Forecasting hypoxia in the Chesapeake Bay and Gulf of Mexico: model accuracy, precision, and sensitivity to ecosystem change

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Abstract

Increasing use of ecological models for management and policy requires robust evaluation of model precision, accuracy, and sensitivity to ecosystem change. We conducted such an evaluation of hypoxia models for the northern Gulf of Mexico and Chesapeake Bay using hindcasts of historical data, comparing several approaches to model calibration. For both systems we find that model sensitivity and precision can be optimized and model accuracy maintained within reasonable bounds by calibrating the model to relatively short, recent 3 year datasets. Model accuracy was higher for Chesapeake Bay than for the Gulf of Mexico, potentially indicating the greater importance of unmodeled processes in the latter system. Retrospective analyses demonstrate both directional and variable changes in sensitivity of hypoxia to nutrient loads.

Keywords: model-data comparison, coastal systems, nitrogen loading, eutrophication

1. Introduction

Ecological models are increasingly moving from heuristic to applied, and this movement requires rigorous analysis and optimization of accuracy, precision, and sensitivity to system change. Ecological systems are subject to sporadic changes caused by internal dynamics (Bronmark \textit{et al} 2010), shifts in drivers (climate (Scheffer and van Nes 2007), human inputs (Goosby \textit{et al} 2001, Rabalais \textit{et al} 2002a)), invasive species (Higgins and Zanden 2010), and other factors. Some of these changes can be included in models explicitly, but others are beyond the scope of most modeling activities. These unmodeled changes and processes are generally parameterized through key model coefficients, and because systems change, those parameterizations are subject to change, therefore it is important for model calibrations to reflect the current state of the system.

Ecosystems are also subject to relatively high ‘random’ short-term variability (e.g. weather) that does not necessarily reflect directional change. Robust model parameterization thus also requires sufficiently long time frames to capture the range of system variability to both detect mean behavior and undertake reasonable uncertainty analysis. There is a potential tension between the goals of providing high accuracy and high precision and between the challenges of incorporating information about both random variability and long-term system changes. So, it is important to develop model calibration approaches that optimize model performance (accuracy, precision) in the face of systems that are both undergoing directional change and are highly variable.

Models of varying degrees of complexity have been informative tools in understanding the controls on hypoxia occurrence in river-impacted coastal areas (Peña \textit{et al} 2010). Hypoxia, low oxygen concentrations in bottom waters, occurs...
when decomposition rates exceed those of oxygen diffusion and mixing. Hypoxia is a widespread and increasing phenomenon (Diaz and Rosenberg 2008, Zhang et al 2010) that can lead to widespread ecosystem changes including altered biogeochemical cycles (Kemp et al 2005, Turner et al 2008), fish kills (Diaz and Rosenberg 2008), decreased or displaced fish production (Rabalais and Turner 2001), and decreased value to human use through recreation and fisheries harvest losses (Renaud 1986).

Two major river-impacted coastal hypoxic areas of the United States occur in the Gulf of Mexico (GOM) along the Louisiana–Texas coasts and in Chesapeake Bay (CB). Hypoxia has been heavily studied in these areas (Justić et al 1993, Bierman et al 1994, Rabalais et al 1994, 1998, Boesch et al 2001, Rabalais and Turner 2001, Hagy 2002, Rabalais et al 2002a, 2002c, Childs et al 2002, Rabalais et al 2004, Kemp et al 2005, Rabalais 2006, Walker and Rabalais 2006, Scully 2010, etc), due in part to concern over potential fisheries impacts (Renaud 1986, Rabalais and Turner 2001), and management goals have been set to limit hypoxia severity. Models have been used successfully in both systems to explore the underlining causes of hypoxia and to make specific management recommendations (Cerco and Cole 1993, Rabalais et al 2002b, Justić et al 2003, Scavia et al 2003, Hagy et al 2004, Turner et al 2005, Scavia et al 2006, Turner et al 2006, Justić et al 2007, Rabalais et al 2007, Turner et al 2008, Greene et al 2009, Penta et al 2009, Wang and Justić 2009, Bianchi et al 2010, Liu et al 2010, Liu and Scavia 2010, Peña et al 2010). Models and empirical data indicate that hypoxia in these systems is caused by a combination of nutrient-driven, mostly nitrogen, production of phytoplankton organic matter; decomposition; freshwater-driven stratification of the water-column; and storm mixing. Management recommendations have generally focused on control of nitrogen loading to these systems due to evidence that it is an important driver of hypoxia and its susceptibility to management compared to other drivers. However, phosphorus load control has also been addressed (Boesch et al 2001, Environmental Protection Agency (EPA) Science Advisory Board (SAB) 2007, Mississippi River/Gulf of Mexico Watershed Nutrient Task Force 2008).

