Parameter sensitivity and identifiability for a biogeochemical model of hypoxia in the northern Gulf of Mexico[☆]

Marcus W. Beck*

USEPA National Health and Environmental Effects Research Laboratory, Gulf Ecology Division, 1 Sabine Island Drive, Gulf
Breeze, FL 32561

John C. Lehrter

Dauphin Island Sea Lab, University of South Alabama, Dauphin Island, AL 36528

Lisa L. Lowe

Lockheed Martin IS & GS - Civil supporting the USEPA, Research Triangle Park, NC 27709

Brandon M. Jarvis

USEPA National Health and Environmental Effects Research Laboratory, Gulf Ecology Division, 1 Sabine Island Drive, Gulf Breeze, FL 32561

Abstract

This study addresses quantitative limitations of coupled hydrodynamic-ecological models by evaluating parameter sensitivity and identifability of a zero-dimensional (0-D) unit of a larger spatio-temporal model of hypoxia on the Louisiana continental shelf of Gulf of Mexico. A systematic framework is used to infer larger trends in dissolved oxygen dynamics over time, having implications for understanding factors that contribute to environmental conditions that are detrimental to aquatic resources. In particular, we focus on issues of parameter identifiability using local sensitivity analyses to provide quantitative descriptions of numerical constraints on model precision. The sensitivity of state variables differed considerably with parameter changes, although most variables were responsive to changes in parameters that influenced planktonic growth rates and less sensitive to physical or chemical parameters. Variation in sensitivity had a direct correspendence with identifiability, such that only small subsets of the complete parameter set were characterized as having unique effects on the model output. As a result, we provide a set of parameter selection heuristics that can be used to identify parameters for model calibration that depend on relative sensitivity and ecological categories within the biogeochemical equations. Although these concerns have been expressed in the literature, they are rarely explicitly addressed or included in evaluations of water quality models.

Keywords: Coastal General Ecosystem Model (CGEM), Gulf of Mexico (GOM), Hypoxia, Identifiability, Sensitivity

[☆]Version: Thu Mar 9 13:12:38 2017 -0600, 70c8da3c6d4aa0e50231198ae90027a250ce2ae5

^{*}Corresponding author

Email addresses: beck.marcus@epa.gov (Marcus W. Beck), jlehrter@disl.org (John C. Lehrter), lowe.lisa@epa.gov (Lisa L. Lowe), jarvis.brandon@epa.gov (Brandon M. Jarvis)

1. Introduction

Hypoxia formation in bottom waters of coastal oceans occurs primarily from excess nutrient inputs from land-based sources (Justíc et al., 1987; Diaz and Rosenberg, 1995; Howarth et al., 1996). These events are detrimental to aquatic organisms and have significant negative effects on economic resources derived from coastal ecosystems (Lipton and Hicks, 2003; Diaz and Rosenberg, 2011). An understanding of the biological, physical, and chemical processes that contribute to the growth of hypoxic areas is a critical concern for mitigating and preventing these negative impacts. Numerical ecosystem models are important tools that synthesize knowledge of ecosystem processes that contribute to hypoxia formation and for predicting the effects of proposed management activities or future scenarios (Scavia et al., 2004; Hagy and Murrell, 2007; Pauer et al., 2016). Unlike statistical models with more generic structures, simulation and process-based models include explicit descriptions of relevant processes that are constrained by empirical or observational 11 data relevant to the system of interest (e.g. Omlin et al., 2001b; Eldridge and Roelke, 2010). These models 12 are often coupled with hydrodynamic grids to provide spatially-explicit representations of patterns in three 13 dimensions (Warner et al., 2005; Zhao et al., 2010; Ganju et al., 2016). Combined hydrodynamic and bio-geo-chemical models have been developed specifically to describe hypoxic conditions on the Louisiana continental shelf (LCS) in the northern Gulf of Mexico (GOM) (Fennel et al. 2013; Obenour et al. 2015; 16 Pauer et al. 2016, Lehrter et al. in press). This area drains a significant portion of the continental United 17 States through the Mississippi-Atchafalaya River Basin (MARB) and is the second largest hypoxic area in 18 the world (Rabalais et al., 2002). Understanding processes that contribute to the frequency and duration of 19 hypoxic events remains a critical research goal for the region.

The development and application of a model represents a tradeoff between characteristics expected from 21 the output or provided by the structural components. An idealized model is sufficiently generalizable across 22 systems, provides results that are precise given the inputs, and includes components that are realistic descrip-23 tions of actual processes (Levins, 1966). Given that these characteristics cannot be simultaneously achieved, models are developed in partial dependence of reality and theoretical constructs, completely separate from both, or dependent on one or the other (Morrison and Morgan, 1999; Ganju et al., 2016). These challenges are analogous to the well-known bias-variance tradeoff in statistical models that balances the competing 27 objectives of over- and under-fitting to an observed dataset. Process-based models are more commonly imbalanced between reality and theory, such that most are over-parameterized in an attempt to completely describe reality (Denman, 2003; Nossent and Bauwens, 2012; Petrucci and Bonhomme, 2014). Quantitative limitations of over-parameterization are analogous to degrees of freedom in standard statistical models as free parameters cannot be numerically estimated when constrained to an observed dataset (Kirchner, 2006). 32 More importantly, over-parameterization can limit use across systems outside of the data domain and impose uncertainty in model predictions as realistic values for every variable may not be known or inaccurately applied from existing studies (Durand et al., 2002; Refsgaard et al., 2007; Wade et al., 2008).

Model precision can be evaluated relative to the effects of initial conditions or the observed data used

for calibration, changes in parameter values, or variation in the structural components (i.e., observational, parameter, or structural uncertainty) (Beck, 1987). Evaluating effects of parameter changes is by far the most common and simplest approach. Although sensitivity analyses should be integrated with model development, parameters are often evaluated post-hoc as a form of 'damage control' for further calibration. This approach 40 is sometimes called inverse modelling where results from sensitivity analyses are used to guide calibration or fit of the developed model to observations (Soetaert and Petzoldt 2010, or confronting models with data, 42 sensu Hilborn and Mangel 1997). Parameter sensitivity analysis combined with inverse modelling necessarily involves questions of parameter 'identifiability'. Redundancies in parameter effects lead to unidentifiable models where calibration is empirically impossible (i.e., standard algorithms will not converge) or parameter 45 values may be non-unique leading to the right answer for the wrong reason (Kirchner, 2006). Unidentifiable parameter sets have effects on model output that can be undone or compensated for by alteration of other 47 parameters. Identifiability issues are not foreign to hypoxia or eutrophication models (Omlin et al., 2001a; Estrada and Diaz, 2010; Mateus and Franz, 2015), although there is a clear need for greater integration of these concepts in practice (Fasham et al., 2006). This study describes a parameter sensitivity and identifiability analysis of a zero-dimensional (0-D) unit 51 of a larger spatial-temporal model of hypoxia dynamics on the LCS. The objectives were to provide a 52 statistical approach that demonstrates numerical limitations of parameter sets for model calibration and provide a framework for selecting parameters within the identifiability constraints. The specific goals were to 1) identify the parameters that have the greatest influence on state variables using local sensitivity analysis, 2) quantify the identifiability of subsets of the total parameter space based on sensitivity, 3) and provide a set of heuristics for choosing parameters based on sensitivity, identifiability, and parameter categories. In 57 addition to dissolved oxygen (O₂), other state variables that were evaluated included ammonium, chlorophyll a (chl-a), irradiance, nitrate, particulate organic matter (POM), dissolved organic matter (DOM), and phosphorus. In general, we provide empirical results to support the assumption that models are generally 60 over-parameterized and only a finite and smaller subset of the complete parameter set can be optimized. Although concerns of identifiability constraints have been shown in the literature, rulesets are not readily 62 available that define how parameters should be selected given tradeoffs betwen precision, sensitivity, and

65 2. Methods

66 2.1. Model description

Hypoxic events, defined as <2 mg L⁻¹ of O₂ (< 64 mmol m⁻³), occur seasonally in bottom waters in the northern GOM. The hypoxic area averages 15,540 km² annually (1993-2015) with minimum concentrations observed from late spring to early fall. Seasonal variation is strongly related to carbon and nutrient export from the MARB (Lohrenz et al., 2008; Bianchi et al., 2010), whereas hydrologic variation, currents, and wind patterns can affect vertical salinity gradients that contribute to the formation of hypoxia (Wiseman et al.,

identifiability, especially for biogeochemical models of ecosystem processes.

1997; Paerl et al., 1998; Obenour et al., 2015). The Coastal General Ecosystem Model (CGEM) was developed
to describe hypoxia dynamics on the LCS and includes elements from the Navy Coastal Ocean Model
(Martin, 2000) for hydrodynamics and a biogeochemical model with multiple plankton groups, water-column
metabolism, and sediment diagenesis (Eldridge and Roelke, 2010). The hydrodynamic component of CGEM
provides a spatially-explicit description of hypoxia using an orthogonal grid with an approximate horizontal
resolution of 1.9 km² and twenty equally-spaced vertical sigma layers on the shelf. The biogeochemical
component includes equations for 36 state variables including six phytoplankton groups (with nitrogen and
phosophorus quotas for each), two zooplankton groups, nitrate, ammonium, phosphate, dissolved inorganic
carbon, oxygen, silica, and multiple variables for dissolved and particulate organic matter from different
sources.

The core unit of CGEM is FishTank, a 0-D model that implements the biogeochemical equations in 82 Eldridge and Roelke (2010) and does not include any advection, mixing, or sediment diagenesis. Although FishTank was developed for specific application in CGEM, it can easily be applied to other hydrodynamic grids. Results are based on time-dependent differential equations that describe energy flow between phytoplankton and zooplankton groups given nutrient uptake rates, organic matter inputs and losses, inherent optical properties, and temperature (Penta et al. 2008; Eldridge and Roelke 2010, see appendix in Lehrter 87 et al. in press). A total of 108 equations are estimated at each time step to return values for each of the 36 state variables described by the model. In addition to the initial conditions, 251 parameter values for each of the equations are also supplied at model execution. Values for each of the parameters were based on estimates from the literature, field or laboratory-based measurements, or expert knowledge in absence of the former. As such, a sensitivity analysis of parameter values is warranted given that, for example, literature 92 or field-based estimates may not apply under all scenarios or expert knowledge is not completely certain (Refsgaard et al., 2007).

