

Mapping the spatial distribution of the biomass and filter-feeding effect of invasive dreissenid mussels on the winter-spring phytoplankton bloom in Lake Michigan

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Summary

1. The effects of the invasive bivalves *Dreissena polymorpha* (zebra mussel) and *Dreissena rostriformis bugensis* (quagga mussel) on aquatic ecosystems, including Lake Michigan, are a topic of current interest to scientists and resource managers. We hypothesized that the winter-spring phytoplankton bloom in Lake Michigan is reduced at locations where the fraction of the water column cleared per day by *Dreissena* filter

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feeding approached the net growth rate of phytoplankton, when the water column was not stratified. To test this hypothesis, we compared the spatial distribution of *Dreissena* filter feeding intensity (determined from geostatistical modelling) to the spatial distribution of chlorophyll (determined from satellite remote sensing).

2. To map the spatial distribution of *Dreissena* biomass and filter-feeding intensity, we developed a geostatistical model based on point observations of mussel biomass measured in Lake Michigan in 1994/95, 2000, 2005 and 2010. The model provided fine-scale estimates of the spatial distribution of biomass for the survey years and provided estimates, with their uncertainty, of total biomass lake-wide and within sub-regions. The approach outlined could be applied more generally to map the distribution of benthic biota in lakes from point observations.

3. Total biomass of *Dreissena* in Lake Michigan, estimated from the geostatistical model, increased significantly over each five-year period. The total biomass in units of 10^6 kg ash-free dry mass (AFDM) (with 90% confidence interval) was 6 (4-8) in 1994/95, 18 (14-23) in 2000, 408 (338-485) in 2005, and 610 (547-680) in 2010. From 1994/95 to 2005, increases were observed in all regions of the lake (northern, central and southern) and in all depth zones (< 30, 30-50, 50-90, and > 90). However, from 2005 to 2010, for depths of < 50 m, biomass declined in the northern region, remained constant in the central region, and increased in the southern region; biomass continued to increase in all three lake regions for depths > 50 m.

4. The filter feeding intensity of *Dreissena* exceeded the benchmark spring phytoplankton growth rate of 0.06 d^{-1} in 2005 for depths < 50 m (lake-wide). In 2010, the filter-feeding impact exceeded 0.06 d^{-1} within depths < 90 m (lake-wide), which greatly increased the spatial area affected relative to 2005. A regression analysis indicated a significant relationship between the reduction in satellite-derived chlorophyll concentration (pre *D. r. bugensis* period to post *D. r. bugensis* period) and spatially co-located filter-feeding intensity (fraction of water column cleared per day) during periods when the water column was not stratified (December to April).

Introduction

The dreissenid mussels, *Dreissena polymorpha* (zebra mussel) and *Dreissena rostriformis bugensis* (quagga mussel), are prolific invaders of aquatic ecosystems. *Dreissena* originated in the Ponto Caspian region, and spread to many aquatic systems in Europe in the 19th and 20th centuries (Vanderploeg *et al.*, 2002; Karatayev, Burlakova & Padilla, 2014). The larval mussels were carried to North America in ship ballast water (Brown & Stepien, 2010) and adults were first reported in the Laurentian Great Lakes in 1988, subsequently spreading through the Mississippi River system and to lakes and reservoirs in western North America (Benson, 2014). The spread of *Dreissena* is facilitated by their pelagic larval stage and the ability of adults to attach to hard substrata, including boats and ships, traits that are lacking in most freshwater bivalves native to North America (Vanderploeg *et al.*, 2002; Karatayev, Burlakova & Padilla, 2014).

Dreissena polymorpha and *D. r. bugensis* share many similarities; however, *D. r. bugensis* is much better adapted to deep water. Karatayev *et al.* (2011) reviewed the course of invasion of *D. polymorpha* and *D. r. bugensis* across water bodies in Europe and North America and found that the two species often coexist in shallow systems although, in deep lakes, *D. r. bugensis* typically displaces *D. polymorpha* to become the dominant species within 10 y after it invades. Most of Lake Michigan is sufficiently deep (75% of the lake area is ≥ 30 m) to provide an advantage to *D. r. bugensis* over *D. polymorpha*. Consistent with the pattern observed in other systems, *D. r. bugensis* has displaced *D. polymorpha* in Lake Michigan: *D. polymorpha* was alone in 1994/95, *D. r. bugensis* began to invade northern Lake Michigan in 2000 (first found in 1997; Nalepa *et al.*, 2001), and total *Dreissena* biomass consisted almost exclusively of *D. r. bugensis* in 2005 (Nalepa, Fanslow & Pothoven, 2010) and 2010 (Fig. 1: Nalepa *et al.*, 2014).

As an ecosystem engineer, *Dreissena* modifies aquatic systems in several ways, having substantial ecological (Karatayev, Burlakova & Padilla, 2002; Vanderploeg *et al.*, 2002) and economic effects (Connelly *et al.*, 2007). *Dreissena* alters benthic substrata by creating aggregates of living mussels ('druses') and deposits of shells. *Dreissena* may create additional food and habitat for invertebrates in the littoral zone, but quagga mussels in deep water may outcompete other invertebrates, decreasing their diversity and

density (Vanderploeg *et al.*, 2002; Karatayev, Burlakova & Padilla, 2014). The system-wide effect of *Dreissena* in most water bodies is to reduce phytoplankton abundance (chlorophyll concentration) through the direct effect of filter feeding (Karatayev *et al.*, 2014; and works cited therein), although there are some mechanisms through which *Dreissena* can increase primary production under certain circumstances. For instance, *Dreissena* increases water clarity through filter feeding and regenerates dissolved nutrients, which can stimulate growth of attached algae (e.g. *Cladophora*) and macrophytes to nuisance values (Vanderploeg *et al.*, 2002; Hecky *et al.*, 2004; Auer *et al.*, 2010). *Dreissena* may also promote the dominance of cyanobacterial blooms through selective feeding and alteration of the phosphorus cycle, (Vanderploeg *et al.*, 2001; Obenour *et al.*, 2014; Steffen *et al.*, 2014), although such blooms are not presently an issue in the oligotrophic main body of Lake Michigan. In addition, *Dreissena* alters the abundance and distribution of fish and wildlife through effects on benthic habitat and prey species distribution (e.g., Schummer, Petrie & Bailey, 2008). Because of the diverse effects of *Dreissena* on invaded systems, detailed distribution maps and estimates of their lakewide biomass are useful to ecologists and resource managers in measuring and understanding the changes associated with invasion.

To map spatial distribution, or to estimate an aggregated total over given spatial areas, it is necessary to predict abundance for locations where observations are not made. Surveys of the benthos, including dreissenids, usually consist of a limited array of point observations, with the number of points being limited by the considerable cost of data collection and sample processing. Nalepa *et al.* (2009) reported *Dreissena* numerical density and biomass in Lake Michigan, derived from Ponar grab sample surveys conducted in 1994/95, 2000, and 2005; they produced maps of the spatial distribution of *Dreissena* numerical density using natural neighbour spatial interpolation (Sibson, 1981), and acknowledged the tendency of this method to overestimate *Dreissena* density in deep water, where mussel densities were expected to be low. Further, Nalepa *et al.* (2009) estimated lakewide biomass by taking the area-weighted mean of point observations within four broadly-defined, bathymetric depth intervals (≤ 30 , 30–50, 50–90 and > 90 m), but did not account for spatial trends within the depth intervals. Bathymetric depth is an important predictor for the distribution of benthic biota. For example, Auer *et al.*

(2013) reviewed 19 studies that found maxima in the abundance of benthic animals (primarily *Diporeia* spp.) within the Great Lakes at depths between ~30-50 m. Auer *et al.* (2013) collected Ponar grab samples of *Diporeia* in Lake Superior in cross-isobath transects, and identified a depth range (or sediment grain size range) of high abundance (> 95th percentile of profundal abundance); lakewide spatial distribution of abundance was then estimated based upon bathymetry and grain size maps. The geostatistical modelling approach described here offers advantages over previous methods of spatial prediction or interpolation applied to benthos in the Great Lakes: 1) covariates such as bathymetry are readily incorporated, and 2) predictive uncertainty is readily quantified. Here we test the hypothesis that the winter-spring phytoplankton bloom was reduced in locations where *Dreissena* filter feeding intensity approached or exceeded the net growth rate of phytoplankton. Prior to establishment of a large population of *D. r. bugensis* in Lake Michigan, there was a winter-spring phytoplankton bloom, with chlorophyll concentration gradually increasing from February until the onset of stratification in late April-May (Fig. 2; data source: Yousef *et al.*, 2014). The disappearance of the winter-spring phytoplankton bloom in the southern basin of Lake Michigan after establishment of *D. r. bugensis* has been documented by several investigators (Fahnenstiel *et al.*, 2010b; Kerfoot *et al.*, 2010; Vanderploeg *et al.*, 2010; Yousef *et al.*, 2014). The winter-spring bloom is comprised of energy-rich diatoms, and the loss of this food source to pelagic and benthic invertebrates has consequences throughout the food web, including for fish (Madenjian *et al.*, 2006; Nalepa, Fanslow & Lang, 2009; Pothoven, Fahnenstiel & Vanderploeg, 2010). Several lines of evidence implicate filter feeding by *D. r. bugensis* in the disappearance of the winter-spring bloom: 1) loss of the winter-spring bloom was coincident with establishment of a large population of *D. r. bugensis*, 2) the reduction in chlorophyll-a concentration and primary production occurred mainly in the isothermal period when benthic filter feeding can draw down phytoplankton abundance throughout the entire water column (Fahnenstiel *et al.*, 2010b; Kerfoot *et al.*, 2010), and 3) measured clearance rates of *D. r. bugensis* could exceed the spring phytoplankton growth rate, assuming a well-mixed water column (Vanderploeg *et al.*, 2010). Alternate explanations for the loss of the winter-spring bloom, including reduced phosphorus loads, increased zooplankton grazing and climate change were found to be unsatisfactory (Fahnenstiel *et*

al., 2010b). Our study provides a new approach for assessing the relationship between dreissenid filter feeding and the loss of the winter-spring bloom by considering the spatial association between the reduction in chlorophyll (from the pre *D. r. bugensis* period to the post *D. r. bugensis* period) and the filter-feeding intensity of *D. r. bugensis*.

