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Sensitivity analysis of an Ocean Carbon Cycle Model in the North Atlantic: an investigation of parameters affecting the air-sea CO_2 flux, primary production and export of detritus

V. Scott¹, H. Kettle², and C. J. Merchant¹

¹School of GeoSciences, The University of Edinburgh, EH9 3JN, UK ²Biomathematics and Statistics Scotland, Edinburgh EH9 3JZ, UK

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Correspondence to: V. Scott (vivian.scott@ed.ac.uk)

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Abstract

The sensitivity of the biological parameters in a nutrient-phytoplankton-zooplankton-detritus (NPZD) model in the calculation of the air-sea CO_2 flux, primary production and detrital export is analysed. The NPZD model is the Hadley Centre Ocean Carbon

- ⁵ Cycle model (HadOCC) from the UK Met Office, used in the Hadley Centre Coupled Model 3 (HadCM3) and FAst Met Office and Universities Simulator (FAMOUS) GCMs. Here, HadOCC is coupled to the 1-D General Ocean Turbulence Model (GOTM) and forced with European Centre for Medium-Range Weather Forecasting meteorology to undertake a sensitivity analysis of its twenty biological parameters. Analyses are per-
- formed at three sites in the EuroSITES European Ocean Observatory Network: the Central Irminger Sea (60° N 40° W), the Porcupine Abyssal Plain (49° N 16° W) and the European Station for Time series in the Ocean Canary Islands (29° N 15° W) to assess variability in parameter sensitivities at different locations in the North Atlantic Ocean. Reasonable changes to the values of key parameters are shown to have a large effect
- on the calculation of the air-sea CO₂ flux, primary production, and export of biological detritus to the deep ocean. Changes in the values of key parameters have a greater effect in more productive regions than in less productive areas. We perform the analysis using one-at-a-time perturbations and using a statistical emulator, and compare results. The most sensitive parameters are generic to many NPZD ocean ecosystem
- ²⁰ models. The air-sea CO₂ flux is most influenced by variation in the parameters that control phytoplankton growth, detrital sinking and carbonate production by phytoplankton (the rain ratio). Primary production is most sensitive to the parameters that define the shape of the photosythesis-irradiance curve. Export production is most sensitive to the parameters that control the rate of detrital sinking and the remineralisation of detritus.



1 Introduction

The ocean absorbs approximately 2 Pg per year of carbon from the atmosphere (Takahashi et al., 2002; Gruber et al., 2009) – around a third of current anthropogenic emissions. The exchange of CO₂ between the atmosphere and ocean is driven by the difference in CO₂ concentration between the air and surface water, which, in turn, is influenced by the uptake of carbon by photosynthesising phytoplankton in the euphotic zone. Phytoplankton and zooplankton mortality, inefficient grazing by zooplankton, and excretion generate organic waste that sinks through the water column (export produc-

- tion). Some of this carbon reaches the ocean bed where it is gradually broken down, either dissolving or remaining locked up in sediments, completing the biological carbon pump. A relatively simple way of modelling this process is to use a four-compartment Nutrient-Phytoplankton-Zooplankton-Detritus (NPZD) model. Such models are widely used in GCMs due to their computational efficiency and typically contain around twenty biological parameters. Many of these parameters represent bulk properties across the
- ¹⁵ whole ocean, and are poorly constrained in value (Fennel et al., 2001; Frenette et al., 1993) leading to large uncertainties in model predictions. Understanding which of these parameters have the greatest influence on model output is important to understanding model results and developing improved models.

In this study we use the Hadley Centre Ocean Carbon Cycle (HadOCC) model of Palmer and Totterdell (2001) which is used to represent the role of the ocean ecosystem in the UK Met Office HadCM3 climate prediction GCM (Gordon et al., 2000) and its faster-running derivative the FAst Met Office and Universities Simulator (FAMOUS) GCM (Smith et al., 2008). HadOCC is an NPZD model that calculates the flow of nitrogen and associated flows of carbon and alkalinity between the four model compart-

²⁵ ments. Twenty free parameters (detailed in Table 1) govern the biological processes of phytoplankton growth and mortality, zooplankton grazing and excretion, and detrital sinking and remineralisation. Analogous parameterisations are used in other NPZD and more complex ocean carbon cycle models (e.g. Waniek, 2003; Popova et al.,



2002)). For this work HadOCC has been coupled to the General Ocean Turbulence Model (GOTM) (Burchard et al. (1999); Kettle and Merchant (2008)) to create a one dimensional (1-D) model (hereafter referred to as HadOCC-GOTM) that can be run at any location and is computationally efficient enough to perform the many model runs required for sensitivity analysis.

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Previous studies have assessed the importance of parameters in ocean ecosystem models at one specific site (e.g. Druon and Le Fèvre, 1999). Here, we analyse three sites that have been used for previous modelling studies (Kettle and Merchant, 2008) in different ocean regimes with different meteorological forcing and nutrient supplies to compare the regional importance of the model parameters. The Central Irminger Sea (hearafter referred to as the CIS) is located at 60° N 40° W between Greenland and Iceland in the North Atlantic subpolar gyre. Nutrients are brought up from deep water by winter storms extending the mixed layer depth allowing a phytoplankton bloom in May/June (Waniek and Holiday, 2006). The Porcupine Abyssal Plain (hearafter called

- the PAP) is located at 49° N 16° W south-east of the British Isles, and south of the main North Atlantic Current Stream. A deep winter mixed layer depth (up to 800 m) enables phytoplankton to bloom around April (Popova et al., 2002). The European Station for Time series in the Ocean Canary Islands (hereafter called ESTOC) lies on the fringes of the North Atlantic's subtropical gyre at 29° N 15° W. It is an oligotrophic site with a
 winter mixed layer depth of around 100 m (Zielinski et al., 2002). Further information on
- all these sites is available from the EuroSITES European Ocean Observatory Network (www.eurosites.info).

In this study we test the sensitivity of three model outputs – net annual air-sea CO_2 flux (mol $CO_2 m^{-2} yr^{-1}$), annual primary production and annual deep export (mgC m⁻² yr⁻¹) to variation in the values of the model parameters. Air-sea CO_2 flux is used in the calculation of absorption of anthropogenic emissions by the ocean, and all three are key processes in climate models. Here we define deep export to be the annual accumulation of detrital material that sinks below the maximum mixed layer depth (MLD) of the water column and is therefore removed from the surface waters. As the mixed



layer at the CIS extends to the deep ocean during the winter, returning sinking detrital material to the surface, the sensitivity of the calculated export to the input parameters at this site is not studied.

- Sensitivity analysis (SA) is the study of how variation in model output can be apportioned to different sources in model input (e.g., Saltelli et al., 2001). SA is performed by varying the value of the parameters across their plausible ranges and recording the resultant change in the model output. Low-sensitivity parameters are those where plausible perturbation to their value does not have a significant effect on the model output, while perturbations of highly sensitive parameters have a large effect on model output. SA qualifies the relative importance of parameter uncertainty in order to better
- output. SA qualifies the relative importance of parameter uncertainty in order to better understand and improve the model. Four steps are required to perform SA. First, the uncertainty in each parameter is assessed and a reasonable range of values for each parameter developed. Second, the parameter ranges are sampled to generate a set of input parameter sets. Third, the model is run for each parameter set and the out-
- ¹⁵ put recorded. Fourth, this information is used to perform a sensitivity assessment. At its simplest, SA is done One-At-a-Time (OAT), with each parameter varied individually while all others remain constant at a some reference value. However, this method assumes that parameter interactions have an insignificant effect on model output. Here we perform both OAT analysis and a more complex global analysis to explore the im-²⁰ pact of each parameter across the full range of all the other parameters.

