Quantifying the impact of ocean acidification on our future climate

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Abstract

Ocean acidification (OA) is the consequence of rising atmospheric \( \text{CO}_2 \), and it is occurring in conjunction with global warming. Observational studies show that OA will impact ocean biogeochemical cycles. Here, we use a coupled carbon-climate Earth System Model under the RCP8.5 emission scenario to evaluate and quantify the first-order impacts of OA on marine biogeochemical cycles and the potential feedback on our future climate over this century. We find that OA impacts have only a small impact on the future atmospheric \( \text{CO}_2 \) (less than 45 ppm) and future global warming (less than a 0.25 K) by 2100. While the climate change feedbacks are small, OA impacts may significantly alter the distribution of biological production and remineralization, which would alter the dissolved oxygen distribution in the ocean interior. Our results demonstrate that the consequences of OA will not be through its impact on climate change, but on how it impacts the flow of energy in marine ecosystems, which may significantly impact their productivity, composition and diversity.

1 Introduction

The oceans have taken up approximately a third of the total fossil fuel \( \text{CO}_2 \) emitted to the atmosphere since the onset of industrialization (140 ± 25 PgC, Khatiwala et al., 2009). This uptake has slowed the rate of global warming, but has led to detectable changes in the ocean chemistry (e.g. Doney et al., 2009). As the \( \text{CO}_2 \) enters the ocean, primarily through sea-air fluxes, it reacts with the sea water reducing the carbonate ion concentration and pH, collectively known as ocean acidification (OA). This uptake of anthropogenic \( \text{CO}_2 \) has reduced the pH of the modern ocean surface waters by 0.1 units or 30% since pre-industrial times (Caldeira, 2005). By the end of this century, under the higher emission scenarios, the projected pH change since pre-industrial times maybe greater than 100% (Orr et al., 2005).
OA has the potential to affect the major biogeochemical (BGC) cycles in the ocean (Gehlen et al., 2011), with potentially significant consequences for our future climate (Matear et al., 2010). In assessing the potential OA impacts on BGC cycles and climate, it is important to recognise that ocean acidification impacts do not occur in isolation, but they are associated with global warming, which may modulate their impacts (e.g. Brewer and Peltzer, 2009). Therefore, to assess the future impacts of OA requires an Earth System Model (ESM) that considers the interactive effects of global warming and OA on BGC cycling (Tagliabue et al., 2011). The goal of this study is to use a coupled-carbon ESM to evaluate and quantify the first-order impacts and climate feedbacks of OA. To do this, we first review the potential for ocean acidification to alter the major BGC cycles. We then use our ESM to assess how these modulations of the BGC by OA alter the projected climate in the current century. From our simulations, we then discuss how the OA modulated BGC changes affect the ocean environment. Finally, we end with a discussion of OA impacts on BGC cycling and climate that required further investigation.

2 Potential impacts on BGC cycles

Ocean acidification has the potential to modify marine BGC cycles in a number of ways, which could alter the future climate. We define the term BGC climate feedback, to denote an oceanic BGC process that may either enhance (positive feedback) or reduce (negative feedback) global warming due to rising greenhouse gases. For this study, we primarily focus on marine BGC processes that are impacted by OA and global warming, which can alter ocean uptake of carbon.

To aid the discussion of the potential impacts of OA on BGC cycles, we separate the impacts into processes that: (1) alter biological production in the photic zone; (2) alter the remineralization of sinking particulate organic and inorganic carbon in the ocean interior. These processes are summarised in Table 1, with an indication of the sign the impact has on the ocean storage of carbon based on published studies.
2.1 Biological production

The rising CO$_2$ in the upper ocean has the potential to affect biological production in several ways:

1. Increase net primary productivity by making photosynthesis more efficient (Rost et al., 2008). Experimental studies have also shown an increase in Particulate Organic Carbon (POC) production in response to increased levels of CO$_2$ (e.g. Zondervan et al., 2002).

2. Alter the stoichiometric nutrient to carbon ratio of the exported particulate organic matter, which enables ocean biology to partially overcome nutrient control on carbon production and export. Mesocosm experiments with natural plankton communities have reported an increased C/N ratio of particulate organic matter under elevated CO$_2$ (Riebesell et al., 2007; Bellerby et al., 2008). In these experiments, the C/N ratio increased from 6.0 at 350 µatm to 8.0 at about 1050 µatm. An increase in the C/N ratio of the export POC would result in increased storage of carbon in the ocean (Oschlies et al., 2008).