The sensitivity of state variables to perturbations of all relevant parameters for the 108 equations was 95 estimated using a five minute timestep from January 1st to December 31st, 2006. Irrelevant parameters were removed for several reasons; parameters were not relevant for the 0-D model (i.e., hydrodynamic 97 parameters), were considered physical constants, or had no effect given initial conditions. Additionally, FishTank includes six phytoplankton and two zooplankton groups to add complexity in community structure and foodweb dynamics. To remove obvious redundancies, the sensitivity analyses were conducted using only 100 one phytoplankton and one zooplankton group. The final set that was evaluated included 51 parameters that 101 were further grouped into one of six categories based on applicable biogeochemical components of the model: 102 optics (n = 4 parameters), organic matter (12), phytoplankton (22), temperature (2), and zooplankton (11). A full description of the model parameters is available as an appendix in Lehrter et al. in press. 104

2.2. Local sensitivity analysis

105

106

A local sensitivity analysis was performed by evaluating the change in state variables following perturbation of each parameter from its original value (Soetaert and Petzoldt, 2010; RDCT (R Development Core

Team), 2016). Parameters were individually perturbed by 50% of the original values and sensitivity S was estimated for each time step i given a change for parameter j as:

$$S_{ij} = \frac{\partial y_i}{\partial \Theta_j} \cdot \frac{w_{\Theta_j}}{w_{y_i}} \tag{1}$$

where the estimate is based on the change in the predicted value for response variable y divided by the change in the parameter Θ_j multiplied by the quotient of scaling factors w for each. The scaling factors, w_{Θ_j} for the parameter Θ_j and w_{y_i} for response variable y_i , were set as the default value of the unperturbed parameter and the predicted value of y_i after perturbation (Soetaert and Petzoldt, 2010). Scaling makes the estimates unitless to compare model sensitivity to parameters and state variables that differ in relative magnitude. Sensitivity values for all j parameters were summarized as a single value across the time series from i = 1 to n as L1:

$$L1 = \sum |S_{ij}|/n \tag{2}$$

All parameters for each of the six equation categories (optics, organic matter, phytoplankton, temperature, and zooplankton) that had non-zero L1 were retained for identifiability analysis.

2.3. Identifiability and selecting parameter subsets

120

121

122

The collinearity index γ provides a measure of potential redundancies in the response of a state variable to changes in parameter values. The index measures the linear dependence between sensitivity functions (i.e., S_i for j parameters) described above for parameter subsets and was estimated from the minimum eigenvector of the cross-product of a selected sensitivity matrix (Brun et al., 2001; Omlin et al., 2001a):

$$\gamma = \frac{1}{\sqrt{\min\left(\text{EV}[\hat{S}^{\top}\hat{S}]\right)}}\tag{3}$$

where γ ranges from one to infinity for perfectly identifiable (orthogonal) or unidentifiable (perfectly collinear) parameter sets. The sensitivity functions were supplied as a matrix \hat{S} with rows i and columns j (eq. (1)) that 125 described deviations of predicted O_2 following perturbations of each parameter. Thus, γ can be estimated 126 from results for any subset of parameter combinations. Sensitivity matrices were first normalized by dividing 127 by the square root of the summed residuals (Omlin et al., 2001a; Soetaert and Petzoldt, 2010). Estimates of γ greater than 10-15 suggest parameter sets are poorly identifiable (Brun et al., 2001; Omlin et al., 2001a), meaning parameter values that maximize precision on a calibration dataset are inestimable by conventional optimization algorithms. An intuitive interpretation of γ is provided by Brun et al. (2001), such that a 131 change in a state variable caused by a change in one parameter can be offset by the fraction $1-1/\gamma$ by the 132 remaining parameters. That is, $\gamma = 10$ suggests the relative change in O_2 for an arbitrary parameter in the selected set can be compensated for by 90% with changes in the other parameters. 134

Parameter selection for model calibration must consider the competing objectives of increased precision with parameter inclusion and reduced identifability as it relates to optimization. An additional challenge is a large number of combinations of parameter sets, which complicates selection given sensitivity differences and desired ecological categories of each parameter (e.g., practitioners may only be interested in optics parameters). Fig. 1 provides a simple graphic of the unique number of combinations that are possible for different subsets of 'complete' parameter sets of different sizes (i.e., n choose k combinations, n!/(k!(n-k)!)). The number of unique combinations increases with the total parameters in the set and is also maximized for moderate selections (e.g., selecting half the total). For example, over 10^{14} combinations are possible by selecting 25 parameters from a set of 50.

A set of heuristics was developed that addresses the tradeoff in model complexity and identifiability 144 given the challenges described above (see also Wagener et al., 2001). These rulesets were developed with the 145 assumption that parameters will be selected with preference for those with high sensitivity and identifability 146 based on $\gamma < 15$ as an acceptable threshold for subsets (e.g., 93% accountability between parameters). 147 Selection heurestics also recognized that parameter categories (i.e., optics, organic matter, phytoplankton, 148 temperature, zooplankton) may have unequal preferences by model users given questions of interest. In 149 all selection scenarios, parameters were selected by decreasing sensitivity starting with the most sensitive 150 until identifiability did not exceed $\gamma = 15$ where selections were 1) blocked within parameter category, 151 2) independent of parameter category, 3) or considering all categories equally. The selection rules produced seven subsets of parameters that could further be used to optimize model calibration. 153

2.4. Model calibration with parameter subsets

3. Results

3.1. Local sensitivity analysis

Local sensitivity analyses showed that O₂ was sensitive to perturbations in 38 of the 51 (75% of total) 157 parameters that were evaluated in FishTank (Fig. 2, Table 1). Within each parameter category, O₂ was 158 sensitive to three parameters for optics (75% of all optic parameters), eight for organic matter (67%), 16 159 for phytoplankton (73%), one for temperature (50%), and 10 for zooplankton (91%). Although O_2 had the 160 greatest sensitivity to parameters in the zooplankton category (as percentage of total), the relative effects 161 varied. Among all parameters, average sensitivity was $L1 = 9.2 \times 10^{-3}$ with values ranging from L1 = 8.34×10^{-8} for QminP (phytoplankton) to 0.05 for umax (phytoplankton). Within categories (excluding 163 temperature with one sensitive parameter), sensitivity ranged from 4.39×10^{-5} (astarOMA) to 7.51×10^{-4} 164 (astar490) for optics, 4.17×10^{-4} (KNH4) to 6.15×10^{-3} (KG1) for organic matter, 8.34×10^{-8} (QminP)165 to 0.05 (umax) for phytoplankton, and 3.69×10^{-5} (ZQp) to 0.05 (ZKa) for zooplankton (Table 1). Average 166 sensitivity values in each category were $L1 = 2.81 \times 10^{-4}$ for optics, 2.17×10^{-3} for organic matter, 0.02 for temperature, 0.01 for phytoplankton, and 0.01 for zooplankton.

Local sensitivity analyses for the additional state variables (ammonium, chl-a, irradiance, nitrate, POM, 169 DOM, and phosphorus) had similar results as O_2 with some exceptions (Fig. 2 and Tables S1 to S7). 170 All variables were sensitive to the same parameters as O₂ (38 of 51 evaluated), although average sensitivity differed between variables. Average L1 ranged from 0.02 for irradiance (Table S3) to 0.71 for DOM (Table S6). 172 All average sensitivity values for the state variables were higher than the average for O_2 ($L1 = 9.2 \times 10^{-3}$). 173 For each variable, L1 ranged from 2.24×10^{-6} (QminP) to 8.49 (mA) for ammonium (Table S1), 1.38×10^{-6} 174 (QminP) to 13.94 (mA) for chl-a (Table S2), 1.92×10^{-7} (QminP) to 0.13 (ZKa) for irradiance (Table S3), 175 6.67×10^{-7} (QminP) to 8.49 (umax) for nitrate (Table S4), 6.41×10^{-5} (KNH4) to 7.22 (mA) for POM (Table S5), 7.41×10^{-5} (KNH4) to 14.25 (mA) for DOM (Table S6), and 8.21×10^{-7} (QminP) to 1.47 (ZKa) 177 for phosphate (Table S7). For the parameter categories, ammonium was most sensitive to phytoplankton 178 parameters (average L1 = 0.8 across all parameters in the category), chl-a to phytoplankton (L1 = 1.14), 179 irradiance to zooplankton (L1 = 0.03), nitrate to zooplankton (L1 = 1.06), POM to temperature (L1 = 0.86), 180 DOM to temperature (L1 = 1.48), and phosphate to zooplankton (L1 = 0.31). Finally, average sensitivity 181 between parameter categories independent of the state variables ranged from 8.38×10^{-3} for optics (average 182 L1 across all variables) to 0.62 for phytoplankton. 183

3.2. Identifiability of parameter subsets and selection rules

202

203

The identifiability analyses suggested that many parameter subsets exceeded the thresholds of $\gamma = 10, 15$. 185 Parameter identifiability for O_2 decreased (increasing γ) at different rates with increasing size of parameter subsets depending on the parameter category or the number of top parameters that were selected (Fig. 3). 187 By category, identifiability was lowest for all combinations of parameter subsets in the phytoplankton (60%) 188 of subsets less than $\gamma = 15, 43\%$ less than $\gamma = 10$) and zooplankton categories (53.1% less than $\gamma = 15, 40\%$ 189 less than $\gamma = 10$), whereas all combinations were identifiable for optics (100% less than $\gamma = 15, 10$) and a 190 majority identifiable for organic matter (91.9% less than $\gamma = 15, 76.5\%$ less than $\gamma = 10$). Identifiability for 191 parameters in the temperature category was not evaluated because O₂ was sensitive to only one parameter 192 (i.e., $\gamma = 1$). Parameter combinations for choosing from the top, top two, top three, and top four parameters 193 in each category together had decreasing identifability with the increasing size of the selection pool (e.g., top 194 one versus top four parameters, Fig. 3). The percentage of parameter subsets that were below the acceptable thresholds for identifability was 100% less than $\gamma = 15, 10$ for the top parameter in each category, 90.6% and 80.7% for the top two, 80.7% and 70.9% for the top three, and 55.8% and 45.7% for the top four. Results for 197 the remaining state variables had similar patterns in identifiability with increasing size of parameter subsets 198 and selection categories, although differences in identifiability between state variables was observed (Fig. 4). 199 Most notably, nitrate was consistently the least identifiable variable (highest overall γ), whereas O_2 was most identifiable. 201

An alternative view of the results in Fig. 3 can be used to demonstrate the effects of parameter selection criteria and number of parameters in the selection pool on identifiability. Fig. 5 shows the percentage of identifiable parameter sets for O_2 using the same selection criteria in Fig. 3, i.e., selection of parameters only

within parameter categories and selection of the top sensitive parameters regardless of category. Fig. 5 is similar to Fig. 3, with the added effect of a chosen γ threshold on identifability. Previous studies have provided only general rules for γ thresholds (Brun et al., 2001; Omlin et al., 2001a), such that exact values for which parameter sets are inestimable likely vary between models and optimization methods. As such, multiple values are shown in Fig. 5 given that actual thresholds could vary in practice. In general, identifiability decreased with the addition of parameters, although the rate of decrease depended on the selected threshold for γ . More conservative values for γ (e.g., $\gamma = 5$) were more sensitive to the number of parameters in a subset, that is, identifiability decreased more quickly with the addition of parameters at lower γ thresholds as compared to higher γ . Notable differences in identifiability were also observed by parameter selection criteria (within categories or top parameters only), which further supports results in Figs. 3 and 4.