Methods

Physical characteristics of Lake Michigan

Lake Michigan is one of the five Laurentian Great Lakes of North America (Fig. 3). It is an oligotrophic lake with a surface area of 57,800 km², a catchment of 118,000 km², a volume of 4,947 km³ and a maximum depth of 281 m (Coordinating Committee on Great Lakes Basic Hydraulic and Hydrologic Data, 1977). It is dimictic, with periods of complete vertical mixing in spring and autumn with summer stratification and variable stratification and ice cover during winter (Wang *et al.*, 2012). Dates of stratification vary from year to year and, in general, the surface mixed layer (SML) varies seasonally and temporally from a few metres deep at the onset of summer stratification (May-June) to 10-20 m in July-August (Beletsky, Schwab & McCormick, 2006). In the autumn, the SML increases due to higher winds and convection (surface cooling), leading into the isothermal period (December-April) when the water column is vertically well-mixed except for intermittent winter stratification when the surface is colder than 4 °C (Beletsky & Schwab, 2001). Surface water temperature in the summer reaches ~20 °C. At depths greater than ~30 m, near-bottom water temperature is ~4 °C for most of the year, except in late summer and autumn when the deepening of the SML may bring warmer water (typically < 10 °C) to the bottom.

Observed dreissenid mussel biomass

Numerical densities of *D. polymorpha* and *D. r. bugensis* were obtained from samples collected with a Ponar grab (sampling area = 0.046 m²) in July-August of 1994/95, 2000, 2005, and 2010 (Nalepa *et al.*, 2008; Nalepa *et al.*, 2014). The number of stations sampled was 90, 157, 160 and 144 in each of the four periods, respectively. Station locations were largely the same in 2000, 2005 and 2010, but differed in 1994/95.

Numerical density was converted to *Dreissena* biomass using length-mass relationships and length frequency distributions (Nalepa *et al.*, 2009; Nalepa *et al.*, 2014). Samples were collected in triplicate at each station and were averaged by station and year for development of the geostatistical model. We report *Dreissena* biomass in units of grams ash-free dry tissue mass (g AFDM). Biomass may be converted to other units through approximate empirical relationships obtained from *D. polymorpha* specimens: 1 g dry tissue mass = 0.88 g AFDM, 1 g dry tissue mass = 0.46 g carbon (Nalepa *et al.*, 1993). We chose to conduct our analysis on total *Dreissena* biomass (sum of *D. polymorpha* and *D. r. bugensis*) because clearance rates of the two species, normalized to biomass, are not significantly different (Vanderploeg *et al.*, 2010), and *D. r. bugensis* dominated total biomass in years for which there were noticeable impacts.

Geostatistical model

We applied a geostatistical modelling framework (Diggle & Ribeiro Jr., 2007; Chiles & Delfiner, 2009) that has been previously used with fisheries benthic trawl surveys (Jardim & Ribeiro Jr, 2007; Jardim & Ribeiro Jr, 2008) and bottom-water hypoxia surveys (Obenour *et al.*, 2013; Zhou *et al.*, 2013). To avoid potential confusion, we emphasize that the geostatistical model predicts dreissenid biomass across space, but it does *not* make predictions across time. The geostatistical model was implemented using the R package (R Core Team, 2012) “geoR” (Ribeiro Jr. & Diggle, 2001).

The following is a brief explanation of the geostatistical modelling framework, which is described in detail elsewhere (Ribeiro Jr. & Diggle, 2001; Diggle & Ribeiro Jr., 2007). Data consist of individual response observations z_i associated with locations x_i , which are a subset of the observation region. In the geostatistical model, the response $Z(x)$ is related to an unobserved stochastic process $S(x)$ called a ‘signal’, which fluctuates around an underlying deterministic trend $\mathbf{F}(x)\beta$. The signal, $S(x)$, is a stationary Gaussian process characterized by a spatial variogram (covariance function) that is typically fitted through variogram analysis or likelihood estimation (Ribeiro Jr. & Diggle, 2001). Here, a spherical covariance function (Ribeiro Jr. & Diggle, 2001), fitted using restricted maximum likelihood (Zimmerman, 2010), was found to produce robust model parameter estimates. Covariance parameters include τ^2 , σ^2 , and ϕ , which are commonly referred to

as the nugget, partial sill and range, respectively. Coincident observations have a covariance of $\sigma^2 + \tau^2$; observations at very small (but non-zero) separation distances have a covariance of σ^2 ; and, as the separation distance increases to the range (ρ), the covariance between observations decreases from σ^2 to zero (and conversely, the variance between observations increases from τ^2 to $\sigma^2 + \tau^2$). For this study, a Box-Cox transformation ($\lambda=0.3$) of the response was found to substantially improve the Gaussian properties of the signal (Box & Cox, 1964; Ribeiro Jr., Christensen & Diggle, 2003). Because the Box-Cox transformation cannot be applied to zero values, an offset of +0.01 g AFDM m⁻² was first applied to all observations, and it was subsequently removed from all model predictions after back-transforming to the original scale.

The deterministic trend component (or ‘drift’), $\mathbf{F}(x)\beta$, characterizes the underlying, large-scale spatial and bathymetric trends in $Z(x)$. Here, $\mathbf{F}(x)$ is a design matrix with covariate elements of the type $f_j(x_i)$, a measurement of the j^{th} covariate at the i^{th} location. Regression parameters, β_j , are estimated through generalized least squares, taking into account the covariance structure of the response. When implemented with covariates, in this way, the geostatistical model is often referred to as a “Universal Kriging” model (Chiles & Delfiner, 2009).

In our model, the design matrix $\mathbf{F}(x)$ included both bathymetric and spatial-coordinate covariates. The depth dependence of *Dreissena* biomass was found to be nonlinear and non-monotonic, with a maximum abundance occurring at an intermediate depth range. To accommodate this pattern within the geostatistical model, we considered two approaches: 1) polynomial depth dependence, and 2) categorical depth variable (the depth range was divided into a series of categorical variables representing different depth intervals). Polynomial functions did not conform well to the observed depth dependence; a second-order polynomial did not capture the asymmetry in the trend around the depth of maximum mussel density, and higher-order polynomials were more likely to produce unrealistic values at depth extremes. In contrast, the categorical depth variable conformed well to observed trends at shallow and deep extremes without being constrained to follow a prescribed functional relationship. Therefore, we selected the categorical depth variable as a predictor in the geostatistical model. A range of depth interval widths (10, 20 and 25

m) was considered for the categorical depth variable, and an algorithm was developed to combine depth intervals systematically, when necessary, to ensure a sufficient number of observations ($n > 9$) within each of the final intervals. Depth intervals with mean biomass < 0.1 g AFDM m^{-2} were not modelled geostatistically because their associated biomass was small and because inclusion of a large number of zero-value observations did not conform well to the Gaussian signal assumption. Thus, the depth covariates, D_k , include categorical variables for each of the k depth intervals included in the geostatistical formulation. Spatial-coordinate covariates were based on X and Y , the UTM easting and northing, respectively; both linear and quadratic spatial trends were considered (using X , Y , X^2 and Y^2). Each of the three sets of depth covariates (based on 10, 20 and 25m candidate depth interval widths), along with all possible combinations of linear and quadratic spatial trends, were evaluated based on a leave-one-out cross-validation and the Bayesian Information Criterion (BIC) (Schwarz, 1978; Ribeiro Jr. & Diggle, 2001). The preferred depth interval width was selected based on comparison of cross-validation skill statistics, including the coefficient of determination (COD) and the percent bias in predicted versus observed results (on the original, untransformed scale). Linear and quadratic trends (with spatial coordinates) were then selected based on the BIC score. Spatial prediction was performed over a 2 km grid covering all of Lake Michigan. Grid and observation coordinates were projected to UTM Zone 16 North to minimize distortion in distance and area calculations. For deterministic trend development, spatial coordinates were converted to units of 10^5 m to avoid scale mismatch issues among covariates. Bathymetry was obtained from the NOAA National Geophysical Data Center (www.ngdc.noaa.gov/mgg/greatlakes/greatlakes.html).

After the model was parameterized, conditional simulations (Ribeiro Jr. & Diggle, 2001; Chiles & Delfiner, 2009) were conducted by sampling from the uncertainty in $S(x)$ and β at prediction grid locations x , thus creating a large number of realizations (1000) of the spatial distribution of $Z(x)$ consistent with the original point observations. Prediction grid locations corresponding to depth intervals excluded from the geostatistical model (as described above) were simulated by randomly sampling from the excluded observations. From this ensemble of realizations, probabilistic estimates of spatially aggregated quantities were then derived (e.g., Obenour *et al.*, 2013). Specifically, the mean, median

and 90% confidence intervals (based on the 5% and 95% quantiles) for biomass were calculated across the entire lake and within specific regions.

