Potential sources of variability in mesocosm experiments on the response of phytoplankton to ocean acidification

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Abstract

Mesocosm experiments on phytoplankton dynamics under high CO\(_2\) concentrations mimic the response of marine primary producers to future ocean acidification. However, potential acidification effects can be hindered by the high standard deviation typically found in the replicates of the same CO\(_2\) treatment level. In experiments with multiple unresolved factors and a sub-optimal number of replicates, post-processing statistical inference tools might fail to detect an effect that is present. We propose that in such cases, data-based model analyses might be suitable tools to unearth potential responses to the treatment and identify the uncertainties that could produce the observed variability. As test cases, we used data from two independent mesocosm experiments. Both experiments showed high standard deviations and, according to statistical inference tools, biomass appeared insensitive to changing CO\(_2\) conditions. Conversely, our simulations showed earlier and more intense phytoplankton blooms in modeled replicates at high CO\(_2\) concentrations and suggested that uncertainties in average cell size, phytoplankton biomass losses, and initial nutrient concentration potentially outweigh acidification effects by triggering strong variability during the bloom phase. We also estimated the thresholds below which uncertainties do not escalate to high variability. This information might help in designing future mesocosm experiments and interpreting controversial results on the effect of acidification or other pressures on ecosystem functions.

Keywords: Variability, Uncertainty Quantification, Mesocosms, Phytoplankton, Ocean Acidification

1 Introduction

Oceans are a sink for about 30% of the excess atmospheric CO\(_2\) generated by human activities [Sabine et al., 2004]. Increasing carbon dioxide concentration in aquatic environments alters the balance of chemical reactions and thereby produces acidity, which is known as ocean acidification (OA) [Caldeira and Wickett, 2003]. Interestingly, the sensitivity of pho-
to autotrophic production of particulate organic carbon (POC) to OA is less pronounced than previously thought. Several studies on CO$_2$ enrichment revealed an overall increase in POC (e.g. Schluter et al., 2014; Eggers et al., 2014; Zondervan et al., 2001; Riebesell et al., 2000), but other studies did not detect CO$_2$ effects on POC abundance (e.g. Jones et al., 2014; Engel et al., 2014) or primary production (Nagelkerken and Connell, 2015). General compilation studies showing controversial results are e.g., Riebesell and Tortell (2011) and Gao et al. (2012). Varying results are found, in particular, in mesocosm experiments, where different treatment levels, i.e., different CO$_2$ concentrations, are applied in parallel repetitions, also known as replicates or sample units. High variance is often present even in the distribution of replicates of the same sample (Paul et al., 2015; Schulz et al., 2008; Engel et al., 2008; Kim et al., 2006; Engel et al., 2005). This variability reveals a severe reduction in the ratio between acidification response signal and the variability in observations, which results in a low signal-to-noise ratio.

Mesocosms enclose plankton communities; therefore they comprise a more realistic experimental set-up compared to batch or chemostat experiments (Riebesell et al., 2008). However, mesocosms also contain a higher number of possible interactions, and thus opportunities for the spread of uncontrolled heterogeneity. Moreover, physiological states vary for different phytoplankton cells and environmental conditions, because of which independent experimental studies with similar but not identical conditions might yield divergent results. In addition, accounting for every possible factor that occurs in the original environment or fine-controlling environmental conditions, initial community structure, and ecophysiological states among replicates might be impractical. Such unresolved ecological details might propagate over the course of the experiment and produce a high standard deviation of the distribution of the replicates of the same treatment level.

Our main working hypotheses on the origins of variability in mesocosm experiments are the following:

– differences among replicates of the same sample can be interpreted as unresolved random variations (named uncertainties hereafter)
uncertainties can amplify during the experiment and generate considerable variability in the response to a given treatment level

which uncertainties are more relevant can be estimated by the decomposition of the variability.

The confirmation of these hypotheses entails two important aspects. First, heuristic exploration of variability would require experiments designed to quantify the sensitivity of mesocosms to variations in potentially relevant factors describing uncertainties in environmental conditions, cell physiology, and community structure. However, this would require high-dimensional multi-factorial set-ups (see Appendix A), which are hard to handle, if at all, even for low number of replicates. Second, statistical inference tools would come to their limitations in estimating treatment effects. Repeated measures of relevant ecophysiological data (e.g., POC) are collected from mesocosm experiments that span a few weeks. If the differences among treatment levels are smaller than those among the replicates of the same treatment level, post-processing statistical analyses typically conclude that there are no detectable effects (Field et al., 2008). In most of the cases, the mean and the variance of the sample are taken as a fair statistical representation of the effect of the treatment level and its variability. However, summary statistics such as the mean and the variance might fail to describe distributions that do not cluster around a central value, i.e., when the data are not normally distributed in the sample. This is because a feature of normally distributed ensembles is that the mean represents the most typical value and departures from that main trend (caused by unresolved factors not directly related to the treatment) very likely cancel out in the calculation of the ensemble average. Actually, this cancellation is the reason for using replicates (Ruxton and Colegrave, 2006), but many circumstances can remarkably lower the likelihood for cancellation, for instance: (i) effects that are sensitive to initial conditions (thus, small initial differences in the replicates of a given sample might become amplified and produce departures that enlarge over the course of the experiment); (ii) non-symmetrically distributed initial conditions in the sample (which might lead to non-symmetrical distribution of the results); and (iii) a low number of replicates, i.e., a sample
size not adapted to the intensity of the treatment effect, the sensitivity of all effects to initial
conditions, and the intended accuracy of the experiment. Each incident decreases the sta-
tistical power and, therefore, might infer misleading conclusions (Miller, 1988; Cohen, 1988;
Peterman, 1990; Cottingham et al., 2005).

A complementary tool for statistical inference to investigate how treatment responses
are masked by uncertainties is a data-supported model analysis based on propagation of
distributions (JCGM, 2008b). In this study, we estimate the effects of ecophysiological un-
certainties by associating the variability in experimental observations to a variational range
from repeated model runs. The margins of the variational range of each factor were thus
confined by the ability of the dynamical model to reproduce the magnitude of the variability
observed in two independent OA mesocosm experiments, namely, Pelagic Enrichment CO₂
Experiment (PeECE II and III). These confidence intervals describe the tolerance thresholds
below which uncertainties do not escalate to high variability in the modeled replicates, and
can serve as an estimator of the tolerance of experimental replicates to such uncertainties.
This information can be important to ensure reproducibility, allowing a comparison between
the results of different independent experiments and increasing confidence regarding the
effects of OA on phytoplankton (Broadgate et al., 2013).

2 Method

Potential sources of variability are estimated following a procedure already applied in sys-
tem dynamics, experimental physics and engineering (JCGM, 2008b). The basic principles
of uncertainty propagation are summarized here using a six-step method (see Fig. 1). Steps
1 and 2 are described in subsection 2.1 and comprise a classical model calibration (using
experimental data of biomass and nutrients) to obtain the reference run representing the
mean dynamics of each treatment level. In this way we found the reference value for the
model factors, i.e., parameters and initial conditions. Steps 3 and 4, described in subsec-
tion 2.2, include the tracked propagation of uncertainties by systematically creating model
trajectories for POC, each one with a slightly different value of a model factor. In steps 5
and 6, we estimated the thresholds of the model-generated variability and the effect of the
uncertainty propagation (also explained in subsection 2.2).