2 Methods

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2.1 Model setup

We use a slightly modified version of HadOCC from that described by Palmer and Totterdell (2001) (see Appendix A for the model equations). HadOCC-GOTM is run at CIS, PAP and ESTOC in the North Atlantic ocean. Initial conditions at each location for alkalinity (Lee et al., 2006), dissolved inorganic carbon (DIC) (Key et al., 2004) are



obtained from the Carbon Dioxide Information Analysis Center (CDIAC), and nitrate profiles (to which the model relaxes to resupply) from the World Ocean Atlas 2005 (Garcia et al., 2006). The number of depth levels used by GOTM are set so that the total depth of the GOTM water column is as close as possible to the actual depth of

the site. To drive the model, meteorological data are taken from ECMWF 40-year Reanalyis (ERA-40) data (Uppala et al., 2005); these comprise air pressure, wind speed, relative humidity, air temperature and total cloud cover, at 6 hourly intervals. Appendix B gives further detail on the resupply of nutrient and meteorlogical forcing.

For each location GOTM-HadOCC is run with one thousand different sets of parameters (as described below). The model is spun up for eight years with yearly repeating meteorological data from 2004 to remove sensitivity to the initial conditions. The three model outputs under investigation (air-sea CO₂ flux, export and primary production) are then taken from the ninth year. This allows output differences between the runs for a given location to be attributed exclusively to the change in the parameter values.

15 2.2 Model parameters: ranges and sampling

We investigate twenty parameters in this study. The model equations in which these parameters are used are detailed in Appendix A. The parameters are listed in Table 1 with the current HadOCC value and values used in other comparable models. For each parameter, we estimate its likely range from expert opinions and the literature values in

Table 1. The parameters are in two distinct categories— positive definite parameters (e.g., detrital sinking velocity), and those with a value confined in the interval 0 to 1 (e.g., the assimilation efficiency of zooplankton feeding).

The parameters that relate directly to phytoplankton growth (K_{nit} , P_{max}^s , α , m_0) are generic to many NPZD and more complex models giving a good resource in determin-

²⁵ ing their possible range. For these parameters the smallest and largest values found in the literature are used for the range (see Tables 1 and 3). Parameterisations of processes such as zooplankton grazing are however less universal so there is less information available to set a likely range. Following consultation with experts who



use HadOCC (personal communication Anderson (2008); Barciela (2008); Hemmings (2008)) we give these parameters a range of the nominal HadOCC value \pm 90% of the difference between the nominal HadOCC value and the closest theoretical limit, in most cases either 0 or 1. To establish a range for Rm_{deep} we use the maximum value for Rm_{shall} as the upper limit for $Rm_{deep}/138.9$ m (the midpoint of the depth level below

for Rm_{shall} as the upper limit for $Rm_{deep}/138.9$ m (the midpoint of the depth level below 100 m is at 138.9 m) from which the 90% range is established as with other parameters above.

From these parameter value ranges one thousand parameter sets are selected using a maximin Latin Hypercube sampling method (Saltelli et al., 2001) which maximizes efficient coverage of the whole parameter space by ensuring selection of parameter values from their full range. The use of the hypercube sampling method gives equivalent parameter space coverage to a randomly sampled selection of parameter value sets a factor of ten greater in size.

These parameter sets are then run in HadOCC-GOTM and the results recorded alongside the appropriate parameter set. The calculated outputs are screened to check for failed runs that have produced wholly unfeasible results created by parameter combinations causing numerical problems (e.g. divison by zero). We use the Takahashi climatology (Takahashi et al., 2002) to screen the CO₂ flux results and set the condition that export and primary production must be greater than or equal to zero. No models runs were found to exceed these checks.

2.3 SA methods

Here, we start with the simplest method of SA – One-at-A-Time (OAT), then use the GEM-SA emulator (see below) to explore the possible effects of interactions between parameters. OAT analysis investigates the effect of changing the value of each parameter across its range while all others remain constant. OAT gives straightforward insight into the effect of parameter value variation on model outputs and the relative importance of parameters, but the amount of the full model parameter space covered is very limited as parameter interactions are not explored (Saltelli et al., 2001).



2.3.1 GEM-SA

In this work, the SA package Gaussian Emulation Machine for Sensitivity Analysis (GEM-SA) www.ctcd.group.shef.ac.uk/gem.html (O'Hagen, 2006), is used to perform and assess a global SA of HadOCC-GOTM. An issue with performing global SA is

- the computational expense of performing large numbers of model runs to generate the input-output data set for study. To reduce this computational cost, GEM-SA uses an input-output data set as a training set to build an emulation of the relationship between the inputs and output in the model, allowing the input-output set, and the corresponding cost of generation, to be smaller. This emulation is a statistical approximation to the ac-
- tual model, which allows the SA to use information from the emulated regions between the points defined by the processed input-output data, as well as the points themselves. Because of this large computational benefit, statistical emulators like GEM-SA are increasinly being used to perform SA in both ecological modelling e.g. Petropoulos et al. (2009) and other research fields such as engineering e.g. Finley et al. (2009).
- ¹⁵ To emulate the relationship between the inputs and the output, GEM-SA interpolates between the points in the training set using a Gaussian process. At each of the points in the training set the emulator gives the same result as that found in the training set – at these points the uncertainty in the fit of the emulator is zero. Between these training set points the emulator gives a "best guess" to the true value of the output that would have
- ²⁰ been calculated if the corresponding parameter value had been explicity used, with the uncertainty of this "best guess" as a normal distribution. This means that uncertainty in the goodness of fit of the emulator increases as the emulated parameter moves further away from the points of the training set. In practice, provided the relationship between the parameter and output is smooth, and a suitably sized training set is used, emulated data are virtually indistinguishable from genuine data.

SA is performed on the emulated model, assessing the contribution of the variance in the value of each parameter to the variance in the model output. Both the individual parameter contribution (without interactions with the other parameters), and the



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total parameter contribution which includes interactions with all other parameters are calculated.

3 Results

OAT results 3.1

- Figures 1, 2 and 3 show the one-at-a-time (OAT) results for each parameter at each of 5 the sites. These are generated by running HadOCC-GOTM with fifty different equally spaced values of each parameter across the parameter ranges shown in Table 3 while all other parameters remain fixed at their orginal value.
- Looking at the results in Figs. 1, 2 and 3, variation in some parameters has a noticeable effect on the output value calculated, while variation in others does not. For 10 CO₂ flux (Fig. 1) individual parameter value variation does not have a huge effect – variation in some parameters (e.g. the maximum rate of photosynthesis P_{max}^{s} and the detrital sinking rate $v_{\rm s}$) has a slight effect on the flux calulated, while other parameters (e.g. F_{messy} , F_{nmp} and F_{zmort}) have a neglible effect. In the case of primary production (Fig. 2) individual parameter variation has a much stronger effect, most notably 15 in the case of the maximum rate of photosynthesis P_{max}^{s} and the initial slope of the photosynthesis-irradiance curve α , as is expected given these parameters role in governing the ability of the phytoplankton to photosynthesise and grow (see Appendix A). For deep export (Fig. 3) the greatest effect is seen for variation in the values of the detrital sinking rate v_s and the deep detrital remineralisation rate Rm_{deep} , again as 20 expected given their role in controlling export and breakdown of detritus in the water
- column.

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Overall, two trends are notable in these OAT results. First, that the effects of individual parameter value variation on the outputs are nearly all monotonic (change only in one direction), the sole exception being the case of the carbon to chlorophyll ratio of phytoplankton θ in the calculation of the primary production (Fig. 2). Second, that while

as expected the output values at the different sites are different, the trends seen in the outputs at each site due to parameter value variation are all similar – i.e. if increase in the value of a parameter causes increase in the output, this effect is seen for all sites. The clearest examples of this are perhaps to be seen in Fig. 2 where increase in the value of P_{max}^{s} and α increases the primary production.