An evaluation of the effects of individual parameters on γ suggested that some parameters had disproportionate effects on identifiability. Based on $\gamma=15$, Fig. 3 suggests that most parameter sets for organic matter were identifiable, regardless of how many parameters were selected (i.e., two through eight). However, some subsets were not identifiable such that identification of one or more redundant parameters that are inflating γ values could provide useful information. Fig. 6 shows an alternative view of identifiability of O_2 with exclusion and inclusion of individual parameters in different sets for the organic matter category. As before, collinearity increases with more parameters in a subset, although the increase varies depending on which parameter was included or excluded from the set. For example, inclusion of KNO3 in a parameter set almost always inflated γ . All parameter subsets that did not include KNO3 were well below $\gamma=15$, suggesting that exclusion of this parameter improves identifiability. Interestingly, the inclusion of some parameters caused a reduction in γ , which contradicts the general rule that more parameters caused reduced identifiability. For example, parameter sets that included KGcdom generally had lower γ values relative to those that excluded the parameter.

Results for each of the three selection heuristics (blocked by parameter category, independent of category, all categories equally) applied to each state variable differed in the number of selected parameters and distribution of parameters within each category (Tables 2 to 4). In general, a corresponendence was observed between the number of parameters that were selected given the threhold of $\gamma = 15$ and relative identifiability between the state variables. As noted above, nitrate was the least identifiable variable (Fig. 4), whereas other variables (e.g., O_2 , irradiance) were more identifiable. The constraints on identifiability between variables were demonstrated with the selection heuristics. For example, heuristics for nitrate typically selected only one or two parameters that met the criteria as compared to more identifiable variables that included several parameters. Overall, the first selection heuristic demonstrated that the number of parameters chosen by parameter category differed independently of the state variables (Table 2). The number of selected parameters averaged across state variables in decreasing order was 4.25 parameters from the phytoplankton category, 3.5 from organic matter, 2.75 from optics, and 2.38 from zooplankton. The second and third selection heuristics (Tables 3 and 4) were similar, although more parameters were generally selected for the third heuristic given

equal importance between categories. 241

4. Discussion

243

244

261

262

263

265

266

267

270

271

272

273

274

4.1. Implications of sensitivity and identifiability analyses

Common goals in the application of biogeochemical models of ecosystem processes are to 1) accurately describe the system by matching predictions with observed data (Reckhow et al., 1990), and 2) provide a means of forecasting ecosystem condition with hypothesized management or environmental scenarios (Clark 246 et al., 2001). Although these objectives are the focus of most applications, the structural components of 247 process-based models should secondarily provide inference into which ecosystem processes and functions are 248 driving observed changes. This latter objective represents a more generic scientific purpose of biogeochemical 249 models that extends beyond the applied benefits of describing and predicting change in a particular system. Modelers often hope to identify universal principles that govern dynamics across systems and the constraint 251 of model parameters to observations provides a means of supporting or refuting hypotheses (Kirchner, 2006). 252 Extension of these principles to test the effects of structural changes and observation uncertainty on model 253 predictions provides further information to support validation of model components. This study provided a simple approach to use the effects of parameter perturbations on model state variables to characterize 255 identifiable parameter subsets that vary by parameter selection criteria. By doing so, we demonstrated 256 that small parameter subsets relative to all sensitive parameters were within the identifiability thresholds 257 described in the literature. The identifiable parameter subsets varied considerably between state variables 258 and the method for parameter selection. In general, these results provide justification for the use of explicit parameter selection heuristics that practitioners should adopt to facilitate model calibration. 260

State variables were most sensitive to parameters in the phytoplankton and zooplankton categories, particularly the maximum growth rates (umax for phytoplankton, Zumax for zooplankton), mortality coefficient for phytoplankton (mA), and the zooplankton half saturation coefficient for grazing (ZKa). An increase in the growth rate of primary producers has the potential to increase oxygen concentration through photosynthetic processes, although increased production of organic matter is balanced with respiration and bacterial decomposition that reduce O2 in the water column. Similarly, increases in zooplankton abundance with increased growth rates causes a reduction in phytoplankton biomass through grazing, which is expected to further deplete pools of organic matter. Most variables were also sensitive to variation in the half-saturation grazing coefficient which moderates the concentration of nutrients that support half the maximum grazing rate. Although the tradeoff between abundance, grazing, and decomposition is complex, the sensitivity of model state variables to parameters that directly control the abundance of primary producers is in agreement with empirical observations of factors that influence hypoxia dynamics on the LCS (Fahnenstiel et al., 1995; Roelke, 2000; Eldridge and Roelke, 2010). The sensitivity of the model output to variation in other parameters that relate to physical and chemical properties of the system was of secondary importance to biological relationships. That is, state variables were sensitive to changes in light and temperature parameters, although to a lesser extent than phytoplankton and zooplankton parameters. As such, the differing sensitivities of state variables to parameters in each of the categories was not unexpected given general ecological relationships that are well understood and described by the model.

277

278

279

280

281

283

284

285

286

287

288

289

290

291

292

293

294

295

298

299

300

302

303

304

305

307

308

309

A general conclusion from the identifiability analyses is that only limited subsets of parameters were identifiable within the constraints of local sensitivity analyses. These results support previous studies that have suggested similarly small subsets of parameters can be identified using traditional calibration schemes (e.g., Wheater et al., 1986; Ye et al., 1997; Omlin et al., 2001a). In addition to CGEM, these conclusions have relevance for many biogeochemical models that include numerous parameters and structural equations to characterize processes in the model domain. A general conclusion is that perhaps a trend towards less complex models could be beneficial given that only a small subset of parameters is identifiable and that ecosystem processes may in fact be sufficiently characterized with few parameters (Ye et al., 1997). Conversely, others have argued that model complexity is not in itself a disadvantage when parsimony is not the sole arbitrator of model structure (Reichert and Omlin, 1997). Over-parameterization can be useful if processes have importance that were not evaluated during model identification. Single objective functions that maximize model precision with identifiable parameters may also provide an incomplete characterization of model worth, which has prompted the development of probability-based models of hypoxia that explicitly include uncertainty in model components (e.g. Obenour et al., 2015). Our results demonstrated that approximately 75% of the evaluated parameters had an effect on the eight state variables, whereas CGEM includes a total of 36 variables and multiple plankton groups, not all of which have immediate concern for understanding hypoxia. The redundancies identified with the sensitivity analyses are only problematic if the primary interest is, for example, O₂ dynamics. Moreover, the proposed selection heuristics provide flexibility for choosing different parameters with the assumption that those chosen depend on the research or management question.

Results from the identifiability analyses provided additional insight into the interactions of parameters in large biogeochemical models. First, identifiability of parameter subsets was not related to the sensitivity of individual variables. As noted above, an identifiable parameter is one that has a unique effect on model predictions that cannot be compensated for or undone by changing other parameters. The magnitude of the effect of a parameter has no bearing on identifiability, which further complicates the selection of parameters for calibration. Although identifiability is the primary limiting factor in choosing a set, the relative sensitivities are more important for the decision to include or exclude individual parameters. Our analysis addressed this challenge by presenting multiple selection criteria for identifiable parameter sets that prioritized the most sensitive parameters during the selection process. Similarly, identifiability was not always related to the number of parameters in a set. Although the overwhelming trend was decreasing identifiability with more parameters, the unique effects of including an individual parameter with an existing set often reduced the γ estimate. For example, Fig. 6 shows that including KGcdom, KO2, or nitmax in parameter sets

more often reduced γ relative to sets that excluded the parameters. These examples demonstrate the complex interactions of parameter changes on variable response, highlighting the need to consider the combined and individual effects of parameters on identifiability. The selection criteria proposed in our above analyses can facilitate parameter selection and also provide diagnostic tools to identify parameters with disproportionate effects on γ .

4.2. Recommendations and conclusions

317

319

320

321

323

324

325

326

328

329

330

331

332

333

334

335

336

338

339

340

341

344

345

An evaluation of sensitivity and identifiability of relevant parameter sets is a preliminary and simplistic approach to improving model predictions. In general, uncertainty analyses that lead to improved models are ultimately expected to increase our understanding of properties defining ecological relationships. The extension of simple parameter sensitivity analyses to the generalization of ecosystem properties requires additional analysis and potential model refinement, at the core of which is the balance between generality and precision. The utility of a specific model depends on the question and objective for application to a specific system. For the above analysis, the FishTank model, as part of the larger CGEM application, was evaluated in the context of hypoxia effects on ecosystem condition and function. At the core of the simple model is a set of biogeochemical equations that characterize the system in relation to planktonic groups, nutrient requirements, and water-column metabolism. Our results have shown that relatively small subsets of parameters are identifiable given the complexity of the model, and as a result, we have provided a general approach to select parameter subsets depending on the ecological context (i.e., selection by parameter category, selection for specific state variables). Thus, the results described above have relevance for further model refinement with the specific goal of better understanding ecological dynamics that moderate hypoxia on the northern GOM. However, the general principles of sensitivity and parameter identifiability have broad applicability beyond this context and we argue that such methods should be more universally applied as an initial approach to quantify numerical constraints of biogeochemical models.

Specific approaches can be used to improve and build on the results presented herein, in addition to the more general considerations above. An evaluation of model precision following calibration with relevant parameter subsets could provide additional information that supports results from the sensitivity analyses. For simplicity, our analysis did not calibrate model parameters and an explicit assumption was that parameter subsets with γ below 10 or 15 were identifiable. To our knowledge, this threshold has not been rigorously evaluated and it is likely to vary between parameter subsets and the chosen calibration method. Variation in parameter estimates given the calibration method and different identifiability thresholds could affect the interpration of model output. Further, how results for the 0-D model extrapolate to the larger three-dimensional model should be evaluated. Although the above analyses were facilitated using the 0-D model (i.e, quick execution times, ease of changing model parameters), lack of physical transport or spatial components potentially limits extrapolation of the results. Variation in parameter estimates at lower dimensions could be magnified as errors propagate to larger scales or site-specific observations for model calibration are not appropriate across a hydrodynamic grid (Harvey, 2000; Lehrter and Cebrian, 2010). The effects of

structural or observational uncertainty should also be evaluated in the context of the larger model. Our simple approach to examine changes in the initial conditions and switches used in the model showed that sensitivity estimates of individual parameters differed relative to the original model configuration, having further implications for characterizing identifiability.

