphorus released by organic matter decay and crustacean zooplankton excretion was concentrated in the upper water column, contributing 100-119% of the phytoplankton phosphorus demand, while phosphorus released by dreissenids and from anoxic sediments was distributed primarily in the hypolimnion during the growing season. Simulated reductions in external phosphorus loads decreased individual phytoplankton groups most at times when they were normally most abundant, e.g., *Microcystis* decreased the most during September. Phosphorus was limiting over the simulation periods, but water temperature and light conditions also played critical roles in phytoplankton succession. While water column phosphorus responded quickly to external phosphorus reduction, pulses of phosphorus (riverine input or sediment resuspension) occurring immediately before the *Microcystis* bloom period could allow it to bloom despite long-term external phosphorus load reduction. Studies are warranted to assess the contribution of seasonal dynamics in phosphorus loading (including sediment resuspension) to *Microcystis* bloom development.
Keywords: Sediment release, Excretion, Eutrophication, Harmful algal bloom, Hypoxia

Introduction

A strong correlation between the concentration of total phosphorus and phytoplankton (hereafter simplified to algae) biomass in freshwater lakes has been documented (Lean, 1973; Scavia and Chapra, 1977; Schindler, 1977; Smith, 1982; Knoll et al., 2003), and phosphorus (P) is the most common limiting macronutrient in freshwater lakes (Schindler, 1977; Arnott and Vanni, 1996; Wetzel, 2001; Wilhelm et al., 2003). Excessive P inputs have dramatically increased water productivity and caused the eutrophication of many lakes (Chapra and Robertson, 1977; Beeton, 2002; Jin, 2003; Schindler, 2012).

Lake Erie was severely eutrophic in the 1960s, resulting from excessive external P loading (Burns and Ross, 1972). Water quality management in Lake Erie demonstrated that control of external P loading provides an effective means of decreasing eutrophication. An external P load reduction program for point sources was carried out in the early 1970s, and soon led to encouraging water quality responses. Not only did total phosphorus concentrations decrease in the water column (Rockwell et al., 1989), but total algal biomass decreased 40% in the western basin by the late 1970s, 65% by the mid-1980s, and both Cyanobacteria and filamentous greens decreased by 80% by the mid-1980s (Makarewicz and Bertram, 1991; Gopalan et al., 1998). Oxygen concentrations increased at the bottom of both the western basin (Krieger et al., 1996) and the central basin (Bertram, 1993; Ludsin et al., 2001).

Recent studies show that dissolved reactive phosphorus loads in some tributaries have increased since 1995 (Baker et al., 2014; IJC, 2014; Scavia et al., 2014) and algal biomass has increased as well (Conroy et al., 2005a). In recent years increases in the frequency and magnitude of *Microcystis* blooms (Michalak et al., 2013; Stumpf et al., 2012) suggest that
climate may be an additional factor triggering the resurgent blooms, which are correlated with an increased soluble phosphorus fraction from agriculturally dominated tributaries (Kane et al., 2014) and the total phosphorus load from spring freshets (Stumpf et al., 2012). The changes in loads are driven by climate-induced variability in precipitation (Scavia et al., 2014) and are accompanied by trends toward warm, calm meteorology during summer, which combine to cause bloom-favorable conditions (Michalak et al., 2013). Consequently, even further reduction in the external P load targets has been recommended (Rucinski et al., 2014; Scavia et al., 2014, and papers in this issue), and the governments of Canada and the United States announced a target of 40% reduction in total phosphorus loads to Lake Erie on February 22, 2016 (USEPA, 2016).

While external loading reduction can reduce symptoms of eutrophication, this result is often delayed by release of internal nutrient loads from years of accumulations and decay of P-rich organic matter in the sediments (Phillips et al., 2005; Turner et al., 2008). Central basin hypoxia (and to a lesser extent in the western basin) can cause sediment ferric phosphate deposits to change to the much more soluble ferrous phosphate form, promoting diffusion of soluble reactive phosphate out of the sediments. Another important internal P source to Lake Erie is excretion by zebra mussels (*Dreissena polymorpha* Pallas) and quagga mussels (*D. rostriformis bugensis* Andrusov). These taxa excrete considerable phosphate (Arnott and Vanni, 1996; James et al., 1997) at rates up to 2.8 mg/m²/d (Conroy et al., 2005b), sufficient to replace the pool of soluble reactive phosphorus (SRP) in the water column in less than 10 days (Conroy et al., 2005b). However, other studies indicate that dreissenid mussels intercept incoming nutrients in the nearshore area and lead to offshore ‘desertification’ (e.g., Hecky et al. 2004). The mussel populations retain a large amount of phosphorus in their body tissue (Mellina et al.,
1995), which can be released into the water with unclear temporal and spatial patterns. An ecosystem with a high cumulative internal P loading rate from these sources can sustain a eutrophic state well after external P loading has decreased. Thus, the amount and spatial distribution of internal P loading strongly affects the efficiency of any external P reduction program.

In this study, we used Zhang et al.’s (2008) model to simulate the spatial distributions of phosphorus throughout Lake Erie’s western, central, and eastern basins during 1997 and 1998. In addition to being the calibration and confirmation years for the model, 1997 and 1998 had higher P loads than the annual target load of 11,000 mt and displayed varying Microcystis bloom tendencies. Phosphorus loads in 1997 were 16,800 mt with no Microcystis bloom, whereas 1998 had lower P loads (12,700 mt) with a moderate Microcystis bloom. We evaluated the effects of different reduction levels (20%, 40%, 60%, and 80%) of external total phosphorus loading on algae in the western basin by comparing the biomass of three algal groups with and without reductions in external P inputs to explore years that experience annual external P loads higher than the annual target load of 11,000 mt. We also evaluated the effects of different reduction levels of external total phosphorus loading on hypoxia in the central basin by comparing the hypolimnetic oxygen concentration and hypoxic area with and without reduction in external P inputs, which was similar to the analyses in the multi-model team reports of the Great Lakes Water Quality Agreement Nutrient Annex 4 (Scavia et al., this issue; Scavia and DePinto, 2015). Model simulations under different P reduction scenarios provide an overview of the P fluxes and fates in the Lake Erie ecosystem during the summer growing season and how they respond to external P loads.

**Methods**
Model Description

A two-dimensional (vertical-longitudinal) hydrodynamic and ecological model, EcoLE, was applied to Lake Erie to simulate the effects of external and internal P loading on the Lake Erie ecosystem. The model is an adaptation of the USACE CE-QUAL-W2 version 2 (Cole and Buchak, 1995), with modifications for large lake hydrodynamics (Boegman et al., 2001), multiple algal groups and dreissenid mussels (Zhang et al., 2008). Hydrodynamics and water quality simulations were calibrated and validated in a previous study (Zhang et al., 2008). EcoLE divides Lake Erie into as many as 65 vertical layers at 1-m intervals and 220 longitudinal segments (2-km wide from west to east). The depths of segments were assigned relative to the Great Lakes Datum (GLD) of 1985. State variables in EcoLE include free water surface elevation, horizontal velocity, vertical velocity, water density, water temperature, suspended solids, dissolved organic matter (DOM), particulate organic matter (POM), diatom-derived particulate organic matter (D-POM), soluble reactive phosphorus (SRP), ammonium, nitrate+nitrite, silicon, dissolved oxygen, algal groups (non-diatom edible algae (NDEA), diatoms, and non-diatom inedible algae (NDIA)), cladocerans and four life stages of copepods (eggs, nauplii, copepodites and adults). Data of water temperature and the water quality state variables mentioned above were taken from the Ohio State University’s Lake Erie Plankton Abundance Study database to initialize, calibrate and verify the model. Other data for external driving forces were provided by various data sources (Zhang et al. 2008).