2.1 Model set-up, data integration, and description of the reference run

In this section, we describe the biological state that was used as reference dynamics. Our
model resolves a minimal set of state variables insofar monitored during experiments that
are assumed to be key agents of the biological dynamics. Model equations are shown in
Table 1. Reference values of the parameters are shown in Table 2. An exhaustive model
documentation is given in Appendix B. The model simulates experimental data from the
Pelagic Enrichment CO$_2$ Experiment (PeECE), a set of 9 outdoor mesocosms placed in
coastal waters close to Bergen (Norway) during the spring seasons of 2003 (PeECE II) and
2005 (PeECE III). In both the experiments, blooms of the natural phytoplankton community
were induced and treated in three replicates for the future, present, and past CO$_2$
conditions (Engel et al., 2008; Schulz et al., 2008; Riebesell et al., 2007, 2008). Experimental data are
available via the data portal Pangaea (doi: 10.1594/PANGAEA.723045 for PeECE II and
doi: 10.1594/PANGAEA.726955 for PeECE III).

Field data of aquatic CO$_2$ concentration, temperature and light were used as direct model
inputs (see Appendix C). Measurements of POC, particulate organic nitrogen (PON), and
dissolved inorganic nitrogen (DIN) were used for model calibration. Although both the ex-
periments differ in their species composition, environmental conditions and nutrient supply,
same parameter set was used to fit PON, POC, and DIN from PeECE II and III (i.e., a total of
54 series of repeated measures over more than two weeks), a feature indicating the model
skills. In addition, the model was validated with another 36 series of biomass and nutrients
data from an independent mesocosm experiment (doi: 10.1594/PANGAEA.840852, data
not shown). The experimental POC and PON data were redefined for a direct comparison
with model results (see Appendix D), since some contributions (e.g., polysaccharides and
goingly exopolymer particles) remain unresolved by our dynamical equations. State
variables of our model comprise carbon and nitrogen contents of phytoplankton, Phy$_C$
and Phy$_N$, and DIN, as representative for all nutrients. The dynamics of non-phytoplanktonic
components, i.e. detritus and heterotrophs (DH), are distinguished by DH\textsubscript{C} and DH\textsubscript{N}. Thus, in our study, POC = Phy\textsubscript{C} + DH\textsubscript{C} and PON = Phy\textsubscript{N} + DH\textsubscript{N}.

The mean cell size in the community, represented as the logarithm of the mean equivalent spherical diameter (ESD), was used as a model parameter. It determines specific ecophysiological features by using allometric relations that are relevant for the computation of subsistence quota, as well as nutrient and carbon uptake rates. Regarding the latter, to resolve sensitivities to different DIC conditions, we used a relatively accurate description of carbon acquisition as a function of DIC and size. It has been suggested by previous observations and models that ambient DIC concentration increases primary production (e.g. Schluter et al., 2014; Rost et al., 2003; Zondervan et al., 2001; Riebesell et al., 2000; Chen, 1994; Riebesell et al., 1993; Riebesell and Tortell, 2011) and mean cell size in the community (Sommer et al., 2015; Eggers et al., 2014; Tortell et al., 2008). While state-of-the-art models such as Artioli et al. (2014) used empirical biomass increase to describe OA effects, we adopted and simplified a biophysically explicit description for carbon uptake from Wirtz (2011), where the efficiency of intracellular DIC transport has been derived as a function of the mean cell size $\ell = \ln(ESD/1\mu m)$ and CO\textsubscript{2} concentration. For very large cells, the formulation converges to the surface to volume ratio, which in our notation reads $e^{-\ell}$. In contrast, the dependence of primary production on CO\textsubscript{2} vanishes for picophytoplankton; the rate limitation by sub-optimal carboxylation then reads:

$$f_{\text{CO2}} = \left( \frac{1 - e^{-a_{\text{CO2}} \cdot \text{CO2}}}{1 + a^* \cdot e^{(\ell - a_{\text{CO2}} \cdot \text{CO2})}} \right).$$

The specific carbon absorption coefficient $a_{\text{CO2}}$ reflects size-independent features of the DIC acquisition machinery (for instance, the carbon concentration mechanisms (Raven and Beardall, 2003)). The coefficient $a^*$ represents carboxylation depletion.

### 2.2 Uncertainty propagation

We considered that uncertainties were only present in the initial set-up of the system; this allowed us to perform a deterministic non-intrusive forward propagation of uncertainty, which
neglects the possible coupling between uncertainties and temporal dynamics unlike in intrusive methods (Chantrasmi and Iaccarino, 2012) involving stochastic dynamical equations with time-varying uncertainties (Toral and Colet, 2014, M. de Castro et al. in preparation). Forward refers to the fact that unresolved differences among replicates simulated as variations of the model control factors are propagated through the model to project the overall variability in the system response, (in contrast to backward methods of parameter estimation where the likelihood of input values is conditioned by the prior knowledge of the output distribution (as, for instance, in Larssen et al. (2006)).

Our approach is based on a Monte Carlo method for the propagation of distributions. It is based on the repeated sampling from the distribution for possible inputs and the evaluation of the model output in each case (JCGM, 2008b). Next, the overall simulated POC variability is compared with that in POC experimental data (i.e., the mean trends of the treatment levels as well as the standard deviations are compared, the former for the calculation of the reference run and the latter for the uncertainty propagation). Among the available experimental data, we favored POC over PON and DIN in the uncertainty propagation analysis since it is usually the target variable of OA effects and shows the highest variability. A variability decomposition with more than one dependent variable (equivalent to a multivariate ANOVA design, for instance) is beyond of the scope of the study. The comparison between simulated and experimental variability in POC helps in the identification of the changes in physiological state and community structure that are the main potential contributors to the variability.

We considered model factors, \( \phi_i \), with \( i = 1, \ldots, N = 19 \), consisting of 14 process parameters and 5 initial conditions for the state variables. Their reference values, \( \langle \phi_i \rangle \), were adjusted to yield model solutions reproducing the mean of each treatment level (steps 1 and 2, Tables 1 and 2). To test our first hypothesis, factor variations representing potential uncertainties are introduced as random values distributed around \( \langle \phi_i \rangle \) with standard deviation \( \Delta \phi_i \). To calculate \( \Delta \phi_i \), we first generate \( 10^4 \) simulations, each one with a different factor value, \( \phi_i \) (steps 3 and 4). The ensemble of model solutions for each factor and treatment level simulates the potential experimental outcomes, hereafter referred to as "virtual repli-
The factor value for each POC trajectory is randomly drawn from a normal distribution around the factor reference value $\langle \phi_i \rangle$ (same distribution is assumed by popular parametric statistical inference tools such as regressions and ANOVA [Field et al. (2008)]). For every treatment level and at every time step, we calculated the ensemble average of the virtual replicates, $\langle POC_{i,mod} \rangle$, and the standard deviation, $\Delta POC_{i,mod}$. Thus, $\Delta \phi_i$ is the standard deviation of the distribution of factor values such as $\Delta POC_{i,mod}$ does not exceed the standard deviation of the experimental POC data, $\Delta POC_{i,exp}$, for any mesocosm at any given time (step 5). The effect of variations of $\phi_i$ on the variability (step 6) is given as follows:

$$\varepsilon_i = \frac{\Delta POC_{i,mod}}{\Delta \phi_i}. \hspace{1cm} (2)$$

This ratio expresses the maximum variability a factor can generate, $\Delta POC_{i,mod}$, relative to the associated range of that factor variations, $\Delta \phi_i$, to ensure that $\Delta POC_{i,mod}$ is the closest to $\Delta POC_{i,exp}$ at any time. In general, $\varepsilon_i$ defines how much of the uncertainty of a dependent variable $Y$ (here $Y=POC$) is explained by the uncertainty of the input factors $\phi_i$, a proxy of which is known as the sensitivity coefficient $c_i = \frac{\partial Y}{\partial \phi_i}$ in the widespread formula to calculate error propagation (Ellison and Williams, 2012), also known as law of propagation of uncertainty (JCGM, 2008a)

$$(\Delta Y)^2 = \sum_{i=1}^{N} c_i^2 \cdot (\Delta \phi_i)^2.$$ 

This expression is based on the assumption that changes in $Y$ in response to variations in one factor $\phi_i$ are independent from those owing to changes in another factor $\phi_j$, and that all changes are small (thus cross terms and higher-order derivatives are neglected). Where no reliable mathematical description of the relationship $Y(\phi_i)$ exists (in our case, only an expression for the rate equation $dPOC/dt$ is known (see Table 1) but not its analytical solution, i.e., POC), $c_i$ can be evaluated experimentally (Ellison and Williams, 2012).
As mentioned in the Introduction and Appendix A, such high-dimensional multi-factorial measurements are costly in mesocosm experiments. Therefore, we obtained equivalent information by numerically calculating $\varepsilon_i$. Such approximations to sensitivity coefficients calculated by our Monte Carlo method of uncertainty propagation correspond to taking all higher-order terms in the Taylor series expansion into account since no linearization is required (see Sections 5.10 and 5.11 and Annex B in JCGM (2008b)). A straightforward extension including the cross terms showing synergistic uncertainties effects, as in an experimental multi-way ANOVA design, requires the assumption of joint distributions for the uncertainty of factors and the calculation of covariance matrices, a considerable effort that is beyond the scope of this paper.

Hereafter, the standard deviation of any given factor, i.e., factor uncertainty, will be given as percentage of the reference values and will be called $\Delta \Phi_i$. The actual factor range is given as $\Delta \phi_i = \frac{\Delta \Phi_i \cdot \phi_i}{100}$. Strong irregularities in the standard deviations of experimental POC data (for instance, small $\Delta \text{POC}^{\text{exp}}$ at day 8 in Fig. 2b), translates to remarkably enhanced or reduced sensitivity coefficients if the model-data comparison would be performed at a daily basis. Therefore, we considered the temporal mean of the standard deviation per phase, i.e., prebloom, bloom, and postbloom. We inferred phases for PeECE II from Engel et al. (2008) and for PeECE III from Schulz et al. (2008) and Tanaka et al. (2008).

To numerically calculate the ensemble of $10^4$ POC trajectories per factor (i.e., the virtual replicates, see Fig. 8), we applied the Heun integration method with a time step of $4 \cdot 10^{-4}$, (about 35 s of experimental time). The number of simulated POC time-series is chosen such as a higher number of model realizations, i.e., a higher number of virtual replicates, will produce the same results (see Adaptive Monte Carlo procedure, Section 7.9. in JCGM (2008b)). We dismissed the negative values that randomly appeared when drawing $10^4$ values from the normal distribution of factor values; this reduction in the number of trajectories did not affect the results.

Environmental data showed low variability among same treatment replicates, (see Fig. 9), suggesting a non-direct relation between variations in environmental factors among replicates and the observed biomass variability. Therefore, we focused in uncertainties in
ecophysiology and community composition and used environmental data as forcing. Perturbations of the similarity among replicates produced by strong changes in environmental conditions (storms, dysfunctional devices, etc.) or by errors in manipulation or sampling procedures are not the scope of this work. After a few decades, the current state-of-the-art of experimental techniques for running plankton mesocosms is advanced. We believe such differences are of low impact or well-understood in terms of their consequences for final outcomes (Riebesell et al., 2010; Cornwall and Hurd, 2015).

Notably, our analysis suggested sufficient (but not necessary (Brennan, 2012)) causes of uncertainties in mesocosm experiments. Variations in model characterization including structural variability (Adamson and Morozov, 2014; Fussmann and Blasius, 2005) or uncertainties in model parametrization (Kennedy and O’Hagan, 2001) or comparisons to different uncertainty propagation methods (M. de Castro et al, in preparation) require further extensive analyses beyond the scope of this study. However, we performed a series of preceding model analyses (including uncertainty propagation) by using slightly different model formulations (data not shown). From these preceding analyses, we found that different model formulations can lead to quantitatively different confidence intervals, but leave the final results qualitatively unchanged.

3 Results

3.1 CO$_2$ effect on POC dynamics

Our model reproduces the means of PON, POC and DIN experimental data per treatment level, i.e., for the future, present, and past CO$_2$ conditions, in two independent PeECE experiments (Figures 2 and 3). For PeECE II, PON is moderately overestimated and post-bloom POC is slightly underestimated. Nonetheless, the model represents the experimental data with similar precision than the means of experimental replicates (see Appendix E). The means of the same treatment replicates and their associated standard deviations are typically used to represent experimental data (see Figure 1b in Engel et al. (2008) for PeECE
II or Figure 8a in Schulz et al. (2008) for PeECE III). The means are in the foundations of the statistical inference tools that did not detect acidification responses for PeECE II and III. However, with our mechanistic model-based analysis, phytoplankton growth in the future CO\textsubscript{2} conditions showed an earlier and elevated bloom with respect to past the CO\textsubscript{2} conditions. The future and past reference trajectories limit the range of the CO\textsubscript{2} enrichment effect, as shown by the dark gray area in Figure 4. POC variability owing to variations in model factors simulating experimental uncertainties is plotted as the light gray area in the figure. Our results suggest that avoiding high POC standard deviations that potentially mask OA effects in experimental data requires the reduction of the factor variations triggering variability during the bloom.

### 3.2 CO\textsubscript{2} effect on uncertainty propagation

The estimation of the tolerance thresholds of the dynamics to uncertainty propagation for the two test-case experiments, per acidification levels and per factor uncertainty, are listed in Table 3. We investigated the potential interaction of the treatment and the uncertainties effects on the tolerance by a linear mixed-effects model with φ\textsubscript{i} as random factor (R Core Team, 2016). The synergistic effect between the factor uncertainty and the treatment levels was found to be non significant (F=2.9 with p=0.06). Therefore, the thresholds do not appear to statistically depend on the treatment level, even when the standard deviation of the measured POC data, △POC\textsuperscript{exp}, for the future and past acidification conditions were, on average, about 70% larger than the standard deviation of the present conditions (POC experimental data in Figs. 2 and 3 are more spread in the future and past concentrations than in the present concentration). Despite the low statistical power of this test (only data from two independent samples, the two PeECE experiments, were available), we still considered the potential lack of CO\textsubscript{2} effect on the uncertainty propagation as sufficient justification to simplify further analysis on variability decomposition by averaging the thresholds and the sensitivity coefficients over treatment levels (see last column in Table 3 and Fig. 7).
### 3.3 Variability decomposition

Our method allows decomposition of POC variability in factor-specific components $\Delta \text{POC}^\text{mod}_i$. The effect of factor variations simulating experimental differences among replicates is classified depending on its nature, intensity and timing (Figures 5 and 7).