- Beck, M. B., 1987. Water quality modeling: A review of the analysis of uncertainty. Water Resour. Res. 23 (8), 1393–1442.
- ³⁵³ Bianchi, T. S., DiMarco, S. F., Jr, J. H. C., Hetland, R. D., Chapman, P., Day, J. W., Allison, M. A., 2010.
- The science of hypoxia in the Northern Gulf of Mexico: a review. Sci. Total. Environ. 408 (7), 1471–1484.
- Brun, R., Reichert, P., Künsch, H. R., 2001. Practical identifiability analysis of large environmental simulation models. Water Resour. Res. 37 (4), 1015–1030.
- clark, J. S., Carpenter, S. R., Barber, M., Collins, S., Dobson, A., Foley, J. A., Lodge, D. M., Pascual, M.,
- Pielke, R., Pizer, W., Pringle, C., Reid, W. V., Rose, K. A., Sala, O., Schlesinger, W. H., Wall, D. H.,
- Wear, D., 2001. Ecological forecasts: an emerging imperative. Sci. 293 (5530), 657–660.
- Denman, K. L., 2003. Modelling planktonic ecosystems: parameterizing complexity. Prog. Oceanogr. 57 (3-4), 429–452.
- Diaz, R. J., Rosenberg, R., 1995. Marine benthic hypoxia: A review of its ecological effects and the behavioural responses of benthic macrofauna. Oceanogr. Mar. Biology 33, 245–303.
- Diaz, R. J., Rosenberg, R., 2011. Introduction to environmental and economic consequences of hypoxia. Int.
 J. Water Resour. Dev. 27 (1), 71–82.
- Durand, P., Gascuel-Odoux, C., Cordier, M. O., 2002. Parameterisation of hydrological models: a review and lessons learned from studies of an agricultural catchment (Naisin, France. Agron. 22 (2), 217–228.
- Eldridge, P. M., Roelke, D. L., 2010. Origins and scales of hypoxia on the Louisiana shelf: importance of seasonal plankton dynamics and river nutrients and discharge. Ecol. Model. 221 (7), 1028–1042.
- Estrada, V., Diaz, M., 2010. Global sensitivity analysis in the development of first principle-based eutrophication models. Environ. Model. Softw. 25, 1539–1551.
- Fahnenstiel, G. L., McCormick, M. J., Lang, G. A., Redalje, D. G., Lohrenz, S. E., Markowitz, M., Wagoner,
- B., Carrick, H. J., 1995. Taxon-specific growth and loss rates for dominant phytoplankton populations from
- the northern Gulf of Mexico. Mar. Ecol. Prog. Ser. 117 (1-3), 229–239.
- Fasham, M. J. R., Flynn, K. J., Pondaven, P., Anderson, T. R., Boyd, P. W., 2006. Development of a robust
- marine ecosystem model to predict the role of iron in biogeochemical cycles: A comparison of results
- for iron-replete and iron-limited areas, and the SOIREE iron-enrichment experiment. Deep. Res. I 53,
- 333-366.
- Fennel, K., Hu, J., Laurent, A., Marta-Almeida, M., Hetland, R., 2013. Sensitivity of hypoxia predictions
- for the northern Gulf of Mexico to sediment oxygen consumption and model nesting. J. Geophys. Res.
- Ocean. 118 (2), 990–1002.

- Ganju, N. K., Brush, M. J., Rashleigh, B., Aretxabaleta, A. L., del Barrio, P., Grear, J. S., Harris, L. A.,
- Lake, S. J., McCardell, G., O'Donnell, J., Ralston, D. K., Signell, R. P., Testa, J. M., Vaudrey, J. M. P.,
- ³⁸⁴ 2016. Progress and challenges in coupled hydrodynamic-ecological estuarine modeling. Estuaries Coasts
- 39 (2), 311–332.
- Hagy, J. D., Murrell, M. C., 2007. Susceptibility of a northern Gulf of Mexico estuary to hypoxia: An analysis
- using box models. Estuar. Coast. Shelf Sci. 74, 239–253.
- Harvey, L. D. D., 2000. Upscaling in global change research. Clim. Chang. 44 (3), 225–263.
- Hilborn, R., Mangel, M., 1997. The Ecological Detective: Confronting Models with Data. Princeton Univer-
- sity Press, Princeton, New Jersey.
- Howarth, R. W., Billen, G., Swaney, D., Townsend, A., Jaworski, N., Lajtha, K., Downing, J. A., Elmgren,
- R., Caraco, N., Jordan, T., Berendse, F., Freney, J., Kudeyarov, V., Murdoch, P., Zhao-Liang, Z., 1996.
- Regional nitrogen budgets and riverine N & P fluxes for the drainages to the North Atlantic Ocean:
- natural and human influences. Biogeochem. 35 (1), 75–139.
- Justíc, D., Legovíc, T., Rottini-Sandrini, L., 1987. Trends in oxygen content 1911–1984 and occurrence of
- benthic mortality in the northern Adriatic Sea. Estuar. Coast. Shelf Sci. 25 (4), 435–445.
- 397 Kirchner, J. W., 2006. Getting the right answers for the right reasons: Linking measurements, analyses, and
- models to advance the science of hydrology. Water Resour. Res. 42 (3), W03S04.
- Lehrter, J. C., Cebrian, J., 2010. Uncertainty propagation in an ecosystem nutrient budget. Ecol. Appl.
- 20 (2), 508–524.
- Lehrter, J. C., Ko, D. S., Lowe, L., Penta, B., In press. Predicted effects of climate change on the severity of
- 402 northern Gulf of Mexico hypoxia. In: Justic et al. (Ed.), Modeling Coastal Hypoxia: Numerical Simulations
- of Patterns, Controls, and Effect of Dissolved Oxygen Dynamics. Springer, New York.
- 404 Levins, R., 1966. The strategy of model building in population biology. Am. Sci. 54 (4), 421–431.
- Lipton, D., Hicks, R., 2003. The cost of stress: low dissolved oxygen and eocnomic benefits of recreational
- striped bass (Morone saxatilis fishing in the Patuxent River. Estuaries 26 (2A), 310–315.
- Lohrenz, S. E., Redalje, D. G., Cai, W. J., Acker, J., Dagg, M., 2008. A retrospective analysis of nutrients
- and phytoplankton productivity in the Mississippi River plume. Cont. Shelf Res. 28 (12), 1466–1475.
- Martin, P. J., 2000. Description of the navy coastal ocean model version 1.0. Tech. Rep. NRL/FR/7322-00-
- 9962, Naval Research Lab, Stennis Space Center, Mississippi.
- 411 Mateus, M. D., Franz, G., 2015. Sensitivity analysis in a complex marine ecological model. Water 7, 2060–
- 412 2081.

- 413 Morrison, M., Morgan, M. S., 1999. Models as mediating agents. In: Morgan, M. S., Morrison, M. (Eds.),
- Models as Mediators. Cambridge University Press, Cambridge, p. 401.
- Nossent, J., Bauwens, W., 2012. Multi-variable sensitivity and identifiability analysis for a complex environ-
- mental model in view of integrated water quantity and water quality modeling. Water Sci. Technol. 65 (3),
- ₄₁₇ 539–549.
- Obenour, D. R., Michalak, A. M., Scavia, D., 2015. Assessing biophysical controls on Gulf of Mexico hypoxia
- through probabilistic modeling. Ecol. Appl. 25 (2), 492–505.
- Omlin, M., Brun, R., Reichert, P., 2001a. Biogeochemical model of Lake Zürich: sensitivity, identifiability
- and uncertainty analysis. Ecol. Model. 141 (1-3), 105–123.
- Omlin, M., Reichert, P., Forster, R., 2001b. Biogeochemical model of Lake Zürich: model equations and
- results. Ecol. Model. 141 (1-3), 77–103.
- Paerl, H. W., Pinckney, J. L., Fear, J. M., Peierls, B. L., 1998. Ecosystem responses to internal and watershed
- organic matter loading: consequences for hypoxia in the eutrophying Neuse River Estuary, North Carolina,
- usa. Mar. Ecol. Prog. Ser. 166, 17–25.
- Pauer, J. J., Feist, T. J., Anstead, A. M., DePetro, P. A., Melendez, W., Lehrter, J. C., Murrell, M. C.,
- Zhang, X., Ko, D. S., 2016. A modeling study examining the impact of nutrient boundaries on primary
- production on the Louisiana continental shelf. Ecol. Model. 328, 136–147.
- Penta, B., Lee, Z., Kudela, R. M., Palacios, S. L., Gray, D. J., Jolliff, J. K., Shulman, I. G., 2008. An
- underwater light attenuation scheme for marine ecosystem models. Opt. Express 16 (21), 16581–16591.
- 432 Petrucci, G., Bonhomme, C., 2014. The dilemma of spatial representation for urban hydrology semi-
- distributed modelling: trade-offs among complexity, calibration and geographical data. J. Hydrol. 517,
- 997-1007.
- Rabalais, N. N., Turner, R. E., Scavia, D., 2002. Beyond science into policy: Gulf of Mexico hypoxia and
- the Mississippi river. Biosci. 52 (2), 129–142.
- 437 RDCT (R Development Core Team), 2016. R: A language and environment for statistical computing, v3.3.1.
- R Foundation for Statistical Computing, Vienna, Austria. http://www.R-project.org.
- Reckhow, K. H., Clements, J. T., Dodd, R. C., 1990. Statistical evaluation of mechanistic water-quality
- models. J. Environ. Eng. 116 (2), 250–268.
- Refsgaard, J. C., van der Sluijs, J. P., Højberg, A. L., Vanrolleghem, P. A., 2007. Uncertainty in the
- environmental modelling process a framework and guidance. Environ. Model. Softw. 22 (11), 1543–1556.

- Reichert, P., Omlin, M., 1997. On the usefulness of over parameterized ecological models. Ecol. Model. 95 (2),
- 444 289–299.
- Roelke, D. L., 2000. Copepod food-quality threshold as a mechanism influencing phytoplankton succession
- and accumulation of biomass, and secondary productivity: a modeling study with management implica-
- tions. Ecol. Model. 134, 245–274.
- Scavia, D., Justic, D., Bierman, V. J., 2004. Reducing hypoxia in the Gulf of Mexico: Advice from three
- models. Estuaries 27 (3), 419–425.
- soetaert, K., Petzoldt, T., 2010. Inverse modelling, sensitivity, and Monte Carlo analysis in R using package
- 451 FME. J. Stat. Softw. 33 (3), 1–28.
- 452 Wade, A. J., Jackson, B. M., Butterfield, D., 2008. Over-parameterised, uncertain 'mathematical marionettes'
- how can we best use catchment water quality models? An example of an 80-year catchment-scale nutrient
- balance. Sci. Total. Environ. 400 (1-3), 52–74.
- Wagener, T., Boyle, D. P., Lees, M. J., Wheater, H. S., Gupta, H. V., Sorooshian, S., 2001. A framework
- for development and application of hydrological models. Hydrol. Earth Syst. Sci. 5 (1), 13–26.
- Warner, J. C., Geyer, W. R., Lerczak, J. A., 2005. Numerical modeling of an estuary: a comprehensive skill
- assessment. J. Geophys. Res. Ocean. 110 (C5), 13.
- Wheater, H. S., Bishop, K. H., Beck, M. B., 1986. The identification of conceptual hydrological models for
- surface water acidification. Hydrol. Process. 1 (1), 89–109.
- 461 Wiseman, W. J., Rabalais, N. N., Turner, R. E., Dinnel, S. P., MacNaughton, A., 1997. Seasonal and
- interannual variability within the Louisiana coastal current: stratification and hypoxia. J. Mar. Syst.
- 12 (1-4), 237–248.
- 464 Ye, W., Bates, B. C., Viney, N. R., Sivapalan, M., Jakeman, A. J., 1997. Performance of conceptual rainfall-
- runoff models in low yielding ephemeral catchments. Water Resour. Res. 33 (1), 153–166.
- Zhao, L., Chen, C., Vallino, J., Hopkinson, C., Beardsley, R. C., Lin, H., Lerczak, J., 2010. Wetland-
- estuarine-shelf interactions on the Plum Island Sound and Merrimack River in the Massachusetts coast.
- ⁴⁶⁸ J. Geophys. Res. 115 (C10), 13.