3. Impact the ability of organisms to calcify (Fabry et al., 2008), and this is anticipated to reduce the production of calcium carbonate. Due to the nature of the carbonate chemistry, reduced calcium carbonate production allows the upper ocean to increase its carbon uptake (Raven, 2005; Heinze, 2004; Hutchins, 2011).

All 3 of these affects would increase the storage of carbon in the ocean and provide a negative feedback to climate change.

2.2 Remineralization of particulate material

Most of the exported POC is remineralized in the upper 1000 m, but $\approx 10\%$ escapes to the deep ocean, where it is remineralized or buried in sediments, and sequestered from the atmosphere on millennium timescales Trull et al. (2001). Analysis of Particulate
Inorganic Carbon (PIC) and POC fluxes at water depths greater than 1000 m suggests a close association between these fluxes (Armstrong et al., 2002). The ratio of PIC production to POC is called the rain ratio (e.g. Archer et al., 2000). OA has the potential to impact the remineralization of sinking particulate material in the following ways:

1. Dissolution of CaCO₃ is an abiotic process driven by thermodynamics, and the rate of dissolution is related to the saturation state of the ocean (Orr et al., 2005). Chemical dissolution can occur when the saturation state of CaCO₃ falls below 1 (or below the lysocline). As the lysocline moves toward the surface with ocean acidification, increased dissolution of CaCO₃ sediments and sinking particles will increase the alkalinity in the ocean (Ridgwell et al., 2009). This potentially increases the supply of alkalinity to the upper ocean increasing CO₂ uptake, which acts as a negative climate feedback.

2. Armstrong et al. (2002) proposed that CaCO₃ acts as a carrier for transporting POC to the deep ocean by ballasting the POC thereby increasing its sinking speed. It is also hypothesised that the association between CaCO₃ and POC might protect the latter from bacterial degradation (Armstrong et al., 2002). If deepwater POC fluxes are controlled by CaCO₃, then a decrease in the CaCO₃ production would result in a decreased POC transport to the deep ocean. Consequently, POC would remineralise at shallower depths and therefore the net efficiency of the biological pump would decrease, resulting in a positive climate change feedback.

3. The remineralisation length scale of POC is dependent on microbial activity. It has been hypothesised that ocean acidification may increase the rate of microbial activity (Weinbauer et al., 2011), leading to more rapid remineralization of sinking POC, which would result in less carbon sequestration, and a positive climate change feedback.

In summary, changing the strength of the biological pump can have a feedback on climate change.
3 Earth system model and simulations

3.1 Earth system model

For the simulations presented here a coupled carbon-climate earth system model was used (Mk3L-COAL). Mk3L-COAL includes a climate model, Mk3L (Phipps et al., 2011), coupled to a biogeochemical model of carbon, nitrogen and phosphorus cycles on land (CASA-CNP) in the Australian community land surface model, CABLE (Wang et al., 2010; Mao et al., 2011), and an ocean biogeochemical cycle model (Matear and Hirst, 2003; Duteil et al., 2012). The details of the ocean biogeochemical model are summarised in Appendix A.

The atmosphere model has a horizontal resolution of 5.6° by 3.2°, and 18 vertical layers. The land carbon model has the same horizontal resolution as the atmosphere. The ocean model has a resolution of 2.8° by 1.6°, and 21 vertical levels. Mk3L simulates the historical climate well, as compared to the models used for earlier IPCC assessments (Phipps et al., 2011; Pitman et al., 2011). Furthermore, the simulated response of the land carbon cycle to increasing atmospheric CO₂ and warming are consistent with those from the Coupled Model Intercomparison Project Phase 5 (CMIP5) (Zhang et al., 2013). The ocean biogeochemical model was shown to realistically simulate the global ocean carbon cycle (Duteil et al., 2012). Previous studies with this ocean biogeochemical model have shown it to realistically simulate anthropogenic carbon and CFC uptake by the ocean (Matsumoto et al., 2004). We referred to the ocean BGC formulation used in these previous studies as our standard BGC formulation, which is then modified to incorporate OA impacts.