Both systems have also undergone significant ecosystem changes in hypoxia sensitivity to nutrient loads over the last 30 years, such that in both systems the severity of hypoxia for a given nitrogen load is now approximately twice what it was in the early 1980s (Hagy et al 2004, Turner et al 2008, Liu et al 2010, Liu and Scavia 2010). Ongoing research and management scenarios are thus complicated by the need to account for this varying ecosystem sensitivity and by speculation about how the systems will respond as nutrient loads change. Shifts in system sensitivity can appear abrupt when viewed retrospectively (Hagy et al 2004, Environmental Protection Agency (EPA) Science Advisory Board (SAB) 2007, Turner et al 2008, Greene et al 2009, Liu et al 2010), however, because of significant interannual variability, they can be impossible to recognize contemporaneously. This delayed recognition of sensitivity change is a challenge to both short- (annual) and long- (management scenarios) term results, and highlights the need for models and model calibration approaches that optimize model performance in changing and highly variable systems.

In this study we test different model calibration approaches for fitting similar models of the GOM and CB to subsets of historical data that include system changes and periods of high variability. A wide range of modeling approaches, from simple regressions to 3D coupled hydrodynamic–biogeochemical and earth system models have been applied to hypoxia for both management and scientific investigation (Peña et al 2010). More complex models are generally able to resolve finer scale ecological mechanisms and provide process based insight. Simpler models, however, are often better predictors of system state and have proven very useful for management applications (Peña et al 2010). Within this range we use a relatively simple, mechanistically based, model that treats estuary and coastal currents as ‘rivers’ with point source organic matter loads. We selected this model because it has proven useful for management guidance and because the computational simplicity allows the explicit incorporation of uncertainty analysis (Scavia et al 2003, 2004, 2006, Scavia and Donnelly 2007, Stow and Scavia 2009, Liu et al 2010, Liu and Scavia 2010). For a description of this model’s use in the GOM in the context of other modeling approaches, see the recent review by Peña et al (2010).

We test for accuracy, precision, and model sensitivity to system changes by hindcasting parts of the historical dataset. We then compare optimal model calibrations between these two systems and discuss its implications for both ecological interpretation and management. Finally, we use our optimal calibrations to forecast outcomes under different nutrient reduction scenarios.

2. Methods

2.1. Models

We use versions of the Streeter–Phelps (SP) river model (Chapra 1997) developed for CB and the GOM. The model is described in greater depth and its assumptions justified in earlier publications (Scavia et al 2003, 2004, 2006, Scavia and Donnelly 2007, Stow and Scavia 2009, Liu et al 2010, Liu and Scavia 2010). These models share the same basic structure but are adapted to each system. Both models treat the estuary or coastal current as a ‘river’ and calculate longitudinal profiles of dissolved oxygen (DO) concentration downstream of an organic matter (BOD) point source (described below for each system). This organic matter point source is assumed to be proportional to the spring total nitrogen (TN) loading to the system with a proportionality constant equal to the product of the Redfield carbon to nitrogen ratio, the respiration ratio of oxygen consumption per organic carbon, and the dilution of inputs within the receiving water body. Spring TN loads were used because spring loads are the dominant drivers of hypoxia in these systems (Cerco 1995, Scavia et al 2003, Hagy et al 2004, Turner et al 2006).
DO profiles are calculated at steady state, for each location along the profile, DO is calculated by:

\[
DO = DO_0 - \frac{k_d BOD_u F}{k_i - k_d} (e^{-k_i x} - e^{-k_d x}) - D_i e^{-k_i x} \tag{1}
\]

where: \(DO = \) dissolved oxygen (mg l\(^{-1}\)), \(DO_0 = \) oxygen saturation (mg l\(^{-1}\)), \(k_d = \) BOD decay coefficient (1/day), \(k_i = \) reaeration coefficient (1/day), \(BOD_u = \) initial BOD (mg l\(^{-1}\)), \(x = \) downstream distance (km), \(F = \) fraction of BOD sinking below the pycnocline (unitless), \(D_i = \) the initial oxygen deficit (mg l\(^{-1}\)), and \(v = \) net downstream advection (km/day). While in the original SP formulation, \(v\) represents net downstream advection, in this application it also parameterizes the combined effect of horizontal transport and subsequent settling of organic matter from the surface. Therefore, it has no simple physical analog.

The length of the hypoxic zone is summed across the part of the profile with DO at hypoxic levels and converted to a measure of hypoxic area or volume by empirical relationships developed from measurements of the hypoxic area or volume in each system (see below). The model was calibrated by fitting predicted and measured area or volume and minimizing error terms. During calibration, each parameter can be assumed to be either constant across all years or adjusted each year. If a parameter is adjusted each year, we assume that its variability includes the effects of all unmodeled processes.

As in prior applications to CB and the GOM (Stow and Scavia 2009, Liu et al. 2010, Liu and Scavia 2010), the model was calibrated using Bayesian fitting through Markov Chain Monte Carlo methods (Lunn et al. 2000, Gill 2002, Gelman and Hill 2007). All model calibration was conducted in WinBUGS (version 1.4.3), called through R (version 2.6.0, R2WinBUGS, version 2.1-8), using the same methods and inputs described elsewhere (Stow and Scavia 2009, Liu et al. 2010, Liu and Scavia 2010). In prior applications of both models, either \(v\) or \(F\) was allowed to vary by year, and all other parameters were fit as constants across years or determined from empirical data (see below).

Model application to the two systems differed in four ways:

1. The location of the organic matter point source was determined by the geography and physics of each system. In CB, summer surface waters flow seaward and bottom waters flow landward. The primary nutrient input to the modeled area of CB is the Susquehanna River at the head of the bay and most hypoxia occurs in the mid-bay region. Thus, the model origin and organic matter point source were assigned to the lower end of the mid-bay region (220 km down bay from the Susquehanna River mouth) and distance in the model is following the landward flowing bottom water. Organic matter loading was based on Susquehanna River spring TN loading. In the GOM, hypoxia occurs below a westward flowing coastal current along the Louisiana and Texas coasts. Because there are two main nutrient inputs to the GOM (the Mississippi and Atchafalaya Rivers), we model two organic matter point sources, one at the model origin (Mississippi River) and one at 220 km down current (Atchafalaya River). Organic matter is proportional to spring TN load with 50% of the Mississippi River and 100% of the Atchafalaya River TN load assumed to be entrained in the westward current.

2. The initial oxygen deficit (\(D_i\)) was assumed to be 0 in the GOM because there is little oxygen depletion in waters east of the delta. \(D_i\) in CB was estimated each year based on measured bottom-water oxygen concentrations at the model origin and a stochastic term based on measurement variation.

3. In CB, the reaeration coefficient is known to vary with distance down estuary (Hagy 2002). Our model uses this observed variation in distance (\(x\)) and calculates \(k_i = b_x K\) where \(b_x\) is a location specific constant accounting for spatial variation (Scavia et al. 2006) and \(K\) is a fit model parameter scaling reaeration.