Table 1: Sensitivity of O_2 to perturbations of individual parameters. Sensitivities are based on a 50% increase from the initial parameter value, where L1 summarizes differences in model output from the default (see eq. (2)). Parameters that did not affect O_2 are not shown. Parameters are grouped by categories as optics, temperature, phytoplankton, zooplankton, and organic matter.

Description	Parameter	L1
Optics		
Chla specific absorption at 490 nm	a star 490	7.51×10^{-4}
OMZ specific absorption at 490 nm	astarOMZ	4.92×10^{-5}
OMA specific absorption at 490 nm	astarOMA	4.39×10^{-5}
Temperature		
Optimum temperature for growth(C)	$Tref(nospA + nospZ)_{p1}$	0.02
Phytoplankton		
maximum growth rate	umax	0.05
mortality coefficient	mA	0.02
initial slope of the photosynthesis-irradiance relationship	alpha	0.02
edibility vector for Z1	ediblevector(Z1)	0.02
phytoplankton carbon/cell	Qc	0.01
phytoplankton growth respiration coefficient	respg	8.36×10^{-3}
N-uptake rate measured at umax	vmaxN	8.12×10^{-3}
phytoplankton basal respiration coefficient	respb	6.94×10^{-3}
Phytoplankton threshold for grazing, is multiplied by VOLcell	Athresh	4.57×10^{-3}
minimum N cell-quota	QminN	4.32×10^{-3}
P-uptake rate measured at umax	vmaxP	4.27×10^{-3}
coefficient for non-limiting nutrient	aN	4.23×10^{-3}
phytoplankton volume/cell	volcell	4.13×10^{-3}
half-saturation constant for P	Kp	2.9×10^{-3}
half-saturation constant for N	Kn	2.77×10^{-4}
minimum P cell-quota	QminP	8.34×10^{-8}
Zooplankton		
half saturation coefficient for grazing	ZKa	0.05
zooplankton nitrogen/individual	ZQn	0.02
Zooplankton mortality constant for quadratic mortality	Zm	0.02
maximum growth rate of zooplankton	Zumax	0.02
assimilation efficiency as a fraction of ingestion	Zeffic	0.01
proportion of grazed phytoplankton lost to sloppy feeding	Zslop	7.78×10^{-3}
Zooplankton growth-dependent respiration factor	Zrespq	5.32×10^{-3}
Zooplankton biomass-dependent respiration factor	Zrespb	2.96×10^{-3}
zooplankton carbon/individual	ZQc	9.38×10^{-5}
zooplankton phosphorus/individual	ZQp	3.69×10^{-5}
Organic Matter	VI	
turnover rate for OM1A and OM1Z	KG1	6.15×10^{-3}
turnover rate for OM2A and OM2Z	KG2	3.14×10^{-3}
O2 concentration that inhibits denitrification	KstarO2	3.04×10^{-3}
decay rate of CDOM, 1/day	KGcdom	2.98×10^{-3}
half-saturation concentration for O2 utilization	KO2	5.85×10^{-4}
half-saturation concentration for NO3 used in denitrification	KNO3	5.8×10^{-4}
maximum rate of nitrification per day	nitmax	4.99×10^{-4}
NH4 rate constant for nitrification	KNH4	4.17×10^{-4}
THE TANK COMSTAND OF INTERIORISM	1111114	4.11 \ 10

^{*}Temperature parameters apply separately to phytoplankton (p1, one group) or zooplankton (z1, one group), denoted by subscripts

Table 2: Parameter identifiability (as γ , eq. (3)) by category for relevant state variables. Selections followed the first heuristic where parameters were selected within categories from most to least sensitive until $\gamma > 15$. Rank describes the relative parameter sensitivity in each category for each state variable. Duplicate parameters and ranks in the first two columns apply only to γ values in the same row (i.e., parameter ranks vary for each variable).

Optics	Parameter	Rank	Ammonium	Chl-a	O_2	Irradiance	Nitrate	POM	DOM	Phosphate
Section Sect										
astarOMA 2 7.33 5.42 - 5.36 - 7.78 7.87 - 4.73 astarOMA 3 - - 3.87 - - - 4.73 astarOMZ 3 7.58 5.51 - 6.02 - 7.91 7.87 - RG1 1 - - - - 1.0 -<		1	1	1	1	1	1	1	1	1
$ \begin{array}{c c c c c c c c c c c c c c c c c c c $			7.33	5.42	-	5.36	-	7.78	7.87	-
Organic Matter Corganic Ma	astarOMZ	2	-	-	1.39	_	-	-	-	4.73
$ \begin{array}{ c c c c c c c c c c c c c c c c c c c$	astarOMA	3	-	-	3.87	_	-	-	-	10.04
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	astarOMZ	3	7.58	5.51	-	6.02	-	7.91	7.87	-
	Organic Matter									
KStar O	KG1	1	-	-	1	_	_	1	-	1
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	KG2	1	-	-	-	-	-	-	1	-
$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	KGcdom	1	-	1	-	1	-	-	-	-
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	KstarO2	1	-	-	-	-	1	-	-	-
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	nitmax		1	-	-	-	-	-	-	-
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	KG1	2	-	1.12	-	1.93	-	-	-	-
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	KG2	2	-	-	6	-	-	-	-	13.43
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	KGcdom	2	-	-	-	-	-	1.47	1.39	-
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	KNH4		4.03	-	-	-	-	-	-	-
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	KG1		4.09	-	-	-	-	-	-	-
	KG2	3	-	-	-	8.19	-	-	-	-
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	KGcdom	3	-	-	-	-	-	-	-	13.75
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	KO2	3	-	-	-	-	-	14.07	11.96	_
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	KstarO2	3	-	-	6.04	-	-	-	-	_
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	KGcdom	4	4.19	-	6.12	-	-	-	-	-
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	KO2	4	-	-	-	-	-	-	-	14.68
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	KstarO2	4	-	-	-	10.65	-	14.08	-	-
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	KO2	5	9.47	-	8.61	-	-	-	-	-
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	Phytoplankton									
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	mA	1	1	1	-	-	-	1	1	-
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	umax	1	-	-	1	1	1	-	-	1
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	ediblevector(Z1)	2	1.13	1.17	-	-	-	1.15	-	-
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	mA		-	-	1.19	1.29	-	-	-	-
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	Qc		-	-	-	-	11.57	-	-	-
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	umax	2	-	-	-	-	-	-	1.21	-
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	vmaxP		-	-	-	-	-	-	-	7.45
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	alpha		-	-	1.44	1.98	-	-	-	-
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	ediblevector(Z1)	3	-	-	-	-	-	-	2.9	-
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	umax	3	2.73	2.11	-	-	-	3.26	-	-
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	alpha	4	3.55	4.57	-	-	-	-	-	-
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	ediblevector(Z1)	4	-	-	2.09	4.09	-	-	-	-
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	Qc	4	-	-	-	-	-	4.98	-	-
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	vmaxN		-	-	-	-	-	-	4.9	-
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	alpha		-	-		-	-	10.11	-	-
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	Qc		-	-	2.9	-	-	-	-	-
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	vmaxN		8.14	-	-	-	-	-	-	-
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	Athresh	6	11.27	-	-	-	-	-	-	-
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	respg		-	-	3.41	-	-	-	-	-
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	vmaxN	7	-	-	3.97	-	-	-	-	-
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	Zooplankton									
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	$Zar{Ka}$	1	-	-	1	1	1	-	-	1
$egin{array}{cccccccccccccccccccccccccccccccccccc$	Zumax	1	1	1	-	-	-	1	1	-
$egin{array}{cccccccccccccccccccccccccccccccccccc$	ZKa	2	-	4.31	-	-	-	7.3	5.43	-
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	ZQn		-	-	3.18	6.32	9.76	-	-	8.54
$\begin{array}{cccccccccccccccccccccccccccccccccccc$			-	-		-	-	-	-	-
	Zumax		-	-	-	6.93	-	-	-	-
Zumax 4 5.2	Zm		-	-	-	11.86	-	-	-	-
	Zumax	4	-	-	5.2	-	-	-	-	-

Table 3: Parameter identifiability (as γ , eq. (3)) for relevant state variables. Selections followed the second heuristic where parameters were selected independent of category from most to least sensitive (L1, eq. (2)), until $\gamma > 15$. Rank describes the relative parameter sensitivity in each category for each state variable (O: optics, OM: organic matter, P: phytoplankton, T: temperature, Z: zooplankton).

Selections by state variable	Parameter	L1	Rank	γ
Ammonium	1 4141110101		1001111	
1	mA	8.49	1_{P}	1
2	nitmax	1.54	$1_{\rm OM}$	1.16
3	Zumax	1.42	$1_{\rm Z}$	2.9
Chlorophyll			-2	
1	mA	13.94	1_{P}	1
2	Zumax	1.02	$1_{ m Z}$	1.18
Dissolved Oxygen	20,,,,,	1.02	-2	1110
1	umax	0.05	1_{P}	1
2	ZKa	0.05	$1_{\rm Z}$	2.17
3	mA	0.02	$^{-2}_{\mathrm{P}}$	2.31
4	$Tref(nospA + nospZ)_{p1}$	0.02	1_{T}	2.37
5	ZQn	0.02	$2_{\rm Z}$	4.69
6	alpha	0.02	$3_{\rm P}$	4.91
7	Zm	0.02	3z	6.73
8	Zumax	0.02	$4_{ m Z}$	6.81
DOM			-2	
1	mA	14.25	1_{P}	1
2	$Tref(nospA + nospZ)_{p1}$	1.48	1_{T}	1.05
3	umax	1.11	$2_{\rm P}$	2.46
4	Zumax	1.01	$1_{\mathbf{Z}}^{\mathbf{I}}$	2.91
Irradiance				
1	ZKa	0.13	$1_{\mathbf{Z}}$	1
2	umax	0.09	$1_{\rm P}$	4.41
3	ZQn	0.06	$2_{\rm Z}$	7.54
4	mA	0.05	$2_{\rm P}$	8.17
5	KGcdom	0.05	$1_{\rm OM}$	9.44
6	alpha	0.04	$3_{\rm P}$	9.66
7	\overline{Zumax}	0.04	$3_{\mathbf{Z}}$	10.79
Nitrate				
1	umax	8.49	1_{P}	1
Phosphate				
1	ZKa	1.47	$1_{\mathbf{Z}}$	1
2	umax	0.78	$1_{\rm P}$	11.45
3	vmaxP	0.59	$2_{\rm P}$	11.48
4	ZQn	0.5	$2_{\rm Z}$	13.74
POM	-			
1	mA	7.22	1_{P}	1
2	Zumax	0.96	$1_{\mathbf{Z}}$	1.15
3	KG1	0.92	$1_{\rm OM}$	3.87

Table 4: Parameter identifiability (as γ , eq. (3)) for relevant state variables. Selections followed the third heuristic where parameters were selected equally within each category from most to least sensitive (L1, eq. (2)), until $\gamma > 15$. Rank describes the relative parameter sensitivity in each category for each state variable (O: optics, OM: organic matter, P: phytoplankton, T: temperature, Z: zooplankton).