The land model (CABLE) with CASA-CNP simulates the temporal evolution of heat, water and momentum fluxes at the surface, as well as, the biogeochemical cycles of carbon, nitrogen and phosphorus in plants and soils. For this study, we used the spatially explicit estimates of nitrogen deposition for 1990 (Dentener, 2006). The simulated (Zhang et al., 2013) geographic variations of nutrient limitations, major biogeochemical...
fluxes and pools on the land under the present climate conditions are consistent with published studies (Wang et al., 2010; Hedin, 2004).

3.2 Model simulations

The ESM was spun-up under preindustrial atmospheric CO$_2$ (1850: 284.7 ppm) until the simulated climate became stable. Stability was defined as the linear trend of global mean surface temperature over the last 400 yr of the spin-up being less than 0.015 K per century.

For the historical period (1850–2005), the ESM was run using the historical atmospheric CO$_2$ concentrations as prescribed by the CMIP5 simulation protocol. For the historical period two simulations were performed. One, a Control simulation (CTRL) where atmospheric CO$_2$ only affects the carbon cycle and not the radiative properties of the atmosphere (Table 2). In the CTRL simulation there is no climate change due to greenhouse gas warming. Two, a Reference simulation (REF) where atmospheric CO$_2$ affects both the radiative properties of the atmosphere and the carbon cycle.

For the future, we used the RCP8.5 emissions scenario as provided by the CMIP5 (http://cmip-pcmdi.llnl.gov/cmip5/), and let our ESM, which has the full carbon-climate interactions determine the future atmospheric CO$_2$ concentrations (Zhang et al., 2013). The RCP8.5 scenario is the high CO$_2$ emissions scenario used in the IPCC’s Fifth Assessment Report, and in this scenario radiative forcing increases to 8.5 Wm$^{-2}$ by 2100. For comparison, we also did a standard CMIP5 simulation where RCP8.5 atmospheric CO$_2$ concentrations were used rather than letting our ESM determine the future CO$_2$ concentrations from the emission scenario (we called this simulation RCP8.5). As discussed by Zhang et al. (2013), using RCP8.5 emissions rather than concentrations resulted in a slightly warmer world (0.25 K) by 2100, because of reduced land carbon uptake due to nutrient limitation. In all our simulations the vegetation distribution used (Lawrence et al., 2013) remained (unchanged) over the simulation period following the CMIP5 experimental design. Note, the CO$_2$ emissions from land use change were not included in our simulation but were taken into account in the emission estimates used.
in the RCP8.5 emissions scenario. We also neglected changes in anthropogenic N deposition over the simulation period, because of the large uncertainty in the future deposition rate and the small impact it has on net land carbon uptake (Zaehle et al., 2010).

3.3 Ocean acidification impacts on BGC

To assess the potential impacts of OA on BGC cycles, we modified the ocean BGC formulation in the future period (2006–2100). These idealised modifications to the marine BGC cycle are summarised in Table 2 and described in more detail below. The modifications are designed to provide a first-order assessment of the OA impacts discussed in the previous section on the future climate.

The reference simulation (REF), refers to the standard BGC formulation with no OA impacts. With this ocean BGC formulation, the ESM simulates years 1850 to 2100 using historical atmospheric CO$_2$ for 1850 to 2006 and then the RCP8.5 emissions until year 2100. All of the remaining experiments to be discussed were started from year 2006 and use the same RCP8.5 emissions as the REF simulation.

The first OA simulation (EP+) considers the impact of rising CO$_2$ on the C/P ratio of the exported POC. The C/P changes are based on Oschlies et al. (2008), while the PIC export remains unchanged. The modified POC export ($Q_{POC}$) is given by the following equations where CO2 is the atmospheric CO$_2$ concentration and $J(P)$ is the phosphate uptake in the photic zone and $\Delta z$ is the depth of the photic zone.

\[
Q_{POC} = 106 \ J(P) F_o \ \Delta z \\
F_o = 1 + (CO2 - 380) \cdot 2.3/700/6.6, \text{ for } CO2 > 380
\]

At the start of the experiment (2006), the atmospheric CO$_2$ equals 380 ppm and the scaling factor ($F_o$) is equal to 1, which is the value used in the REF experiment.