POC variability during the prebloom phase can be explained mainly by the differences of factors related to subsistence quota, i.e., $Q^*_\text{subs}$ and $\alpha_Q$, in both PeECE II and III experiments (left column in Fig. 5). This suggests that the differences in subsistence quota first intensify the divergence of POC trajectories and then weaken few days later because of the system dynamics. These subsistence parameters only need to vary about 6% and 8% among replicates (see Table 3) to maximize their contribution to the $\Delta \text{POC}^\text{exp}$; thus, their sensitivity coefficients are the highest (see Fig. 7).

Differences in the initial nutrient concentration, DIN(0), mean cell size, $\ell$, and phytoplankton biomass loss coefficient, $L^*$, generate the modeled variability mainly during the bloom (with just about 20% differences among replicates, see Table 3 and second column in Fig. 5) showing high values of the sensitivity coefficient (gray highlight in Fig. 7).

Amplified variability in the postbloom phase (third column in Fig. 5) can be attributed to the uncertainties in the reference temperature $T_{\text{ref}}$ for the Arrhenius equation, Eq. (B2), in sinking loss or export flux, $s$, and in remineralization and excretion, $r^*$. The sensitivity coefficient of $T_{\text{ref}}$ is high, with just about 12% variation. Therefore, even if differences in ambient temperature among replicates of the same sample are negligible (see the low standard deviations in the temperature, Fig. 9), differences in the metabolic dependence on that ambient temperature seems to be relevant in the decay phase. Interestingly, variations among replicates in the physiological dependence on other environmental factors do not show the same relevance (the sensitivity coefficient $\varepsilon_i$ is low for carbon acquisition $a_{\text{CO2}}$ and light absorption $a_{\text{PAR}}$). Generating high divergence during the postbloom requires a strong perturbation of parameters for the description of the non-phytoplanktonic biomass (about 81% of the reference value for sinking and 96% for remineralization and excretion, see Table 3), which translates to a relatively low sensitivity coefficient.
Perturbations of the initial detritus concentration, \(DH_C(0)\) and \(DH_N(0)\) have no impact on the dynamics provided that they remain within reasonable ranges \(\triangle \Phi_i < 100\). In fact, more than tenfold difference among replicates in such non-relevant factors were necessary to achieve a perceptible variability \(\triangle POC_i^{mod}\).

POC variability throughout the bloom phases (right column in Fig. 5) can be attributed to the varying carbon and nitrogen initial conditions, \(Phy_C^\prime\) and \(Phy_N^\prime\), nutrient uptake-related factors, \(V_{\max}^*, \alpha_V,\) and Aff, and protein allocation for photosynthetic machinery, \(f_p\). About the latter, high standard deviations of the tolerance (see Table 3) suggest non-conclusive results.

### 4 Discussion

We used the uncertainty quantification method to decompose POC variability by using a low-complexity model that describes the major features of phytoplankton growth dynamics and fits the mean of mesocosm experimental PeECE II and III data with high accuracy for all \(CO_2\) treatment levels. We confirmed the working hypotheses (Figs. 5,7); in particular, we showed that small differences in initial nutrient concentration, mean cell size and phytoplankton biomass losses are sufficient to generate the experimentally observed bloom variability \(\triangle POC^{exp}\) that potentially mask acidification effects.

#### 4.1 Nutrient concentration

Differences among replicates in the initial nutrient concentration substantially contribute to POC variability, a sensitivity that is, interestingly, not well expressed when varying the initial cellular carbon or nitrogen content of the algae, \(Phy_C^\prime(0)\) and \(Phy_N^\prime(0)\). The relevance of accuracy for the initial nutrient concentration in replicated mesocosms has already pointed in Riebesell et al. (2008). Under a constant growth rate, \(DIN(0)\) determines the timing of nutrient depletion; therefore, differences in the initial nutrient concentration might also translate into temporal variations in the succession of species. We showed that such dependence is noted even in more general dynamics, and that our method can also estimate the varia-
tional range for differences in the initial DIN concentration for experiments with low number of replicates. The standard deviation of DIN(0) in the experimental set-up for PeECE III was 50% of the mean, which is significantly above our tolerance threshold (see Table 3 for initial DIN concentration). Following Riebesell et al. (2007), we considered day 2 as the initial condition, when the measured DIN was $14 \pm 2 \mu\text{mol-C L}^{-1}$, as shown in Table 1. Since $2 \mu\text{mol-C L}^{-1}$ is approximately the 14% of $14 \mu\text{mol-C L}^{-1}$, the variability of replicates at day 2 was about 14%. Therefore, experimental differences in the initial nutrient concentration were similar to the tolerance threshold for the initial DIN established to avoid high variability ($(20 \pm 6)\%$ in Table 3), which represents an explanation for the high divergence observed in POC measurements.

For PeECE II, experimentally measured DIN concentration at day 0 was $10.7 \pm 0.8 \mu\text{mol-C L}^{-1}$ suggesting a 7.5% difference among replicates, which was below our projected tolerance level (7.5 is out of the range $[14, 26]$). The same was noted for day 2, with DIN concentration equal to $8 \pm 0.5 \mu\text{mol-C L}^{-1}$ (Table 1). Our approach showed that differences in initial nutrient concentration in PeECE II were not sufficiently high to trigger the experimentally observed POC variability. Incidentally, phosphate re-addition on day 8 of the experiment established new initial nutrient concentration for the subsequent days. When the dynamics in one replicate significantly diverges from the mean dynamics of the treatment, even if the re-addition occurs at the same time and at the same concentration in all the replicates, the mesocosm with the outlier trajectory will not respond as the others do, and with the addition of a new nutrient condition, the divergence might be further amplified. In the case, nutrient re-addition has the same impact on the systems as variations in the initial conditions of nutrient concentration. Also for PeECE II, variability in POC is about 30% higher than that for PON, as shown in Fig. 2. We attribute the temporal decoupling between C and N dynamics to the break of symmetry among replicates by the nutrient re-addition owing to the strong sensitivity of the system to initial nutrient concentrations and a concomitant change in subsistence N:C quota, which is a sensitive parameter, especially during the pre-bloom phase (Fig. 5 and Fig. 7). Increase of POC:PON ratios under nitrogen deficiency has been observed frequently during experimental studies (e.g. Antia et al. 1963; Biddanda
and Benner (1997) and has been attributed to preferential PON degradation and to intracellular decrease of the N:C ratio (Schartau et al., 2007). Hence, we confirmed that nutrient re-addition during the course of the experiments results in a significant disturbance, as has been previously reported Riebesell et al. (2008), although a complete analysis would require a model that explicitly accounts for other nutrients.

4.2 Mean cell size as a proxy for community structure

We found a limited tolerance to variations in the mean cell size of the community, $\ell$, which has a threshold of about 22% variation (see Table 3). If we consider the averaged mean cell size of PeECE II, $\langle \ell \rangle = 1.6$, and III, $\langle \ell \rangle = 1.8$, from Table 2, we obtain $\langle \ell \rangle = 1.7$. Then the absolute standard deviation is $\Delta \ell = 22 \cdot \frac{1.7}{100} \sim 0.4$. Therefore, our methodology shows that variations within the range limited by $\langle \ell \rangle \pm \Delta \ell$, that is [1.3, 2.1], are sufficient to reproduce the observed experimental POC variability during the bloom. Since $\ell$ is in the log-scale, the corresponding ESD increment is within the variational range $\langle \text{ESD} \rangle \pm \Delta \text{ESD}$, that is, [3.7, 8.1]$\mu m$ (or [25, 285]$\mu m^3$ in volume). These values are easily reached in the course of species succession. This supports studies showing that community composition outweighs ocean acidification (Eggers et al., 2014; Kroeker et al., 2013; Kim et al., 2006).