These results are readily interpreted. Looking at the model equations (see Appendix A), changes in individual parameter values are to be expected to have a monotonic effect on model outputs. Similarly, while the model experiences different forcing at the different sites its overall behaviour in response to parameter variations will stay the same as the mechanisms of the model remain unaltered.

3.2 GEM-SA results

Table 4 details the results of the model runs using the parameter inputs generated by the Latin hypercube sampling of the parameter space (see Sect. 2.2). Comparing these results (in particular the minimum and maximum output values calculated) with

- the OAT results in Figs. 1, 2 and 3 shows that changing the values of multiple parameters has a greater potential effect on the calculated output than changing the value of only one parameter. This is most marked in the case of the CO₂ flux where OAT changes (Fig. 1) have little effect on the flux, while large variation is seen in the CO₂ flux results for the global SA, with in the case of the Porcupine Abyssal Plain (PAP) site
- ²⁰ a change from a minimum of ≈ -5 (outgassing) to a maximum of ≈ 10 (ingassing) mol $CO_2 \text{ m}^{-2} \text{ yr}^{-1}$. This indicates that parameter interactions play a significant role in the output calculations.

Figures 4, 5 and 6 show the SA results from the GEM-SA analysis for the three model outputs at the three sites. The thick bars show the percentage of the output variance attributed to variation in each indivudal parameter, while the thin bars show the output variance attributed to variation in each parameter including interactions with all other parameters.



3.3 Discussion of GEM-SA results

The GEM-SA results show several general trends. The parameters that have a high sensitivity for each output are mostly the same at the different sites – the most obvious exception being θ which has a very high sensitivity at ESTOC but not at the CIS or PAP sites. By extension, parameters that have little sensitivity for the output are mostly the same for the different sites. This is similar to the trend seen for the OAT results in Figs. 1, 2 and 3. Parameters that have a large overall sensitivity (individual and interaction contributions), also mostly have a large individual contribution.

Looking at the GEM-SA results in detail, interactions between each individual possible combination of any two parameters (190 possible pairs of parameters) nearly all contribute to much less than 1% of the output variance, with the majority contributing less than 0.01%. For each parameter, there are 19 possible interactions with the remaining 19 parameters. The sum of these 19 interactions is shown by the difference between the overall contribution of the parameter and the individual contribution (visi-

¹⁵ ble section of the narrow bars in Figs. 4, 5 and 6). This shows that, while the sum of the parameter interactions is important to explaining the output variance, in general, interactions between indivual parameter pairs are not.

Table 5 details all the parameter interactions (9 in total) that contribute greater than 2% to the output variance. These are mostly interactions that might be expected. For primary production, P_{max}^s and α jointly define the photosynthesis-irradiance curve and a significant interaction between these parameters is to be expected. Similarly, as v_s controls the speed at which detritus sinks to the deep, and Rm_{deep} controls the rate at which that detritus breaks down, deep export is likely to influenced by interaction between these parameters.

We now look at the differences in parameter sensitivites for each of the outputs between the different sites. Starting with CO_2 flux, a total of ten different parameters, six at the CIS, five at ESTOC and seven at the PAP have an overall contribution of greater than 5% to the output variance (see Fig. 4). The ocean ecosystem influences the



air-sea CO_2 flux by fixing dissolved inorganic carbon (DIC) through growth and exporting it away from the surface through mortality. Ecosystem processes that increase DIC fixing (e.g. phytoplankton growth) encourage further CO_2 uptake from the atmosphere, while ecosystem processes that release organic carbon back to DIC in the upper ocean (e.g. remineralisation and carbonate formation) restrict CO_2 uptake.

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At the CIS, nutrients and sunlight are in relative abundance, so the value of the maximum rate of photosynthesis P_{max}^s has a very large influence on the CO₂ flux as it controls the amount of DIC taken up by phytoplankton growth at the surface, and hence the ability of the surface ocean to uptake CO₂ from the atmosphere. The initial slope of the photosynthesis-irradiance curve α is also highly influential for the same reason.

At the PAP, while P_{max}^s and α remain sensitive parameters for the reason described above, the sinking rate of detritus v_s is by far the most influential. As shown by the OAT results in Fig. 1, increasing values of v_s increase the CO₂ flux into the ocean, as faster detrital sinking reduces the amount of detritus that breaks down to dissolved inorganic carbon (DIC) near the surface. This appears to be the key process influencing the CO₂ flux at the PAP.

At the ESTOC, the carbon to chlorophyll ratio of phytoplankton θ is found to be by a large margin the most influential parameter. The ESTOC is an oligiotrophic site with limited nutrient availability restricting growth. As a result, changes in the growth rate parameters (P_{max}^{s} and α) have little effect on the DIC uptake by the phytoplankton, but changing the ratio of carbon to chlorophyll in the phytoplankton will change the DIC uptake. Similarly, the rain ratio Y_c which controls the release of organic carbon to DIC by carbonate formation is seen to be influential. However, looking at Table 4, while these parameters are influential in relative terms, the effect of parameter variation on the CO₂ flux at the ESTOC is relatively small compared to that seen at the other sites

(a range of 1.33 mol $CO_2 m^{-2} yr^{-1}$). At a site where biological activity is restricted by low nutrient availability, variation in the parameters governing the ecosystem behaviour do not have a large effect on the uptake of CO_2 flux.



For primary production (Fig. 5) a similar story is seen. At the more productive CIS and PAP sites the phytoplankton growth parameters P_{max}^s and α are the most influential – this can also be seen in the OAT results in Fig. 2. At the less-productive ESTOC, changing the carbon to chlorophyll ratio of phytoplankton θ has a much greater effect

⁵ as, for increasing θ , the same restricted supply of nutrient can produce a greater mass of phytoplankton. Similarly, phytoplankton specific mortality m_0 at the ESTOC is more influential as nutrient limited growth is less able to replace the dead phytoplankton.

Lastly, we consider deep export. As in the case of CO_2 flux and primary production, θ proves highly influential at the ESTOC due to limited nutrient supply. Otherwise, as expected from the OAT results (Fig. 3), the most influential parameters are the detrital sinking rate v_s , and the deep remineralisation rate Rm_{deep} .

4 Discussion

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The results of the OAT analysis and GEM-SA analysis are generally in agreement and as expected from the model structure (see Appendix A). The CO₂ flux is most sensitive to parameters that influence the CO₂ partial pressure (pCO₂) of the ocean surface, and hence the air-sea CO₂ exchange, by altering the DIC content of the sea-surface level. The phytoplankton growth parameters P_{max}^{s} and α influence the ocean surface pCO_{2} through controlling how efficiently phytoplankton fixes dissolved inorganic carbon. At the well-lit surface, P_{max}^{s} is more important in limiting phytoplankton growth, and so has greater influence on the calculation of CO₂ flux than α . The sinking rate of detritus v_{s} controls the rate at which detritus is removed from the sea-surface level

- directly influencing the surface ocean DIC concentration and hence its pCO_2 . The work of Schneider et al. (2008) using the more complex PISCES model, corroborates the importance of the parameterisation of particulate sinking on the calculation of the surface pCO_2 and resulting CO_2 flux. The rain ratio Y_c directly effects the DIC concentration
- by setting the amount of organic carbon released to DIC through carbonate formation. The strong influence of the carbon to chlorophyll ratio θ at the ESTOC arises due to



the relatively low nutrient availability at the site, as discussed in Sect. 3.3 above.

Primary production is sensitive to the parameters that control phytoplankton growth: α , P_{max}^s and m_0 (the phytoplankton-specific mortality rate), as has been found in previous studies such as Druon and Le Fèvre (1999). Deep export is found to be most $_5$ sensitive to the detrital sinking rate v_s , and the deep remineralisation rate Rm_{deep} . Again, high sensitivity to the carbon to chlorophyll ratio θ at the ESTOC arises due to low levels of growth (see Sect. 3.3).