In the second OA experiment (EP++), in addition to the EP+ modification, the export of POC is prescribed to increase as atmospheric CO$_2$ levels rise reflecting the
enhanced production due to enhanced photosynthesis with increased CO\(_2\). In addition the PIC export \((Q_{\text{PIC}})\) is prescribed to decline as CaCO\(_3\) decreases with rising CO\(_2\) (Ridgwell et al., 2009). To achieve this we increase the scaling factor on POC export \((S_{\text{npp}})\) and reduce the rain ratio between PIC and POC export \((r)\) with atmospheric CO\(_2\) as follows.

\[
S_{\text{npp}} = S_{\text{npp}}^0[4.5 \cdot (\text{CO2}/380) − 3.5] \quad (3)
\]

\[
Q_{\text{PIC}} = r \ 106 \ J(P) \ \Delta z \quad (4)
\]

\[
r = r^0 \cdot 1/(9.5 \cdot \text{CO2}/380 − 8.5) \quad (5)
\]

where \(S_{\text{npp}}^0\) and \(r^0\) are the scaling factor and rain ratio respectively, used in the REF experiment. At an atmospheric CO\(_2\) level of 1140 ppm (the approximate value at 2100) the scaling factor on POC export is 10 times and the rain ratio is 1/20 of the REF experiment. Note, that while the scaling factor was increased by 10 times the actual increase in export of POC is still constrained by the availability of phosphate and light, which dramatically reduces the increase in POC export in this experiment.

In the third OA simulation \(\ominus\) min\(+\), the standard BGC formulation is used but both the remineralization of POC and the dissolution of PIC are enhanced. The increased POC remineralization reflects an increasing rate of microbial activity with OA. The increased PIC dissolution reflects the impact of OA on the chemical dissolution of PIC. In the previously discussed experiments, POC and PIC have a prescribed depth profiles (see A10 and A11), which are now modified by the atmospheric CO\(_2\) as follows to give an upper bound estimate of the potential impact of increased POC remineralization and increased PIC dissolution on carbon storage in the ocean. The following equations give the factors used to modify the prescribed depth profiles for POC and PIC from the REF experiment, where \(\text{Remin}_{\text{POC}}^0 = −0.9\) and \(\text{Remin}_{\text{PIC}}^0 = 3500\) in the REF simulation (see Eqs. A10 and A11).

\[
\text{Remin}_{\text{POC}} = \text{Remin}_{\text{POC}}^0[1 + (\text{CO2} − 380)/500] \quad (6)
\]

\[
\text{Remin}_{\text{PIC}} = \text{Remin}_{\text{PIC}}^0 1/[1 + (\text{CO2} − 380)/500] \quad (7)
\]
With an atmospheric CO$_2$ level of 1000 ppm, the depth of remineralization declines by 2.25 from the REF experiment. For POC, this means the POC sinking below 200 m declines from 54 % in REF simulation to 25 %. For PIC, the PIC sinking below 1000 m reduces from 75 % in REF simulation to 37 %.

These BGC changes are an idealised representation of the OA impact on the ocean BGC cycle. They were deliberately chosen to provide extreme perturbations to the BGC cycle and test the potential for these impacts to have a significant consequence on the future projected climate.

4 Results and discussion

We will first discuss the REF simulation by assessing its present-day ocean state and its projected response to the RCP8.5 emission scenario. Then, we will discuss how OA impacts the ocean BGC cycle, and how this in turn impacts the future atmospheric CO$_2$ concentration and climate.

4.1 Historical period

For the historical period, the simulated global surface warming generally agrees with the observed warming (Fig. 1). The observed global surface temperature increase between 1850–1899 and 2001–2005 is 0.76 ± 0.19 K (Trenberth et al., 2007) compared to the simulated increase of 0.57 ± 0.07 K (Zhang et al., 2013). The observed land surface temperature increase from 1850–1899 to 2001–2005 is 1.0 ± 0.25 K (Brohan et al., 2006) compared to a simulated increase of 0.75 ± 0.06 K (Zhang et al., 2013).

Zhang et al. (2013) assessed the simulated anthropogenic CO$_2$ uptake by the land and ocean, and they showed that over the historical period the model realistically reflected the observed estimates. From 1850 to 2005, the total carbon accumulated in the land biosphere is 85 Pg C, which is within the land carbon uptake of 135 ± 85 Pg C (Zhang et al., 2013) calculated from the estimated rates of ocean carbon uptake and at-
mospheric growth. The simulated carbon accumulated in the ocean (118 PgC), agrees with the estimated anthropogenic carbon storage in the ocean of 135 ± 25 PgC (Khatiwala et al., 2009). The simulated land and ocean carbon uptake is also consistent with the latest synthesis for the period 1960 to 2005 (Canadell et al., 2007).