Selections by state variable	Parameter	L1	Rank	γ
Ammonium				
1	mA	8.49	$1_{\rm P}$	1
2	nitmax	1.54	$1_{\rm OM}$	1.16
3	Zumax	1.42	$1_{\rm Z}$	2.9
4	$Tref(nospA + nospZ)_{p1}$	0.79	1_{T}^{2}	3.46
5	astar490	0.03	10	4.25
Chlorophyll			-0	
1	mA	13.94	1_{P}	1
$\overline{2}$	Zumax	1.02	$1_{\rm Z}$	1.18
3	$Tref(nospA + nospZ)_{p1}$	0.6	$1_{ m T}$	2.62
4	KGcdom	0.07	$1_{\rm OM}$	3.24
5	astar490	0.02	$1_{\rm O}$	5.98
Dissolved Oxygen	400	0.02	-0	
1	umax	0.05	1_{P}	1
$\overline{2}$	ZKa	0.05	$1_{\mathbf{Z}}$	2.17
3	$Tref(nospA + nospZ)_{p1}$	0.02	1_{T}	2.29
4	KG1	6.15×10^{-3}	1_{OM}	3.85
5	astar490	7.51×10^{-4}	1 _O M	3.89
6	mA	0.02	$^{10}_{ m 2p}$	4.42
7	ZQn	0.02		5.22
DOM	ZQn	0.02	2_{Z}	3.22
1	mA	14.25	1_{P}	1
2	$Tref(nospA + nospZ)_{p1}$	1.48	1_{T}	1.05
3	Zumax	1.01	$1_{ m Z}$	2.61
4	KG2	0.94		3.39
5			$1_{\rm OM}$	
5 6	astar 490	0.04	$1_{\rm O}$	4.46
7	$umax \ ZKa$	1.11 0.88	$^{2}_{P}$	$6.02 \\ 9.21$
Irradiance	ZNa	0.00	2_{Z}	9.21
	ZKa	0.12	1_	1
$\frac{1}{2}$		0.13	$1_{ m Z}$	$\frac{1}{4.41}$
3	umax VC - 1	0.09	1_{P}	
3 4	KGcdom	0.05	1_{OM}	4.5
	$Tref(nospA + nospZ)_{p1}$	0.03	1_{T}	4.5
5	astar490	0.02	$1_{\rm O}$	6.9
6	ZQn	0.06	$2_{\rm Z}$	10.63
7	mA	0.05	2_{P}	11.21
8	KG1	3.96×10^{-3}	$2_{\rm OM}$	14.65
9	astarOMA	1.47×10^{-3}	$2_{\rm O}$	14.72
Nitrate		0.40	1	
1	umax	8.49	1_{P}	1
Phosphate	ZV.	1 47	1	1
1	ZKa	1.47	1_{Z}	1
2	umax	0.78	1_{P}	11.45
3	$Tref(nospA + nospZ)_{p1}$	0.16	1_{T}	13.71
4 POM	KG1	0.14	1_{OM}	14.64
POM	4	7 00	1	1
1	mA	7.22	1_{P}	1
2	Zumax	0.96	1_{Z}	1.15
3	KG1	0.92	1_{OM}	3.87
4	$Tref(nospA + nospZ)_{p1}$	0.86	1_{T}	3.93
5	astar 490	0.03	$1_{\rm O}$	5.81

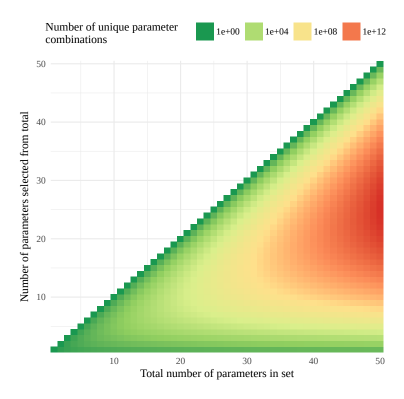


Fig. 1: Examples of unique parameter combinations from different parameter sets and number of selected parameters. The number of combinations are shown for increasing numbers of selected parameters from the total in the set, where 50 parameter sets are shown each with one through 50 total parameters. Note that the number of unique combinations is shown as the natural-log.



Fig. 2: Sensitivity values (L1, eq. (2)) of all state variables to changes in a 50% increase in parameter values. Parameters are grouped by category: optics, organic matter, phytoplankton, zooplankton, temperature, and zoplankton. See Table 1 for L1 values for O_2 and Tables S1 to S7 for the other state variables.

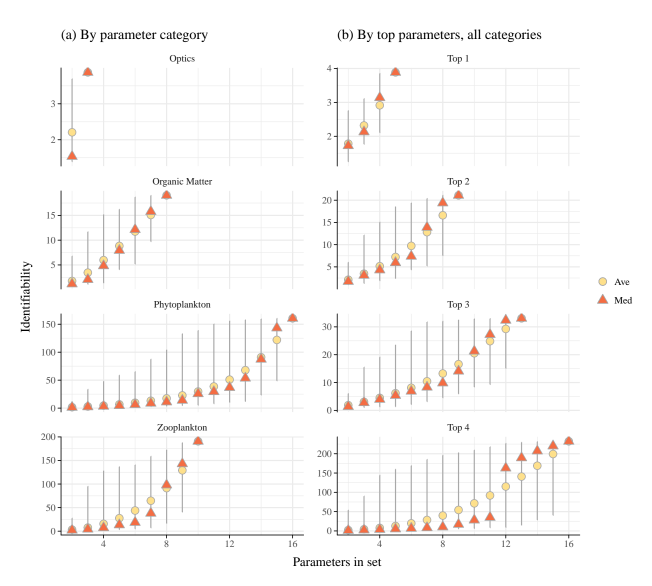


Fig. 3: Identifiability (as γ , eq. (3)) of parameter subsets for O_2 . Plots in (a) show identifiability by parameter categories and (b) shows identifiability by selecting the top 1 through 4 parameters in all categories. Lines represent identifiability ranges for the possible combinations given the number of parameters in the set. The temperature category is not shown because O_2 was sensitive to only one parameter (i.e., $\gamma = 1$).

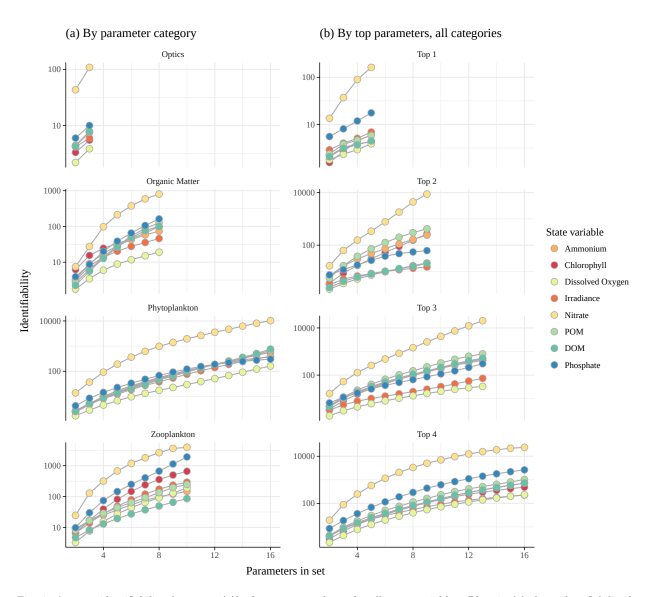


Fig. 4: Average identifiability (as γ , eq. (3)) of parameter subsets for all state variables. Plots in (a) show identifiability by parameter categories and (b) shows identifiability by selecting the top 1 through 4 parameters in all categories. Identifiability was averaged for all combinations in a parameter set to evaluate relative difference between state variables. The temperature category is not shown because all state variables were sensitive to only one parameter (i.e., $\gamma = 1$).

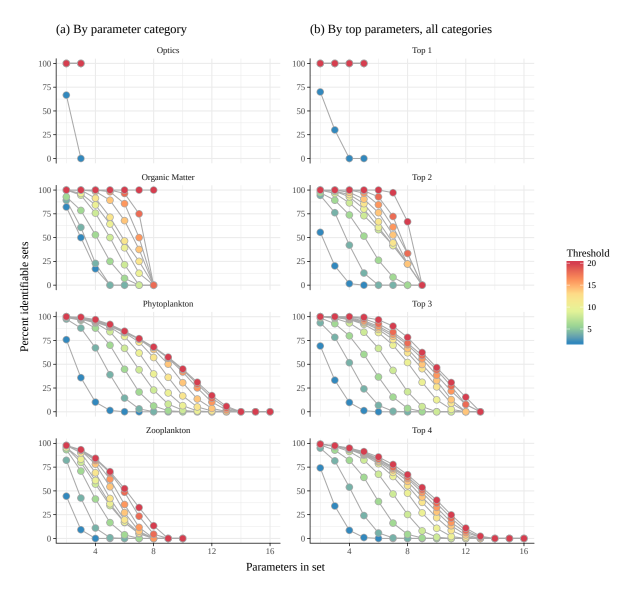


Fig. 5: Percent of identifiable parameter sets for O_2 at different γ thresholds, selection criteria, and total number of parameters in the set. Thresholds varied from $\gamma=2$ to 20 such that sets with γ below a threshold were considered identifiable relative to the value. Plots in (a) show percent of identifiable sets by selecting parameters within categories and (b) shows percent identifiable by selecting from the top 1 through 4 parameters in all categories. Percent identifiable was based on all sets in Fig. 3.