As can be seen from the literature resources used to identify the parameter ranges in Tables 1 and 2, these influential parameters are all generic to many NPZD type ocean ecosystem models. The less generic HadOCC parameterisations and corresponding parameters such as the zooplankton feeding parameters F_{injest} , F_{messy} , F_{nmp} and F_{zmort} are mostly found to have weak influence. This is encouraging for the comparison of the results of different NPZD models, as it indicates that provided the parameterisations of phytoplankton growth and detrital export are similar, differences in the parameterisation of other processes may not be as critical.

While the more complex GEM-SA approach reveals some greater detail about the sensitivity of the parameters, its results agree closely with those seen in the OAT analysis. As seen in the GEM-SA results (Figs. 4, 5 and 6) individually important parameters remain the most important when parameter interactions are explored. While parame-

- ter interactions remain important in the calculation of the model outputs, changes to the value of individual parameters have a greater overall effect. Overall, these results show that the relatively simple form of an NPZD model is robust to assumptions about its behaviour based on its equations, and that parameter sensitivity can be assessed using simple methods.
- ²⁵ For use in GCMs, NPZD models like HadOCC are tuned to match measured bulk properties of the ocean ecosystem. While these sensitivity analysis results show that only certain widely used parameters may need to be tuned, they also indicate that changes to these parameters that might arise from the alteration of the marine environment by athropogenic activity can have a large effect on fundamental ocean



biogeochemical processes. This is most clearly seen in the OAT results shown in Figs. 1, 2 and 3 where changing highly sensitive parameters over a reasonable range of values is seen to have a large effect on the model outputs, particularly at the more productive CIS and PAP sites.

- ⁵ The ocean ecosystem is subject to change in its environment due to the anthropogenic decrease in ocean pH arising from increased CO₂ absorbtion by the ocean (Orr et al., 2005; Key et al., 2004), and the warming of the ocean (Doney, 2006). The effects of these environmental changes on the ocean ecosystem and its biogeochemical behaviour remain uncertain (Doney et al., 2009b; Cao and Caldeira, 2008). Having
- only one generic phytoplankton, zooplankton, nutrient and detritus compartment, controlled by a limited set of basic parameters, NPZD models such as HadOCC will not capture the effect of environmental change on the ecosystems behaviour as currently formulated. This is perhaps most clearly apparent in the use of a fixed value for the rain ratio Y_c. Ocean acidification is likely to have a detrimental effect on carbonate
 producing phytoplankton, possibly changing the global rain ratio (Doney et al., 2009a),
- and this effect is not captured in HadOCC. Models that contain greater flexibility have capacity for response to environmental changes.

The development of more advanced and flexible ocean biogeochemistry models is focussed on the use of phytoplankton functional types (PFTs) each with their own as-

- sociated parameters to separately represent the behaviours of different types of phytoplankton (Le Quéré et al. (2005), Hood et al. (2006)). There is much debate on the development of PFT models (e.g., Hood et al. (2006), Anderson (2005), Flynn (2006) and Le Quéré (2006)) due to issues of poorly understood ecology and lack of data. Selection of PFT groups and parameterisation of their characteristics is subject to the
- ²⁵ limitations of current knowledge, but does enable exploration of the possible effects of environmental change on ocean ecosystem behaviour. The changes seen in the model outputs arising from changes in the values of 'bulk property' parameters demonstrate that more flexible representations of the ocean ecosystem are needed to be able to predict the ocean ecosystem's response to climate change and potential feedbacks.



5 Summary and conclusion

We have shown that reasonable variation in the values of the biological parameters used in the HadOCC NPZD ocean biogeochemistry model have a large effect on the calculation of three fundamental outputs for biogeochemical modelling and climate pre-

diction (air-sea CO₂ flux, export and primary production). The parameters of greatest importance are generally found to be those that are generic to most NPZD models. While parameter interactions are found to influence the output calculation, changes to the values of indvidual parameters have a much greater effect than interaction with other parameters. The results of a simple one-at-a-time sensitivity analysis are not
 overturned by the results of a more complex global sensitivity analysis.

Parameters that control phytoplankton growth (the maximum photosynthetic rate P_{max}^s and the initial slope of the photosynthesis-irradiance curve α), and the parameters that control the rate of sinking of detritus and the formation of carbonate (the rain ratio) have the greatest influence on the calculation of the air-sea CO₂ flux. The cal-

- ¹⁵ culation of the primary production is most influencd by the values of the phytoplankton growth parameters P_{max}^s and α , and the specific mortality rate of the phytoplankton. Deep export (export to below the maximum annual mixed layer depth) is most influenced by the values of the sinking rate of detritus, and the remineralisation rate below the upper ocean. In a nutrient depleted oligiotrophic site, the ratio of carbon to chloro-
- ²⁰ phyll of phytoplankton θ is the most influential parameter, but changes in its (and other parameter values) have much less absolute effect on the calculated outputs than in more productive regions.

The parameters of NPZD models used in GCMs are tuned to the large-scale properties of the ocean ecosystem. These large scale properties have the potential to change

²⁵ due to anthropogenic modification of the marine environment (Siegel and Franz, 2010). While NPZD models can be successfully tuned to reproduce the measured large-scale ocean ecosystem properties, their most influential parameters are not formulated to respond to large-scale changes in the marine environment. This potentially limits the



power of NPZD models in GCMs to provide insight into the role of the ocean ecosystem in future climates.

Appendix A

5 Model equations

The NPZD model equations are described below. For more detail on these please refer to the original paper on HadOCC (Palmer and Totterdell, 2001). The model parameters are listed in Table 1.

Zooplankton graze on both phytoplankton and detritus. If *h* is the grazing rate per unit food concentration then the losses to phytoplankton and detritus are $H_P = hP$ and $H_D = hD$ respectively, where

$$h = \frac{B_Z Z}{F_{\text{tot}}} g_{\text{max}} \frac{F^2}{F^2 + K_F^2}$$
(A1)

and $F = \max(0, F_{tot} - F_{th})$, where $F_{tot} = B_P P + B_D D$, $F_{th} = 0.1$ and K_F is the half saturation constant for grazing. The factors $B_P = \frac{14.01+12.01\Theta_P}{14.01+12.01\Theta_{Red}} = 1$ (where Θ_{Red} is the Redfield ratio), $B_Z = \frac{14.01+12.01\Theta_Z}{14.01+12.01\Theta_{Red}}$ and $B_D = \frac{14.01+12.01\Theta_D}{14.01+12.01\Theta_{Red}}$ are used to adjust for the different nitrogen content per unit biomass in zooplankton, phytoplankton and detritus. Phytoplankton P (mMol N m⁻³)

$$\left. \frac{\partial P}{\partial t} \right|_{\text{bio}} = I_{\text{lim}} N_{\text{lim}} P - M_P - H_P - \eta P \tag{A2}$$

where M_P is phytoplankton mortality. This is assumed to result from viral infection, so the specific rate increases with population density such that $M_P = mP^2$ where $m = m_0$. However, if the phytoplankton concentration falls below the threshold of $P \le$



