Title....

1

2

Marcus W. Beck¹, John C. Lehrter¹

¹USEPA National Health and Environmental Effects Research Laboratory Gulf Ecology Division, 1 Sabine Island Drive, Gulf Breeze, FL 32561 Phone: 850-934-2480, Fax: 850-934-2401

Emails: beck.marcus@epa.gov, lehrter.john@epa.gov

Version Date: Fri Jun 24 16:57:50 2016 -0500

3 Abstract

Bio-geo-chemical models are useful tools in environmental sciences that can guide management and policy-making. Consequently, significant time and resources are spent developing these models in system-specific contexts. The optimization of model parameters to maximize precision, including transferability of these models to different systems, are fundamental concerns in the development and application of these tools. This study describes quantitative limitations of coupled hydrodynamic-ecological modelling by contrasting numeric and ecological certainty with a systematic framework for characterizing parameter sensitivity and identifability. We evaluate a simple bio-geo-chemical model that is the 1-dimensional unit of a larger spatio-temporal model of hypoxia on the Louisiana continental shelf of Gulf of Mexico as an example. Results from analysis of the 1D model are 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. We argue that quantitative and ecological certainty in model calibration are often at odds and the practitioner must explicitly choose model components to optimize given tradeoffs between the two. We further conclude that numerically optimal parameter sets for models of hypoxia are often small subsets of the complete parameter set because of redundancies in the unique effects of paramater perturbations on model output. As a result, we demonstrate that use of a model for inference into ecological mechanisms of observed or predicted changes in hypoxic condition can be potentially misguided in the absence of quantitative descriptions of identifiability. Although these concerns have been expressed in the literature, they are rarely explicitly addressed or included in model evaluations. In addition to immediate implications for regional models, we provide a framework for describing the effects of parameter uncertainty and identifiability that can be applied to similar models to better inform environmental management.

1 Introduction

10

11

12

13

14

15

16

17

18

19

20

21

22

23

24

25

26

27

28

31

32

- 1. Simulation/biogeochemical/process-based models overview, contrast with statistical models
- 2. What models seek to provide generality, precision, realism Levins (1966), there is a

- tradeoff so models are 1) developed in partial independence and dependence on the world
 and theory, 2) function autonomously from both, or 3) represent both at the same time, from
 Morrison and Morgan (1999), cited in Ganju et al. (2016). This is similar to the
 bias-variance tradeoff for statistical models, e.g., overparameterization of a model makes it
 very biased as it fits the data (the world) exactly, tradeoff between sensitity and error with
 changes in model complexity (more complexity is less error but increasing sensitivity)
 described in Snowling and Kramer (2001)
- 3. How is model performance/uncertainty evaluated regarding what they should provide structural, observational, parameter Beck (1987)?
- 4. Parameter uncertainy as low-hanging fruit can do post-hoc and from inner to outer level of
 complexity, parameter uncertainty is the most common, e.g., marine ecological model
 Mateus and Franz (2015), but stopped short, global sensitivity analysis of eutrophication
 model Estrada and Diaz (2010)
 - 5. Challenges related to uncertainty similar to degrees of freedom, identifiability definition from Brun et al. (2001) and need to evaluate identifiability Fasham et al. (2006), Omlin et al. (2001) did a similar analysis with freshwater biogeochem model

46

47

48

This study describes a parameter sensitivity analysis to evaluate identifiability for a
bio-geo-chemical model of hypoxia for the northern Gulf of Mexico (GOM). We evaluate a
simple 1-dimensional unit of a larger spatial-temporal model to explore relationships between
multiple parameter sets and hypoxia dynamics on the Louisiana continental shelf (LCS). The
study also provides a general framework for sensitivity analysis and parameter identifiability that
can be used on similar mechanistic models. Specifically, an assumption is that models are
generally over-parameterized and only a finite and smaller subset of the larger parameter set can
be optimized for a given research question or dataset. We provide explicit guidance for choosing
such subsets of the parameter space given constraints on identifiability as directly related to
sensitivity analyses. The specific objectives are to 1) identify the parameters that have the
greatest influence on dissolved oxygen using local sensitivity analysis, 2) quantify the
identifiability of subsets of the total parameter space based on sensitivity, 3) provide a set of

heuristics for choosing parameters based on sensitivity and parameter categories with the larger mechanistic model, including extension to other state variables, and 4) discuss implications for hypoxia formation in coastal regions, including management strategies for nutrient reduction and use of mechanistic models to inform decision-making. The 'optimum' parameter space is defined as the chosen subset that represents the maximum number of identifiable parameters. Here, 'optimum' is both a qualitative description based on a research question or management goal and a quantitative objective based on numerical optimization criteria for fitting model output to a 67 calibration dataset. These results can be used to refine existing models or guide application of models to novel contexts, such as downscaling or application to new environments.