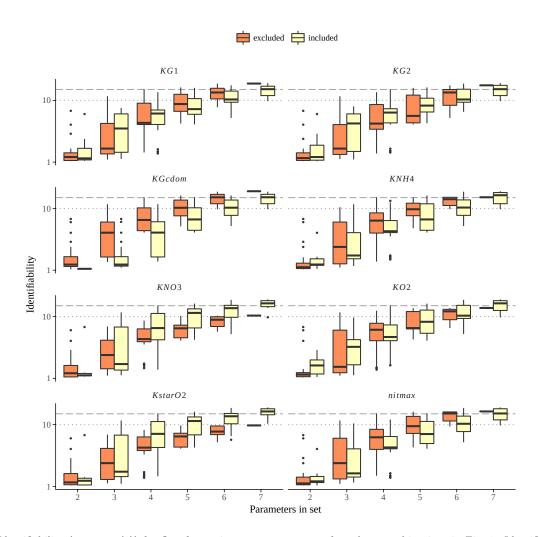


Fig. 6: Identifiability (as γ , eq. (3)) for O_2 of organic matter parameters for subset combinations in Fig. 3. Identifiability is evaluated for subsets that excluded and included the parameters at the top of each plot. Identifiability of including all eight parameters is in Fig. 3. Grey lines indicate potential thresholds at $\gamma = 10, 15$ for maximum acceptable identifiability.

Table S1: Sensitivity of ammonium to perturbations of individual parameters. Sensitivities are based on a 50% increase from the initial parameter value, where L1 summarizes differences in model output from the default (see eq. (2)). Parameters that did not affect ammonium are not shown. Parameters are grouped by categories as optics, temperature, phytoplankton, zooplankton, and organic matter.

Description	Parameter	L1
Optics		
Chla specific absorption at 490 nm	a star 490	0.03
OMA specific absorption at 490 nm	astarOMA	1.63×10^{-3}
OMZ specific absorption at 490 nm	astarOMZ	1.5×10^{-3}
Temperature		
Optimum temperature for growth(C)	$Tref(nospA + nospZ)_{p1}$	0.79
Phytoplankton		
mortality coefficient	mA	8.49
edibility vector for Z1	ediblevector(Z1)	1.32
maximum growth rate	umax	0.65
initial slope of the photosynthesis-irradiance relationship	alpha	0.6
N-uptake rate measured at umax	vmaxN	0.46
Phytoplankton threshold for grazing, is multiplied by VOLcell	Athresh	0.29
coefficient for non-limiting nutrient	aN	0.17
phytoplankton growth respiration coefficient	respg	0.16
phytoplankton basal respiration coefficient	respb	0.15
half-saturation constant for P	Kp	0.14
phytoplankton volume/cell	volcell	0.14
minimum N cell-quota	QminN	0.1
P-uptake rate measured at umax	vmaxP	0.1
phytoplankton carbon/cell	Qc	0.03
half-saturation constant for N	Kn	0.01
minimum P cell-quota	QminP	2.24×10^{-6}
Zooplankton		
maximum growth rate of zooplankton	Zumax	1.42
assimilation efficiency as a fraction of ingestion	$Zef\!\!f\!\!f\!\!ic$	0.76
half saturation coefficient for grazing	ZKa	0.74
zooplankton nitrogen/individual	ZQn	0.62
Zooplankton mortality constant for quadratic mortality	Zm	0.5
proportion of grazed phytoplankton lost to sloppy feeding	Zslop	0.3
Zooplankton growth-dependent respiration factor	Zrespg	0.22
Zooplankton biomass-dependent respiration factor	Zrespb	0.16
zooplankton phosphorus/individual	ZQp	1.07×10^{-3}
zooplankton carbon/individual	ZQc	1.44×10^{-4}
Organic Matter		
maximum rate of nitrification per day	nitmax	1.54
NH4 rate constant for nitrification	KNH4	0.66
turnover rate for OM1A and OM1Z	KG1	0.07
decay rate of CDOM, 1/day	KGcdom	0.07
half-saturation concentration for O2 utilization	KO2	0.06
O2 concentration that inhibits denitrification	KstarO2	0.05
turnover rate for OM2A and OM2Z	KG2	0.03
half-saturation concentration for NO3 used in denitrification	KNO3	7.55×10^{-3}

Table S2: Sensitivity of chl-a to perturbations of individual parameters. Sensitivities are based on a 50% increase from the initial parameter value, where L1 summarizes differences in model output from the default (see eq. (2)). Parameters that did not affect chl-a are not shown. Parameters are grouped by categories as optics, temperature, phytoplankton, zooplankton, and organic matter.

Description	Parameter	L1
Optics		
Chla specific absorption at 490 nm	a star 490	0.02
OMA specific absorption at 490 nm	astarOMA	1.45×10^{-3}
OMZ specific absorption at 490 nm	astarOMZ	1.13×10^{-3}
Temperature		
Optimum temperature for growth(C)	$Tref(nospA + nospZ)_{p1}$	0.6
Phytoplankton	- 71	
mortality coefficient	mA	13.94
edibility vector for Z1	ediblevector(Z1)	0.95
maximum growth rate	umax	0.85
initial slope of the photosynthesis-irradiance relationship	alpha	0.62
N-uptake rate measured at umax	vmaxN	0.53
phytoplankton growth respiration coefficient	respq	0.26
Phytoplankton threshold for grazing, is multiplied by VOLcell	Athresh	0.25
phytoplankton basal respiration coefficient	respb	0.24
coefficient for non-limiting nutrient	aN	0.17
half-saturation constant for P	Kp	0.14
P-uptake rate measured at umax	vmaxP	0.12
phytoplankton volume/cell	volcell	0.1
minimum N cell-quota	QminN	0.07
phytoplankton carbon/cell	Qc	0.02
half-saturation constant for N	$\check{K}n$	0.01
minimum P cell-quota	QminP	1.38×10^{-6}
Zooplankton		-
maximum growth rate of zooplankton	Zumax	1.02
half saturation coefficient for grazing	ZKa	0.85
assimilation efficiency as a fraction of ingestion	Zeffic	0.57
zooplankton nitrogen/individual	ZQn	0.52
Zooplankton mortality constant for quadratic mortality	Zm	0.41
proportion of grazed phytoplankton lost to sloppy feeding	Zslop	0.23
Zooplankton growth-dependent respiration factor	Zrespg	0.17
Zooplankton biomass-dependent respiration factor	Zrespb	0.14
zooplankton phosphorus/individual	ZQp	1.29×10^{-3}
zooplankton carbon/individual	ZQc	7.55×10^{-5}
Organic Matter		
decay rate of CDOM, 1/day	KGcdom	0.07
turnover rate for OM1A and OM1Z	KG1	0.03
turnover rate for OM2A and OM2Z	KG2	0.01
O2 concentration that inhibits denitrification	KstarO2	0.01
half-saturation concentration for O2 utilization	KO2	3.35×10^{-3}
half-saturation concentration for NO3 used in denitrification	KNO3	1.19×10^{-3}
maximum rate of nitrification per day	nitmax	3.4×10^{-5}
NH4 rate constant for nitrification	KNH4	2.97×10^{-5}

Table S3: Sensitivity of irradiance to perturbations of individual parameters. Sensitivities are based on a 50% increase from the initial parameter value, where L1 summarizes differences in model output from the default (see eq. (2)). Parameters that did not affect irradiance are not shown. Parameters are grouped by categories as optics, temperature, phytoplankton, zooplankton, and organic matter.

Description	Parameter	L1
Optics		
Chla specific absorption at 490 nm	a star 490	0.02
OMA specific absorption at 490 nm	astarOMA	1.47×10^{-3}
OMZ specific absorption at 490 nm	astarOMZ	1.34×10^{-3}
Temperature		
Optimum temperature for growth(C)	$Tref(nospA + nospZ)_{p1}$	0.03
Phytoplankton	· · ·	
maximum growth rate	umax	0.09
mortality coefficient	mA	0.05
initial slope of the photosynthesis-irradiance relationship	alpha	0.04
edibility vector for Z1	ediblevector(Z1)	0.04
N-uptake rate measured at umax	vmaxN	0.03
Phytoplankton threshold for grazing, is multiplied by VOLcell	Athresh	0.02
coefficient for non-limiting nutrient	aN	0.01
phytoplankton growth respiration coefficient	respg	0.01
half-saturation constant for P	Kp	0.01
P-uptake rate measured at umax	vmaxP	9.48×10^{-3}
phytoplankton basal respiration coefficient	respb	9.38×10^{-3}
phytoplankton volume/cell	volcell	8.1×10^{-3}
minimum N cell-quota	QminN	5.75×10^{-3}
phytoplankton carbon/cell	Qc	3.78×10^{-3}
half-saturation constant for N	Kn	9.81×10^{-4}
minimum P cell-quota	QminP	1.92×10^{-7}
Zooplankton		
half saturation coefficient for grazing	ZKa	0.13
zooplankton nitrogen/individual	ZQn	0.06
maximum growth rate of zooplankton	Zumax	0.04
Zooplankton mortality constant for quadratic mortality	Zm	0.04
assimilation efficiency as a fraction of ingestion	$Zef\!fic$	0.03
proportion of grazed phytoplankton lost to sloppy feeding	Zslop	0.02
Zooplankton growth-dependent respiration factor	Zrespg	0.01
Zooplankton biomass-dependent respiration factor	Zrespb	9.67×10^{-3}
zooplankton phosphorus/individual	ZQp	9.34×10^{-5}
zooplankton carbon/individual	ZQc	1.99×10^{-5}
Organic Matter		
decay rate of CDOM, 1/day	KGcdom	0.05
turnover rate for OM1A and OM1Z	KG1	3.96×10^{-3}
turnover rate for OM2A and OM2Z	KG2	9.88×10^{-4}
O2 concentration that inhibits denitrification	KstarO2	7.2×10^{-4}
half-saturation concentration for O2 utilization	KO2	3.54×10^{-4}
half-saturation concentration for NO3 used in denitrification	KNO3	6.18×10^{-5}
maximum rate of nitrification per day	nitmax	1.72×10^{-6}
NH4 rate constant for nitrification	KNH4	1.48×10^{-6}

Table S4: Sensitivity of nitrate to perturbations of individual parameters. Sensitivities are based on a 50% increase from the initial parameter value, where L1 summarizes differences in model output from the default (see eq. (2)). Parameters that did not affect nitrate are not shown. Parameters are grouped by categories as optics, temperature, phytoplankton, zooplankton, and organic matter.