Dreissenid mussels were modeled here as external forces, grazing on algae and excreting phosphate and ammonia back into the system. Zhang et al. (2008) used the depth-dependent dreissenid density estimations by Jarvis et al. (2000). However, due to the seasonal hypoxia, low mussel densities were recorded in the deep central basin (Burlakova et al., 2014; Jarvis et
In this study, we used Jarvis et al.’s density (135 ind/m²) to populate deep water areas (>22 m) in the central basin. The remaining settings for mussel simulations were kept as in Zhang et al. (2008). We compared the simulation results of before and after the change (see Supplementary Materials A for details).

The simulation periods were chosen based on availability of field data, and ran from May 10 - September 30, 1997 (model calibration) and June 10 - October 30, 1998 (model confirmation) (Figure 1). Using hydrodynamics coefficients based on the Boegman et al. (2001) calibration, our previous modeling efforts with EcoLE focused on the simulation of water temperature, phosphorus and nitrogen, and biomass of algal groups and zooplankton groups over the summer growing season (Zhang et al., 2008). The simulated surface and bottom water temperatures showed good agreement with field observations, and the simulated values of biological and nutrient state variables also matched well with field measurements. Taken together, the simulation results of Boegman et al.’s (2008a) model detailing the interaction between hydrodynamics and dreissenid impacts, and the agreement of state variables (algae, N+N, NH₄ and SRP) between field observations and model predictions (Boegman et al., 2008b), and the simulation results of dissolved oxygen in the western central basin (Conroy et al. 2011) and in the central basin (Scavia and DePinto, 2015), we consider our current model to be a valid analytical tool, which we use herein to study the processes involved in phosphorus recycling in the Lake Erie system under varying external P inputs.

External Phosphorus Loads

We estimated seasonal external P loads from wastewater treatment plants (WWTP) and non-point sources entering via rivers (Table 1, Figure 1). Phosphorus load from rivers was
measured as soluble reactive phosphorus (SRP) and total phosphorus (TP). SRP was input into EcoLE directly. Although a big fraction of TP from tributaries is inorganic mater, the current model doesn’t simulate the dynamics of inorganic particulate phosphorus. So other phosphorus (TP minus SRP) was converted into phosphorus-containing organic matter assuming organic matter contains 1% phosphorus (Bowie et al., 1985) and was input into EcoLE as organic matter. Because only TP data were available from WWTP, 76.9% of TP was considered as soluble phosphorus and 23.1% was considered as organic matter phosphorus (Young et al., 1982). The external P was assumed to be well mixed in its entrance model cells (i.e., specific segments and layers) and reached other cells by physical or biochemical transport that varied from location to location within the lake. The total external P loads over the simulation periods were calculated as the sum of the products of discharge flow and P concentration in the discharge from each tributary. See Supplementary Materials B for calculation details. To calculate their accumulative spatial distributions over the simulation periods, we turned off all the biological and chemical processes in the model, and the resultant spatial distribution was fully determined by physical transport and mixing processes.

Internal Phosphorus Loads

Two internal P sources were considered in the model: anoxic sediment release and dreissenid mussel excretion. For comparison, we also considered two phosphorus cycling processes in the water column: crustacean zooplankton excretion (combined excretion from cladocerans and copepods) and organic matter decay. The phosphorus (as SRP) released from these sources was involved in the SRP dynamics indiscriminately. However, to depict the spatial distribution of phosphorus and track the amount of phosphorus from each source, the
four sources were recorded as extra state variables. The resultant spatial distributions from physical mixing processes determined the potential availability from each P source to algal growth.

Anoxic sediment release

Under normoxic conditions, sediments hardly release any phosphorus to the overlying water, whereas they release phosphorus dramatically under anoxic conditions (c.f., Mortimer, 1941, 1971; Burns and Ross, 1972; Di Toro and Connolly, 1980). Lam et al., (1987) assumed that anoxia occurred at 1.5 mg O₂/L (Chapra and Canale, 1991) in the lower layer of their two-layer model, while Burns et al. (2005) considered bottom water anoxic if the epibenthic dissolved oxygen concentration declined to 1.0 mg/L. We adopted Lame et al.’s approach, and assumed that no phosphorus was released when DO concentrations of the bottom water were above 1.0 mg/L, while a constant release rate was used, 0.0044 g P/m²/d (Lam et al., 1987), when DO concentrations were below 1.0 mg/L. This DO threshold for anoxia was lower than Lam et al.’s (1987), because our deepest water layer was thinner than theirs.

Dreissenid mussel excretion

Zebra mussels first invaded Lake Erie in the late 1980s, but have been more or less replaced by quagga mussels recently (Stoeckmann, 2003, Patterson et al., 2005), such that by 1998, 84.4 % of mussels in the eastern basin, 99.7% in the central basin were quagga mussels, but only 36.9% in the western basin (Jarvis et al., 2000). We assume therefore, for simplicity, that during the 1997-1998 periods mussels in the western basin were 100% zebra mussels, whereas those in the central and the eastern basins were 100% quagga mussels. Zebra mussels and quagga mussels have different weight-specific phosphorus excretion rates (Conroy et al.,...
2005b) and the phosphorus excretion of a mussel population in a model cell over the growing season was calculated as the sum of the products of individual excretion rates and numbers of mussels. See Supplementary Materials B for the calculation details.

Crustacean zooplankton excretion

We simulated copepods and cladocerans as the two crustacean zooplankters in the lake, using Fennel and Neumann’s (2003) stage-structured population model for copepods and a generic bioenergetic model for cladocerans (Zhang et al., 2008). We used a ratio of phosphorus to dry weight ($\delta_{p-cop}$ for copepods, $\delta_{p-clad}$ for cladocerans) to convert maintenance cost to phosphorus excretion (Andersen and Hessen 1991). See Supplementary Materials B for the calculation details.

Organic matter decay

The organic matter pools (dissolved organic matter (DOM), particulate organic matter (POM) and (diatom-derived particulate organic matter (D-POM)) in the water column had temperature-specific decay rates, which were converted to phosphorus release rates by a ratio of phosphorus to organic matter mass of 0.01 (Bowie et al., 1985). Although phosphorus in DOM is dissolved phosphorus (DP), it is not available directly to algal growth and becomes available through DOM degradation.

Basin-wide phosphorus budget
The phosphorus content of four phosphorus pools: soluble reactive phosphorus (SRP), dissolved phosphorus (DP), particulate phosphorus (PP) and crustacean phosphorus (CP), were estimated basin wide and over the growing season (Figure 2). DP referred to the phosphorus in dissolved organic matter; PP referred to the phosphorus in particulate organic matter and algae; whereas CP included the phosphorus content of cladocerans and copepods. The pathway from SRP to PP included phosphorus uptake by algae, while PP to SRP included phosphorus excreted by algae and released by POM decay. Our model had a traditional crustacean grazing food web. However, by including organic matter, the model implicitly also includes the microbial food web (Debruyn et al., 2004).

The average mass of each phosphorus pool during the simulation period was calculated as the average sum of the products of the model cell volume and the concentration of phosphorus content of a constituent mentioned above in the cell for all model cells across the whole basin. Daily basin-wide phosphorus fluxes between state variables, mussels and sediment were calculated as the daily average of a sum of products of the phosphorus transfer rate from one constituent to another within a model cell and the cell volume for all model cells across the whole basin. See Supplementary Materials B for the calculation details. To evaluate the effects of reduction in external P loads on the in-lake P dynamics, we calculated the percent changes in the basin-wide P budget for different P reduction scenarios from the no-reduction scenario.