Methods 2 70

72

2.1 **Model description** 71

Low concentrations of dissolved oxygen occur seasonally on the LCS in the northern GOM. These hypoxic events, defined as <2 mg L⁻¹ (<64 mmol m⁻³ of O₂), occur in bottom waters and are caused by nutrient inputs from the Mississippi-Atchafalaya River Basin (MARB) that drains a significant portion of the continental United States. Nutrient-stimulated primary production in surface waters increases biological oxygen demand in bottom waters as sinking organic matter is decomposed. 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 factors related to hydrologic variation and wind patterns can affect vertical salinity 80 gradients that contribute to the formation of hypoxia (Wiseman et al. 1997). 81 Three-dimensional numerical simulation models have been developed to better understand 82 factors contributing to hypoxia and to predict the effects of management actions or climate 83 scenarios on future patterns (Fennel et al. 2013, Pauer et al. 2016, Lehrter et al. in review). This 84 study evaluates a recently developed hydrodynamic and ecological model that describes 85 horizontal and vertical transport and mixing of state variables relevant for hypoxia. The Coastal General Ecosystem Model (CGEM) includes elements from the Navy Coastal Ocean Model (Martin 2000) to describe hydrodynamics on the LCS 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 dynamics using an orthogonal grid with an approximate horizontal resolution of 1.9 km²
and twenty equally-spaced vertical sigma layers on the shelf (depth ≤ 100 m, with additional
hybrid layers at deeper depths). The biogenical component includes equations for 33 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 model is run for a set period of time at a given time step using atmospheric and hydrological
boundary conditions described in cites....

FishTank, parameters and parameter categories, how they compare to other models

2.2 Local sensitivity analysis

Methods from Reichert and Vanrolleghem (2001), Soetaert and Petzoldt (2010)

Sensitivity of the model to parameter changes is evaluated using the difference in model results before and after perturbing each parameter value. Each parameter is perturbed by the same value as a percentage of the whole. The default value (as 1 + 1e-8 proportion, default value for 'tiny' argument in 'sensFun'). A sensitivity value S is estimated for each time step i given a set value for parameter j as:

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

where the estimate depends 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, are set as the default value of the unperturbed parameter and the predicted value of y_i after perturbation (Soetaert and Petzoldt 2010). The scaling ensures the estimates are unitless such that the relative magnitudes provide a comparison for model sensitivity to parameter changes that may vary in scale. The FME package summarizes sensitivity as L1 and L2 across the time series:

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

$$L2 = \sqrt{\sum \left(S_{ij}^2\right)/n} \tag{3}$$

The mean, minimum, and maximum S_{ij} values for each parameter are also reported. In general, positive mean sensitivity estimates generally indicate that a parameter has a positive effect on the model results for a given increase in the parameter. However, the effect can change over time so a plot of the 'sensitivity function' should be viewed which shows the difference from 117 the values of the response variable before and after changing a parameter value. Note that the perturbation factor in 'sensFun' depends on the default value of each parameter such perturbations are not consistent for parameter values less than or greater than the perturbation factor. This can produce results that are inconsistent for different levels of perturbation. A custom function was keeps the perturbation factors constant regardless of the magnitude of the default values. Sensitivity for each parameter using the custom function are estimated using the above equations from the FME package. All parameters that are considered 'sensitive' should be further evaluated by plotting the predictions before and after perturbation and across the total range of the parameter.

Each parameter was perturbed by 50% using 'sensfun' to identify model sensitivity. Parameters that produced different model results from the default values are shown below, by category.

The parameters that have the greatest effect on the model by category (optics, organics, phytoplankton, zooplankton) are as follows.

Plotting the raw values from the sensitivity analysis provides a visual assessment of changes.

2.3 **Identifiability**

115

116

119

120

121

122

124

125

126

127

129

130

131

132

133

134

135

137

138

139

Identifiability describes the ability to estimate a parameter in relation to variation among the remaining parameters. A parameter is identifiable if all parameters within the set can be uniquely estimated based on the observed data. Parameters that are unidentifiable typically produce similar model outputs for a given relative perturbation, i.e., the effect of altering one parameter can be undone by altering one or more other parameters. Model calibration will not converge for parameters sets that are unidentifiable. Identifiability is estimated from the minimum eigenvector of the cross-product of a model's sensitivity matrix (Brun et al. 2001, Omlin et al. 2001):

$$\gamma = \frac{1}{\sqrt{\min\left(\text{EV}[\hat{S}^{\dagger}\hat{S}]\right)}} \tag{4}$$