 $0.01 \text{ mMol N m}^{-3}$ the natural mortality is switched off (m = 0) as the population is assumed to be too small to transmit infection. The nutrient limitation on phytoplankton concentration is given by

$$N_{\rm lim} = \frac{N}{N + K_{\rm nit}}.$$

⁵ The light limitation I_{lim} is estimated using the daily-averaged version of the spectrallyaveraged parameterisation by Anderson (1993) such that $I_{\text{lim}} = 1 - \exp\left(\frac{-\alpha_{\text{max}}}{P_{\text{max}}^{s} \Theta N_{\text{lim}}} \alpha^{\#} I\right)$ where $\alpha_{\text{max}} = 2.602\alpha$, the derivation of $\alpha^{\#}$ is detailed by Anderson (1993) and I is photosynthetically active radiation (Wm⁻²). Zooplankton Z (mMol N m⁻³)

$$\left. \frac{\partial Z}{\partial t} \right|_{\text{bio}} = F_{\text{injest}} (\beta_P H_P + \beta_D H_D) - M_Z \tag{A4}$$

where zooplankton mortality $M_Z = \mu_1 Z + \mu_2 Z^2$ Detritus *D* (mMol N m⁻³)

$$\frac{\partial D}{\partial t}\Big|_{\text{bio}} = \frac{\Theta_P}{\Theta_D} (1 - F_{\text{nmp}}) M_P + \frac{\Theta_Z}{\Theta_D} (1 - F_{\text{zmort}}) M_Z$$

$$+ \frac{\Theta_P}{\Theta_D} a_{PD} H_P + (a_{DD} - 1) H_D - \lambda D$$
(A5)

where $\Theta_P = 6.625$, $\Theta_Z = 5.625$ and $\Theta_D = 7.5$ are the C:N ratios of phytoplankton, zooplankton and detritus respectively. $a_{PD} = (1 - F_{messy})(1 - F_{injest}) + (1 - \beta_P)F_{injest}$, $a_{DD} = (1 - F_{messy})(1 - F_{injest}) + (1 - \beta_D)F_{injest}$ and $\lambda = Rm_{shall}$ above 100 m and $\lambda = Rm_{deep}$ below 100 m. Dissolved inorganic nitrogen N (mMol N m⁻³)

$$\frac{\partial N}{\partial t}\Big|_{\text{bio}} = \left\{F_{\text{nmp}} + \left(1 - \frac{\Theta_P}{\Theta_D}\right)(1 - F_{\text{nmp}})\right\}M_P + \eta P + \left\{F_{\text{zmort}} + \left(1 - \frac{\Theta_Z}{\Theta_D}\right)(1 - F_{\text{zmort}})\right\}M_Z + \left(1 - \frac{\Theta_Z}{\Theta_D}\right)(1 - F_{\text{zmort}})\left\{F_{\text{zmort}} + \left(1 - \frac{\Theta_Z}{\Theta_D}\right)(1 - F_{\text{zmort}})\right\}M_Z + \left(1 - \frac{\Theta_Z}{\Theta_D}\right)(1 - F_{\text{zmort}})\left\{F_{\text{zmort}} + \left(1 - \frac{\Theta_Z}{\Theta_D}\right)(1 - F_{\text{zmort}})\right\}M_Z + \left(1 - \frac{\Theta_Z}{\Theta_D}\right)(1 - F_{\text{zmort}})\left\{F_{\text{zmort}} + \left(1 - \frac{\Theta_Z}{\Theta_D}\right)(1 - F_{\text{zmort}})\right\}M_Z + \left(1 - \frac{\Theta_Z}{\Theta_D}\right)(1 - F_{\text{zmort}})\left\{F_{\text{zmort}} + \left(1 - \frac{\Theta_Z}{\Theta_D}\right)(1 - F_{\text{zmort}}\right)\right\}M_Z + \left(1 - \frac{\Theta_Z}{\Theta_D}\right)(1 - F_{\text{zmort}})\left\{F_{\text{zmort}} + \left(1 - \frac{\Theta_Z}{\Theta_D}\right)(1 - F_{\text{zmort}}\right)\right\}M_Z + \left(1 - \frac{\Theta_Z}{\Theta_D}\right)(1 - F_{\text{zmort}})\left\{F_{\text{zmort}} + \left(1 - \frac{\Theta_Z}{\Theta_D}\right)(1 - F_{\text{zmort}}\right)\right\}M_Z + \left(1 - \frac{\Theta_Z}{\Theta_D}\right)(1 - F_{\text{zmort}})\left\{F_{\text{zmort}} + \left(1 - \frac{\Theta_Z}{\Theta_D}\right)(1 - F_{\text{zmort}}\right)\right\}M_Z + \left(1 - \frac{\Theta_Z}{\Theta_D}\right)(1 - F_{\text{zmort}})\left\{F_{\text{zmort}} + \left(1 - \frac{\Theta_Z}{\Theta_D}\right)(1 - F_{\text{zmort}}\right)\right\}M_Z + \left(1 - \frac{\Theta_Z}{\Theta_D}\right)(1 - F_{\text{zmort}}\right)\left\{F_{\text{zmort}} + \left(1 - \frac{\Theta_Z}{\Theta_D}\right)(1 - F_{\text{zmort}}\right)\right\}M_Z + \left(1 - \frac{\Theta_Z}{\Theta_D}\right)(1 - F_{\text{zmort}}\right)$$

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(A3)

+
$$F_{\text{messy}}(1 - F_{\text{injest}})(H_P + H_D)$$

+ $\left(1 - \frac{\Theta_P}{\Theta_D}\right)a_{PD}H_P$
+ $\lambda D - I_{\text{lim}}N_{\text{lim}}P$

Dissolved inorganic Carbon C (mMol m⁻³)

$$5 \quad \frac{\partial C}{\partial t} = \Theta_P F_{nmp} M_P + \Theta_P \eta P + \Theta_Z F_{zmort} + F_{messy} (1 - F_{injest}) (\Theta_P H_P + \Theta_D H_D) + F_{injest} \{ (\Theta_P - \Theta_Z) \beta_P H_P + (\Theta_D - \Theta_Z) \beta_D H_D \} + \Theta_D \lambda D - (1 - Y_c) \Theta_P I_{lim} N_{lim} P Alkalinity, A (mMol m-3)$$

¹⁰
$$\frac{\partial A}{\partial t} = -2Y_c \Theta_P I_{\text{lim}} N_{\text{lim}} P - \frac{\partial N}{\partial t}$$

Appendix B

Model drivers: nutrient supply and meteorology

The replenishment of nutrient in the upper ocean by mixing depends partly on lateral advection from upwelling regions and estuarine regions. To represent this additional input from outside the 1-D GOTM column, the nutrient in HadOCC-GOTM is relaxed below the productive depth to a nutrient profile taken from Levitus et al. (1993). The productive depth is defined to be the greatest depth at which both light and nutrient are available in sufficient quantities for photosynthesis to take place – i.e. the shallower of the mixed layer depth or euphotic depth. The modelled nutrient profile (*N*) is relaxed



to the fixed Levitus et al. (1993) profile (N_{Lev}) by reducing the difference between them by 1/60 per days worth of timesteps (Δt_d) such that the replenished nutrient

$$N_{\rm new} = \frac{N + (N_{\rm Lev} - N)\Delta t_d}{60}$$

HadOCC-GOTM is forced with meterological data for air pressure, wind speed, relative humidity, air temperature and total cloud cover at six hourly intervals. This data is taken from the ECMWF 40-year Re-analysis (ERA-40) dataset (Uppala et al., 2005). These are the meteorological processes that influence the rate of air-sea CO₂ transfer (wind, humidity, pressure and temperature and the amount of light (cloud cover) that reaches the ocean surface and is hence available for phytoplankton photosynthesis.

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 (CASIX).