External P loaded to each of the three Lake Erie basins and the exchanges between basins were calculated to complete the P budget. The net P exchanges between connected basins were the sum of the net P fluxes of NDEA-P, NDIA-P, diatom-P, SRP, DOM-P and POM-P. The P exchanges at the intersection between basin separation lines (Figure 1) were
considered as exchanges between basins. Both horizontal advection and diffusion were included.

Effects of phosphorus reduction on algal community in the western basin

To test the effects of a decrease in external P loads on different algal groups for years 1997 and 1998, we reduced the concentrations of SRP, DOM and POM (constituents in tributary loads that contain phosphorus, no algal data were available for the tributaries) by 20%, 40%, 60% and 80% in tributaries without altering flows over the simulation period, which resulted in 20%, 40%, 60% and 80% reductions in external P loads while preserving the seasonal dynamics of the P inputs.

The three modeled categories of algae in Lake Erie were non-diatom edible algae (NDEA), non-diatom inedible algae (NDIA), and diatoms. NDEA included algae of Chlorophyta, Cryptophyta and Pyrrophyta, and were dominated by genera of *Chlamydomonas*, *Oocystis*, *Chroomonas*, *Cryptomonas*, *Rhodomonas*, and *Gymnodinium*. NDIA included algae of Cyanophyta, Pyrrophyta, filamentous Chlorophyta, and were dominated by *Microcystis*. Diatoms were dominated by *Melosira*, *Fragilaria*, and *Cyclotella*. The basin-wide percent differences in the total algal biomass under different levels of external P reduction were calculated as:

\[
\frac{B_{\text{lowExtP},t} - B_{\text{ExtP},t}}{B_{\text{ExtP},t}} \times 100
\]

Where \(B_{\text{lowExtP},t}\) was basin-wide total algal biomass at time step \(t\) under different reduced levels of external P conditions, while \(B_{\text{ExtP},t}\) was under no reduction in the external P loads. We did not provide the percent change in each algal group because when the biomass was low the percent change could overemphasize the effects. Instead, we provided the time series of daily basin-
wide algal biomass over the simulation periods for each algal group. We calculated the changes in the monthly-averaged algal biomass for each algal group and the total algal biomass for September, when the peak of Microcystis blooms occurred (Bridgeman et al., 2012; Wynne et al., 2010).

Factors influencing the dynamics of algal biomass

In order to analyze in detail the processes that could affect the dynamics of different algal groups, we picked a representative model cell that was located in the western basin close to the Maumee River (segment 5 at 1 m depth), which should show strong effects from changes in the external P loads. We output daily-averaged limiting factors, net growth rates (d^{-1}) (gross growth rate minus excretion rate, mortality rate and settling rate), daily net growth (g DW/m^3/d) (the product of net growth rates and algal biomass), and crustacean zooplankton consumptions (g DW/m^3/d) of different algal groups for this model cell under different levels of external P reductions. In the western basin, all mussels were located on the bottom layer (Zhang et al. 2008), so no mussel grazing occurred in this cell.

Effects of phosphorus reduction on hypoxia in the central basin

Sediment oxygen demand (SOD) is expressed as a function of oxygen concentration and temperature (Lucas and Thomas 1972, Lam et al. 1987).

\[
SOD = SOD_{\text{max}} \times \frac{\Phi_{DO}}{\Phi_{DO} + O_h} \theta^{(T-20)}
\]

where \(SOD_{\text{max}}\) is maximum sediment oxygen demand at 20°C, g O_2 m^{-2} d^{-1}, \(O_h\) is oxygen concentration half-saturation constant, and \(\Phi_{DO}\) is the DO concentration in the bottom layer right above sediment. In addition to changes in external P loads, we adjusted the
maximum SOD ($SOD_{\text{max}}$) for different TP load scenarios according to the empirical relationship between TP loads and SOD developed by Rucinski et al. (2014; Scavia and DePinto, 2015). To calculated hypoxic area (area with DO <2 mg/L), we used the Zhou et al.'s method (Zhou et al. 2013), which was based on the average of bottom dissolved oxygen concentration (DO) measured by U.S. Environmental Protect Agency every fall from 10 monitoring stations. Our bottom DO (1 - 3m above the sediment) was averaged among model segments that corresponding to the 10 monitoring stations from August 1 to September 30, 1998, a year that had an annual TP phosphorus load close to that of 2008. We then converted these DO concentrations to hypoxic area using Zhou et al.'s empirical relationship for each TP reduction scenario.

**Results**

*External phosphorus loads*

The total phosphorus loaded into Lake Erie from tributaries during the simulation periods was primarily from the Maumee and the Detroit Rivers, whereas loads from waste water treatment plants were relatively small (Table 1). Temporally, most of the external P loads occurred during early summer (Julian day 131-160) in 1997, while a big fraction of external P load occurred during summer 1998 (Julian day 190-240) (Figure 3). Spatially, the model depicts that most of the external P loads, under pure advection and physical mixing processes, were concentrated in the western and west-central basins (Figure 4), and contributed little to the phosphorus concentrations in the east-central and eastern basins.

*Spatial distribution of internal phosphorus loads*
We used specific state variables to track the phosphorus from each of the two internal P sources and the two water column cycling processes. The tracked phosphorus was accumulated in the water column and distributed fully by the physical mixing process. Here, we report the resultant phosphorus concentration distributions from different P sources at the end of September.

Anoxic sediment released phosphorus primarily at the bottom of the central basin over the simulation period (Figure 5a). Sediment phosphorus was trapped in the hypolimnion by thermal stratification and showed little influence on the epilimnion phosphorus concentration during the growing season. Because sediment only releases phosphorus when it is under anoxic conditions, the sediment phosphorus release profiles also reflect the simulated oxygen conditions on the lake bottom.

Dreissenid mussels excreted a total amount of phosphorus higher than that of zooplankton in the western basin (Table 2). Dreissenid excretion resulted in 5-20 μg P/L in the upper water in the western and the west-central basins, while it resulted in less than 5 μg P/L in the upper water in the rest of the lake (Figure 5b). Very low dreissenid excretion occurred in the central basin, which experiences seasonal hypoxia. In the stratified eastern basin, most of the phosphorus was concentrated in the lower water column.

Vertically, crustacean excretion was concentrated in the water column with depths of 20 m and shallower in 1997 (Figure 5c). Horizontally, crustacean excretion contributed the highest fluxes in the segments close to the Maumee River mouth (segments 1-11) and in the western central basin. By the end of September, crustacean excretion resulted in a concentration in the water column as high as 20 μg P/L in 1997 (Figure 5c). The phosphorus excreted by crustaceans in the western basin between segments 12 and 24 (east of the segments influenced
by the Maumee River) was extremely low, which illustrates the influence of the Detroit River’s high flows and low phosphorus content. The phosphorus released by organic matter in 1997 was also concentrated in segments 1-11 that were close to the Maumee River and in the upper water column of the central and eastern basins, where it resulted in up to 50 µg P/L by the end of September (Figure 5d).

Spatial distributions of these four P sources in 1998 were similar to those of 1997 (Figure 6). However, the zooplankton excretion of phosphorus in 1998 contributed little to the deep water phosphorus content in the eastern basin compared to 1997 (Figures 5c and 6c). Organic matter decay resulted in a lower phosphorus concentration in eastern Lake Erie compared to that in 1997 (Figures 5d and 6d), consistent with the lower external nutrient loads in 1998 than in 1997.