4.3 Phytoplankton loss

Another major contributor to POC variability during the bloom phase is phytoplankton biomass loss, $L^*$. With a standard deviation of about 20% (Table 3), uncertainties in $L^*$ generate variability larger than the model response to OA, in particular at the end of the growth phase and the beginning of the decay phase. Unresolved details in phytoplankton loss rate include, among others, replicate differences in cell aggregation or damage by collisions, mortality by virus, parasites, and morphologic malformations, or grazing by non-filtered mixotrophs or micro-zooplankton.
4.4 Consequences for the design of mesocosm experiments

In our simulations, the CO$_2$ level affected the intensity and timing of the bloom (Figure 4). Thus, the slope of the growth phase can be regarded as a suitable target variable to detect OA effects. Moreover, our model analysis revealed a low signal-to-noise ratio. The ability to distinguish the treatment effect from noise depends on the experimental design, the strength of the treatment, and the variability that it is not explained by the treatment. When the signal-to-noise ratio is as low as it is shown by our simulations, a large experimental sample size is needed to avoid to incur in a Type II error (Field et al. 2008). In particular, we can assume a two sample two sided balanced t-test with two treatment levels as in Figure 4. The maximum difference between means equal to approximately 5µmol-C L$^{-1}$ (see for instance PeECE III at day 10) and the variability $\Delta$POC$^{mod}$ approximately equal to 4µmol-C L$^{-1}$. If we aim for a statistical power of 0.8, i.e., a 80% chance of observing a statistically significant result with that experimental design, the required number of replicates per treatment would be 11 (R Core Team 2016), which is unpractical in mesocosm experiments. With n=3 replicates, the chance declines to only 20%.

We provided an estimation for the uncertainty thresholds that can be used for improving future sampling strategies with low number of replicates, i.e. n=3. Tolerances shown in Table 3 can be used to quantify how much replicates similarity can be compromised before the variability of the outcomes outweighs potential acidification effects. Some tolerances indicate maximal variations in observable quantities, such as nutrient concentration and community composition. These model results suggest that a better control of such dissimilarities among replicates can help maintain the variability below the range of the acidification effect, especially during the bloom.

Strategies to reduce $\Delta$POC$^{mod}$ should similarly apply to lower $\Delta$POC$^{exp}$. For example, model results turned out to be very sensitive to variations in mean logarithmic cell size. Variations of this factor during the initial filling of the mesocosms may already generate divergent responses in POC so that a potential CO$_2$ signal becomes difficult to detect, if at all. To determine spectra of cell sizes (or mean of logarithmic cell size) of the initial plank-
ton community prior to CO$_2$ perturbation would be a possibility to countervail this difficulty. The decision of which mesocosm to select for which kind (i.e. intensity) of perturbation may then be adjusted according to similarities in initial plankton community structure. For example, we may consider some number of available mesocosms that should become subject to two different CO$_2$ perturbations. We may first select two mesocosms that reveal the greatest similarity with respect to their initial size spectra and assign them to the two different CO$_2$ treatments. Likewise, from the remaining mesocosms we again chose those two mesocosms that show closest similarity between their size spectra. Those two are chosen to become subject to the two different CO$_2$ perturbations. The selection procedure could be repeated until all mesocosms have been assigned to either of the two CO$_2$ treatments. Thus, mesocosms with similar initial conditions are assured to become subjected to different CO$_2$ perturbations. This reduces a mixture of random effects due to variations in initial conditions and CO$_2$ effect and it will likely facilitate data analysis in experiments with low number of replicates. Mesocosms may then be first analysed pairwise (similar initial conditions) with respect to differences in CO$_2$ response.

In addition, our analysis results help interpreting non-conclusive results and provide plausible explanations for the negative results for the detection of potential acidification effects (Paul et al., 2015; Schulz et al., 2008; Engel et al., 2008; Kim et al., 2006; Engel et al., 2005). Thus, our study also suggests the limitation of the statistical inference tools commonly used to assess the statistical significance of effect detectability.

Finally, we found the same main contributors to POC variability for all the treatment levels, even if experimental variability is about 70% higher in the mesocosms where the carbon chemistry was manipulated. In particular, the heterogeneity of variance measured in future levels is larger than under the other acidification conditions (see fluctuations of the standard deviations of CO$_2$ concentrations, Fig. 9). These differences in biomass variability among treatment levels are not explained by uncertainties in our model factors. They might have been originated by the irregularities in the CO$_2$ aeration (Riebesell et al., 2008; Cornwall and Hurd, 2015), however, further analyses need to be conducted to determine potential sources of differences in variability.
5 Conclusions

Our model projections indicated that phytoplankton responses to OA were mainly expected to occur during the bloom phase, presenting a higher and earlier bloom under acidification conditions. Moreover, we found that amplified POC variability during the bloom that potentially reduces the low signal-to-noise ratio can be explained by small variations in the initial DIN concentration, mean cell size, and phytoplankton loss rate.

The results of the model-based analysis can be used for refinements of experimental design and sampling strategies. We identified specific ecophysiological factors that need to be confined in order to ensure that acidification responses do not become masked by variability in POC.

With our approach we reverse the question of how experimental data can constrain model parameter estimates and instead determine the range of variability in experimental data that can be explained by modeling with variational ranges bounding uncertainties of specific control factors. We tested the hypothesis whether small differences among replicates have the potential to generate higher variability in biomass time-series than the response that can be attributed to the effect of CO$_2$. Therefore, we conclude that modeling studies that integrate data from acidification experiments should resolve physiological regulation capacities at cellular and community levels. In fact, modeling the propagation of uncertainties revealed cell size to be a major contributor to phytoplankton biomass variability. This suggests the use of adaptive size-trait based dynamics since such approaches allow for the resolution of ecophysiological trait shifts in non-stationary scenarios (Wirtz, 2013). The role of intracellular protein allocation can also be clarified by using a trait-based approach, since our results about the impact of its variations were non-conclusive.

In this study, we established a foundation for further model-based analysis for uncertainty propagation that can be generalized to any kind of experiments in biogeosciences. Extensions comprising time-varying uncertainties by introducing a new random value for parameters at every time step or including covariance matrices, showing the simultaneous interaction of variations in two factors, can be straightforward implemented (M. de Castro et
Finally, we believe that an explicit description of uncertainty quantification is essential for the interpretation and generalization of experimental results.

Appendix A: Model representation of replicates

Heuristic exploration of the potential origins of the observed variability uses statistical inference tools such as a multi-way repeated measures ANOVA exploring which independent factors are contributing the most to the standard deviations. Such approaches have the advantage of accounting for interacting effects between combinations of factors (and not only for the synergistic effects of each factor and acidification, as in our model-based approach, see Sec. 3). However, the realization through and experimental set-up would require a high-dimensional multi-factorial experiment extremely difficult to perform (Fig. 8). For three acidification levels, the minimum number of factor levels (i.e., high and low), minimum number of sample units (i.e., duplicates), and the same number of factors we analyze here, (i.e., N=19), the total number of mesocosms would be 3 x 2 x 2 x 19 = 228. The possibility of simulating a high number of replicates is one of the unique strengths of modeling. For each factor, we simulate possible realizations of the same acidification level with slight variations of the factor reference value (simulating differences in physiological states and community structure). We generated model solutions for $10^4$ normally distributed factor values i.e., in total, 3 acidification levels x 19 factors x $10^4$ virtual replicates for PeECE II and III experiments. Examples of 50 virtual replicates with uncertainty in initial nutrient concentration are shown in Figure 8 and examples of 10 virtual replicates with uncertainty in phytoplankton biomass losses are shown in Figure 1, both numerically calculated for low CO$_2$ conditions in PeECE III.