References

Anderson, T.: A spectrally averaged model of light penetration and photosynthesis, Limnol. Oceanogr., 38, 1403–1419, 1993. 1994

Anderson, T.: Plankton functional type modelling: running before we can walk?, J. Plankton

20 Res., 27, 1073–1081, 2005. 1991

Anderson, T.: personal communication, 2008. 1983

 Anderson, T. and Pondaven, P.: Non-redfield carbon and nitrogen cycling in the Sargasso Sea: pelagic imbalances and export flux, Deep-Sea Res. Pt. I, 50, 573–591, 2003. 2009
 Anderson, T., Ryabchenko, V., Fasham, M., and Gorchakov, V.: Denitrification in the arabian

sea: A 3D ecosystem modelling study, Deep Sea Res. Pt. I, 54, 2082–2119, 2007. 2009



(B1)

- Baklouti, M., Faure, V., Pawlowski, L., and Sciandra, A.: Investigation and sensitivity analysis of a mechanistic phytoplankton model implemented in an new modular numerical tool (eco3m) dedicated to biogeochemical modelling, Prog. Oceanogr., 71, 34–58, 2006. 2009
 Barciela, R.: personal communication, 2008. 1983
- ⁵ Burchard, H., Bolding, K., and Ruiz Villarreal, M.: GOTM, a general ocean turbulence model, Theory, applications and test cases, Research article, European Commission, 1999. 1980
 Cao, L. and Caldeira, K.: Atmospheric CO₂ stabilization and ocean acidification, Geophys. Res. Lett., 35, L19609, doi:10.1029/2008GL035072, 2008. 1991
 - Chuck, A., Tyrell, T., Totterdell, I., and Holligan, P.: The oceanic response to carbon emissions
- over the next century: investigation using three ocean carbon cycle models, Tellus B, 57(1), 70–86, 2005. 2009
 - Cloern, J., Grenz, C., and Videgar-Lucas, L.: An empirical model of the phytoplankton chlorophyll:carbon ratio- the conversion factor between productivity and growth rate, Limnol. Oceanogr., 40, 1313–1321, 1995. 2009
- ¹⁵ Doney, S.: Oceanography: Plankton in a warmer world, Nature, 444, 695–696, 2006. 1991 Doney, S., Fabry, V., Feely, R., and Kleypas, J.: Ocean acidification: The other CO₂ problem, Annual Review of Marine Sciences, 169–192, 2009a. 1991
 - Doney, S., Tilbrook, B., Roy, S., Metzl, N., Le Quéré, C., Hood, M., Feely, R., and Bakker, D.: Surface ocean CO₂ variability and vulnerability, Deep Sea Res. Pt. II, 56, 504–511, 2009b. 1991
 - Druon, J.-N. and Le Fèvre, J.: Sensitivity of a pleagic ecosystem model to variations of process parameters within a realistic range, J. Mar. Syst., 19, 1–26, 1999. 1980, 1990, 2009

20

- Fennel, K., Losch, M., Schröter, J., and Wenzel, M.: Testing a marine ecosystem model: sensitivity analysis and parameter optimisation, J. Mar. Syst., 28, 45–63, 2001. 1979, 2009
- ²⁵ Finley, S., Sidwell, V., Marzocca, P., and Willmert, K.: Application of variance based sensitivity analysis to blade outer air seals, Advanced Modeling and Optimization, 11, 2009. 1984
 - Flynn, K.: . Reply to horizons article 'Plankton functional type modelling: running before we can walk?' II, putting trophic functionality into plankton functional types, J. Plankton Res., 28, 873–875, 2006. 1991
- Frenette, J.-J., Demers, S., Legendre, L., and Dodson, J.: Lack of agreement among models for estimation the photosynthetic parameters, Limnol. Oceanogr., 38(3), 679–687, 1993. 1979
 Fujii, M. and Chai, F.: Modelling carbon and silicon cycling in the equatorial pacific, Deep-Sea Res. Pt. II, 54, 496–520, 2007. 2009



Fujii, M., Ikeda, M., and Yamanka, Y.: Roles of biogeochemical processes in the oceanic carbon cycle described with a simple coupled physical biogeochemical model, J. Oceanogr., 61, 803–815, 2005. 2009

Garcia, H., Locarnini, R. A., Boyer, T., and Antonov, J.: World ocean atlas 2005, volume 4:

- Nutrients: phosphate, nitrate, silicate, Research article, NOAA Atlas NESDIS, 2006. 1982 Geider, R., MacIntyre, H., and Kana, T. M.: Dynamic model of phytoplankton growth and acclimation: responses of a balanced growth rate and the chlorophyll *a*: carbon ratio to light, nutrient-limitation and temperature, Mar. Ecol. Prog-Ser., 148, 187–200, 1997. 2009
- Geider, R., MacIntyre, H., and Kana, T. M.: A dynamic regulatory model of phytoplanktonic acclimation to light, nutrients, and temperature, Limnol. Oceanogr., 43(4), 679–694, 1998. 2009
 - Gordon, C., Cooper, C., Senior, C., Banks, H., Gregory, J., Johns, T., Mitchell, J., and Wood, R.: The simulation of SST, sea ice extents and ocean heat transports in a version of the Hadley Centre coupled model without flux adjustments, J. Clim. Dyn., 16, 147–168, 2000. 1979
- Gruber, N., Gloor, M., Mikaloff-Fletcher, S., Doney, S., Dutkiewicz, S., Follows, M., Gerber, M., Jacobson, A., Joos, F., Lindsay, K., Menemenlis, D., Mouchet, A., Muller, S., Sarmiento, J., and Takahsahi, T.: Oceanic sources, sinks, and transport of atmospheric CO₂, Global Biogeochem. Cy., 23, GB1005, doi:10.1029/2008GB003349, 2009. 1979 Hemmings, J.: personal communication, 2008. 1983
- Hood, R., Laws, E., Armstrong, R., Bates, N., Brown, C., Carlson, C., Chai, F., Doney, S., Falkowski, P., Feely, R., Friedrichs, M., Landry, M., Moore, J., Nelson, D., Richardson, T., Salihoglu, B., Schartau, M., Toole, D., and Wiggert, J.: Pelagic functional group modeling: Progress, challenges and prospects, Deep Sea Res. Pt. II, 53, 459–512, 2006. 1991 Kawamiya, M., Kishi, M., and Suginohara, N.: An ecosystem model for the North Pacific em-
- ²⁵ bedded in a general circulation model part I: Model description and characteristics of spatial distributions of biological variables, J. Mar. Syst., 25, 129–157, 2000. 2009
 - Kettle, H. and Merchant, C.: Modeling ocean primary production: sensitivity to spectral resolution of attenuation and absorption of light, Prog. Oceanogr., 78, 135–146, 2008. 1980, 2009
- ³⁰ Key, R., Kozyr, A., Sabine, C., Lee, K., Wanninkhof, R., Bullister, J., Feely, R., Millero, F., Mordy, C., and Peng, T.-H.: A global ocean climatology: Results from Global Data Analysis Project (GLODAP), Global Biogeochem. Cy., 18, GB4031, doi:10.1029/2004GB002247, 2004. 1981, 1991

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Le Quéré, C.: Reply to horizons article "Plankton functional type modelling: running before we can walk?" I abrupt changes in marine ecoystems? J. Plankton Res., 28, 871–872, 2006. 1991

Le Quéré, C., Harrison, S., Prentice, C., Buitenhuis, E., Aumont, O., Bopp, L., Claustre, H.,

Da Cunha, L., Geider, R., Giraud, X., Klaas, C., Kohfeld, K., Legendre, L., Manizza, M., Platt, T., Rivkin, R., Sathyendranath, S., Uitz, J., Watson, A., and Wolf-Gladrow, D.: Ecosystem dynamics based on plankton functional types for global ocean biogeochemistry models, Glob. Change Biol., 11, 2016–2040, 2005. 1991

Lee, K., Tong, L., Millero, F., Sabine, C., Dickson, A., Goyet, C., Park, G.-H., Wanninkhof,

R., Feely, R., and Key, R.: Global relationships of total alkalinity with salinity and temperature in surface waters of the world's oceans, Geophys. Res. Lett., 33(19), L19605, 5 pp., doi:10.1029/2006GL027207, 2006. 1981

Levitus, S., Conkright, M., Reid, J., Najjar, R., and Mantyla, A.: Distribution of nitrate, phosphate and silicate in the world oceans, Prog. Oceanogr., 31, 245–273, 1993. 1995, 1996