**Basin-wide phosphorus budgets**

Particulate phosphorus (PP) was the largest phosphorus pool, with an average of 2143 mt in the lake in 1997 and 2366 mt in 1998 (Tables 2-3). SRP in the central and eastern basins was similar to, or larger than dissolved organic phosphorus (DP), but was much lower than DP in the western basin. The zooplankton phosphorus pool was the smallest phosphorus pool among the four, and was less than one-tenth of the other particulate phosphorus pool.

The model showed that the most active phosphorus pathway was from SRP to PP, due to the active phosphorus uptake by algae in all three basins for both years (Tables 2-3). Algal uptake could deplete the western basin SRP pool within 2 days, while the phosphorus regenerated daily in the water column by organic matter and zooplankton excretion was 82-85% of the western basin algal demand, with 26-36% from DP, 30-43% from PP, and 16% from crustaceans. Dreissenid mussels could provide 19-27% of the algal P demand each day. In the
central and eastern basins, the P regenerated daily in the water column by organic matter and zooplankton excretion was 100-119% of the algal demand, with 78-97% from organic matter decay. Dreissenid mussels could provide 7% of algal P demand in the central basin and 24-27% in the eastern basin, but it only becomes available to algae after fall turnover starts. The sums of organic matter decay, zooplankton excretion, mussel excretion and sediment release were similar to the total algal uptake in all three basins, indicating fast recycling of phosphorus in the water column (Tables 2-3).

In the western basin, the daily external P loads were three times higher than P sedimentation in 1997, but slightly smaller than P sedimentation in 1998. Phosphorus pathways within the water column (e.g., SRP→PP and PP→SRP) and P exchange between western basin and central basin were greater in 1998 than those in 1997, which suggested that when nutrient loads were lower, phosphorus recycled faster within the water column. The total P input to water column (sum of external P, sediment release and mussel excretion) were higher than P loss to sediment in both years. Western basin sediment release indicated that anoxic conditions occurred during the simulation period (Loewen et al. 2007). External loads to the central and eastern basins were small compared to their internal loads, while sediment release was the largest P input to the water column for the central basin and mussel excretion was the largest P input to the water column for the eastern basin. The difference between external loading and withdrawals from the Niagara River and Welland canal showed that the lake retained 77-82% of the total external P loads; some of which will be lost to sedimentation.

Under the four P reduction scenarios, large changes in P pools and fluxes occurred in the western basin, less in in the central basin, and essentially no change in the eastern basin for both years (Tables 4-5). The SRP pool in 1997 was reduced by less than the percent reduction in
external P loads (Table 4), but decreased more in line with the reduction of the external loads in 1998 (Table 5). In the western basin, decreases in the water column fluxes were much larger than those on the bottom, while mussel excretion increased due to improved oxygen conditions (Tables 4-5). However, our simulations were limited by fixed mussel excretion rates, while in reality mussel P excretion should be a function of phosphorus ingested from grazed algae (Vanderploeg et al., in review), which, in turn, should be a function of the external loads.

Consistent between the two years, most phosphorus pools and fluxes decreased proportional to the reduction of the external P loads, except zooplankton biomass decreased by percentages higher than the reductions in the external P loads.

Effects of phosphorus reduction on the algal community in the western basin

Upon simulation of reductions of 20%, 40%, 60%, and 80% of external P loading in the western basin, the total algal biomass decreased more under higher reduction scenarios. The total algal biomass decreased by up to 65% during late June of 1997 and leveled off thereafter, while it decreased gradually by 88% from June to October in 1998 (Figure 7). NDIA decreased more than other algal groups during September under most TP reduction scenarios, except when the TP reduction amount was large (>60%) in 1997 and NDIA biomass was already low (Figure 8).

A clear algal succession pattern occurred in 1997, where diatoms peaked in early June when temperature was still cool and optimal for diatom growth, then NDEA became dominant with favorable high water temperatures, followed by an increase in NDIA in August (Figures 9a,
In 1998, nutrient loads during summer boosted the growth of NDIA (Figures 3, 9b), while NDEA and diatoms decreased due to competition for light (Figure 11b).

Phosphorus loading reduction simulations predict large decreases in the algal groups with the highest biomass. NDEA decreased during its peak time in late June-July (Julian days of 170-220) by as much as 93% in 1997 (in the 80% reduction scenario) and as much as 85% in 1998. Diatoms showed a large decrease during the period when water temperature was optimal for diatom growth and were less affected by the P reduction during hot periods (Figures 9, 10). NDIA biomass was low in 1997 and decreased by 20% until later in August, when NDIA biomass was high and the decrease in biomass became larger (by up to 69%) with phosphorus loading reductions, while NDIA was abundant in 1998 and decreased by up to 97%.

Factors for the dynamics of algal biomass

For the representative model cell close to the Maumee River in the western basin (segment 5 and at 1m depth), we output detailed calculation results relevant to algal dynamics, including limiting factors of water temperature, light, nitrogen, phosphorus, silicon for diatoms, algal daily respiration, excretion, senescence mortality, sedimentation and gross growth rate (d⁻¹), plus crustacean zooplankton grazing mortality on NDEA and diatoms.

Different algal groups had different water temperature preferences, with diatoms favoring cooler temperature, NDIA favoring warm temperatures and NDEA in between (Figure 10). The temperature influence on algal growth was not affected by the reduction of external P loads. Nitrogen was not a limiting factor for any of the three groups, and silicon was not a limiting factor for diatoms over the simulation periods in this model cell. However, the effects of light and phosphorus on algal groups were changed with different levels of P reduction.
Phosphorus loads can strongly decrease the light conditions for the growth of NDEA and Diatoms, especially when the P loading events were followed with fast growth of NDIA (Figures 11, 12a). In 1998, after an increase of nutrients around Julian day 220, NDIA had a strong growth pulse, and the light conditions for NDEA and Diatoms replaced phosphorus to become a limiting factor. Reduction in P loads led to decreases in NDIA and increases in light conditions for NDEA and Diatoms. This explains why NDEA biomass was higher under P reductions later during the simulation period of 1998 (Figure 9b).

Although algal daily respiration, excretion, and senescence mortality rates were much smaller than the gross growth rate, especially when gross growth rates were high, under lower gross growth rates, net growth rates can be negative (Figure 12a). Although NDEA and Diatoms had similar net growth rates (d⁻¹), their individual realized daily growth (g DW/m³/d) differed from each other and from the curves of the net growth rates (d⁻¹, Figure 12b-c). The realized daily growth rates were defined by the current algal biomass. NDIA biomass was very low at the beginning of simulations, and gained little biomass daily before Julian day 200 even with a big growth rate around Julian day 170 (Figure 12a). However, nutrient loads after Julian day of 220 obviously enhanced the growth of NDIA for both 1997 and 1998, especially in 1998 (Figure 12a). Sedimentation and crustacean grazing mortality on NDEA and diatoms were proportional to the current biomass.

Effects of phosphorus reduction on hypoxia in the central basin

With no TP reduction, the average hypolimnetic DO concentration in the central basin in 1998 was 1.6 mg/L, and the hypoxic area was 6435 km² (Figure 13). Hypoxia in the central
basin improved with decreased TP load, until TP load was decreased below 4000 MT with hypoxia declining rapidly with TP reduction.