Appendix B: Definition of relative growth rate

Relative growth rate $\mu$ is calculated from the primary production rate by subtracting respiration and mortality losses as follows: $\mu = P - R - L$. 
Primary production

Primary production rate reflects the limiting effects of light, dissolved inorganic carbon (DIC), temperature and nutrient internal quota as follows:

\[ P = P_{\text{max}} \cdot f_{\text{PAR}} \cdot f_{\text{CO}_2} \cdot f_T \cdot f_Q \cdot f_p. \]  \hspace{1cm} (B1)

\( P_{\text{max}} \) is the maximum primary production rate, (Table 2). Specific light limitation \( f_{\text{PAR}} \) depends on light and \( \text{CO}_2 \). For the attenuation coefficient \( a_z \), we consider that in coastal regions light intensity is typically reduced to 1\% of its surface value at 5 m (Denman and Gargett, 1983) and we obtained \( a_z = 0.75\text{m}^{-1} \). Next, \( \text{PAR} \) experienced by cells at mixed layer depth (MLD = 4.5 m, Engel et al. (2008)), was calculated from the level of radiation at the water surface, \( \text{PAR}_0 \) (see Appendix C), following an exponential decay described by the Lambert-Beer law

\[ \text{PAR} = \text{PAR}_0 \int_{0}^{\text{MLD}} e^{-a_z \cdot z} dz. \]

The relationship between photosynthesis and irradiance can be formulated by referring to a cumulative one-hit Poisson distribution (Ley and Mauzerall, 1982; Dubinsky et al., 1986). With the temperature and carbon acquisition dependence, it yields

\[ f_{\text{PAR}} = \left( 1 - e^{-a_{\text{PAR}} \cdot \text{PAR}} \right) \left( 1 + \frac{a_{\text{PAR}} \cdot \text{PAR}}{P_{\text{max}} \cdot f_{\text{CO}_2} \cdot f_T} \right), \]

where \( a_{\text{PAR}} \) is the effective absorption related to the chloroplast cross-section and saturation response time for receptors (Geider et al. (1998a) Wirtz and Pahlow (2010)); the carbon acquisition term \( f_{\text{CO}_2} \) is described in Section 2.1 Eq. (1).

\( f_T \) is the temperature dependence. We considered that all metabolic rates depend on protein folding that increases with rising temperature following the Arrhenius equation (Scalley
and Baker, 1997) as described in Geider et al. (1998b) or Schartau et al. (2007)

$$\text{f}_T = e^{\frac{E_a}{T} \left(1 - \frac{1}{T_{\text{ref}}} \right)},$$  \hspace{1cm} (B2)

with activation energy $E_a = \frac{T_{\text{ref}}^2}{10} \cdot \log(Q_{10})$ as in Wirtz (2013), where we used $Q_{10} = 1.88$ for phytoplankton (Eppley, 1972; Brush et al., 2002) and $T_{\text{ref}}$ was the mean measured temperature (see Appendix C).

The allometric factor $\alpha_Q$ quantifies the scaling relation of subsistence quota and cell size. We used Droop dependency on nutrient N:C ratio (Droop, 1973), which has been recently mechanistically derived (Wirtz and Pahlow, 2010; Pahlow and Oschlies, 2013)

$$f_Q = \left(1 - \frac{Q_{\text{subs}}}{Q} \right)$$

where $Q = \frac{\text{Phy}_N}{\text{Phy}_C}$. Its lower reference, the subsistence quota $Q_{\text{subs}} = Q^*_{\text{subs}} \cdot e^{-\alpha_Q \cdot \ell}$, is considered size-dependent and reflects a lower protein demand for uptake mechanisms in large cells (Litchman et al., 2007).

The last term in Eq. (B1) accounts for an energy allocation trade-off in phytoplankton cells: protein allocation for photosynthetic compounds such as RuBisCo and pigments, $f_p$, versus allocation for nutrient uptake, $f_v$, expressed by $f_p + f_v = 1$ (Wirtz and Pahlow, 2010; Pahlow and Oschlies, 2013). We simplified the detailed partition models by setting the trait fractions as constant.

**Respiratory cost and nutrient uptake rates**

Efforts related to nutrient uptake $V$ are represented by a respiration term. Other expenses such as biosynthetic costs are neglected (Pahlow, 2005). Respiration rate is then calculated as $R = \zeta \cdot V$ where $\zeta$ expresses the specific respiratory cost of nitrogen assimilation (Raven, 1980; Aksnes and Egge, 1991; Pahlow, 2005). For simplicity, our model merges the set of
potentially limiting nutrients (e.g. P, Si and N) to a single resource only, that is DIN. We follow Aksnes and Egge (1991) as described in Pahlow (2005) for the maximum uptake rate

\[ V_{\text{max}} = \frac{1}{\frac{1}{V_{\text{max}}} + \frac{1}{f_T + \text{Aff} \cdot \text{DIN}}} \]

comprising the maximum uptake coefficient \( V_{\text{max}} \) and nutrient affinity \( \text{Aff} \). In addition to a temperature dependence of nutrient uptake as reported by Schartau et al. (2007), we assumed that respiratory costs decrease with increasing cell size (Edwards et al., 2012), which leads to an allometric scaling in nutrient uptake (Wirtz, 2013) with exponent \( \alpha_V \). We also accounted for the static proteins allocation trade-offs between photosynthetic machinery, \( f_p \), and nutrients uptake, \( f_v = 1 - f_p \). Thus, the nutrient uptake term yields

\[ V = (1 - f_p) \cdot V_{\text{max}} \cdot e^{-\alpha_V \cdot \ell}. \]

**Loss rates**

To describe the loss rate of phytoplankton biomass, we used a density-dependent term

\[ L = L^* \cdot (\text{Phy}_C + \text{DH}_C). \]

The resulting matter flux increases the biomass of detritus and heterotrophs (DH), and a fraction of it becomes a part of the remineralizable pool. A temperature-dependent remineralization term (Schartau et al., 2007)

\[ r = r^* \cdot f_T \]

describes any kind of DIN production, such as hydrolysis and remineralization of organic matter, excretion of ammonia directly by zooplankton, and rapid remineralization of fecal pellets produced also by the zooplankton. The other fraction of the non-phytoplanktonic biomass is removed by settling with a rate related to the sinking coefficient, \( s \), shown in
Tables 1 and 2. Our model was calibrated with experimental data from enclosed mesocosms where aquarium pumps ensured mixing. Therefore, we assumed that sufficiently wealthy organisms could achieve neutral buoyancy (Boyd and Gradmann, 2002), and thus sinking might not have directly affected the phytoplankton biomass.

Appendix C: Forcings

We used measured aquatic CO$_2$ and temperature per mesocosm and ambient PAR, as model inputs (see Fig. 9). For the two PeECE experiments, the photon flux density was measured by the Geophysical Institute of the University of Bergen. To calculate the surface radiation inside the mesocosms, PAR$_0$, we followed (Schulz et al., 2008) and considered that 80% of incident PAR passed through the gas tight tents, of which up to 15% penetrated to approximately 2.5 m depth, the center of the mixed surface layer in PeECE III. The daily carbon dioxide data were interpolated and PAR signal was filtered by singular spectrum analysis to avoid sudden changes that could be detrimental for the performance of the numerical calculation, since the Heun method requires differentiable functions.