Dreissena filter-feeding intensity, F_C

We calculated the filter feeding intensity (F_C , day⁻¹) by *Dreissena* as $F_C = B \times C / d$, where B is the biomass, C is the biomass-specific volume of water cleared by filter-feeding per unit time, and d is the local bathymetric depth (Vanderploeg *et al.*, 2010). We used $C = 12 \text{ mL (mg AFDM)}^{-1} \text{ h}^{-1}$ at 3 °C, based on experiments with *D. r. bugensis* from Lake Michigan feeding on *Cryptomonas*, a preferred food of *Dreissena* and a representative alga of the Lake Michigan winter-spring phytoplankton assemblage (Vanderploeg *et al.*, 2010).

The quantity F_C may be thought of as the first-order rate coefficient for phytoplankton mortality due to dreissenid grazing under conditions of a vertically well-mixed water column. Vanderploeg *et al.* (2001; 2002; 2010) used F_C to predict the effect of mussels on summer and winter assemblages of phytoplankton, based on the observation that dreissenids filter a broad range of particle sizes. Higgins & Vander Zanden (2010) used F_C as a predictor of impact in meta-analyses of *Dreissena* across systems. Vanderploeg *et al.* (2010) related F_C to an average net phytoplankton growth rate of 0.06 d⁻¹ during spring isothermal conditions in Lakes Michigan, Erie, Huron and Ontario (Fahnenstiel *et al.*, 2000) as a benchmark to estimate the ability of *Dreissena* populations to affect the winter-spring phytoplankton bloom.

Satellite-derived chlorophyll concentration

SeaWiFS (Sea-viewing Wide Field-of-view Sensor) data were downloaded from NASA's Ocean Colour data archive (<http://oceancolor.gsfc.nasa.gov>). Level2 (L2) images were used for our analysis with ~ 1 kilometre (km) pixel resolution for final products. SeaDAS7 software was used to process and map the acquired data to UTM (Zone 16 North) projection. SeaWiFS L2 chlorophyll maps were produced using the NASA OC4 algorithm, with methods fully described in Yousef *et al.* (2014). Band ratio algorithms for chlorophyll, such as OC4, are suitable for waters in which chlorophyll is the main colour producing agent, and are known to suffer from artifacts due to interference from other constituents of surface waters, including suspended mineral

particles and coloured dissolved organic matter. Kerfoot *et al.* (2008) found good agreement between SeaWiFS-derived chlorophyll and *in situ* measurements in Lake Michigan ($R^2 = 0.874$). Comparison of SeaWiFS OC3 chlorophyll retrieval (a band ratio algorithm similar to OC4) to *in situ* chlorophyll measurements indicated that OC3 produced acceptable retrievals for open waters of Lake Michigan (Shuchman *et al.*, 2013, their Table 3). Comparison of SeaWiFS OC4 chlorophyll to several hundred *in situ* chlorophyll measurements in the Great Lakes indicated good agreement up to chlorophyll concentration of $3 \mu\text{g L}^{-1}$ with increasing bias at higher chlorophyll concentration (Lesht, Barbiero & Warren, 2013, their Fig. 4). In order to avoid potential artifacts introduced by optically complex waters, we excluded areas from our analysis with chlorophyll concentrations $> 3 \mu\text{g L}^{-1}$ in any given SeaWiFS image. Chlorophyll concentrations $> 3 \mu\text{g L}^{-1}$ were limited to nearshore areas and to eutrophic Green Bay. In addition, we excluded areas shallower than 15 m to avoid artifacts caused by bottom reflectance. Monthly mean chlorophyll concentration was determined by averaging the cloud-free pixels of each daily satellite image over the month. We excluded monthly mean images with $< 50\%$ spatial coverage from the analysis, which mainly occurred in November to January due to greater cloud cover in these months.

Regression of chlorophyll reduction on filter feeding intensity

A regression analysis was conducted to test the hypothesis that the reduction in chlorophyll concentration (before and after the invasion of *D. r. bugensis*), ΔChl , was spatially associated with the filter feeding intensity, F_C , during the isothermal period (when the water column is vertically well-mixed). To account for seasonal trends in chlorophyll concentration when estimating ΔChl , we compared chlorophyll concentration within the same month for different years. In our regression analysis, we accounted for three sources of uncertainty in the regression parameter estimates: 1) spatial and interannual variation in pre- and post-invasion chlorophyll, 2) parameter estimation by least-squares regression, and 3) estimation of dreissenid biomass spatial distribution by the geostatistical model. These sources of uncertainty were addressed by means of a sampling-based Monte Carlo method in which we sampled from the ensembles of pre-invasion years (1998-2001), post-invasion years (2005-2010), and

realizations of mussel spatial distribution from the geostatistical model. Samples of dreissenid biomass distribution for post-mussel years were determined through linear interpolation between the geostatistical conditional simulations of biomass distribution for 2005 and 2010. Each Monte Carlo sample produced a map of ΔChl and a map of post-invasion F_C on the 2 km prediction grid, and 1000 such samples were created in total. To avoid influence of spatial autocorrelation on the regression, ΔChl was averaged lake-wide within intervals of F_C . The ten intervals of F_C were selected to give an equal number of pixels within each interval (~1000). Thus, each sample produced 10 data points on which the regression was conducted. A two-segment, piecewise linear regression was selected. Regression parameter estimation was conducted using the R package “segmented” (Muggeo, 2003; Muggeo, 2008). The Monte Carlo sampling method resulted in an ensemble of 1000 regression parameter estimates. To account for uncertainty in estimation of the parameters by regression, the ensemble was enlarged to 10,000 by sampling ($n = 10$) from the uncertainty (i.e. standard error) of each of the 1000 parameter estimates. The final parameter estimate and confidence interval was taken from the percentiles of the ensemble of parameter estimates.

Results

The maps of dreissenid mussel biomass (Fig. 4) produced by the geostatistical model show the spatial patterns of the invasion of Lake Michigan by *D. polymorpha* and *D. r. bugensis*. The geostatistical model expands upon the benthic survey data of Nalepa *et al.* (2014) by predicting biomass in locations that were not sampled to provide a distinct visualization of the ring of high dreissenid mussel biomass that developed around the lake within an intermediate-depth band in 2005, with further expansion of the depth range in 2010. The categorical depth intervals (Fig. 5) and spatial coordinates were found to be significant explanatory variables in models for each period (Table 1).

The ability to quantify the spatial prediction uncertainty is an important outcome of the geostatistical modelling approach applied here. The standard deviation of the conditional simulations is an indicator of the spatial prediction uncertainty (Fig. 4, bottom).

Throughout the lake, standard deviations are comparable in magnitude to the best

estimates of mussel biomass (Fig. 4), reflecting the fact that variance increases with increasing biomass (motivating the Box-Cox transformation of the observations). The sum of $\tau^2 + \sigma^2$, representing the total variance of the observations around the covariate trend, was lowest in 1994-95 and increased substantially in later periods, consistent with the substantial increase in the variance of observations apparent in Fig. 5. This variability does not have a high degree of spatial correlation, as the range of spatial correlation (ϕ) was only 14.0 and 2.5 km in 1994-95 and 2010, respectively. The range of spatial correlation was larger in 2000 and 2005, but in these years the greatest portion of the variance was uncorrelated (i.e., $\sigma^2 < \tau^2$). Due to the limited spatial correlation in the fitted model, proximity to sampling locations does not appear to substantially affect predictive uncertainty (Fig. 4, bottom), and although sampling effort increased from 1994-95 to later periods, there was no net reduction in predictive uncertainty.

In order to summarize how biomass varied spatially and through time, biomass estimates developed through conditional simulations were aggregated lakewide and for the northern, central, and southern regions of the lake (Fig. 3), and for the same depth intervals used by Nalepa *et al.* (2009). Lakewide total biomass increased significantly for each five-year period from 1994/95 to 2010 (Table 2), despite the fact that biomass declined or remained constant in some regions in the period 2005 to 2010 (Fig. 6). The highest biomass (30-40 g AFDM m⁻²) occurred in the northern region in 2005 in the 30-50 m depth range, and subsequently declined in 2010 (Fig. 6). However, biomass increased continuously over the study period in the southern region for all depth ranges, and at depths > 50 m over all regions.

The spatial distribution of dreissenid filter feeding intensity, F_C , (Fig. 7, top) was similar to that of mussel biomass (Fig. 4), but was intensified in shallower areas, as expected from the appearance of d in the denominator of the F_C relationship. In 2005 and 2010, April chlorophyll (Fig. 7, bottom; Yousef *et al.*, 2014) was greatly reduced lakewide, relative to 2000 and earlier (Figs 2 & 7), indicating near disappearance of the winter-spring phytoplankton bloom. A white contour line was added to the chlorophyll plots in Fig. 7 to indicate the region in which F_C exceeded the benchmark net growth rate of phytoplankton during the winter-spring bloom in Lake Michigan of 0.06 d⁻¹; and we would expect the winter-spring bloom to be greatly reduced in this region. In 2005, and

to a greater extent in 2010, low chlorophyll concentrations are evident within the white contour lines (Fig. 7), providing observational evidence for a direct impact of dreissenid mussel filter feeding on the winter-spring bloom in Lake Michigan.