Discussion

The cause-effect relationship between external P loads and harmful algal blooms (e.g., *Microcystis* blooms) in Lake Erie has been reevaluated and confirmed recently (Kane et al., 2014; Obenour et al., 2014; Stumpf et al., 2012; Verhamme et al., 2016 this issue). In addition to the increasing soluble phosphorus in external P loads, the ongoing changes in land use, agricultural practices and climate change also favor the re-eutrophication of Lake Erie (Michalak et al., 2013; Scavia et al., 2014). The phosphorus loading target that once improved water quality in the 1970-80s will not be good enough to maintain water quality now and in the future (Scavia et al., 2014). A new external P loading target of a 40% decrease from current levels was announced as part of an adaptive management plan (US EPA, 2016). Our study provides some scientific insight and support for this further P reduction. Using a process-oriented mathematical model to depict the spatial distribution of both external and internal loads, we estimated the sizes of basin-wide P pools and pathways to determine the relative importance of different P sources and availability to algal growth. We also simulated responses of three algal groups to different levels of phosphorus reduction, which advances most of the research on eutrophication of Lake Erie that only addressed single model algal group (IJC 2014).

External phosphorus loading

Our model showed that externally loaded phosphorus accumulated in the water column of western Lake Erie in both years, which was consistent with Schwab et al.’s (2009) study that
used a 3-D hydrodynamics model and simulated the external P distribution over a whole year for 1994. This long retention time may help to explain how phosphorus loaded into Lake Erie in early spring can be a determining factor to predict *Microcystis* blooms later in summer and fall (e.g., Stumpf et al., 2012).

Although our model does not include all tributaries along the lake shore, the major tributaries have been included (Bolsenga and Herdendorf, 1993; Dolan, 1993; Schwab et al., 2009). Furthermore, considering the overwhelming loads from the Maumee River and the Detroit River, our model captures the general character of the external loading to Lake Erie during each summer growing season. Model performance would be improved if P loading data from the Grand River, Ontario, were available (Boegman et al., 2008 a, b), as this river is the major source of external nutrient loading to the eastern basin. Due to field data limitations, our simulation periods were from May to September in 1997, and from June to October in 1998, which missed the high P loadings from the Maumee River during January through April (or May in 1998), when a large fraction of the annual loads entered the lake (Figure 3). The effects of these loads should be captured with our model initial conditions; however, they will not be reduced in the load reduction scenarios.

**Internal phosphorus loading**

The uncertainty in the relationship between external P loads and *Microcystis* blooms (Obenour et al., 2014) calls for more studies on unknown factors that could play an important role in *Microcystis* dynamics, such as internal P loading (Kane et al., 2014). Our results show that P recycling within the upper water column contributes the major portion to algal demands, with organic matter decay as the primary contributor. Phosphorus pools and pathways in the
water column responded faster to the reduction of external P loads than pathways to the sediment.

Burns and Ross (1972) estimated that the phosphate regeneration rate under oxygenated conditions in the central basin of Lake Erie was 22 µmoles P m⁻² d⁻¹ (0.0007 g m⁻² d⁻¹); while the anoxic regeneration rate was 245 µmoles P m⁻² d⁻¹ (0.0076 g m⁻² d⁻¹). Lam et al. (1983) assumed that the release rate was 0.0044 g m⁻² d⁻¹ under anoxic conditions. We took Lam et al.’s approach and ignored the phosphorus release under oxygenated conditions. Thus, phosphorus release by sediments occurred mainly in the central basin, where seasonal hypoxia occurs. Our estimates should be considered conservative with a relative low anoxic release rate, in addition to the ignorance of the small but significant normoxic release in Lake Erie (Matisoff et al. 2016). Sediment release was not available to algae during the stratification period in the central and eastern basins, but will be available after the fall overturn when it may support algal growth. This production will further enhance the net organic matter sedimentation to the bottom, and fuel sediment oxygen demand in the central basin in the next year and delay the reduction of hypoxia in the central basin expected from any external P reduction (IJC 2014). Our study supports Burns et al.’s (2005) finding that the oxygen depletion rate in the hypolimnion of the central basin was correlated with the previous year’s load of total phosphorus. A long-term simulation (e.g., several years) of the ecosystem (including October to May periods) is needed in order to quantitatively estimate this delay in response.

Numerous studies have focused on dreissenid P excretion and its ecological impacts (e.g., Mellina et al., 1995; Arnott and Vanni, 1996; Bierman et al., 2005; Conroy et al., 2005b). Our model showed a zebra mussel population excretion rate lower than Mellina et al.’s (1995) estimates, because the mussel density (220×10³ ind/m²) in Mellina et al.’s study was much lower than the density in our study.
higher than that in our study (3 to $6 \times 10^3$ m$^{-2}$). Our estimates of mussel P excretion were also at the lower range of Arnott and Vanni’s (1996) estimates, but mussel excretion still contributed 19-27% of algal demands in the western basin, with an even higher contribution under lower external loads. Mussel excretion as an internal P source would become more important with increasing mussel populations (Zhang et al., 2011), and a recent survey showed that the dreissenid mussel biomass in western Lake Erie tripled from 1998 to 2011 (Karatayev et al., 2014). Mussel excretion is also temperature-dependent (Johengen et al. 2013), while constant excretion rates in our model were measured at the high end of the bottom temperatures. Thus, we may overestimate the mussel phosphorus excretion in this aspect.

Zooplankton excretion is an important phosphorus source in lakes (Hudson et al., 1999; Vanni, 2002; Conroy et al., 2005b). Zooplankton excreta and organic matter release are readily available to algae and dominate the P supply supporting algae production (Carpenter and Kitchell, 1984; Scavia et al., 1988). Our model suggests crustacean excretion provided up to 22% of the algal P demand, which was consistent with Boegman et al.’s (2008b) estimates that zooplankton excretion supported 26% of algal P uptake in 1994. Vanni (2002) estimated zooplankton excretion supports as much as 58% of the primary producer P demand, higher than our estimates.

Phosphorus release from the sediment was less than the P sedimentation losses, which indicates that extra P is accumulated in the lake. However, this study didn’t simulate sediment resuspension, which may bring a large amount of phosphorus from sediment into the water column in the western basin (Matisoff and Carson, 2014; T. Johengen, University of Michigan, personal communication). Increases in epilimnnion phosphorus in the central basin after fall
overturn due to up-mixing of hypolimnion phosphorus and sediment resuspension were well documented for the fall of 1970, and the resultant high concentration lasted for several months (Burns, 1976; Lam and Jaquet, 1976). Hawley and Eadie (2007) calculated that the top 20-40 mm sediment in the central basin was subject to erosion and deposition on an annual basis. However, large storms (e.g., the November 1940 storm) can resuspend up to 2 m of sediment (Lick et al., 1994). Thus if the settled organic matter is not buried deep enough on the bottom or there are strong storms, sediment resuspension during storm events can fuel major algal production in the water column. With storms becoming much stronger with climate change, this factor will become more and more important, causing a delay in oligotrophication despite any P loading management plan. However, the phosphorus input from sediment resuspension may be significantly less bioavailable (Matisoff and Carson, 2014) and less influential to algal growth (LimnoTech reports XXX).