- ¹⁵ O'Hagen, A.: Bayesian analysis of computer code outputs: A tutorial. Reliability Engineering & System Safety, 91, 1290–1300, 2006. 1984
 - Orr, J., Fabry, V., Aumont, O., Bopp, L., Doney, S., Feely, R., Gnanadesikan, A., Gruber, N., Ishida, A., Joos, F., Key, R., Lindsay, K., Maier-Reimer, E., Matear, R., Monfray, P., Mouchet, A., Najjar, R., Plattner, G., Rodgers, K., Sabine, C., Sarmiento, J., Schlitzer, R., Slater, R.,
- I.J., T., Weirig, M., Yamanaka, Y., and Yool, A.: Anthropogenic ocean acidification over the twenty-first century and its impact on calcifying organisms, Nature, 437, 681–686, 2005.
 1991
 - Pahlow, M., Vézina, F., Casault, B., Maass, H., Malloch, L., Wright, D., and Lu, Y.: Adaptive model of plankton dynamics for the North Atlantic, Prog. Oceanogr., 76, 151–191, 2008. 2009
 - Palmer, J. and Totterdell, I.: Production and export in a global ecosystem model, Deep-Sea Res. Pt. I, 48, 1169–1198, 2001. 1979, 1981, 1993
 - Pätsch, J., Kühn, W., Radach, G., Santana Casiano, J., Gonzalez Davila, M., Neuer, S., Freudenthal, T., and Llinas, O.: Interannual variability of carbon fluxes at the North Atlantic
- ³⁰ Station ESTOC, Deep-Sea Res. Pt. II, 49, 253–288, 2002. 2009

25

Petropoulos, G., Wooster, M., Carlson, T., Kennedy, M., and Scholz, M.: A global Bayesian sensitivity analysis of the 1d SimSphere soil-vegetation-atmospheric transfer (SVAT) model using Gaussian model emulation, Ecol. Modell., 220, 2427–2440, 2009. 1984



- Popova, E., Lozano, C., Srokosz, M., Fasham, M., Haley, P., and Robinson, A.: Coupled 3D physical and biological modelling of the mesoscale variability observed in North-East Atlantic in spring 1997: biological processes, Deep-Sea Res. Pt. I, 49, 1741–1768, 2002. 1979, 1980, 2009
- 5 Saltelli, A., Chan, K., and Scott, E.: Sensitivity Analysis, John Wiley & Sons Ltd, 2001. 1981, 1983
 - Schneider, B., Bopp, L., and Gehlen, M.: Assessing the sensitivity of modeled air-sea CO₂ exchange to the remineralization depth of particulate organic and inorganic carbon, Global Biogeochem. Cy., 22, GB3021, 13 pp., doi:10.1029/2007GB003100, 2008. 1989
- ¹⁰ Siegel, D. and Franz, B. A.: Century of phytoplankton change, Nature, 466, 569–571, 2010. 1992
 - Smith, R., Gregory, J., and Osprey, A.: A description of the FAMOUS (version XDBUA) climate model and control run, Geoscientific Model Development Discussions, 1, 147–185, 2008. 1979
- Takahashi, T., Sutherland, S., Sweeny, C., Poisson, A., Metzl, N., Tilbrook, B., Bates, N., Wanninkhof, R., Feely, R., Sabine, C., Olafsson, J., and Nojiri, Y.: Global sea-air CO₂ flux based on climatological surface ocean pCO₂, and seasonal biological and temperature effects, Deep-Sea Res. Pt. II, 49, 1601–1622, 2002. 1979, 1983

Tjiputra, J., Polzin, D., and Winguth, A. M. E ..: Assimilation of seasonal chlorophyll and nu-

- trient data into an adjoint three-dimensional ocean carbon cycle model: Sensitivity analysis and ecosytem parameter optimization, Global Biogeochem. Cy., 21, GB1001, 13 pp., doi:10.1029/2006GB002745, 2007. 2009
 - Uppala, S., Kallberg, P., Simmons, A., Andrae, U., da Costa Bechtold, V., Fiorino, M., Gibson, J., Haseler, J., Hernandez, A., Kelly, G., Li, X., Onogi, K., Saarinen, S., Sokka, N., Allan, R.,
- Andersson, E., Arpe, K., Balmaseda, M., Beljaars, A., van de Berg, L., Bidlot, J., Bormann, N., Caires, S., Chevallier, F., Dethof, A., Dragosavac, M., Fisher, M., Fuentes, M., Hagemann, S., Holm, E., Hoskins, B., Isaksen, L., Janssen, P., Jenne, R., McNally, A., Mahfouf, J.-F., Morcrette, J.-J., Rayner, N., Saunders, R., Simon, P., Sterl, A., Trenberth, K., Untch, A., Vasiljevic, D., Viterbo, P., and Woollen, J.: The ERA-40 re-analysis, Quarterly Journal Royal
 Meteorological Society, 131, doi:10.1256/gj.04.176, 2005. 1982, 1996
 - Waniek, J.: The role of physical forcing in initiation of spring blooms in the northeast atlantic, J. Mar. Syst., 39, 57–82, 2003. 1979

Waniek, J. and Holiday, P.: Large-scale physical controls on phytoplankton growth in the central



Irminger Sea, part II: model study of the physical and meterological preconditioning, J. Mar. Syst., 59, 219–237, 2006. 1980, 2009

- Zielinski, O., Llinás, O., Oschlies, A., and Reuter, R.: Underwater light field and its effect on a one-dimensional ecosystem model at station ESTOC north of the Canary Islands, Deep-Sea
- 5 Res. Pt. II, 49, 3529–3542, 2002.
 1980, 2009





Fig. 1. OAT plots for CO₂ flux (y-axis) in mol CO₂ m⁻² yr⁻¹ for each of the sites. CIS green, PAP blue and ESTOC red. The y-axis scale is the same in all plots for easy comparison of the relative effect of variation in the value of each parameter (x-axis) on the CO₂ flux. The effect of individual parameter variation on the CO₂ flux is generally small, with changes to the CO₂ flux greatest for the parameters P_{max}^s , α , g_{max} , v_s and θ .











Fig. 3. OAT plots for export (y-axis) in mgC m⁻² yr⁻¹ for each of the sites. PAP blue and ESTOC red. The y-axis scale is the same in all plots for easy comparison of the relative effect of variation in the value of each parameter (x-axis) on the export. Changes to the value of v_s and Rm_{deep} have the greatest effect on the export calculated.





Fig. 4. GEM-SA results of parameter effect on the calculated CO₂ flux. Wide bars show the percentage of output variance attributable to variation in each individual parameter, narrow bars show total percentage of output variance attributable to variation in each parameter including interactions with all other parameters. CIS green, PAP blue and ESTOC red.

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Fig. 5. GEM-SA results of parameter effect on the calculated production. Wide bars show the percentage of output variance attributable to variation in each individual parameter, narrow bars show total percentage of output variance attributable to variation in each parameter including interactions with all other parameters. CIS green, PAP blue and ESTOC red.

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Fig. 6. GEM-SA results of parameter effect on the calculated deep export. Wide bars show the percentage of output variance attributable to variation in each individual parameter, narrow bars show total percentage of output variance attributable to variation in each parameter including interactions with all other parameters. PAP blue and ESTOC red.

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 Table 1. HadOCC parameters and literature values (references and additional information in Table 2).