4. In CB, the volume of water with \(DO < 2\) mg l\(^{-1}\) is determined each year, so the model hypoxia cutoff is set to 2 mg l\(^{-1}\) when determining length (\(L\)), and volume (\(V\)) is calculated using the empirical relationship \(V = 0.00391 L^2\) (Scavia et al. 2006). In the GOM, hypoxia is reported as the area of hypoxic bottom water, with measurements taken just above the sediment water interface. Because the model simulates the entire sub- pycnocline layer and because available DO profiles show that when near-bottom DO is 2 mg l\(^{-1}\), average sub-pycnocline DO approaches 3 mg l\(^{-1}\), the GOM model hypoxia cutoff is set to 3 mg l\(^{-1}\). Hypoxic area (\(A\)) is calculated using the empirical relationship \(A = 38.835 L\) (Scavia and Donnelly 2007).

2.2. Data sources

We use spring total nitrogen (MT TN/d) loading data from the USGS to drive both models. Average January through May TN loads from the Susquehanna River (at Conowingo, MD gauging station) are used for CB and May TN loads from the Mississippi (at St Francisville) and Atchafalaya (at Melville) Rivers are used for the GOM (USGS 2007, 2009, 2010).

Model calibrations and tests are conducted using empirically measured hypoxic area (GOM) or volume (CB). GOM hypoxic area has been interpolated from near-bottom DO measurements collected by shelf-wide cruises in late July or early August (Rabalais et al. 2002b, Rabalais 2009). Cruises have been conducted yearly since 1985 with the exception of 1989. Because the measured hypoxic area was potentially impacted by tropical storms in 1996, 1998, 2003, and 2005, we removed these years from both our calibration and test datasets (see Turner et al. 2008) because the model is incapable of accounting for these extreme conditions. Such tropical storms can disrupt water-column stratification, mixing oxygenated water downward, and thus temporarily break the link between production and hypoxia observed in non-storm years. CB hypoxic volume is determined from DO profiles taken on four cruises in July and August each year since 1984 and sporadically before then (Hagy et al. 2004, Chesapeake Bay Program (CBP) 2008). We use the July CBP cruise data from the consistent record since 1984 for model calibration and testing.

Initial oxygen deficit (\(D_i\)) for CB was based on average July bottom-water oxygen concentration measured at stations in the mid-bay region (Scavia et al. 2006, Chesapeake Bay...
2.3. Hindcast and forecast tests

We tested several model calibration algorithms to optimize model performance. Each test used a calibration dataset and a test dataset. Model performance was measured by assessing precision, accuracy, robustness, and sensitivity to system change, although some tests focus on a subset of these measures. Precision was assessed as the size of the coefficient of variation and the 95% credible interval (CI) of the model prediction. Accuracy was assessed as the percentage of observations in the test dataset that fell within the 95% CI of the model prediction. It is expected that this value could differ from 95% because the test dataset contains observations for years that are not included in the model calibration and thus the model is predicting outside its statistical sample. Robustness was based on the impact of individual years in the calibration data set on calibrated parameter values. Sensitivity to system change was assumed to be maximized when few, recent years of calibration data were used because averaging across larger numbers of data points decreases the impact of any given point on the average.

To test the impact of increasing the number of years in the calibration set on model precision and robustness (Test 1), calibrations began with the first three years of data and we progressively added years for successive calibrations (calibration dataset). To test precision, we used each calibrated model to predict hypoxia for each year in the full dataset (test dataset) and the average CV and 95% CI were calculated. To test robustness, we examined variation in parameter values over time from each calibration test. Model accuracy was quantified by calculating the percentage of observed hypoxic areas or volumes that fell within the model’s 95% CI for that year’s prediction. We repeated this test (Test 2) beginning with the three most recent years and adding years in reverse order.