To assess the realism of the ocean carbon simulation we compare: (1) dissolved oxygen at 500 m; (2) surface aragonite saturation state; (3) aragonite lysocline depth; and (4) surface phosphate with the observations using a Taylor diagram (Fig. 2).

For dissolved oxygen at 500 m, the REF simulation is highly correlated with the observations (Boyer et al., 2009) (Fig. 2 and Table 3) with the simulation slightly over-estimating the average oxygen concentration at 500 m. In general, the simulated dissolved oxygen levels at mid-depths are over-estimated, but the simulated thickness of suboxic water (defined as oxygen concentrations less than 5 µmolL\(^{-1}\)) was about double the observed value.

For surface aragonite saturation state, the REF simulation is highly correlated with the aragonite saturation state calculated from the observations (Key et al., 2004), with the simulation slightly under-estimating the average saturation state (Fig. 3 and Table 3). In the ocean interior, the REF simulation has a similar pattern for the depth of the aragonite lysocline to the observations. In general, the simulated lysocline depth is a little deeper than observed, except in the Western Pacific where the simulated values are several hundred meters too deep (Fig. 3 and Table 3).

For surface phosphate, the REF simulation is highly spatially correlated with the observations (Boyer et al., 2009), but with slightly greater spatial variability (Fig. 2 and Table 3). The simulated mean surface phosphate concentration is also slightly less than observed.

In summary, the REF simulation generally reflects the present-day ocean state (correlation coefficient with the observations greater than 0.8). However, the important discrepancies with the observations are that the suboxic water is too thick and the lysocline in the Western Pacific subtropical and equatorial water is too deep. These
model biases need to be considered when assessing the projected regional changes associated with OA.

4.2 Future OA impacts on ocean BGC

4.2.1 Carbon climate feedbacks

The addition of the OA impacts on the ocean BGC alters the atmospheric CO$_2$ by less than 45 ppm by 2100 (Fig. 4). Enhanced POC export increases ocean carbon uptake, and atmospheric CO$_2$ drops by almost 43 ppm by 2100. While the enhanced remineralization of the PIC and POC reduces carbon uptake, and atmospheric CO$_2$ increases by about 18 ppm by 2100. The combination of enhanced POC export with the shoaling of POC remineralization and PIC dissolution reduces atmospheric CO$_2$ by 38 ppm. The atmospheric CO$_2$ change in the COMB experiment is greater than the difference between EP++ and Remin+ simulations because the combination of larger export with shallower recycling of POC is more efficient at storing carbon in the ocean than when the two changes are considered separately. For the global climate, the OA impacts of these small atmospheric CO$_2$ changes causes the global surface temperature to deviate by less than 0.25 K (Fig. 4).

4.2.2 Changes in BGC fields

While the OA impacts on the ocean carbon storage and climate by the end of this century are small, we now investigate whether the BGC fields in the ocean are significantly altered by OA. Key ocean BGC fields that have been shown to be impacted by global warming and OA are aragonite saturation state, lysocline depth, export production, dissolved oxygen and volume of suboxic water. We focus on how the OA impacts each of these fields relative to the REF simulation.
4.2.3 Aragonite saturation state

For the surface aragonite saturation state, there are only subtle differences among the simulations (Fig. 5). As expected, all simulations show a dramatic reduction in the surface aragonite saturation state by 2100, with the surface water poleward of approximately 40° S and 40° N being under-saturated with respect to aragonite (Fig. 5). At low latitudes, by 2100 the maximum aragonite saturation state from all simulations is less than 2.75, a level of saturation state that historically corals are not found (Guinotte et al., 2003). In all simulations, the upwelling region of the Eastern Equatorial Pacific shows a minimum in the tropical aragonite saturation state.

For the depth of the aragonite lysocline, all simulations show a dramatic shoaling in the polar regions (Fig. 6). In all simulations, the eastern tropical and subtropical Pacific lysocline is less than 100 m deep. Similarly, all simulations show lysocline depths of less than 100 m in the Indian Ocean. It is only in the equatorial western Pacific that the lysocline depths remain greater than 1000 m by the 2100. Note, this is the region, where the REF simulation over-estimated the lysocline depth by several hundred meters and in the OA experiments this region retains its resilience to change. This probably reflects a bias in the model, and it is anticipated that the region would show much greater shoaling by the end of the century (Bopp et al., 2013).