Description	Parameter	L1
Optics		•
Chla specific absorption at 490 nm	a star 490	0.02
OMZ specific absorption at 490 nm	astarOMZ	1.27×10^{-3}
OMA specific absorption at 490 nm	astarOMA	1.19×10^{-3}
Temperature		
Optimum temperature for growth(C)	$Tref(nospA + nospZ)_{p1}$	0.3
Phytoplankton		_
maximum growth rate	umax	8.49
phytoplankton carbon/cell	Qc	0.89
initial slope of the photosynthesis-irradiance relationship	alpha	0.7
edibility vector for Z1	ediblevector(Z1)	0.33
mortality coefficient	mA	0.27
Phytoplankton threshold for grazing, is multiplied by VOLcell	Athresh	0.2
N-uptake rate measured at umax	vmaxN	0.19
coefficient for non-limiting nutrient	aN	0.13
phytoplankton growth respiration coefficient	respg	0.11
phytoplankton volume/cell	volcell	0.1
P-uptake rate measured at umax	vmaxP	0.1
half-saturation constant for P	Kp	0.09
minimum N cell-quota	QminN	0.09
phytoplankton basal respiration coefficient	respb	0.07
half-saturation constant for N	Kn	7.06×10^{-3}
minimum P cell-quota	QminP	6.67×10^{-7}
Zooplankton		
half saturation coefficient for grazing	ZKa	7.59
zooplankton nitrogen/individual	ZQn	1.17
Zooplankton mortality constant for quadratic mortality	Zm	0.7
maximum growth rate of zooplankton	Zumax	0.34
proportion of grazed phytoplankton lost to sloppy feeding	Zslop	0.26
assimilation efficiency as a fraction of ingestion	Zeffic	0.25
Zooplankton growth-dependent respiration factor	Zrespg	0.17
Zooplankton biomass-dependent respiration factor	Zrespb	0.1
zooplankton carbon/individual	ZQc	3.8×10^{-3}
zooplankton phosphorus/individual	ZQp	8.59×10^{-4}
Organic Matter		
O2 concentration that inhibits denitrification	KstarO2	0.78
half-saturation concentration for NO3 used in denitrification	KNO3	0.07
decay rate of CDOM, 1/day	KGcdom	0.04
half-saturation concentration for O2 utilization	KO2	0.03
turnover rate for OM1A and OM1Z	KG1	0.02
turnover rate for OM2A and OM2Z	KG2	0.01
maximum rate of nitrification per day	nitmax	9.96×10^{-3}
NH4 rate constant for nitrification	KNH4	9.87×10^{-3}

Table S5: Sensitivity of POM to perturbations of individual parameters. Sensitivities are based on a 50% increase from the initial parameter value, where L1 summarizes differences in model output from the default (see eq. (2)). Parameters that did not affect POM are not shown. Parameters are grouped by categories as optics, temperature, phytoplankton, zooplankton, and organic matter.

Description	Parameter	L1
Optics		
Chla specific absorption at 490 nm	a star 490	0.03
OMA specific absorption at 490 nm	astarOMA	1.73×10^{-3}
OMZ specific absorption at 490 nm	astarOMZ	1.49×10^{-3}
Temperature		
Optimum temperature for growth(C)	$Tref(nospA + nospZ)_{p1}$	0.86
Phytoplankton		
mortality coefficient	mA	7.22
edibility vector for Z1	ediblevector(Z1)	0.9
maximum growth rate	umax	0.89
phytoplankton carbon/cell	Qc	0.67
initial slope of the photosynthesis-irradiance relationship	alpha	0.67
N-uptake rate measured at umax	vmaxN	0.45
phytoplankton growth respiration coefficient	respg	0.29
phytoplankton basal respiration coefficient	respb	0.24
Phytoplankton threshold for grazing, is multiplied by VOLcell	Athresh	0.22
minimum N cell-quota	QminN	0.21
coefficient for non-limiting nutrient	aN	0.14
half-saturation constant for P	Kp	0.11
phytoplankton volume/cell	volcell	0.1
P-uptake rate measured at umax	vmaxP	0.09
half-saturation constant for N	Kn	0.01
minimum P cell-quota	QminP	7.35×10^{-4}
Zooplankton	-	
maximum growth rate of zooplankton	Zumax	0.96
half saturation coefficient for grazing	ZKa	0.79
assimilation efficiency as a fraction of ingestion	Zeffic	0.54
zooplankton nitrogen/individual	ZQn	0.49
Zooplankton mortality constant for quadratic mortality	Zm	0.39
proportion of grazed phytoplankton lost to sloppy feeding	Zslop	0.27
Zooplankton growth-dependent respiration factor	Zrespq	0.16
Zooplankton biomass-dependent respiration factor	Zrespb	0.12
zooplankton carbon/individual	ZQc	9.64×10^{-3}
zooplankton phosphorus/individual	ZQp	1.06×10^{-3}
Organic Matter		
turnover rate for OM1A and OM1Z	KG1	0.92
decay rate of CDOM, 1/day	KGcdom	0.07
half-saturation concentration for O2 utilization	KO2	0.04
O2 concentration that inhibits denitrification	KstarO2	0.02
turnover rate for OM2A and OM2Z	KG2	0.01
half-saturation concentration for NO3 used in denitrification	KNO3	3.72×10^{-3}
maximum rate of nitrification per day	nitmax	6.98×10^{-5}
NH4 rate constant for nitrification		6.41×10^{-5}
half-saturation concentration for NO3 used in denitrification maximum rate of nitrification per day	KNO3	$\begin{array}{c} 3.72 \times 10^{-3} \\ 6.98 \times 10^{-5} \end{array}$

Table S6: Sensitivity of dissolved organic matter to perturbations of individual parameters. Sensitivities are based on a 50% increase from the initial parameter value, where L1 summarizes differences in model output from the default (see eq. (2)). Parameters that did not affect dissolved organic matter are not shown. Parameters are grouped by categories as optics, temperature, phytoplankton, zooplankton, and organic matter.

Description	Parameter	L1
Optics		
Chla specific absorption at 490 nm	astar 490	0.04
OMA specific absorption at 490 nm	astarOMA	2.48×10^{-3}
OMZ specific absorption at 490 nm	astarOMZ	2.04×10^{-3}
Temperature		
Optimum temperature for growth(C)	$Tref(nospA + nospZ)_{p1}$	1.48
Phytoplankton	J(1 1)P1	
mortality coefficient	mA	14.25
maximum growth rate	umax	1.11
edibility vector for Z1	ediblevector(Z1)	0.94
N-uptake rate measured at umax	vmaxN	0.86
initial slope of the photosynthesis-irradiance relationship	alpha	0.85
phytoplankton carbon/cell	Qc	0.67
phytoplankton growth respiration coefficient	respq	0.36
phytoplankton basal respiration coefficient	respb	0.29
coefficient for non-limiting nutrient	aN	0.25
minimum N cell-quota	QminN	0.24
Phytoplankton threshold for grazing, is multiplied by VOLcell	Athresh	0.22
half-saturation constant for P	Kp	0.2
P-uptake rate measured at umax	vmaxP	0.14
phytoplankton volume/cell	volcell	0.1
half-saturation constant for N	Kn	0.02
minimum P cell-quota	QminP	4.37×10^{-3}
Zooplankton	-	
maximum growth rate of zooplankton	Zumax	1.01
half saturation coefficient for grazing	ZKa	0.88
assimilation efficiency as a fraction of ingestion	Zeffic	0.58
zooplankton nitrogen/individual	ZQn	0.54
Zooplankton mortality constant for quadratic mortality	Zm	0.41
Zooplankton growth-dependent respiration factor	Zrespg	0.17
Zooplankton biomass-dependent respiration factor	Zrespb	0.13
proportion of grazed phytoplankton lost to sloppy feeding	Zslop	0.12
zooplankton carbon/individual	ZQc	0.04
zooplankton phosphorus/individual	ZQp	1.69×10^{-3}
Organic Matter		
turnover rate for OM2A and OM2Z	KG2	0.94
decay rate of CDOM, 1/day	KGcdom	0.1
half-saturation concentration for O2 utilization	KO2	0.04
turnover rate for OM1A and OM1Z	KG1	0.04
O2 concentration that inhibits denitrification	KstarO2	0.03
half-saturation concentration for NO3 used in denitrification	KNO3	3.16×10^{-3}
maximum rate of nitrification per day	nitmax	8.44×10^{-5}
NH4 rate constant for nitrification	KNH4	7.41×10^{-5}

Table S7: Sensitivity of phosphate to perturbations of individual parameters. Sensitivities are based on a 50% increase from the initial parameter value, where L1 summarizes differences in model output from the default (see eq. (2)). Parameters that did not affect phosphate are not shown. Parameters are grouped by categories as optics, temperature, phytoplankton, zooplankton, and organic matter.

Description	Parameter	L1
Optics		
Chla specific absorption at 490 nm	astar 490	9.01×10^{-3}
OMZ specific absorption at 490 nm	astarOMZ	5.21×10^{-4}
OMA specific absorption at 490 nm	astarOMA	5.13×10^{-4}
Temperature		
Optimum temperature for growth(C)	$Tref(nospA + nospZ)_{p1}$	0.16
Phytoplankton		
maximum growth rate	umax	0.78
P-uptake rate measured at umax	vmaxP	0.59
edibility vector for Z1	ediblevector(Z1)	0.25
initial slope of the photosynthesis-irradiance relationship	alpha	0.23
mortality coefficient	mA	0.2
N-uptake rate measured at umax	vmaxN	0.18
Phytoplankton threshold for grazing, is multiplied by VOLcell	Athresh	0.13
coefficient for non-limiting nutrient	aN	0.11
phytoplankton growth respiration coefficient	respg	0.09
phytoplankton volume/cell	volcell	0.06
phytoplankton basal respiration coefficient	respb	0.06
minimum N cell-quota	QminN	0.04
half-saturation constant for P	Kp	0.03
half-saturation constant for N	\overline{Kn}	6.97×10^{-3}
phytoplankton carbon/cell	Qc	6.68×10^{-3}
minimum P cell-quota	QminP	8.21×10^{-7}
Zooplankton		
half saturation coefficient for grazing	ZKa	1.47
zooplankton nitrogen/individual	ZQn	0.5
Zooplankton mortality constant for quadratic mortality	Zm	0.35
maximum growth rate of zooplankton	Zumax	0.26
assimilation efficiency as a fraction of ingestion	Zeffic	0.19
proportion of grazed phytoplankton lost to sloppy feeding	Zslop	0.15
Zooplankton growth-dependent respiration factor	Zrespg	0.1
Zooplankton biomass-dependent respiration factor	Zrespb	0.06
zooplankton phosphorus/individual	ZQp	6.43×10^{-3}
zooplankton carbon/individual	ZQc	3.38×10^{-5}
Organic Matter		
turnover rate for OM1A and OM1Z	KG1	0.14
turnover rate for OM2A and OM2Z	KG2	0.06
decay rate of CDOM, 1/day	KGcdom	0.02
half-saturation concentration for O2 utilization	KO2	0.01
O2 concentration that inhibits denitrification	KstarO2	7.29×10^{-3}
half-saturation concentration for NO3 used in denitrification	KNO3	1.19×10^{-3}
maximum rate of nitrification per day	nitmax	2.7×10^{-5}
NH4 rate constant for nitrification	KNH4	2.64×10^{-5}