Effects of phosphorus reduction on the algal community in the western basin

Our simulation results show a decrease in P loads is an effective way to control algae, especially the NDIA such as *Microcystis*, and supports the newly announced reduction target in external P load to Lake Erie. Our study shows the dynamics of different algal groups (NDEA, diatoms and NDIA) over the growing seasons and their interactions. In Lake Erie NDIA was dominated by *Microcystis*, so we parameterized the NDIA group as *Microcystis* (Zhang et al. 2008), and we focus on *Microcystis* in this discussion. Diatoms prefer cooler water temperatures and experience higher sinking rates due to their heavy silica frustules, and are often abundant during spring and fall. Thus diatoms do not compete with *Microcystis* severely. However, NDEA has a similar temperature and phosphorus niche to that of *Microcystis*, and showed
strong competition interactions with *Microcystis* in our model. *Microcystis* outcompeted NDEA in August and September by decreasing light conditions for NDEA growth. *Microcystis* is less affected by light conditions than its competitors given its capacity of maintaining growth under relative lower light intensity (Reynolds and Walsby, 1975) and by buoyancy regulation (Belov and Giles, 1997). A pulse of nutrient input is often associated with high turbidity (riverine input or sediment resuspension) that can significantly limit NDEA growth. This light condition may be accompanied with the fast growth of *Microcystis* boosted by the P loads when temperature is optimal, as was the case in 1998, when NDEA lost in competition with *Microcystis*. The competition for phosphorus between NDEA and *Microcystis* is not bilaterally equal. *Microcystis* can regulate its buoyancy to locate itself in a water layer with high phosphorus content, and store phosphorus intracellularly for later growth after moving into a water layer with good light conditions (Harke et al., 2016). Once *Microcystis* dominates the community, NDEA loses the competition until temperatures become cool or nutrients are depleted and *Microcystis* declines (Wynne et al., 2010). Another disadvantage to NDEA for this competition is that they are food for zooplankton and mussels, while *Microcystis* is not (Aleya et al., 2006). Although the output of the representative model cell did not show that zooplankton grazing mortality causes significant decreases in NDEA, zooplankton does have a high grazing capacity and causes a late June clear-water phase in many lakes (e.g., Wu and Culver, 1991). Zooplankton grazing may thus contribute to a community shift to *Microcystis* dominance. Dreissenid mussels’ selective filtration also promotes *Microcystis* blooms (e.g., Vanderploeg et al. 2012).  

The two simulation years in our model had contrasting P loads and *Microcystis* bloom sizes, with high P loads in 1997 but no *Microcystis* bloom and lower P loads in 1998 accompanied by a moderate *Microcystis* bloom, thus the total amount of external P load is not
the sole determining factor of Microcystis blooms. The timing of external loads can be important, e.g., high loading during July and August of 1998 boosted the growth of Microcystis. The output from the representative model cell shows that several consecutive P input pulses occurred immediately before the increase in Microcystis biomass, which highlights the importance of timing of nutrient inputs (IJC, 2014). Similarly, Michalak et al. (2013) hypothesized that a strong resuspension event immediately preceding bloom onset was one of the ideal conditions for bloom development. Although they assumed this event will bring more over-wintering Microcystis cells from the sediment to the water column, it was also associated with a big P input from the sediment into the water column. Unfortunately, without a clear definition of ‘strong,’ they rejected this hypothesis later on. Year 1997 could be an outlier in statistical models for prediction of Microcystis bloom using external P loads (e.g., Stumpf et al., 2012). However, it does provide us a unique chance to use a different approach to study Microcystis blooms and identifies some critical factors that can be incorporated into statistical models and improve their predictions and reduce uncertainty.

Effects of phosphorus reduction on hypoxia in the central basin

Hypoxia has become much stronger (expressed as areal extent and duration) in recent years (Scavia et al., 2014). Our modeling results showed that hypoxia’s response to external P loading was similar to that reported by the Annex 4 Resemble Modeling Group (Scavia and DePinto, 2015; Scavia et al., this issue). However, the simulated hypoxic area tended to be larger in 1998 under TP load similar to those in their focus year (2008), which is likely a result of differences in meteorology and resulting thermal structure (Rucinski et al., 2014).

Conclusion
We used a 2-D water quality model to study the effects of external and internal P loads on algal dynamics. Our results support the use of a phosphorus reduction program to improve water quality in Lake Erie. Our study indicates that reduction in external phosphorus would result in fast and large decreases in algal biomass and Microcystis blooms in the western basin. However, several phosphorus input pulses during Microcystis developing time can dramatically increase Microcystis and consequently decrease the light condition for NDEA and limit NDEA growth, promoting a rapid community shift to Microcystis dominance. Central basin hypoxia would likely take a longer time to respond the external phosphorus reduction. Additional factors, such as the spatial distribution of external P loads, the contribution of internal phosphorus loads and sediment resuspension events, may delay the occurrence of improved water quality, in part because sediment resuspension and transportation are less responsive to changes in external phosphorus. More studies are needed of long-term cycling of phosphorus in the lake, including the processes in sediment-water interactions.

Acknowledgments

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Nitrogen and phosphorus excretion by detritivorous gizzard shad in a reservoir


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Table 1. TP and SRP loads (metric tons per season) from tributaries and Waste Water Treatment Plants (WWTP) in 1997 and 1998. Although the seasonal data are listed here, daily/monthly loads were collected and linearly interpolated into every-30- min inputs in the model. *data from D. Dolan, University of Wisconsin-Green Bay, Green Bay, WI, USA, personal communication.

<table>
<thead>
<tr>
<th></th>
<th>1997</th>
<th>1998</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(May 10-Sep.30)</td>
<td>(Jun.10 – Oct. 30)</td>
</tr>
<tr>
<td></td>
<td>TP</td>
<td>SRP</td>
</tr>
<tr>
<td>Maumee River</td>
<td>1221</td>
<td>203</td>
</tr>
<tr>
<td>Toledo WWTP</td>
<td>29</td>
<td>3</td>
</tr>
<tr>
<td>Detroit River</td>
<td>1166</td>
<td>207</td>
</tr>
<tr>
<td>Sandusky River</td>
<td>275</td>
<td>31</td>
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<tr>
<td>Total</td>
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<td>464</td>
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Table 2. Basin-wide P cycling in 1997, shown as concentrations in major pools, and rates of transfer between pools. DP: P in dissolved organic matter; PP: P in phytoplankton and detritus; CP: P in crustaceans; DP→SRP: phosphorus release due to decomposition of dissolved organic matter; PP→SRP: phosphorus excretion by phytoplankton and release due to decomposition of detritus; CP→SRP: phosphorus excretion by crustaceans; MP→SRP: phosphorus excretion by dreissenids; SED→SRP: phosphorus release by sediments under anoxic conditions; PP→SED: phosphorus loss to sediment due to sedimentation of PP. External loading: P loading from tributaries of Lake Erie; Withdrawals: total phosphorus (SRP+DP+PP) loss through the Welland Canal and the Niagara River; Exchange between basins: net total phosphorus transported by horizontal currents between basins. All values are an average over the simulation period from May 10 to September 30, 1997.

<table>
<thead>
<tr>
<th></th>
<th>WB</th>
<th>CB</th>
<th>EB</th>
<th>Total</th>
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<td>511.0</td>
<td>1250.9</td>
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<td>841.2</td>
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<tr>
<td>PP</td>
<td>176.1</td>
<td>1533.5</td>
<td>433.3</td>
<td>2142.9</td>
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<tr>
<td>CP</td>
<td>14.6</td>
<td>143.8</td>
<td>38.7</td>
<td>197.1</td>
</tr>
<tr>
<td><strong>P pathways</strong> mt P/d</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>DP→SRP</td>
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<td>33.3</td>
<td>10.2</td>
<td>52.5</td>
</tr>
<tr>
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<td>52.1</td>
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<td>71.9</td>
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<tr>
<td>CP→SRP</td>
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<td>22.3</td>
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<td>31.1</td>
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<td>MP→SRP</td>
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<td>6.6</td>
<td>6.3</td>
<td>19.5</td>
</tr>
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<td>10.2</td>
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<td>23.2</td>
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<td>13.8</td>
<td>3.7</td>
<td>22.7</td>
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<tr>
<td><strong>External loading</strong> mt P/d</td>
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<td>3.1</td>
<td>0.2</td>
<td>18.9</td>
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<tr>
<td><strong>Exchange between basins</strong> mt P/d</td>
<td>-5.9</td>
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<td>3.7</td>
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<tr>
<td><strong>Withdrawals</strong> mt P/d</td>
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Table 3. As Table 2, but for June 10 through October 30, 1998.