Because the above comparisons are confounded by overlap between the calibration and test datasets, they were used only to narrow the range of years for which a more complete test was conducted. In these tests of accuracy, forecasts were conducted using 3, 5, and 7 year windows of calibration data (Test 3, range selected based on the results of Tests 1 and 2, see below). Precision was quantified by the CV of the hypoxia forecast in the year following each calibration and the average of these CVs across all calibrations using the same window size. The accuracy of these calibrations was tested by forecasting hypoxia in the year following each calibration window and calculating the percentage of observed hypoxic areas or volumes that fell within the forecast’s 95% CI across all calibrations using that window (test dataset).

To prepare for forecasts where all model coefficients are to be held constant, we tested two methods of parameter calibration. In prior work with this model (Liu et al 2010, Liu and Scavia 2010), \( v \) was estimated as the year-specific term \( v_i \) and then forecasts used the mean and standard deviation of \( v_i \) through time, ignoring the Bayesian fit parameter distribution. We compared this approach with one that estimated all parameters (including \( v \)) as constant distributions through time such that forecasts could be conducted directly from the calibrated distributions (Test 4). For both methods (using \( v_i \) and \( v \)) in CB, the parameter \( D_i \) which is not calibrated but calculated from observed values for calibration years, was estimated using the average and variation in the previously observed values.

2.4. Response curves

Response curves of predicted hypoxic area or volume versus spring TN load, were constructed in the same way as hindcasts and forecasts but using evenly spaced spring TN loads spanning the observed range rather than exact historical loads (Scavia et al 2006, Liu et al 2010, Liu and Scavia 2010). As in prior publications (Liu et al 2010, Liu and Scavia 2010), response curves were constructed using 50% CIs to better constrain conditions in typical or average years.

3. Results

3.1. Full dataset calibration

When the model is fit to the entire CB and GOM datasets, allowing \( v_i \) to vary in each year and then averaging \( v_i \) for hindcasts, hindcasting accuracy is high (100% and 80% of the observations are within the 95% CI of the hindcast, respectively, see right-most symbols associated with the full dataset in figures 1(a) and (c)). These percentages differ from 95% because of additional variability added to the model when the parameter \( v_i \), and the parameter \( D_i \) in the CB model, are averaged across years. Model precision, as measured by the CV of the predicted hypoxic region, is better for CB (33%) than for the GOM (41%) (see right-most symbols associated with the full dataset in figures 1(a) and (c)). Model parameters have mean values of \( F = 0.91, k_d = 0.14, K = 0.58, v_A = 2.5 \) (where \( A \) indicates the average across years), \( D_A = 1.2 \) for CB; and \( F = 0.51, k_d = 0.006, k_i = 0.012, v_A = 0.64 \) for the GOM (see right-most symbols associated with the full dataset in figures 1(b) and (d)). These values are consistent with previously published model calibrations and, as in prior calibrations, estimated process rates based on these parameters are consistent with observed rates (Scavia et al 2006, Scavia and Donnelly 2007, Liu et al 2010, Liu and Scavia 2010).

3.2. Effect of number of years in the calibration

The precision and accuracy of the models calibrated to the full datasets represent a goal for calibrations using sub-datasets. However, they may not represent the best overall model calibration because using the full dataset ignores temporal trends and regime shifts within the system and thus sacrifices model sensitivity to system change. It also confounds calibration and test datasets, causing a possible overestimation of model accuracy in predicting novel conditions. So, we tested model performance by calibrating to subsets of data...
with increasing number of years. Beginning at the start of each dataset (Test 1), adding years causes a rapid improvement for the CB model performance in both precision and accuracy until about 5 years of data are used (figure 1(a)). Beyond this point, precision and accuracy asymptote toward values of the full dataset. Similarly, CB parameter values were highly variable in the beginning until about 5 years of data were used (figure 1(b)). Beyond 5 years, there was little change in values, despite known changes in system behavior, indicating that model calibration to long datasets loses sensitivity to these changes. These same patterns of precision, accuracy, and parameter variability were observed when calibration was started using the three most recent years and adding years in reverse order (Test 2, data not shown). Thus, using about 5 years of calibration data seems to offer an optimal combination of model precision, accuracy, and robustness, while avoiding a loss of sensitivity to system change in this system.