Overall the OA impacts only make subtle changes to the overall dramatic reduction in surface aragonite saturation state and lysocline depth projected with the RCP8.5 emission scenario.

4.2.4 Export production

For export of POC from the upper ocean, in the REF simulation there is a small global reduction (Fig. 7), which is mostly confined to the Equatorial Pacific, Indian Ocean and North Atlantic Oceans (Fig. 8). The export of PIC from the upper are also shown in Fig. 7, with all experiments showing a decline global export in the future except for REMIN+. In the REMIN+ simulation, the increase in POC export is associated
with an increase in PIC export. Interestingly, the different OA experiments show large regional differences in export production (Fig. 8), with EP++ and REMIN+ substantially increasing export production in the Southern Ocean. However, for the global integrated value, OA impacts all cause an increase in POC export production, with the greatest increase occurring in the COMB projection.

The large range in the export production response of the different OA experiments is consistent with previous studies (e.g. Tagliabue et al., 2011). The simulations reveal that OA impacts on export production are much greater than their impacts on climate change. Such behaviour demonstrates the consequence of OA will not be through its impact on climate change, but on how it impacts the flow of energy in marine ecosystems. These changes may have significant affects on marine ecosystems and their productivity, biodiversity, and our future ability to exploit them as a food resource.

4.2.5 Dissolved oxygen

While the OA impacts had a small effect on atmospheric CO$_2$ levels, the export production varied dramatically amongst the simulations (Fig. 8), which has the potential to alter dissolved oxygen levels in the ocean. With climate change, the REF simulation shows a decline in the mid-water dissolve oxygen levels in the North Pacific, Equatorial Pacific and Southern Oceans (Fig. 9). The oceanic oxygen levels are expected to decline under global warming (e.g. Matear et al., 2000; Bopp et al., 2002), because surface warming lowers the sea surface oxygen concentrations, enhances stratification, reduces ventilation of the thermocline, and reduces thermohaline circulation, which all tend to decrease the supply of oxygen to the ocean interior. The reduce oxygen supply to the ocean interior is also linked to increased residence time of water at depth, thereby enhancing biological oxygen consumption in the ocean interior.

The REF simulation projects a global decline in total dissolved oxygen inventory of 1.8% between 2006 and 2100 (Fig. 10b) and 2.5% between 1850 and 2100. The projected small decline is comparable to other ESMs projections, which show a small decrease of 2% to 4% by the end of 2100 with climate change (Cocco et al., 2013).
decline in dissolved oxygen in the major ocean basins is also reflected in a change in the thickness of the suboxic water (defined as water with dissolved oxygen of less than 5 µmol L\(^{-1}\)) (Fig. 10a). In the REF simulation, there is a projected 5% decline in suboxic water by 2100 (Fig. 10a). This is consistent with the multi-model analysis (Cocco et al., 2013), which showed the projected total volume of suboxic waters remain relatively unchanged (±5%) by the end of 2100 with climate change. As discussed by Cocco et al. (2013), the projected oxygen changes in the upper 1000 m of the ocean displayed a complex regional pattern with both positive and negative trends reflecting the complex interactions between changes in circulation, biological production, biological remineralization, and temperature.

The inclusion of OA impacts that could either increase POC export from the upper ocean (Fig. 7) and/or reduced its depth of remineralization may substantially decrease oxygen levels in the ocean interior. For the total oxygen inventory in the ocean, the simulations reveal that increased POC export causes a decline in oxygen while the shoaling of POC remineralization has little impact (Fig. 10b).

The EP+ simulation showed an increase in the thickness in the suboxic water of 17% by 2100 (Fig. 10). Similar simulations where the C/N ratio of exported POC matter is increased with OA have projected reduce dissolved oxygen levels in the ocean and comparable values for the increase in the volume of suboxic water to the EP+ projection (Oschlies et al., 2008; Tagliabue et al., 2011).