Appendix D: Definition of POC

The applied model equations attribute phytoplankton, detritus, and herbivorous heterotrophs to particulate organic matter. Measurements of particulate organic carbon also include some fractions of large bacterioplankton, carnivorous zooplankton, as well as extracellular gel particles such as transparent exopolymer particles. These additional organic contributions to POC measurements are not explicitly resolved in our model. Therefore, for comparisons between simulation results and observations, we redefine the raw data from PANGAEA, named POC’ hereafter (dots in Figs. 2, 3 and 5 represent the already modified POC data). We used data of transparent exopolymer particles, TEP, from Egge et al. (2009) for PeECE III, such as here POC = POC’ - TEP. For PeECE II, TEP data were not available. We used POC = POC’ - POC", where POC" is the difference between particle abundance,
PA, of the Coulter Counter measurements and the Flow Cytometry data in Engel et al. (2008):

\[ \text{POC}'' = \beta \cdot (\text{PA}_{\text{Coulter Counter}} - \text{PA}_{\text{Flow Cytometry}}) \].

The scaling parameter \( \beta = 0.000065 \, \mu\text{mol-C}^{-1} \, \text{L} \) was tuned to provide reductions between 40 and 50\% from total POC, in agreement with the adjustments of PeECE III.

Appendix E: Residuals of the model-data fit

For the model-data fit shown in Figs. 2 and 3, we calculated the cumulative residuals E and M (Table 4) with respect to the mean of experimental replicates per treatment, time and mesocosm. For experimental data, residuals E were calculated as follows:

\[ E = \sum_{\text{treat,rep,day}} \left| Y_{\text{treat,rep,day}}^{\exp} - \langle Y_{\text{treat,day}}^{\exp} \rangle \right| / \eta \]

and for model results, residuals M were calculated as follows:

\[ M = \sum_{\text{treat,rep,day}} \left| Y_{\text{treat,rep,day}}^{\text{mod}} - \langle Y_{\text{treat,day}}^{\exp} \rangle \right| / \eta \]

with \( \eta = 9 \) being the total number of mesocosms. High residuals entail high deviation from the trend. In the case of E, this is the deviation from the mean of the treatment (typically used in statistical inference tools), and in the case of M, the deviation from the model reference run. When both E and M values are comparable, we can infer that the quality of both representations is similar (see Table 4). Thus, conclusions inferred from both approaches are based on equally valid assumptions.

Author contributions. K.W., M.S., and M.M.C. developed the model code; M.M.C performed the simulations and prepared the manuscript, which was revised by K.W. and M.S.
Acknowledgements. We thank Sabine Mathesius for the PAR and temperature data for both the PeECE II and III experiments and Kaela Slavik for the English edition of the preliminary version of the manuscript. We acknowledge our two anonymous reviewers for their helpful comments and suggestions. This work is a contribution to the National German project Biological Impacts of Ocean Acidification (BIOACID) and it is also supported by the Helmholtz society via the program PACES.

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Eggers, S. L., Lewandowska, A. M., Barcelos e Ramos, J., Blanco-Ameijeiras, S., Gallo, F., and Matthiessen, B.: Community composition has greater impact on the functioning of ma-


Figure 1. Variability decomposition method based on uncertainty propagation (summary of the basic principles given in Sections 5.1.1 and 5.6.2. and Annex B in JCGM (2008b)).
Figure 2. Solid lines show reference runs for POC, PON and DIN simulating the mean of the replicates per treatment level, with different colors for the three experimental CO$_2$ set-ups. Dots are replicated data from the Pelagic Enrichment CO$_2$ Experiment (PeECE II) for newly produced POC and PON, i.e. starting values at day 2 were subtracted from subsequent measurements as in Riebesell et al. (2007).
Figure 3. As in Fig. 2 for PeECE III.
Figure 4. Reference simulations of POC for high CO₂ (red) and low CO₂ (blue) conditions bind the range of acidification effect (dark gray) according to our model projections. Light gray area shows the limits of the overall simulated POC variability, $\Delta POC^{\text{mod}}$. Inlay graph display the signal-to-noise ratio (black solid lines), i.e., the ratio between the variance of the acidification effect and the variance of the overall variability.
Figure 5. POC variability decomposition per factor, $\triangle POC_{i}^{\text{mod}}$ for PeECE II. Shaded areas are limited by the standard deviation of $10^4$ simulated POC time-series (see Sec. 2), around the mean trajectory of the ensemble (solid line). The timing of the amplification of the variability determines four separated kinds of behavior: factor uncertainties generating variability during the prebloom, bloom, postbloom or at irregular phase (see Sec. 3).
Figure 6. As Fig. 5 for PeECE III.
Figure 7. Sensitivity coefficients $\varepsilon_i$, Eq. (2) of factors $\phi_i$ listed in Tables 1 and 2 for different bloom phases in two OA independent mesocosm experiments. Factors whose uncertainties potentially mask acidification effects (Fig. 4) by triggering variability during the bloom (Figs. 5 and 6) are highlighted.
Figure 8. The exploration of the sources of variability in an experiment with a multi-way repeated measures ANOVA design with 3 acidification levels requires a multi-factorial high-dimensional set-up (left panel). Alternately, we numerically simulate the biomass dynamics with $10^4$ virtual replicates, each one with a different normally distributed factor value (right panel). Uncertainty propagation relates the dispersion of the factor values with the dispersion of the POC trajectories. As an example, we plot results of POC variability in 50 virtual replicates of PeECE III at low acidification with uncertainty in initial nutrient concentration. Mesocosm drawing from [Scheinin et al. (2015)].
**Figure 9.** Environmental data from PeECE II and III are taken as model inputs. Error bars denote the standard deviation of the same treatment replicates.
**Table 1.** States variables and their dynamics.

<table>
<thead>
<tr>
<th>State variable</th>
<th>Dynamical equation</th>
<th>Init. cond.</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>phytoplankton carbon</td>
<td>$\frac{d\text{Phy}_C}{dt} = (P-R-L) \cdot \text{Phy}_C$</td>
<td>2.5</td>
<td>$\mu\text{mol-C L}^{-1}$</td>
</tr>
<tr>
<td>phytoplankton nitrogen</td>
<td>$\frac{d\text{Phy}_N}{dt} = V \cdot \text{Phy}_C - L \cdot \text{Phy}_N$</td>
<td>0.4</td>
<td>$\mu\text{mol-N L}^{-1}$</td>
</tr>
<tr>
<td>nutrient concentration</td>
<td>$\frac{d\text{DIN}}{dt} = r \cdot \text{DH}_N - V \cdot \text{Phy}_C$</td>
<td>$8 \pm 0.5$ (*)</td>
<td>$\mu\text{mol-N L}^{-1}$</td>
</tr>
<tr>
<td>detritus and heterotrophs C</td>
<td>$\frac{d\text{DH}_C}{dt} = L \cdot \text{Phy}_C - (s \cdot \text{DH}_C + r) \cdot \text{DH}_C$</td>
<td>0.1</td>
<td>$\mu\text{mol-C L}^{-1}$</td>
</tr>
<tr>
<td>detritus and heterotrophs N</td>
<td>$\frac{d\text{DH}_N}{dt} = L \cdot \text{Phy}_N - (s \cdot \text{DH}_N + r) \cdot \text{DH}_N$</td>
<td>0.01</td>
<td>$\mu\text{mol-N L}^{-1}$</td>
</tr>
</tbody>
</table>

* PeECE II, ** PeECE III
Table 2. Parameter values used for the reference run, \( \langle \phi_i \rangle \). All values are common to both PeECE II and III experiments, only the mean temperature (determined by environmental forcing) and the averaged cell size in the community are different since different species composition succeeded in the experiments (*Emiliania huxleyi* was the major contributor to POC in PeECE II [Engel et al. 2008] but also diatoms significantly bloomed during PeECE III [Schulz et al. 2008]).