Determining optimal calibration for the GOM involves a greater compromise between precision and sensitivity. As in CB, adding calibration years (Test 1) causes a rapid improvement in model precision and accuracy (figure 1(c)). However, precision and accuracy continue to improve until about 15 years of calibration data. The model also maintained more sensitivity to system change with the addition of calibration years for longer datasets than the CB model (figure 1(d)). Parameter values, especially \( F \) and \( k_r \), continue to change as years are added up to at least 15 years of calibration data. When calibration started using the three most recent years and years were added in reverse order (Test 2, data not shown), model accuracy was high (100% of observations in the 95% CI) using even 3 years of data and remained at this level as years were added. Model precision improved quickly until about 5 years and then saturated, and parameter values varied in a similar pattern. Though results were more mixed.

**Figure 1.** (Test 1) Average CV of predicted hypoxic volume or area and % observations in 95% CI using (a) a test dataset of Chesapeake Bay hypoxic volume from 1984–2008 and calibration datasets of increasing numbers of years starting in 1984 and (c) a test dataset of Gulf of Mexico hypoxic area from 1985–2009 and calibration datasets of increasing numbers of years starting in 1985. Parameter estimates (panels (b) and (d); means, with 50% CI bars for \( v_A \)) corresponding to the model calibrations used in panels (a) and (c). Note that results from calibrating to the full datasets are shown as the right-most symbols in each graph.
for the GOM, with continued improvement in model precision and accuracy beyond the first 5 years of calibration data in the forward though not the backward calibration tests, we decided to further test models using 5 years of calibration data because using 15 or 20 years lost sensitivity to system changes which have been observed on shorter time scales (Turner et al. 2008, Liu et al. 2010).

3.3. Moving window calibrations: case 1—averaging $v_i$

Previous applications of this model estimated $v_i$ for each year in a calibration dataset and then averaged it for forecasts. So we first test the moving window calibrations (Test 3) with this method and then compare it below to the case where $v$ is estimated as a constant over the calibration widow period (Test 4). Tests with 3, 5, and 7 year moving windows showed little difference in precision (CV) when forecasting CB (figure 2(a)) or GOM (figure 3(a)) hypoxia in the year following the calibration window and no overall improvement in precision using larger windows. A change in precision could indicate over or under fitting the model, but this does not seem to be taking place. For all window sizes, parameter values changed over time; however, parameter variability was highest using the smallest (3 year) window (figures 2(b) and 3(b)). This increased variability indicates higher model sensitivity to system state because more of the underlying variability is reflected in the parameters. Model accuracy was high for all window sizes in CB and decreases with window size in the GOM. In CB the per cent of observations in the 95% CI is 100%, 95%, and 100% for the 3, 5, and 7 year calibration periods, respectively, compared to 93%, 80%, and 73% for these calibrations in the GOM. Thus, the results support the use of a 3 year moving window.
Compared to using the full dataset, 3 year moving window calibrations resulted in the same accuracy for CB (100%) and improved accuracy for GOM (95% versus 80% of observations within the 95% CI). Average model precision (CV) for the 3 year moving window calibration was slightly poorer in CB (39% versus 33%), but was slightly improved in the GOM (30% versus 41%). Using a moving window allows the model precision to vary over time based on recent system variability. Precision is higher (lower CV) during periods of system stability, such as the late 1990s in the GOM, and lower (higher CV) following regime shifts (figure 3(a)).

### 3.4. Moving window calibrations: case 2—constant $v$

Fitting $v_i$ for each year and then averaging it for forecasts introduces arbitrary variation into the model. As an alternative, we tested moving window calibrations of 3, 5, and 7 years fitting all parameters, including $v$, as constants (Test 4).