The geostatistical model results allow us to estimate when the filter feeding intensity, F_C , first exceeded benchmark phytoplankton growth rate within specific regions and depth ranges. In 2000, F_C was much less than 0.06 d^{-1} over most of Lake Michigan, but slightly exceeded that value in the northern region (Fig. 8). In 2005, F_C exceeded 0.06 d^{-1} in all regions for < 30 and $30\text{-}50$ m depth ranges. In 2010, compared to 2005, F_C declined in the northern region, remained constant in the central region, and continued to increase in the southern region. In the $50\text{-}90$ m depth range, F_C first exceeded 0.06 d^{-1} on a lakewide basis in 2010, resulting in a large expansion of the area impacted by dreissenid grazing (Fig. 7).

To test our hypothesis that the post-invasion reduction in chlorophyll, ΔChl , was spatially associated with *Dreissena* filter feeding intensity, F_C , we conducted a regression analysis. The two-segment, piecewise linear regression model used (Fig. 9) has four parameters, each of which may be interpreted to provide information relevant to our hypothesis: 1) γ_0 , the y-axis intercept, ΔChl at $F_C = 0$, represents the reduction in chlorophyll at locations that were not locally affected by *Dreissena* filter feeding, 2) γ_1 , the slope of the line segment from $F_C = 0$ to $F_C = \gamma_b$, is expected to be negative if ΔChl is spatially associated with F_C , 3) γ_b , the breakpoint value of F_C for the piecewise regression, is expected to be near the benchmark phytoplankton growth rate of 0.06 d^{-1} , and 4) γ_2 , the slope of the line segment for $F_C > \gamma_b$, is expected to have a slope of zero because the local impact of *Dreissena* filter feeding is expected to saturate at F_C greater than the phytoplankton growth rate.

Results of the regression analysis indicated significant negative values of γ_0 for February, April to August, and December (Fig. 10, top), indicating a significant post-invasion reduction in chlorophyll concentration even at locations where local filter feeding intensity was near zero. Significant negative γ_1 values were found for December to April (Fig. 10, bottom), indicating that filter feeding had a significant localized effect up to some threshold value in F_C (γ_b). As expected, these significant local effects

occurred in months when the lake was relatively well-mixed vertically. Also consistent with our hypothesis was the finding that γ_1 values were not significantly different from zero for months associated with summer stratification (May to November), with the exception of June, which had a significant positive value of γ_1 . For months associated with the winter-spring bloom (February to April), γ_b was 0.03 d^{-1} (0.00 to 0.08, 95% CI for the three months combined), not significantly different from the benchmark phytoplankton growth rate of 0.06 d^{-1} . Finally, γ_2 was not significantly different from zero (95% CI does not include zero) for any month, indicating that the reduction of chlorophyll due to the local effect of *Dreissena* filter feeding was saturated for $F_C > \gamma_b$.

Discussion

Before the invasion of Lake Michigan by *D. r. bugensis*, a late winter to spring (March to May) phytoplankton bloom was typical (Fahnenstiel & Scavia, 1987; Fahnenstiel *et al.*, 2000; Kerfoot *et al.*, 2008; Kerfoot *et al.*, 2010). Lake Michigan was often vertically well-mixed during the spring bloom (Fahnenstiel *et al.*, 2000), resulting in spatial patterns of winter-spring chlorophyll concentrations related to bathymetry: higher chlorophyll in moderate depth regions where SML-mean light exposure is higher, and low chlorophyll in the deep central basins where SML-mean light exposure is least (Fig. 7; 1998, 2000).

Initiation of the winter-spring phytoplankton bloom in lakes and oceans is sensitive to the thickness of the surface mixed layer which, in combination with the light attenuation coefficient, controls the mean light exposure of phytoplankton cells (Sverdrup, 1953; Fahnenstiel *et al.*, 2000; Siegel, Doney & Yoder, 2002). In order for the winter-spring phytoplankton bloom to begin, the population growth rate must exceed loss processes, which include phytoplankton respiration, excretion, sinking, and losses to heterotrophic grazing; thus the critical light level to initiate the winter-spring bloom (i.e. compensation irradiance) increases with increasing losses of phytoplankton to heterotrophic grazing (Sverdrup, 1953; Siegel, Doney & Yoder, 2002). If grazing pressure on phytoplankton is sufficiently high, then the compensation irradiance may not be exceeded until the

thickness of the surface mixed layer has been reduced by summer stratification, and the winter-spring bloom may not occur at all.

While the disappearance of the winter-spring phytoplankton bloom after the dreissenid invasion has been shown previously for the southern basin of Lake Michigan (Fahnenstiel *et al.*, 2010b; Kerfoot *et al.*, 2010; Vanderploeg *et al.*, 2010; Yousef *et al.*, 2014), our refined maps of dreissenid biomass spatial distribution, covering all of Lake Michigan, show the spatial coherence of the mussel filter feeding intensity with reduced chlorophyll concentrations. There was a significant reduction in chlorophyll caused by the local impact of dreissenid filter feeding (Fig. 10, $\gamma_1 < 0$) in winter and spring, when Lake Michigan is often well-mixed to the bottom. The local filter-feeding impact saturated when the fraction of the water column cleared per day by *Dreissena* was near the benchmark spring phytoplankton growth rate of 0.06 d^{-1} . In addition to the local filter-feeding impact, a post-invasion reduction in chlorophyll occurred lake-wide, including locations that were not locally affected by dreissenid grazing (Fig. 10, $\gamma_0 < 0$).

The significant positive γ_1 value that occurred in June was surprising because it might be interpreted to suggest greater chlorophyll concentration at locations where dreissenid filter feeding intensity was greatest. Direct, local effects of dreissenid filter feeding would not be expected in June when stratification cuts off benthic filter feeders from surface chlorophyll. One possible explanation for the higher June chlorophyll concentration in locations affected by dreissenid filter feeding is an indirect effect through altered nutrient cycling; uptake of available phosphorus may have been delayed by suppression of the winter-spring bloom so that available phosphorus in June was greater post- than pre-invasion. Additional data would be needed to test this hypothesis. June was also unique for having the largest post-invasion reduction in chlorophyll (Fig. 10, top) that was not related to local filter-feeding impacts (negative γ_0 value). The large lake-wide reduction in June chlorophyll after the invasion is likely to be a cumulative effect of the suppression of the winter-spring bloom over the preceding months.

The mid-lake reef provided an interesting test of the ability of the geostatistical model to predict the occurrence of *Dreissena* at locations where observations were not available in the Ponar survey data set. The mid-lake reef has a substratum consisting of carbonate

rock, and separates the southern and central regions of Lake Michigan (Fig. 3). This reef area could not be sampled in the surveys owing to the inability of the Ponar grab to collect a sample from a hard substratum. The geostatistical model predicted the occurrence of *Dreissena* on the mid-lake reef complex based on the suitable depth habitat of the reef (depths of 40 - 100 m). Surveys and images taken with a remotely operated underwater vehicle (ROV) indicated that *D. r. bugensis* was scarce on the reef complex in 2002 but had densely colonized it by 2006 (Houghton, Paddock & Janssen, 2014), which is consistent with the spatial distributions shown in Fig. 4.

Given the rapid expansion of *D. r. bugensis* over the 15-year sampling period, and the decline in phytoplankton, it is interesting to consider whether there is any indication that this species has approached a limited carrying capacity in Lake Michigan over the most recent period (2005 to 2010). Note that *D. r. bugensis* first invaded Lake Michigan in the northern region (first found in 1997) and subsequently spread southwards (Nalepa *et al.*, 2001; Nalepa, Fanslow & Lang, 2009). In the < 30 m depth interval, biomass converged toward 10 g AFDM m⁻², decreasing in the northern, increasing in the southern and remaining constant in the central region. In a similar manner, biomass converged toward 20-30 g AFDM m⁻² within the 30-50 and 50-90 m depth intervals. The observation that biomass declined or remained constant in some northern regions, while biomass increased in the south but had not yet reached the peak values observed in the north, would be consistent with carrying capacities of ~10 g AFDM m⁻² in the < 30 m depth range and 20-30 g AFDM m⁻² in the 30-90 m depth range. Since biomass was still increasing in the > 90 m depth range lake-wide, and this range comprises a large portion of total lake area (43%), further monitoring will be needed to reveal the ultimate carrying capacity for *Dreissena* in Lake Michigan.

We expect spatial patterns in *Dreissena* biomass in Lake Michigan to continue to shift in relation to variable population growth and responses to environmental conditions (self-induced or otherwise). As illustrated, biomass can be broadly different depending on lake region and depth. Since dreissenids have such profound impacts on water quality, nutrient and energy cycling, and the abundance of other species, not only in Lake Michigan (Fahnenstiel *et al.*, 2010a) but in other bodies of water (Higgins & Vander Zanden, 2010), detailed distribution maps based on benthic surveys and the geostatistical

modelling approach outlined here provide a valuable tool for assessing and interpreting further impacts at both whole lake and regional scales.

Supporting information

Electronic data set: Lake Michigan *Dreissena* biomass values from the geostatistical model on the 2-km prediction grid are provided in the text files

“Rowe_etal_LakeMichiganDreissenaBiomass_X.txt”, where X is the year (1994-95, 2000, 2005, 2010), and the *Dreissena* biomass observed values are provided in the file “Nalepa_DreissenidLakeMichigan.txt”.