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<th>EB</th>
<th>Total</th>
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<td>1998</td>
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<tr>
<td>P pools</td>
<td>mt P</td>
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<tr>
<td>SRP</td>
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<td>1061.5</td>
<td>826.1</td>
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<td>1492.5</td>
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<td>19.6</td>
<td>175.4</td>
</tr>
<tr>
<td>P pathways</td>
<td>mt P/d</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>DP→SRP</td>
<td>8.8</td>
<td>29.5</td>
<td>9.8</td>
<td>48.1</td>
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<tr>
<td>PP→SRP</td>
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<td>51.2</td>
<td>15.4</td>
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<td>21.2</td>
<td>2.0</td>
<td>28.6</td>
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<tr>
<td>MP→SRP</td>
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<td>6.8</td>
<td>6.3</td>
<td>19.4</td>
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<td>Withdrawals</td>
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<td>-2.8</td>
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Table 4. Percent changes in the phosphorus pools and pathways under different reduction levels in the external phosphorus loads at the end of September of 1997.

<table>
<thead>
<tr>
<th></th>
<th>1997</th>
<th>WB</th>
<th></th>
<th></th>
<th>CB</th>
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<tr>
<td></td>
<td></td>
<td>20%</td>
<td>40%</td>
<td>60%</td>
<td>80%</td>
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<td>40%</td>
<td>60%</td>
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<td></td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>SRP</td>
<td>-7</td>
<td>-12.1</td>
<td>-18</td>
<td>-29</td>
<td>0</td>
<td>-0.6</td>
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<td>-11.2</td>
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<tr>
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<td>-18.9</td>
<td>-33.1</td>
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<tr>
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<td>-58.6</td>
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<td>-11.7</td>
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<td>-1.3</td>
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<td>-9.9</td>
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Table 5. Percent changes in the phosphorus pools and pathways under different reduction levels in the external phosphorus loads at the end of September of 1998.

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P pathways

|       |       |       |       |       |       |       |       |       |
|-------|-------|-------|-------|-------|-------|-------|-------|
|       | DP→SRP |       |       |       |       |       |       |       |
|       | PP→SRP |       |       |       |       |       |       |       |
|       | CP→SRP |       |       |       |       |       |       |       |
|       | MP→SRP |       |       |       |       |       |       |       |
|       | SED→SRP|       |       |       |       |       |       |       |
|       | SRP→PP |       |       |       |       |       |       |       |
|       | PP→SED |       |       |       |       |       |       |       |

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Figure 1. Locations of tributaries, depth contours of 5, 22, 30 and 50m, and separations of the Lake Erie western (WB), central (CB) and eastern (EB) basins.

Figure 2. Basin-wide P pools and cycling pathways incorporated in EcoLE. SRP: soluble reactive phosphorus; DP: P in dissolved organic matter; PP: P in phytoplankton and detritus; CP: crustacean phosphorus content; PP→SRP: phosphorus excretion by phytoplankton and release due to decomposition of detritus; CP→SRP: phosphorus excretion by crustaceans; MP→SRP: phosphorus excretion by dreissenids; SED→SRP: phosphorus release by sediments under anoxic conditions; PP→SED: phosphorus loss to sediment due to sedimentation of PP.

Note that these pools and pathways were simulated in each model cell, but are aggregated here to represent basin-wide estimates. External loading: TP loading from tributaries of Lake Erie; Withdrawals: total phosphorus (DP+PP) loss through the Welland Canal and the Niagara River; Exchange between basins: net total phosphorus transported by horizontal currents between basins.

Figure 3. Total daily phosphorus loads into Lake Erie from the Maumee, Detroit, Sandusky and the Cuyahoga rivers. Data are from the National Center for Water Quality Research, Heidelberg University, Tiffin, Ohio. Note that most of the annual loading occurs by early June in 1997, but 1998 had two later high loading events, one in June and the other in July - August.

Figure 4. Comparison of the results of physical transport and mixing processes in the EcoLE model on the spatial distribution of TP entering the lake by external
loading during the two years in the study, without involvement of chemical and
biological processes. The isopleths show the distribution of TP externally loaded
by the end of the two simulation periods (30 September 1997 and 30 October
1998). The x axis represents the 220 2-km long west-to-east spatial segments of
the model, whereas the y-axis represents depth in the lake (and the 1-m thick
layers of the model). Each segment and layer combination constitutes one cell in
this two-dimensional model.

Figure 5. Comparison of the spatial distribution of phosphorus from each of the internal
sources and the water column cycling processes over the simulation period of
1997, as results of physical transport and mixing processes. The isopleths show
the distribution of internally loaded phosphorus on September 30, 1997.

Figure 6. Comparison of the spatial distribution of phosphorus from each of the internal
sources and the water column cycling processes over the simulation period of
1998, as results of physical transport and mixing processes. The isopleths show
the distribution of internally loaded phosphorus on October 30, 1998.

Figure 7. Percent change of total algal biomass over the simulation periods for 1997 and
1998 under different levels of external phosphorus loading reduction (decrease by
20, 40, 60 and 80%) from model runs with no reduction. The dotted lines indicate
a 50% decrease in algal biomass.

Figure 8. September-averaged biomass for total algae, and three algal groups for 1997
and 1998 under different levels of external phosphorus loads to the western basin.
Biomass was scaled to the biomass with no phosphorus reduction. The horizontal
line represents a 50% reduction relative to model runs with no reduction.
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Figure 10. Limiting factor of water temperature (dimensionless) on the growth of three algal groups in a typical model cell close to the Maumee River in the western basin (segment 5 and 1m below the water surface) for a) 1997 and b) 1998.

Figure 11. Limiting factors (dimensionless) calculated based on light and phosphorus for NDEA and diatoms in the typical model cell for a) 1997 and b) 1998 under different phosphorus reduction scenarios (no reduction, reduction of 20%, 40%, 60% and 80%). NDEA and diatoms were affected by light and phosphorus in the same way.

Figure 12. Daily net growth rate (d⁻¹) and net growth (g DW /m³/d) over the simulation period of 1998 for a) non-diatom inedible algae NDIA, b) non-diatom edible algae NDEA, and c) diatoms under different phosphorus reduction scenarios (no reduction, reduction of 20%, 40%, 60%, and 80%). Note the different scales on the y-axes.

Figure 13. Relationships between total TP load to the western and central basins and hypolimnetic DO (black line) and hypoxic area (gray line) for model year 1998. The reference line indicates the hypolimnetic DO concentration of 4 mg/L, and a threshold hypoxic area of 2000 km².
Figure 1.
Figure 2.
Figure 3.

![Graph showing TP loads over Julian days for 1998 and 1997.](Image)
Figure 4.
Figure 5.

a) Anoxic sediment release

b) Dreissenid mussel excretion

c) Crustacean excretion

d) Organic matter decay
Figure 6.

1099 a) Anoxic sediment release

1101 b) Dreissenid mussel excretion

1103 c) Crustacean excretion

1105 d) Organic matter decay
Figure 7.
Figure 8.

a) 1997

![Graph showing WB annual TP load (MT) vs. Scaled September averaged biomass (%).]  
- Total
- Diatoms
- NDEA
- NDIA

b) 1998

![Graph showing WB annual TP load (MT) vs. Scaled September averaged biomass (%).]  
- Total
- Diatoms
- NDEA
- NDIA
Figure 9.

a) 1997

Non-Diatom Edible Algae (NDEA)

Diatoms

Non-Diatom Inedible Algae (NDIA)

Basin-wide biomass (mt DW)

Julian days
Non-Diatom Edible Algae (NDEA)

Diatoms

Non-Diatom Inedible Algae (NDIA)

Basin-wide biomass (mt DW)

Julian days
Figure 10.