With 3, 5, and 7 year moving window calibrations, CB hypoxia forecast accuracy is lower than expected. Accuracy is highest for the 3 year window and decreases with increasing window size (82%, 70%, and 68% of observed hypoxic volume were within the model 95% CI, figure 4(c) compares forecast and observed hypoxia for the 3 year moving window calibration). There was very little difference in precision (CV) among window sizes (figure 4(a)) and no overall improvement in precision using larger windows. For all window sizes, parameter values changed over time; however, as in prior tests, parameter variability was highest using the smallest (3 year) window (figure 4(b)), indicating the highest model sensitivity to system state.

Test for the GOM resulted in lower accuracy, with 73%, 68%, and 46% of the observed hypoxic areas within the model's 95% CI for 3, 5, and 7 year windows, respectively (figure 5(c) compares forecast and observed hypoxia for the 3 year moving window calibration). There was very little difference in precision (CV) among window sizes (figure 5(a)), no overall improvement in precision using larger windows, and parameter values changed over time with the highest variability associated with the smallest (3 year) window (figure 5(b)).

The CV of predicted hypoxic area or volume varies with time in all moving window calibrations. However, the average
CV is improved by calibrating with a constant \( v \) in both systems. Using the 3 year window, the CV for CB is improved from 39% to 33% and for GOM from 30% to 18%, compared to moving window tests averaging \( v_i \). This is a substantial improvement in model precision. This increase in model precision is accompanied by a decrease in model accuracy. However, because the increased variability introduced into the model by averaging year-specific \( v_i \) is not related to a specific mechanism or known ecological process, the lower forecasting accuracy is likely a better representation of true model performance. The SP model is a vast simplification of nature and the accuracy cost of using this model (95% − 82% = 13% for CB and 22% for the GOM) reflect unmodeled variation in these systems.

4. Discussion

The dataset for the GOM included years in three distinct and previously observed system states with varying sensitivity to hypoxia formation (Environmental Protection Agency (EPA) Science Advisory Board (SAB) 2007, Turner et al 2008, Greene et al 2009, Liu et al 2010). Similar changes in system state have been observed in CB, however, data limitations prevented us from including the historic CB system state (Hagy et al 2004, Liu and Scavia 2010) in the current model tests. Model accuracy was poorer for the GOM than for CB and one of the reasons could be the attempt to calibrate the model across multiple system states. Including multiple states is minimized when using short calibration windows and improved model accuracy in the shortest windows are a result. GOM model accuracy for the 3 year moving window calibration, fitting \( v \) as a constant, is further improved to 78% when excluding calibrations that overlap more than one system state. Though such exclusions can only be identified post facto, this further supports use of shorter calibration windows to minimize including multiple system states.

Initial comparisons, adding years to the calibration dataset, starting from the oldest (figure 1) or most recent (data not shown) data, indicated that 5 or more years...
of calibration data were needed to optimize precision and accuracy. However, more extensive comparisons using 3, 5, and 7 year calibration datasets and employing more robust accuracy measures (by forecasting only the year following a calibration window) show that model performance is optimized with 3 year calibrations. These tests showed less of a tradeoff between model precision and accuracy than was expected. Model precision did not differ among window lengths in any comparison and in fitting either year-specific or constant $v_i$. Model accuracy was either higher in the shortest window or did not change with window size. As expected, the shortest window calibrations were the most adaptive to system change. This responsiveness is seen in both the improved model accuracy in the changing GOM and in increased parameter variability in both systems.

The tradeoff between precision and accuracy existed for both models calibrated using year-specific or constant $v_i$. We believe that fitting and averaging year-specific $v_i$ introduces an artifact that improves model accuracy by arbitrarily reducing precision. Thus, we propose that the optimal calibration for annual forecasting is to use a rather short (3 year) recent dataset treating all parameters as constants.