where γ ranges from one to infinity for perfectly identifiable (orthogonal) or unidentifiable (perfectly collinear) results for a set of parameters in the sensitivity matrix S. Identifiability can be estimated for any combination of parameters for a model. Values less than 10-15 indicate parameters are generally identifiable. The FME package provides the 'collin' function to estimate identifiability of output from the 'sensFun'. For reasons described above, a custom function was created to determine identifiability from model residuals, where the residuals were based on a relative perturbation of the model parameters. A comparison of results of identifiability using the custom functions and those from the FME package were generally in agreement. Standard output from the identifiability function is shown below, where identifiability is displayed as a matrix with increasing combinations of subsets of the parameters.

The identifiability functions evaluate the ability to identify all subsets from pairwise to all parameters in the subset. As an example, the identifiability of pairwise combinations for each of the categories are shown below.

Pairwise identifiability of the top two most sensitive parameters in each category is shown, first as a table showing all unique combinations from two to all parameters and second as a figure showing pairwise combinations.

2.4 Identifying subsets and testing calibration

Model calibration must consider the competing objectives of parameter inclusion and identifability. Estimating parameters for novel datasets increases model complexity and the ability to identify parameters decreases with the inclusion of parameters to estimate. The above analyses indicated that estimating all of the parameters was impossible because of redundancies in the model output. Calibration to novel datasets should balance the competing objectives of identifiability while including parameters for which model output is most sensitive. The plots below show different scenarios of parameter inclusion for model calibration. Reading each plot from left to right can be interpreted as including additional parameters, where each parameter is

ranked by relative sensitivity. The inset in each plot shows the identifiability of including parameters, up to a maximum where additional inclusion exceeds the identifability threshold of 169 fifteen. The scenarios for including parameters all begin with the parameters that have the greatest 170 effect on the model output. The inclusion of additional parameters depends on the scenario. The 171 first scenario selected parameters by decreasing sensitivity within each category (i.e., four 172 separate models calibrated for optics, organics, phytoplankton, or zooplankton), the second 173 scenario selected parameters by sensitivity regardless of category, and the third scenario selects 174 parameters by sensitivity with equal representation between categories. Boxes in each plot 175 represent a unique parameter and the numbers in each box represent the relative rank of a 176 parameter's sensitivity to the model output within each category. 177

The following plots show identifiability with the addition of parameters.

3 Results

178

181

182

183

184

185

186

187

188

189

190

191

192

193

194

195

196

4 Discussion

Questions specific to GOM - what initial conditions are important? How many phytoplankton groups do we need (e.g., related to structural uncertainty)?

How does the assimilation of additional parameters (e.g., other state variables) during calibration influence the conclusions?

How does uncertainty translate to what a model should provide (generality v precision)? The first step - find out what can be optimized but then do not overfit....

What about structural uncertainty - does sensitivity of a model to variation in a parameter imply parameter uncertainty and/or structural uncertainty?

A final point about optimization with identifiable parameter sets - optimization to fit the data still does not ensure a correct model. Failing in one way can be over-compensated by another feature, e.g., the parameter set that is optimized (see Flynn (2005), p. 1207, third paragraph)

Omlin et al. (2001) state that the sensitivity, identifiability, estimation process is iterative (p. 113), need to rinse and repeat for proper calibration.

How to improve identifiability - get more/better observed data, include obs from other state variables in RSS minimization (eqn q in Omlin et al. (2001))

Alternative methods for uncertainty analysis - bayesian, MCMC, nonlinear

calibration-constrained optimization (Gallagher and Doherty 2007)

¹⁹⁸ References

- Beck MB. 1987. Water quality modeling: A review of the analysis of uncertainty. Water Resources Research, 23(8):1393–1442.
- Bianchi TS, DiMarco SF, Jr JHC, Hetland RD, Chapman P, Day JW, Allison MA. 2010. The science of hypoxia in the Northern Gulf of Mexico: a review. Science of the Total Environment, 408(7):1471–1484.
- Brun R, Reichert P, Künsch HR. 2001. Practical identifiability analysis of large environmental simulation models. Water Resources Research, 37(4):1015–1030.
- Eldridge PM, Roelke DL. 2010. Origins and scales of hypoxia on the Louisiana shelf: importance of seasonal plankton dynamics and river nutrients and discharge. Ecological Modelling, 221(7):1028–1042.
- Estrada V, Diaz M. 2010. Global sensitivity analysis in the development of first principle-based eutrophication models. Environmental Modelling and Software, 25:1539–1551.
- Fasham MJR, Flynn KJ, Pondaven P, Anderson TR, Boyd PW. 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-Sea Research I, 53:333–366.
- Fennel K, Hu J, Laurent A, Marta-Almeida M, Hetland R. 2013. Sensitivity of hypoxia predictions for the norther Gulf of Mexico to sediment oxygen consumption and model nesting.