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Table 1. Geostatistical covariate trend model, skill statistics, and covariance parameters for each year. Percent bias and coefficient of determination (COD) were determined by leave-one-out cross validation. The covariates are the categorical depth variables D_k , and spatial coordinates X and Y (UTM easting and northing, respectively). The covariance parameters τ^2 and σ^2 are the nugget and partial sill (for Box-Cox transformed *Dreissena* biomass) and ϕ is the range of spatial correlation (km).

Year	Covariates	Pct. bias	COD	τ^2	σ^2	ϕ
1994-95	$D_k + X + Y$	-9.3	0.27	0.0	0.7	14.0
2000	$D_k + X$	-17.9	0.13	1.4	0.5	96.1
2005	$D_k + X + Y + X^2 + Y^2$	-0.4	0.27	4.5	2.8	29.3
2010	$D_k + X + Y + Y^2$	2.4	0.46	0.4	2.5	2.5

Table 2: Mean and percentiles of the Lake Michigan total *Dreissena* biomass from the conditional simulations, millions of kg ash-free dry mass.

Year	Mean	5	25	50	75	95
1994-95	6	4	5	6	7	8
2000	18	14	17	18	20	23
2005	408	338	376	406	437	485
2010	610	547	582	607	636	680

Figure captions

Fig. 1. Summary of the *Dreissena* survey data (Ponar grab samples, Nalepa *et al.*, 2014), showing the displacement of *D. polymorpha* by *D. r. bugensis* over the study period. The top panel shows the percentage of stations at which each species was detected. The bottom panel shows boxplots of biomass at the sampling stations. Boxplots indicate the 5, 25, 50, 75 and 95th percentile of the observations, and the mean is indicated by a symbol. The number of stations sampled in each period is given below the plot.

Fig. 2. Monthly lake-wide mean surface chlorophyll concentration from SeaWiFS remote sensing imagery for the period 1998 through 2010 (Yousef *et al.*, 2014). Vertical dashed lines indicate April of each year.

Fig. 3. Map of Lake Michigan showing bathymetry contours and regions that were used to summarize results. The inset in the right panel shows the location of Lake Michigan in North America.

Fig. 4. Spatial distribution of mean (top row) and standard deviation (bottom row) of dreissenid mussel biomass from geostatistical conditional simulations. Observed values are shown as circles, in the same colour bar scale.

Fig. 5. Boxplots of observed dreissenid mussel biomass as related to station depth where samples were collected. The bathymetric depth intervals that were used to create the categorical depth variable in the covariate trend model are indicated for each year in the study period (vertical lines). The depth ranges shaded in grey had minimal biomass, and were treated separately from the geostatistical model. Boxplots are defined as in Fig. 1. Note that the y-axis scale differs among years.

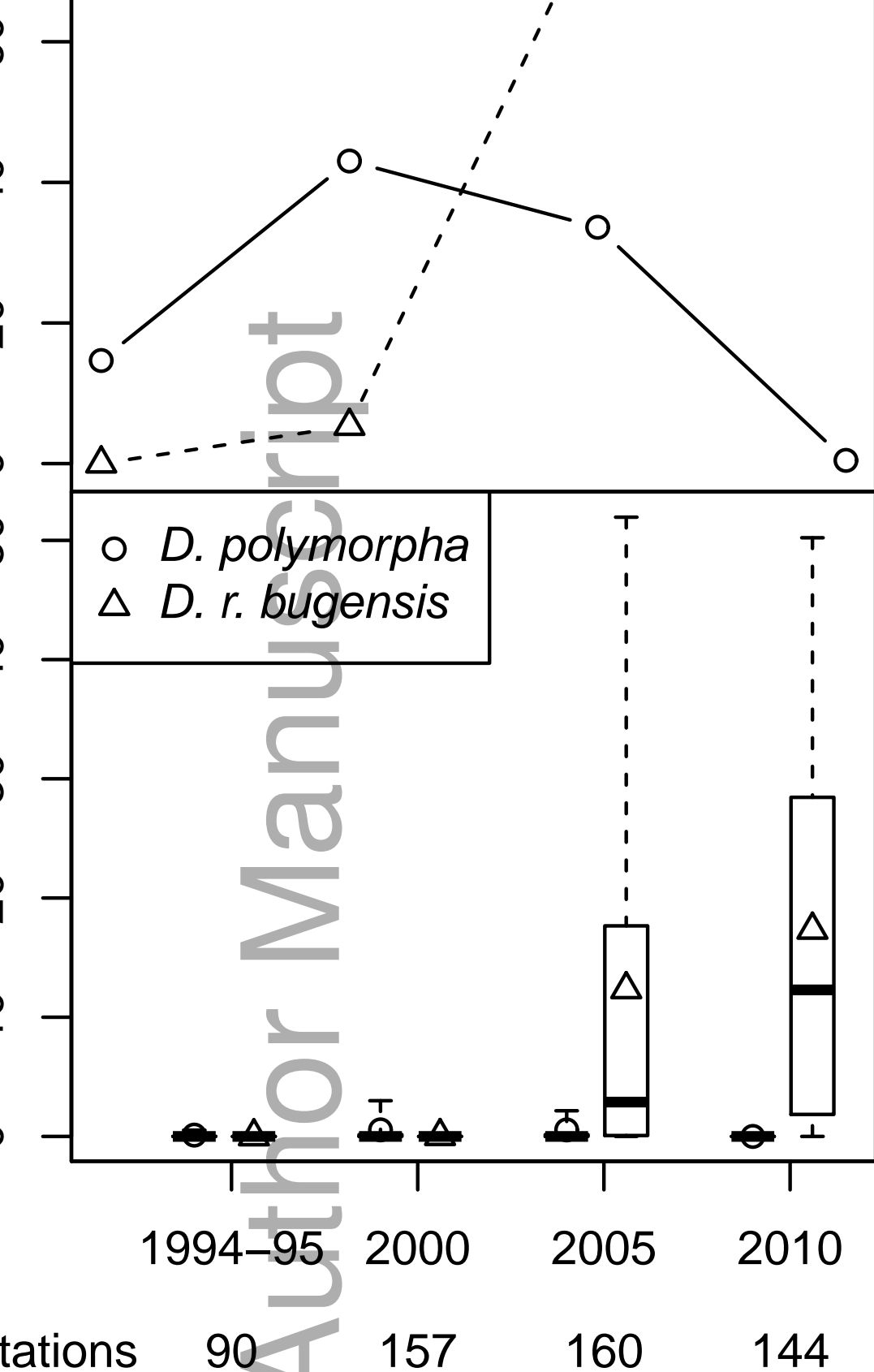
Fig. 6. Mean dreissenid mussel biomass within regions of Lake Michigan indicated in Fig. 3. Boxplots indicate the 5, 25, 50, 75, and 95th percentiles of the 1000 values obtained from the conditional simulations.

Fig. 7. Spatial distribution of the fraction of the water column cleared per day by *Dreissena* filter feeding, F_C (top row), and the mean April chlorophyll concentration for corresponding years from the SeaWiFS satellite (Yousef *et al.*, 2014) (bottom row). 1998 was the earliest year available for SeaWiFS chlorophyll, so it was paired with 1994/95 F_C . The black and white contour line indicates the 0.06 day⁻¹ contour of the fraction cleared per day, which is the benchmark phytoplankton growth rate during the winter-spring bloom (Vanderploeg *et al.*, 2010). Areas shaded in grey were excluded from the analysis by quality screening criteria applied to the satellite-derived chlorophyll data (see Methods).

Fig. 8. Fraction of the water column cleared per day by *Dreissena* filter feeding, F_C , summarized over the regions indicated in Fig. 3. The dashed line indicates the 0.06 day⁻¹ level of fraction cleared per day. Boxplots are defined as in Fig. 8.

Fig. 9. An example of fitting a piecewise linear regression model of spatially-associated values of filter feeding intensity and the change in chlorophyll concentration (ΔChl , pre *D. r. bugensis* period to post *D. r. bugensis* period). In this example, the month is April, the pre *D. r. bugensis* year is 1999, and the post *D. r. bugensis* year is 2009. Each point in the plot represents the mean ΔChl for an interval of F_C (~1000 2-km pixels per interval; point shown at mean value of F_C for each interval).