Parameter	Description	Units	Had-OCC value	Literature values
K _{nit}	N half saturation constant	$\rm mMolNm^{-3}$	0.1	$\begin{array}{c} 0.01 {-} 0.5, \ 0.82, \ 0.85^1, \ 0.5^2, \ 0.001, \ 0.003 \ {}^3, \\ 0.5^4, \ 0.25^5 \end{array}$
P_{\max}^s	Maximum photosynthesis rate	d ⁻¹	1.5	3–5.1 ³ , 1.0 ⁵ , 0.25, 0.1–4.1 ⁷ , 0.01–1.71 ¹
α	Initial slope of P–I curve	mgC mg Chl ⁻¹ h ⁻¹ (Wm ⁻²) ⁻¹	0.09	$\begin{array}{llllllllllllllllllllllllllllllllllll$
η	Phyto. respiration rate	d ⁻¹	0.05	0.001 ⁹ , 0.0–0.025 ³
<i>m</i> ₀	Phytospecific mortality rate	d ⁻¹ (mMol N m ⁻³) ⁻¹	0.05	$0.1-0.25, 0.03^1, 0.05^5, 0.05^{10}, 0.008^{11}, 0.035^{12}, \text{ conversion}^{26}$
μ_1	Linear zoo. mortality rate	d ⁻¹	0.05	0.03–0.05 ⁶ , 0.2 ¹³
μ_2	Depth dependant zoo. mortality	d ⁻¹ (mMol N m ⁻³) ⁻¹	0.3	0.2 ¹⁴
K _F	Zoo. grazing half saturation	$\rm mMolNm^{-3}$	0.5	0.4–0.5 ¹⁵ , 1.0 ¹² , 0.5 ⁵
g_{\max}	Zoo. maximum grazing rate	d ⁻¹	0.8	0.06–1.9, 1.1, 2.0 ¹ , 0.1–0.5, 0.9 ⁶ , 2.0 ¹⁶
β_P	Zoo. assimilation effiency on phyto.	-	0.9	0.75 ¹⁰ , 0.75, 0.46, 0.76 ¹⁷ , 0.7–0.95 ¹⁸
β_D	Zoo. assimilation efficiency on det.	-	0.65	0.75^{10} , and others in β_P above.
F _{ingest}	Zoo. injestion fraction	-	0.77	-
F _{messy}	Zoo. messy feeding fraction	-	0.1	0.23 ¹⁹ , 0.02 ⁹
F _{nmp}	Phyto. mortality frac- tion to labile N & C	-	0.01	-
Fzmort	Zoo. mortality fraction to nutrient	-	0.67	-
Vs	Detrital sink rate	$m d^{-1}$	10	32.0 ²⁰ , 5.0 ² , 3.0 ²⁰ , 20.0 ²¹ , 24.7 ⁹
Rm _{shall}	Shallow (> 100 m) remineralisation rate	d ⁻¹	0.1	0.05–0.1 ²² , 0.05 ²³
Rm _{deep}	Deep (< 100 m) remineralisation rate	$m d^{-1}$	8.58	see $Rm_{\rm shall}$ above
Y _c	Rain ratio	-	0.013	0.15, 0.25, 0.08–0.12 ⁷ , 0.16 ²⁴ , 0.05–0.25 ²⁵
Θ	C:Chl ratio	_	40	10–333 ²⁷



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Footnote Number	Citation and information	ēr	Ocean Cark
1	Waniek and Holiday (2006)		Mode
2	Zielinski et al. (2002)		
3	Geider et al. (1998)	\Box	V. Scott
4	Baklouti et al. (2006)	SC	
5	Fennel et al. (2001)	S	
6	Druon and Le Fèvre (1999)	S.	
7	Chuck et al. (2005)	no	Title P
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14	Waniek and Holiday (2006) single zoo, mortality rate		Tables
15	Kettle and Merchant (2008)	SO	
16	Anderson and Pondaven (2003)		
17	Waniek and Holiday (2006) single assimilation parameter	<u>s</u> .	
18	Druon and Le Fèvre (1999) single assimilation parameter	0 N	
19	Anderson and Pondaven (2003) to DOM		
20	Anderson et al. (2007)	ap	
21	Kawamiya et al. (2000))er	Back
22	Druon and Le Fèvre (1999) at all depths	ſ	Dack
23	Waniek and Holiday (2006) at all depths		E 11 C
24	Fujii and Chai (2007)		Full Scree
25	Fujii et al. (2005)		
26	Phytoplankton concentration is assumed to be $1 \text{ mMol N} \text{ m}^{-3}$ for nurposes of conversion	SO	
20	Cloern et al. (1995)	Ŭ.	Printer-friend
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Table 2 References and additional information for Table 1



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Table 3. HadOCC parameters: Ranges used to perform the set of runs for the sensitivity analysis. Ranges are taken from either upper and lower literature values (where available), or from 90% of HadOCC value to closest theoretical limit each way (CTL).

Parameter	lower limit	upper limit	Source
K _{nit}	0.01	0.85	literature
P_{\max}^s	0.01	5.1	literature
α	0.023	1.2	literature
η	0.005	0.095	90% of HadOCC to CTL
m ₀	0.008	0.25	literature
μ_1	0.03	0.2	literature
μ_2	0.03	0.57	90% of HadOCC to CTL
K _F	0.4	1.0	literature
g_{max}	0.06	2.0	literature
β_P	0.46	0.95	literature
β_D	0.46	0.75	lower limit from β_P literature
F _{ingest}	0.63	0.977	90% of HadOCC to CTL
Fmessy	0.02	0.23	literature
Fnmp	0.001	0.019	90% of HadOCC to CTL
F _{zmort}	0.373	0.967	90% of HadOCC to CTL
V_s	3.0	32.0	literature
Rm _{shall}	0.05	0.1	literature
Rm _{deep}	3.8	13.36	90% to Rm_{shall} limit (see text)
Y _c	0.013	0.25	literature
Θ	10.0	333.0	literature



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Table 4. Results of GEM-SA model runs for net annual air-sea CO_2 flux (mol $CO_2 m^{-2} yr^{-1}$), primary production (mgC m⁻² yr⁻¹) and annual export below maximum MLD (mgC m⁻² yr⁻¹). MADM is the median absolute deviation from the median scaled such that for a normal distributon it is equal to the standard deviation σ .

output	location	control	min	max	median	MADM	mean	σ
CO_2 flux mol CO_2 m ⁻² yr ⁻¹	CIS	14.98	13.19	17.22	15.73	0.68	15.52	0.88
	PAP	1.81	-4.96	9.78	2.50	1.61	2.62	2.11
	ESTOC	0.70	-0.16	1.17	0.24	0.16	0.29	0.23
Primary production mgC m ⁻² yr ⁻¹	CIS	1.44×10 ⁵	0.0	5.68×10^{7}	1.85×10 ⁴	7.27×10^{4}	1.86×10 ⁵	9.60×10 ⁴
	PAP	1.37×10 ⁵	0.0	5.93×10 ⁵	1.90×10 ⁵	7.96×10 ⁴	1.92×10 ⁵	1.01×10 ⁵
	ESTOC	5.96×10 ⁴	0.0	2.43×10 ⁵	1.94×10 ⁴	2.50×10 ⁴	3.34×10^{4}	3.87×10 ⁴
Export mgC m ⁻² yr ⁻¹	PAP ESTOC	3.84×10 ³ 1.24×10 ³	0.087 1.10×10 ⁻³	1.42×10 ⁵ 9.60×10 ³	1.69×10 ⁴ 976.42	1.99×10 ⁴ 1.01×10 ³	2.55×10 ⁴ 1.45×10 ³	2.66×10^4 1.5×10^3

output	location	parameters	% output variance
CO ₂ flux	CIS	_	_
-	PAP	αV_s	2.08
	ESTOC	$m_0 \theta$	3.68
		$Y_c \theta$	3.53
Primary production	CIS	$P_{\max}^s \alpha$	3.25
	PAP	$P_{\max}^{s} \alpha$	3.13
	ESTOC	$m_0 \theta$	5.24
Export	PAP	v _s Rm _{deep}	6.64
	ESTOC	$v_s R m_{deep}$	2.51
		$v_s \theta$	3.67

Table 5. GE