### 4.1. Forecasts

Using the 3 year window calibrations, we developed load–hypoxia response curves for CB and the GOM for different periods in the historical record (figure 6). The GOM has undergone two shifts in sensitivity between the earliest (1986–8) and most recent (2005–7) calibration periods, and the resulting increase in sensitivity can be seen across the TN range.

Though the primary CB regime shift appears to have occurred before the start of our dataset (Hagy et al. 2004, Kemp et al. 2005, Scavia et al. 2006, Conley et al. 2009), CB appears to be undergoing a gradual increase in sensitivity to nutrient loads from 1983 through at least 2005 (Liu and Scavia 2010, Scully 2010). This increasing sensitivity is reflected in our parameter estimates (figure 4(b)) and in response curves using 1988–90 and 2002–4 calibration datasets (figure 6). Between these periods, hypoxia sensitivity increased, especially at high TN loads. Parameter values for recent years trend back toward, and even beyond, those earlier in the dataset. Accordingly, the most recent (2005–2007) response curve shows decreased sensitivity. At high TN loads, the curve resembles the 1988–90 ‘low sensitivity’ case, and it appears to have even lower sensitivity at lower TN loads.

These changes are driven mostly by changes in the parameters $F$, $v$, $K$, and $D_i$. $F$ and $v$ increase between the first two periods and then decrease again before the final period. Sensitivity analyses (not shown) indicate that increases in $F$ tend to increase sensitivity at all TN loads while increases in $v$ increases sensitivity more at high TN loads. At the same time $K$ decreased between the first two time periods while remaining relatively unchanged between the second and third. Like increases in $v$, decreases in $K$ tend to increase sensitivity at high TN loads. Finally, the measured parameter $D_i$ remained constant between the first two periods but decreased between the second and third. Decreases in $D_i$ lead to decreases in sensitivity at low TN loads. This measured decrease in DO deficit in recent years may indicate a release from oxygen stress further down bay.

Using models calibrated with the three most recent years, or in the case of the GOM the three most recent years that were not impacted by severe tropical storms, provides a consistent method for annual forecasts that is relatively robust to regime shifts and changes in system sensitivity. However, changes in system sensitivity still pose a significant challenge for developing long-term scenarios—that is, in setting nutrient load targets, which response curve is most appropriate? Such long-term forecasts require assumptions about the future system sensitivity to hypoxia formation. Will
the system continue to follow the most recent curve, will it revert to a former sensitivity (as may be happening in CB), or will it become even more sensitive? We suggest this public policy challenge is best met with ensemble modeling using the family of response curves with curve selection weighted based on expert judgment and acceptable risk. For example, if a given hypoxia level were deemed ecologically or socially unacceptable, any response curve that predicted hypoxia above this level at certain nutrient loadings could be weighted higher over that loading range based on the precautionary principle. Alternately, evidence of system recovery to a lower sensitivity state could shift the weighting of curves toward those with lower sensitivity while still maintaining some weight on other observed curves.

The model presented here is primarily focused on forecasts of interest for hypoxia management and has also been used to explore system level trends in hypoxia sensitivity (Liu et al. 2010). Both simpler and more complex models have also been applied to the GOM and CB systems and each model type yields different insights into the physical, watershed, and biological controls of hypoxia as well as its impacts on individual organisms, food-webs, and biogeochemistry (Peña et al. 2010).

5. Conclusions

The forecasting ability of a simple hypoxia model with Bayesian incorporation of parameter uncertainty and variability for GOM and CB was optimized by calibration to short (3 year), recent datasets. This calibration window approach was used to assess the tradeoff between incorporating adequate system variability into model parameterization and the ability to track gradual (in CB) and abrupt (in the GOM) ecosystem changes in hypoxia sensitivity to nutrient loads. We propose use of this moving window calibration method for future short-term (annual) forecasts. The underlying changes in system sensitivity pose a great challenge to the long-term forecasting and additional work, using Bayesian weighting among families of models or incorporation of more complex model features, coupled with climate models, is likely needed.

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