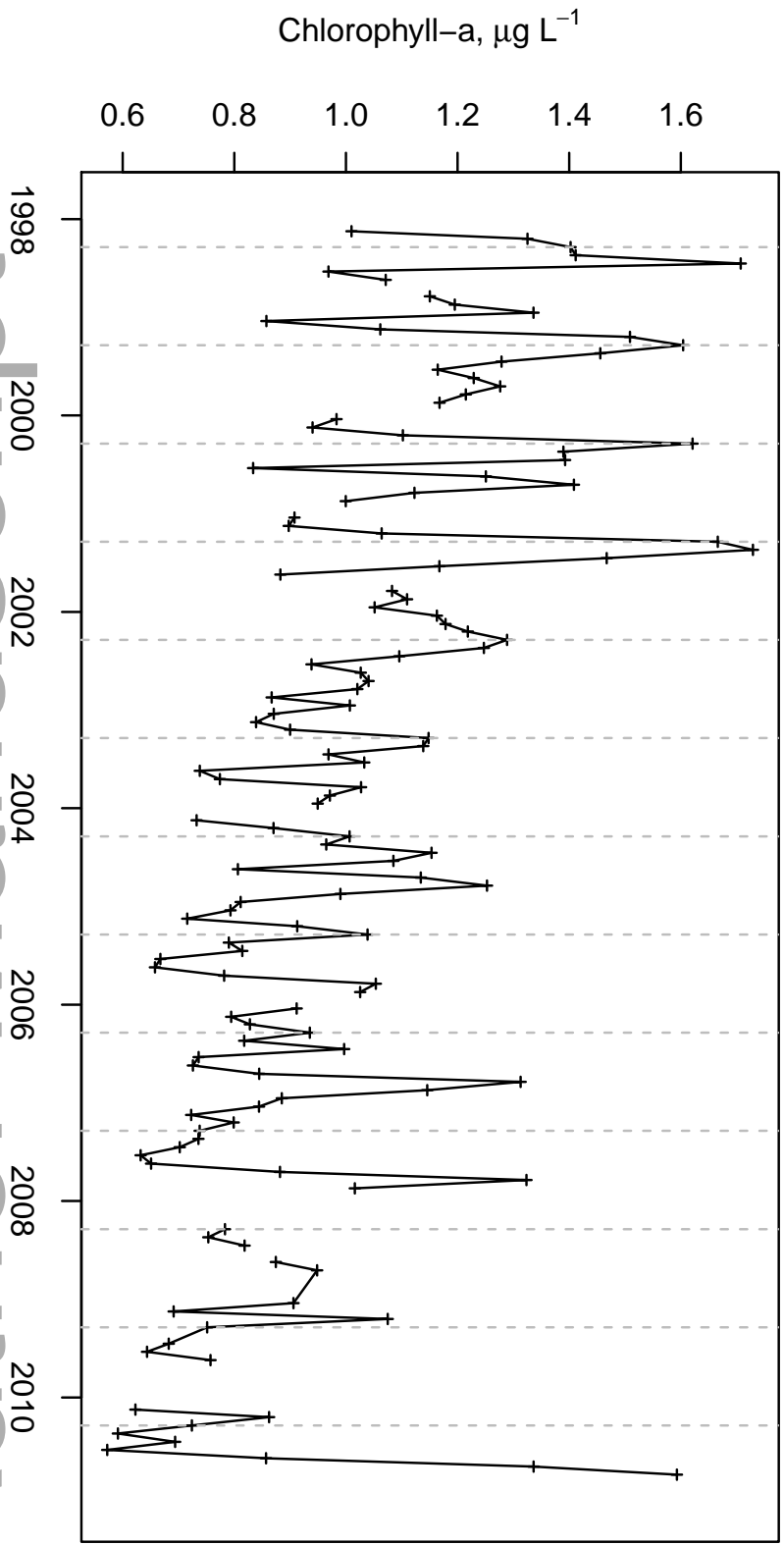
Fig. 10. Parameter estimates by month for the piecewise linear regression of ΔChl on F_C (one example shown in Fig. 9). The parameter γ_0 is the y-axis intercept (ΔChl at $F_C = 0$) and γ_1 is the slope of the line segment between $F_C = 0$ and the breakpoint value of F_C . Each bar represents the median of the Monte Carlo ensemble of parameter estimates, and the number is the frequency of occurrence of negative values in the ensemble. Bars shaded in black indicate a significant difference from zero (95% CI, based on percentiles of the ensemble, did not include zero). Negative γ_0 indicates a lake-wide post-invasion reduction in chlorophyll independent of the spatial distribution of *Dreissena* filter feeding. Negative γ_1 indicates a post-invasion reduction in chlorophyll that is spatially associated with *Dreissena* filter feeding intensity, F_C .



1994-95 2000 2005 2010

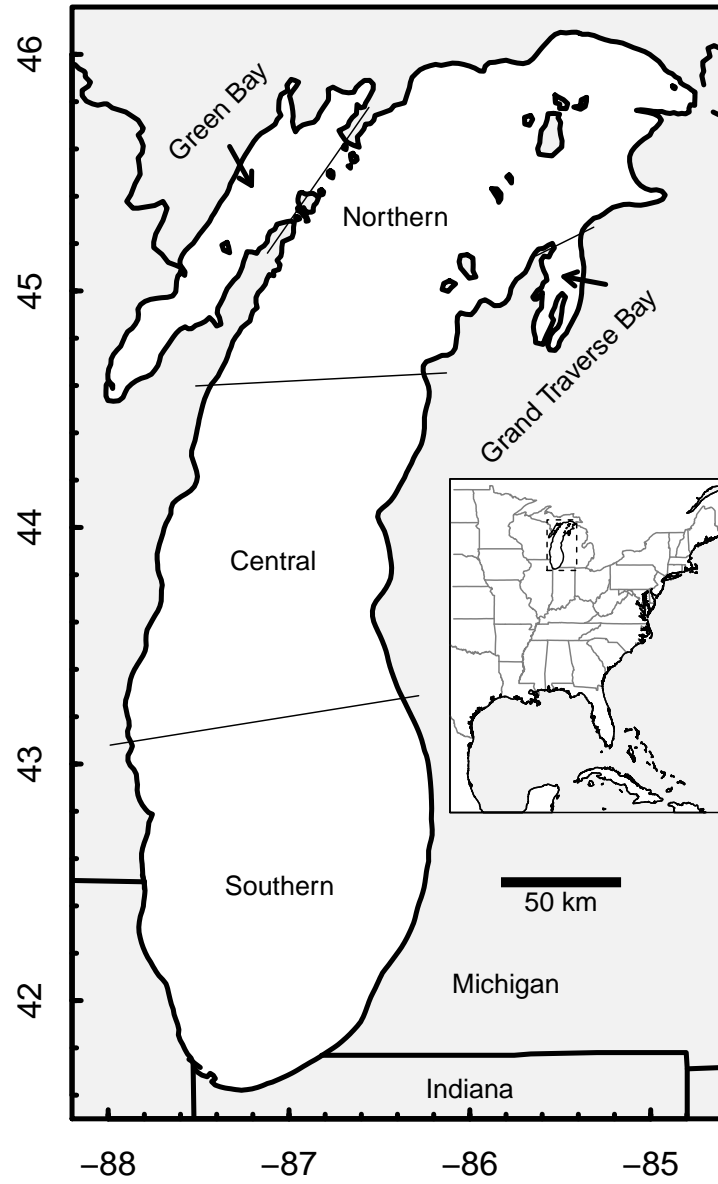
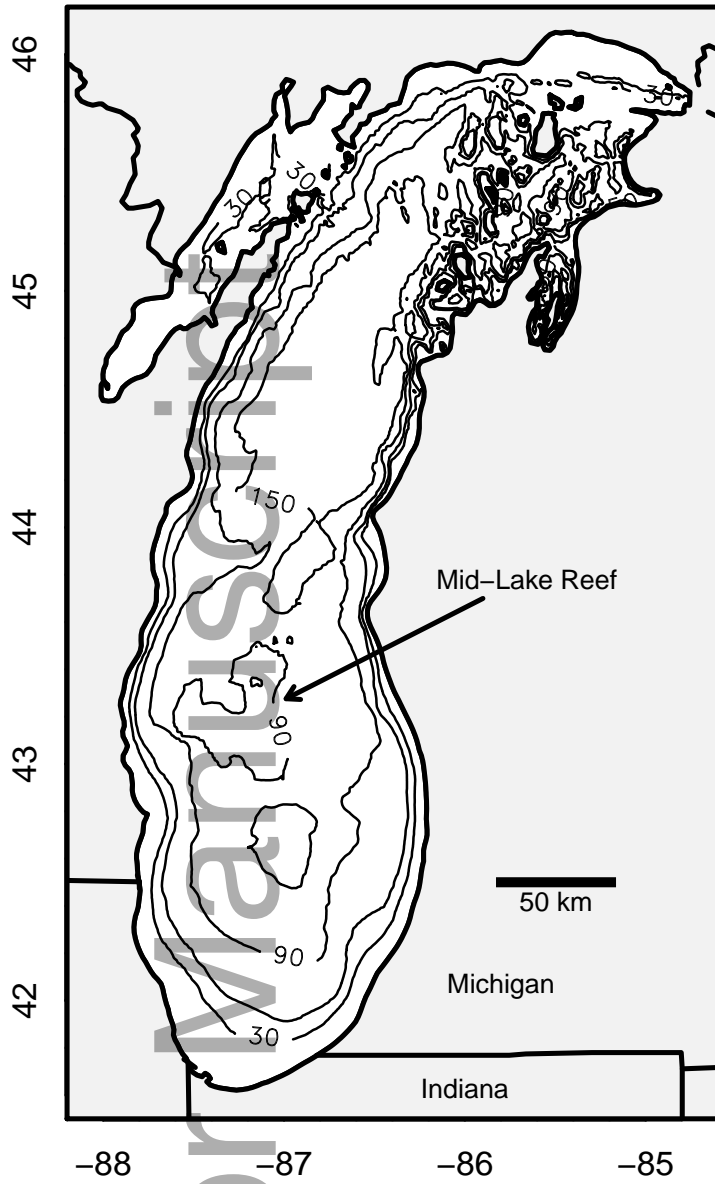
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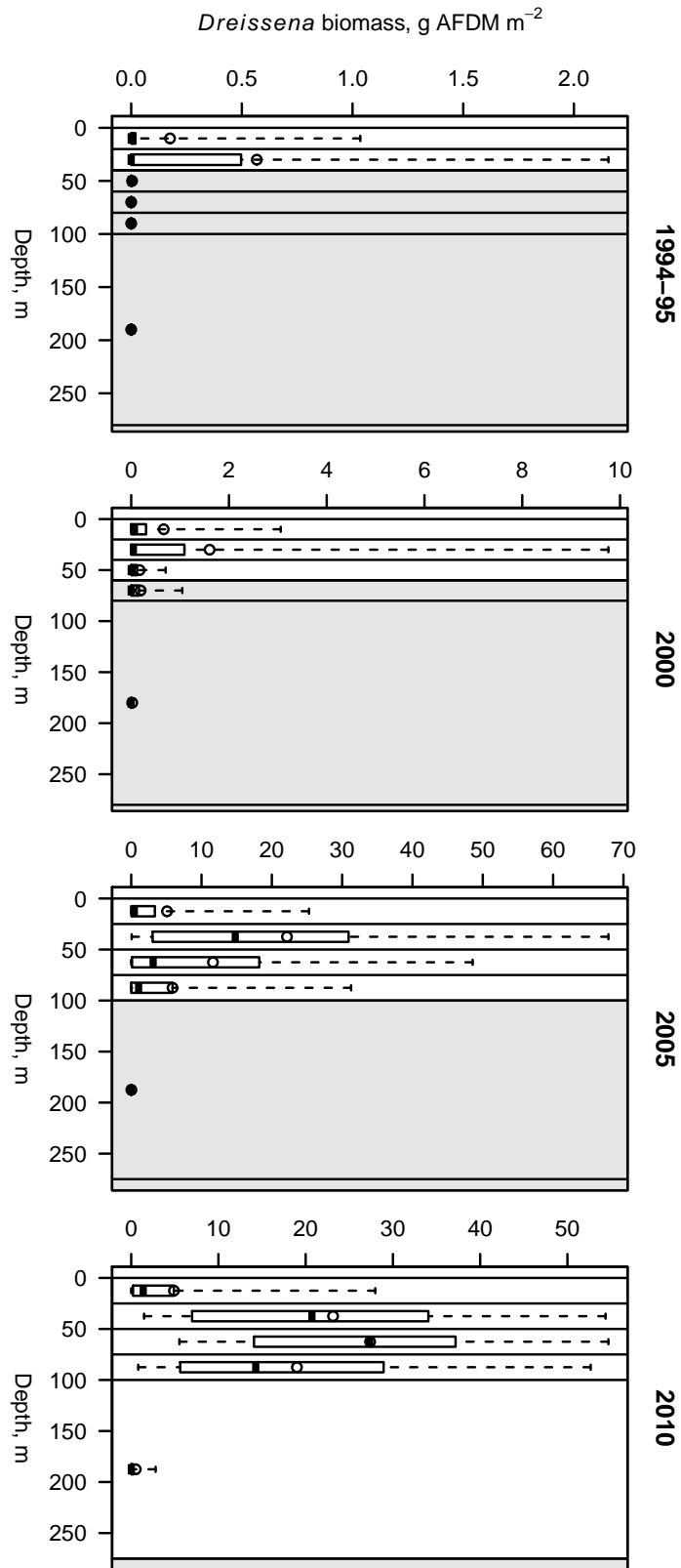
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Latitude