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Units</th>
<th>Variable</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>( a_{\text{CO}_2} )</td>
<td>0.15</td>
<td>( (\mu\text{mol-C})^{-1}\text{L} )</td>
<td>( \text{Phy}_C )</td>
<td>this study</td>
</tr>
<tr>
<td>( a_{\text{PAR}} )</td>
<td>0.7</td>
<td>( \mu\text{mol phot}^{-1}\text{m}^2 )</td>
<td>&quot;</td>
<td>this study</td>
</tr>
<tr>
<td>( a^* )</td>
<td>0.15</td>
<td>( \mu\text{m}^{-1} )</td>
<td>&quot;</td>
<td>this study</td>
</tr>
<tr>
<td>( P_{\text{max}} )</td>
<td>12</td>
<td>( \text{d}^{-1} )</td>
<td>&quot;</td>
<td>this study</td>
</tr>
<tr>
<td>( Q_{\text{subs}}^* )</td>
<td>0.33</td>
<td>( \text{mol-N (mol-C)}^{-1} )</td>
<td>&quot;</td>
<td>this study</td>
</tr>
<tr>
<td>( \alpha_Q )</td>
<td>0.4</td>
<td>-</td>
<td>&quot;</td>
<td>this study</td>
</tr>
<tr>
<td>( \zeta )</td>
<td>2</td>
<td>( \text{mol-C (mol-N)}^{-1} )</td>
<td>&quot;</td>
<td>Raven (1980)</td>
</tr>
<tr>
<td>( \ell )</td>
<td>1.6</td>
<td>( \text{Ln(ESD/1\mu m)} )</td>
<td>&quot;</td>
<td>Phy(_C), Phy(_N), DIN</td>
</tr>
<tr>
<td></td>
<td>1.8</td>
<td>-</td>
<td>&quot;</td>
<td>PeECE II data</td>
</tr>
<tr>
<td>( f_p )</td>
<td>0.4</td>
<td>-</td>
<td>&quot;</td>
<td>this study</td>
</tr>
<tr>
<td>( V_{\text{max}}^* )</td>
<td>0.5</td>
<td>( \text{mol-N (mol-C d)}^{-1} )</td>
<td>&quot;</td>
<td>this study</td>
</tr>
<tr>
<td>( \text{Aff} )</td>
<td>0.2</td>
<td>( (\mu\text{mol-C d})^{-1}\text{L} )</td>
<td>&quot;</td>
<td>this study</td>
</tr>
<tr>
<td>( \alpha_V )</td>
<td>0.45</td>
<td>-</td>
<td>&quot;</td>
<td>Edwards et al. (2012)</td>
</tr>
<tr>
<td>( L^* )</td>
<td>( 11 \cdot 10^{-3} (\mu\text{mol-C d})^{-1} )</td>
<td>Phy(_C), Phy(_N) and DH(_C), DH(_N)</td>
<td>this study</td>
<td></td>
</tr>
<tr>
<td>( r^* )</td>
<td>1.5</td>
<td>( \text{d}^{-1} )</td>
<td>DH(_C), DH(_N)</td>
<td>this study</td>
</tr>
<tr>
<td>( s )</td>
<td>10</td>
<td>( L(\mu\text{mol-C d})^{-1} )</td>
<td>&quot;</td>
<td>this study</td>
</tr>
<tr>
<td>( T_{\text{ref}} )</td>
<td>8.3</td>
<td>Celsius</td>
<td>Phy(_C), Phy(_N) and DIN, DH(_C), DH(_N)</td>
<td>PeECE II data</td>
</tr>
<tr>
<td></td>
<td>10.1</td>
<td>Celsius</td>
<td></td>
<td>PeECE III data</td>
</tr>
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</table>
Table 3. Tolerance of mesocosms experiments to differences among replicates, given as a percentage of the reference factor value listed in Tables 1 and 2. According to our model projections, above these thresholds the simulated variability, $\Delta POC_{mod}^i$, exceeds the observed variability, $\Delta POC_{exp}^i$. Main contributors to the simulated variability during the bloom are highlighted in bold (see Sec. 3).

<table>
<thead>
<tr>
<th>factor $\phi_i$</th>
<th>$\Delta \Phi_i$ (%)</th>
<th>averaged tolerance (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>PeECE II</td>
<td>Present</td>
</tr>
<tr>
<td>PhyC(0) initial phyto C biomass</td>
<td>68</td>
<td>49</td>
</tr>
<tr>
<td>PhyN(0) initial phyto N biomass</td>
<td>26</td>
<td>19</td>
</tr>
<tr>
<td>DIN(0) initial DIN</td>
<td>20</td>
<td>28</td>
</tr>
<tr>
<td>aCO2 carbon acquisition</td>
<td>89</td>
<td>46</td>
</tr>
<tr>
<td>aPAR light absorption</td>
<td>&gt;100</td>
<td>&gt;100</td>
</tr>
<tr>
<td>Pmax maximum photosyn. rate</td>
<td>27</td>
<td>18</td>
</tr>
<tr>
<td>Qsubs subsistence quota offset</td>
<td>6</td>
<td>5</td>
</tr>
<tr>
<td>αQ Qsubs allometry</td>
<td>9</td>
<td>7</td>
</tr>
<tr>
<td>ℓ size Ln(ESD/1 µm)</td>
<td>25</td>
<td>20</td>
</tr>
<tr>
<td>fP fraction of protein in photosyn. machinery</td>
<td>92</td>
<td>75</td>
</tr>
<tr>
<td>Vmax* maximum nutrient uptake</td>
<td>13</td>
<td>11</td>
</tr>
<tr>
<td>Aff nutrients affinity</td>
<td>39</td>
<td>31</td>
</tr>
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<td>αV Vmax allometry</td>
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<td>11</td>
</tr>
<tr>
<td>L* phytoplankton losses</td>
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<td>30</td>
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<td>r* DIN remineralization</td>
<td>73</td>
<td>99</td>
</tr>
<tr>
<td>s DH sinking</td>
<td>&gt;100</td>
<td>&gt;100</td>
</tr>
<tr>
<td>Tref reference temperature</td>
<td>17</td>
<td>12</td>
</tr>
</tbody>
</table>
### Table 4. Cumulative residuals for PeECE III.

<table>
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<th></th>
<th>Y</th>
<th>E</th>
<th>M</th>
<th>units</th>
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<tbody>
<tr>
<td>POC</td>
<td>35.1</td>
<td>37.4</td>
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<td>$\mu$mol-C L$^{-1}$</td>
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<tr>
<td>PON</td>
<td>6.0</td>
<td>9.1</td>
<td></td>
<td>$\mu$mol-N L$^{-1}$</td>
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<tr>
<td>DIN</td>
<td>6.7</td>
<td>9.2</td>
<td></td>
<td>$\mu$mol-N L$^{-1}$</td>
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