 Journal of Geophysical Research: Oceans, 118(2):990–1002.
- Flynn KJ. 2005. Castles built on sand: dysfunctionality in plankton models and the inadequacy of dialogue between biologists and modellers. Journal of Plankton Research, 27(12):1205–1210.
- Gallagher M, Doherty J. 2007. Parameter estimation and uncertainty analysis for a watershed model. Environmental Modelling and Software, 22(7):1000–1020.
- Ganju NK, Brush MJ, Rashleigh B, Aretxabaleta AL, del Barrio P, Grear JS, Harris LA, Lake SJ, McCardell G, O'Donnell J, Ralston DK, Signell RP, Testa JM, Vaudrey JMP. 2016. Progress and challenges in coupled hydrodynamic-ecological estuarine modeling. Estuaries and Coasts, 39(2):311–332.
- Lehrter JC, Ko DS, Lowe L, Penta B. In review. Predicted effects of climate change on the severity of northern Gulf of Mexico hypoxia. In: Justic et al., editor, Modeling Coastal Hypoxia: Numerical Simulations of Patterns, Controls, and Effect of Dissolved Oxygen Dynamics. Springer, New York.
- Levins R. 1966. The strategy of model building in population biology. American Scientist, 54(4):421–431.

- Lohrenz SE, Redalje DG, Cai WJ, Acker J, Dagg M. 2008. A retrospective analysis of nutrients and phytoplankton productivity in the Mississippi River plume. Continental Shelf Research, 28(12):1466–1475.
- Martin PJ. 2000. Description of the navy coastal ocean model version 1.0. Technical Report NRL/FR/7322-00-9962, Naval Research Lab, Stennis Space Center, Mississippi.
- Mateus MD, Franz G. 2015. Sensitivity analysis in a complex marine ecological model. Water, 7:2060–2081.
- Morrison M, Morgan MS. 1999. Models as mediating agents. In: Morgan MS, Morrison M, editors, Models as Mediators, page 401. Cambridge University Press, Cambridge.
- Omlin M, Brun R, Reichert P. 2001. Biogeochemical model of Lake Zürich: sensitivity, identifiability and uncertainty analysis. Ecological Modelling, 141(1-3):105–123.
- Pauer JJ, Feist TJ, Anstead AM, DePetro PA, Melendez W, Lehrter JC, Murrell MC, Zhang X, Ko
 DS. 2016. A modeling study examining the impact of nutrient boundaries on primary
 production on the Louisiana continental shelf. Ecological Modelling, 328:136–147.
- Reichert P, Vanrolleghem P. 2001. Identifiability and uncertainty analysis of the River Water Quality Model No. 1 RWQM1. Water Science and Technology, 43(7):329–338.
- Snowling SD, Kramer JR. 2001. Evaluating modelling uncertainty for model selection. Ecological Modelling, 138:17–30.
- Soetaert K, Petzoldt T. 2010. Inverse modelling, sensitivity, and Monte Carlo analysis in R using package FME. Journal of Statistical Software, 33(3):1–28.
- Wiseman WJ, Rabalais NN, Turner RE, Dinnel SP, MacNaughton A. 1997. Seasonal and
 interannual variability within the Louisiana coastal current: stratification and hypoxia. Journal
 of Marine Systems, 12(1-4):237–248.

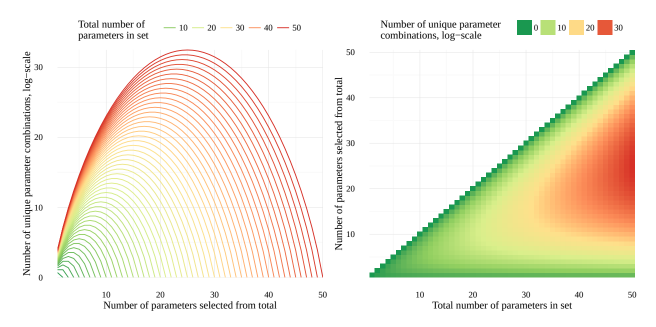


Fig. 1: Number of unique combinations that can occur by selecting parameters for optimization from different parameter sets. The number of combinations are shown for increasing numbers of selected parameters from the total in the set, where 100 parameter sets are shown each with one through 100 total parameters. For example, there are six unique parameter combinations that result from choosing two parameters from a set. Note that the number of unique combinations is shown as the natural-log. The same information is viewed differently in each plot.