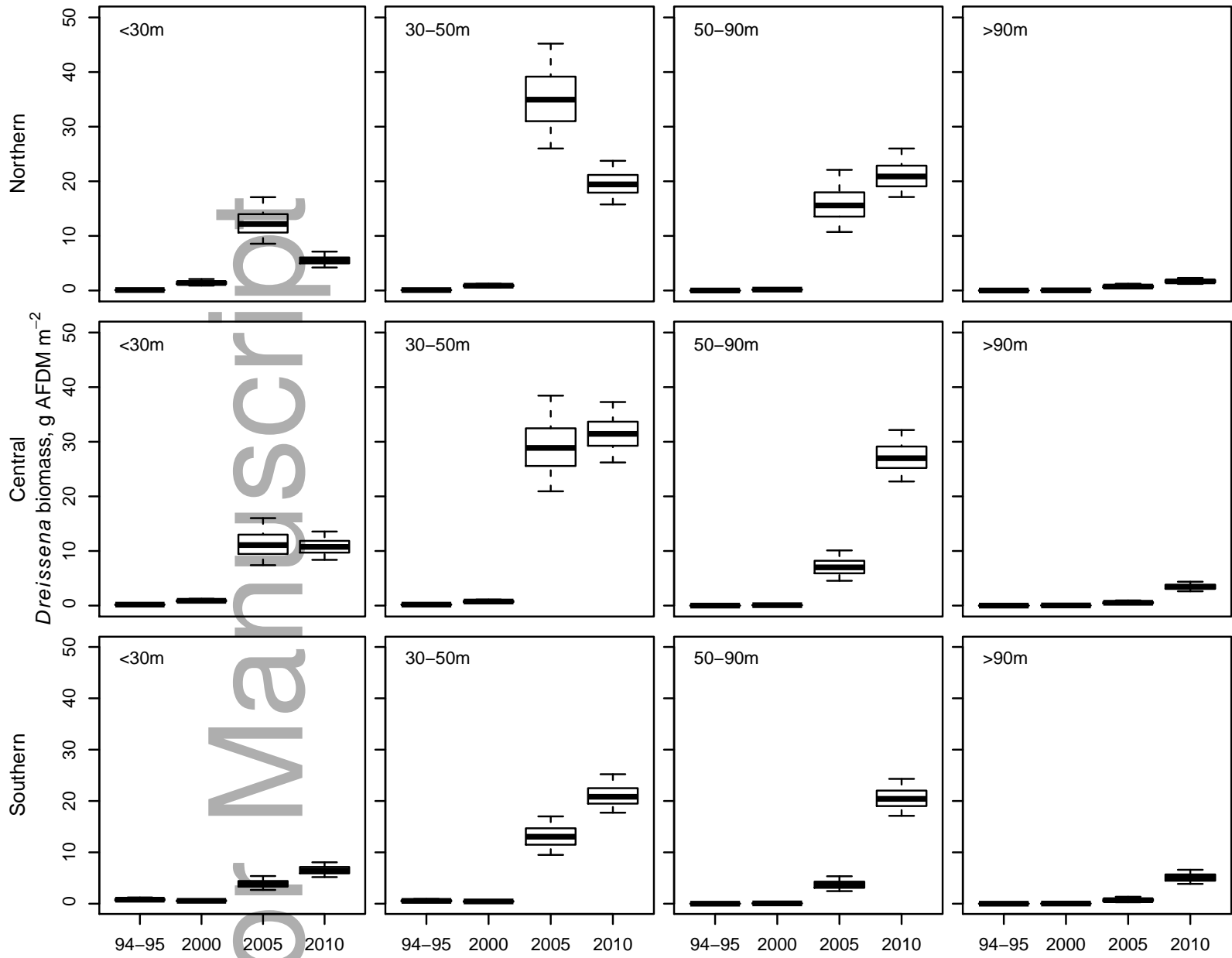


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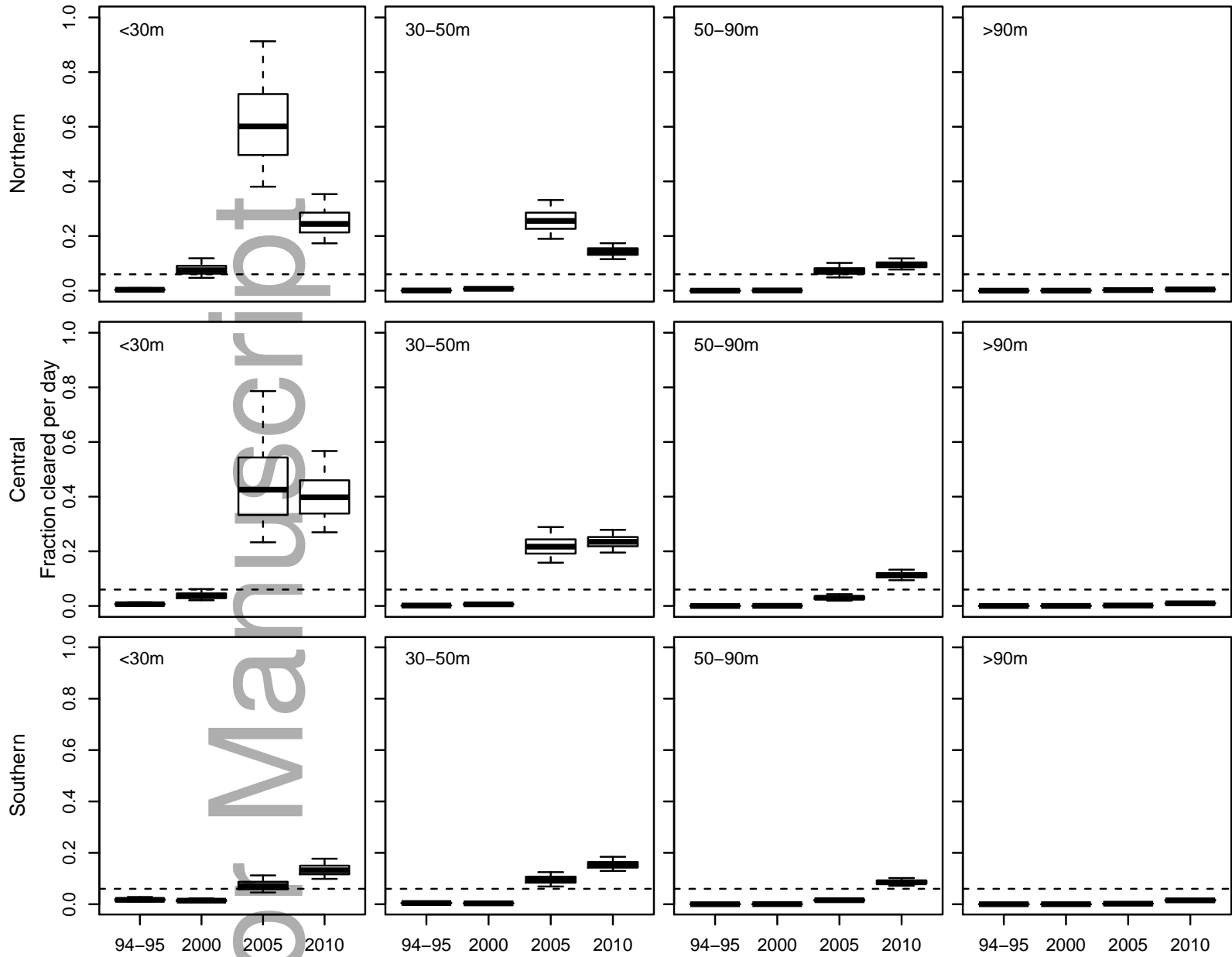
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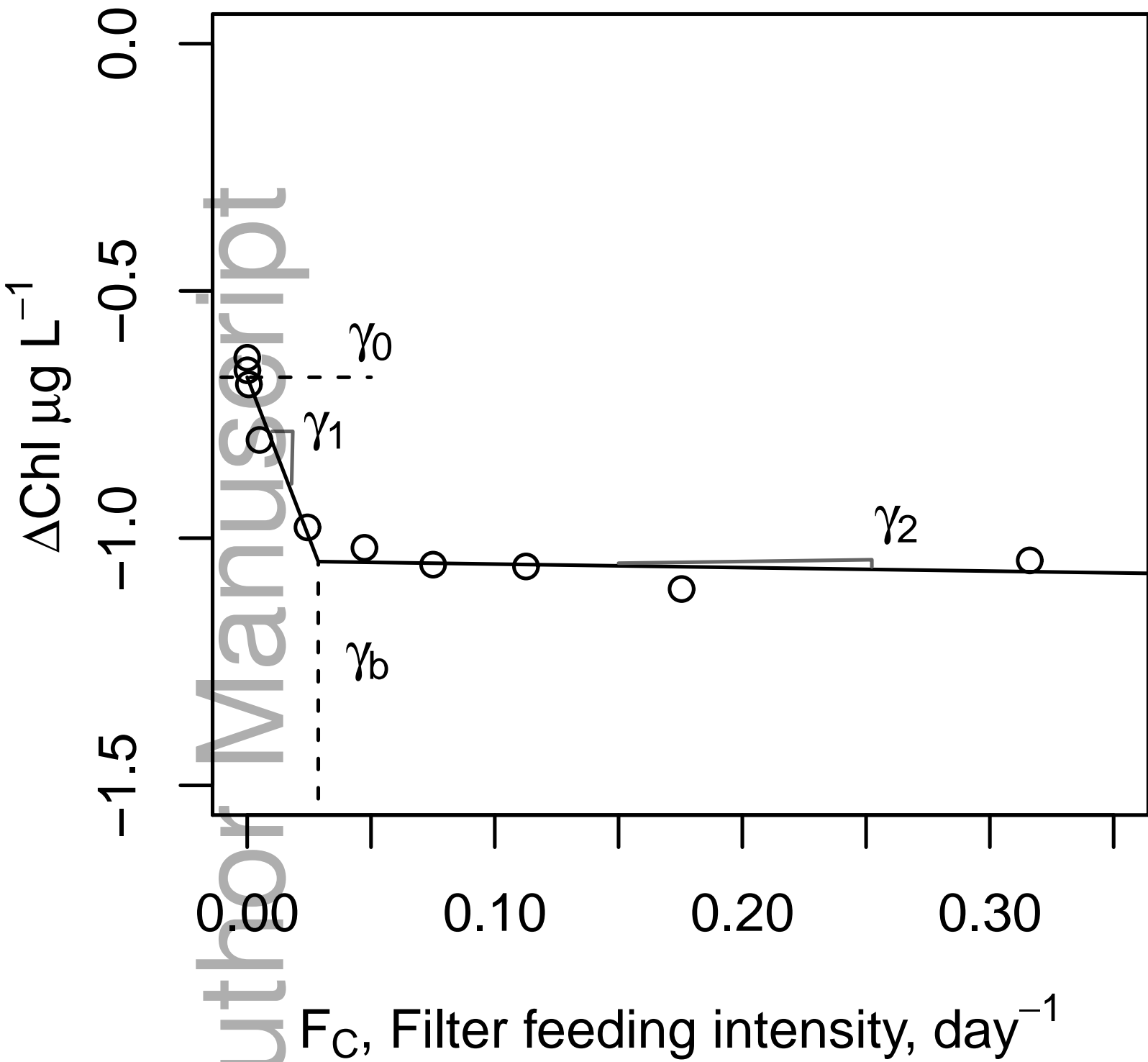
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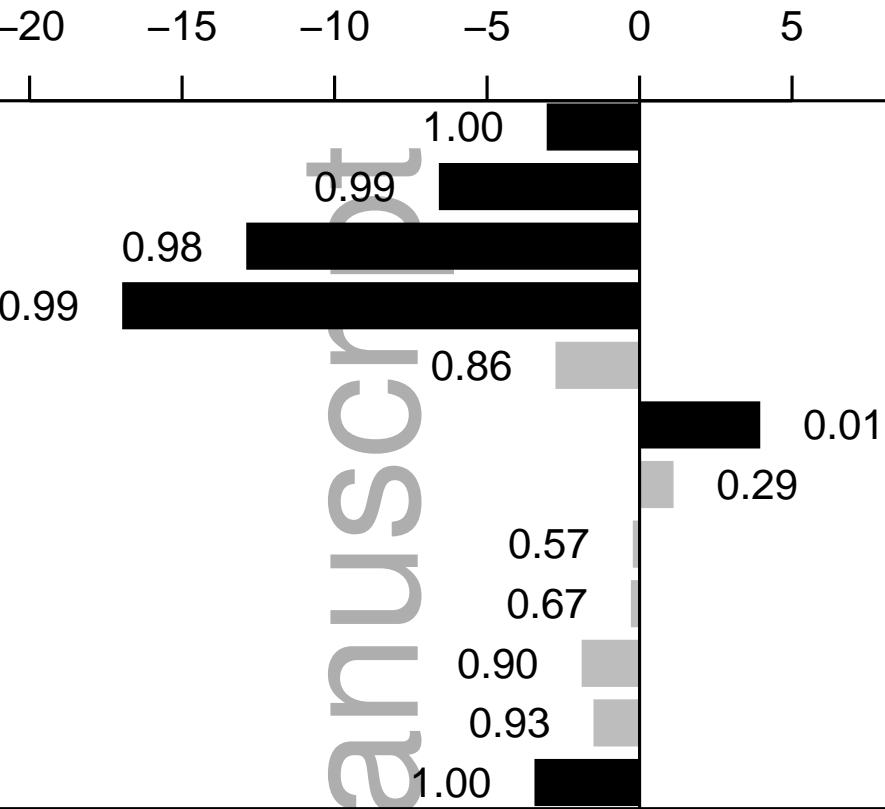