- Limiting factor: Temperature
- Julian days: 140 160 180 200 220 240 260 280 300
- 1997 and 1998 graphs
- NDEA, Diatoms, NDIA lines
Figure 11.

(a) 1997

(b) 1998

Limiting factor fractions

Julian days

Light

Phosphorus

No reduction

20%

40%

60%

80%
Figure 12.

a) NDIA

![Graph of net growth rate (d^-1) and daily net growth (g DW/m^3/d) vs. Julian days.](image)
b) NDEA

[Graph showing net growth rate (d⁻¹) and daily net growth (g DW/m³/d) over Julian days.]
c) Diatoms

![Graph showing net growth rate and daily net growth over Julian days]

- Net growth rate (d\(^{-1}\))
- Daily net growth (g DW/m\(^3\)/d)
- Lines representing different reduction percentages: No reduction, 20%, 40%, 60%, and 80%
Figure 13.

WB + CB annual TP load (MT)

Hypolimnetic DO (mg/L)

Hypoxic area (km²)

WB + CB annual TP load (MT)
A. Density of dreissenid mussels in the hypoxia zone of central basin

Zhang et al. (2008) used the depth-dependent dreissenid density estimations by Jarvis et al. (2000). However, due to the seasonal hypoxia, very low mussel densities were recorded in the deep central basin. In this study, we estimated mussel density at the bottom of the central basin according to studies by Jarvis et al. (2000), Patterson et al. (2005), and field data from IFYLE (2005). Jarvis et al. (2000) reported 135 ind m$^{-2}$ at a deep-water station (24.2 m). Patterson et al. (2005) showed that most of the deep-water stations had densities within range of 0-500 ind m$^{-2}$. IFYLE (2005) recorded a range of 0-260 ind m$^{-2}$. Since Jarvis et al.’s study included data from our simulation years (1997-1999) and their density values were well within the ranges of other two studies, we used Jarvis et al.’s density to populate deep water areas (>22 m) in the central basin. The remaining model settings for mussel simulations were kept as in Zhang et al. (2008). We compared the simulation results of before and after these density changes, and found that they decreased the modeled mussel excretion (about 50%) in the central basin. Sediment phosphorus release also decreased by 9-15% because the smaller dreissenid mussel population had lower oxygen consumption and led to improved dissolved oxygen conditions on the bottom, while sediment phosphorus release only happened under anoxic conditions. Consequently, the SRP pool had decreased by 16-18%. Other state variables and process rates of phosphorus dynamics were largely unaffected (<2%, but most <0.5%) compared to those before the density change due to seasonal thermal stratification (Table S1).
Table S1. The percent changes in the P pools and P pathways of simulation with new dreissenid mussel density (Standard), compared to simulations with high dreissenid mussel density in the central basin with depths >22m. Values of ‘0’ indicated the changes were less than 0.5%. * indicate very low values of sediment P release from the eastern basin and the changes would be misleading.

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P pathways

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B. Calculations of phosphorus pools and pathways.

External Phosphorus Loads

The total external phosphorus loads ($L$, g P) over the simulation periods were estimated by:

$$ L = \sum_i \sum_j Q_{ij} C_{ij} (t_{j+1} - t_j) $$

where, $Q_{ij}$ was the flow rate of the source $i$ at time $j$ (m$^3$ s$^{-1}$). $C_{ij}$ was the total phosphorus concentration (SRP and organic matter phosphorus) of the source $i$ at the time $j$ (g P m$^{-3}$) or the monthly mean phosphorus concentration of source $i$ when the instantaneous concentrations were not available.

Crustacean zooplankton excretion

The total crustacean P excretion ($f_{CP}$) during the growing season (g P m$^{-3}$) was calculated as:

$$ f_{CP} = \sum x (\delta_{p-clad} r_{clad} m_{cladoceran} + \sum_{i} (\delta_{p-cop} r_{cop} m_{i})) $$

where, $\delta_{p-cop}$ was the ratio of phosphorus to dry weight for copepods, $\delta_{p-clad}$ was the ratio of phosphorus to dry weight for cladocerans. $r$ was the respiration rate (s$^{-1}$), $T$ was the duration of a growing season, and $m$ was biomass (g m$^{-3}$).

Dreissenid mussel excretion

Zebra mussels and quagga mussels have different weight-specific phosphorus excretion rates, $ZMP$ and $QMP$ ($\mu$g P mg$^{-1}$ DW d$^{-1}$) (Conroy et al. 2005).
\[ \log_{10}(ZMP) = 0.506[\log_{10}(W_{zm})] - 1.172 \]
\[ \log_{10}(QMP) = 0.297[\log_{10}(W_{qm})] - 1.195 \]

Where, \( W_{zm} \) and \( W_{qm} \) are the dry weights (mg) of individual 10-mm mussels.

Thus, the phosphorus excretion of a mussel population (\( f_{zmP} \) for zebra mussels; \( f_{qmP} \) for quagga mussels) in a model cell over the growing season (g P m\(^{-3}\)) was calculated as the sum of the products of individual excretion rates and numbers of mussels of the two species.

\[ f_{zmP} = \sum (N_{zm}(W_{zm}ZMP)) / V \]
\[ f_{qmP} = \sum (N_{qm}(W_{qm}QMP)) / V \]

Where, \( V \) is the volume of the corresponding model cell, m\(^3\).

Basin-wide phosphorus budget

The phosphorus content of four phosphorus pools: soluble reactive phosphorus (SRP), dissolved phosphorus (DP), particulate phosphorus (PP) and crustacean phosphorus (CP), and of the four internal phosphorus sources were estimated basin wide and over the growing season (Figure 2). DP referred to the phosphorus in dissolved organic matter; PP referred to the phosphorus in particulate organic matter and algae; whereas CP included the phosphorus content of cladocerans and copepods. The pathway from SRP to PP included phosphorus uptake by algae, while PP to SRP included phosphorus excreted by algae and released by POM decay. Our model had a traditional
crustacean grazing food web. However, by including organic matter, the model
implicitly also includes the microbial food web (Debruyn et al. 2004).

The average mass of each phosphorus pool during the simulation period (g P,
converted to metric tons P later) was calculated by
\[ \frac{\sum_i \sum_j C_{ij}\ V_{ij}}{N}, \]
where \( C_{ij} \) was the concentration of phosphorus content of a constituent mentioned above
in the model cell \((i,j)\) and at time \(t\), in g P m\(^{-3}\). \( V_{ij} \) was the water volume of the
corresponding model cell \((i,j)\), m\(^3\), at time \(t\). \( N \) was the total number of simulation time
steps (about 30 mins) during a simulation period.

Daily basin-wide phosphorus fluxes (\( F \), g P d\(^{-1}\), converted to mt P d\(^{-1}\) later)
between state variables and mussels and sediment were calculated by:
\[ \frac{\sum_i \sum_j \sum_i q_{ij}\ V_{ij}}{N}, \]
where, \( q_{ij} \) was the phosphorus transfer rate from one constituent to another within a
model cell \((i,j)\) at time \(t\), g P m\(^{-3}\) s\(^{-1}\). For example, \( q_{ij} \) of the phosphorus from CP to SRP
is the total phosphorus excreted by crustacean zooplankton per m\(^3\) in the model cell \((i,j)\)
at time \(t\).


IFYLE (International Field Years on Lake Erie), 2005. database http://www.glerl.noaa.gov/ifyle/