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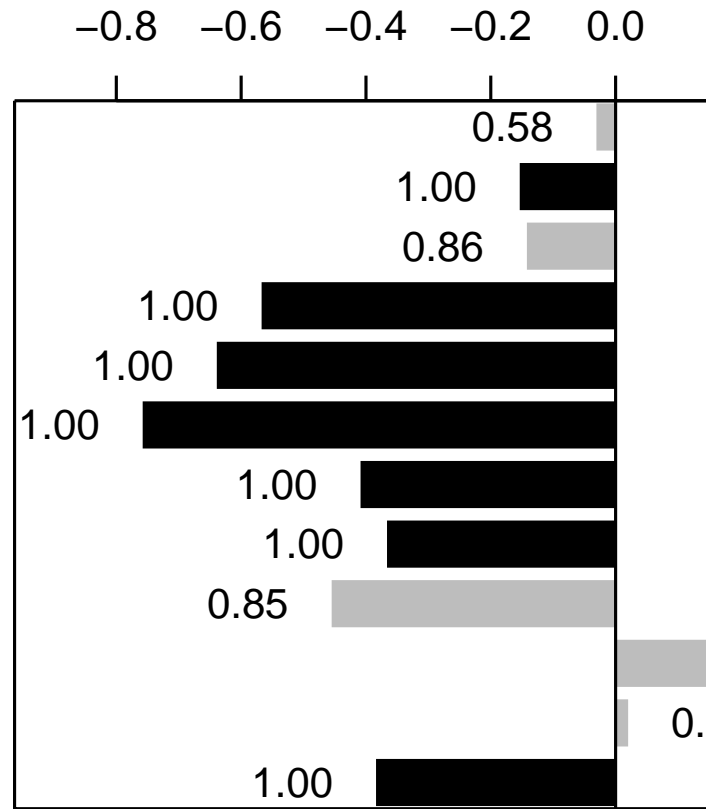


fwb12653-fig-0009.eps

γ_1 , slope of F_C vs ΔChl



γ_0 , ΔChl at $F_C = 0$



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fwb12653-fig-0010.eps