Oschlies et al. (2008) showed the volume of suboxic water is very sensitive to small changes in the remineralization of POC in the ocean interior. Our simulations confirm this result, but also show the volume of suboxic water is sensitive to where OA increases POC export. With the shoaling of POC remineralization (REMIN+) there is a reduction in the volume of suboxic water. By confining POC remineralization to the upper ocean, the equatorial Pacific has less suboxic water, because this water is now more influenced by air–sea gas exchange and there is a global decline in the volume of suboxic water (Fig. 10). Similarly, the EP++ simulation with increased export production shows the greatest decline in total oxygen inventory (Fig. 10b), but a decline
in the volume of suboxic water by 2100 (Fig. 10a). The COMB simulation, with its increased POC export and shoaling of the POC remineralization also projects a decline in the volume of suboxic water. However, COMB simulation projects the development of suboxic zone in the North Pacific and Southern Ocean where the combination of increased export production, shoaling of depth of POC remineralization, and increased stratification with global warming allows the develop of suboxic water in these regions.

The large uncertainty in the potential changes in POC export and remineralisation with OA, at present make it difficult to project the potential consequences of OA on dissolved oxygen levels with confidence, and this makes it a critical issue for further investigation.

5 Summary and perspectives

Ocean acidification is the inevitable consequence of rising atmospheric CO$_2$ and occurs in conjunction with global warming. Published studies have hypothesised the potential of OA to impact biogeochemical cycling in the ocean. However, to date no studies have combined these impacts to quantify the integrated impacts on ocean biogeochemistry and the feedbacks to the future climate. Here, we explore the integrated consequences of these OA induced changes using a coupled carbon-climate ESM and some first-order representations of the potential OA impacts on marine biogeochemical cycles. A key result of this study is that OA does not significantly alter the total carbon stored in the ocean, with the potential changes in atmospheric CO$_2$ levels (45 ppm maximum) being small compared to the future concentration projected with the RCP8.5 emission scenario by the end of this century (more than 1000 ppm). The small impact on the future atmospheric CO$_2$ means the OA impacts will have only a minor feedback on projected global warming, and our simulations suggest that by 2100 the global averaged surface temperature would be altered by less than 0.25 K. Therefore, while the simulations do project significant global warming ($\approx$ 3 K by 2100), the inclusion of OA impacts on the marine BGC cycle did not significantly alter the projected
changes. Consistent with both the small impact on carbon storage in the ocean and on global warming, the inclusion of OA impacts did not significantly alter the projected trajectory of future ocean acidification (e.g. surface aragonite state and lysocline depth). While the differences that occur by including OA impacts on biogeochemistry are small, we emphasize with the RCP8.5 scenario by 2100 there will be significant changes in ocean acidification, that will impact the marine ecosystem. All polar surface waters will be under-saturated with respect to aragonite, and the maximum surface aragonite saturation state in the tropics will be less than 2.75, a value below which coral reefs are not historically found.

Where OA has the potential to have a significant impact is on the POC and PIC export from the upper ocean. We emphasize the consequence of these changes on marine ecosystems is highly uncertain and needs further study. While deliberately conceived to be large, the changes in PIC and POC export did not significantly change the future depth of the lysocline, however, they did significantly change the regional export production and the interior oxygen levels.

The inclusion of OA impacts that could either increase POC export from the upper ocean or reduced its depth of remineralization could substantially decrease oxygen levels in the ocean interior. However, the large variability in potential changes in POC export with OA, at present make it difficult to confidently assess the consequences of OA on dissolved oxygen levels and therefore this is another important issue to addressed. The decline in oxygen with the rising CO₂ could also have consequences for marine organisms with high metabolic rates. Global warming, lower oxygen and higher CO₂ levels represent physiological stresses for marine aerobic organisms that may act synergistically with ocean acidification (Portner and Farrell, 2008). Understanding how OA and global warming impacts marine organisms warrants further investigation.

While CO₂ is the most important greenhouse gas modulated by the ocean, other greenhouse gases may be altered by OA. The next two most important greenhouse gas produced in the ocean are methane (CH₄) and nitrous oxide (N₂O), and their production in the ocean is linked to the remineralization of organic matter in low oxygen water
The decline in the interior oxygen levels should be associated with increased production of both these gases (Glessmer et al., 2009). However, it is expected the response of CH$_4$ and N$_2$O would be less than the CO$_2$ impact shown here. Consistent with this conclusion, Schmittner et al. (2008) used an ESM climate change projection run until year 4000 to show the expansion of suboxic water doubled N$_2$O production in the ocean. The net result was an increase in atmospheric concentrations by 60 ppb that caused a warming of about 0.25 K, a small change given the length of their simulation.

Enhanced dinitrogen (N$_2$) fixation by cyanobacteria occurs under elevated $p$CO$_2$ concentrations (Hutchins et al., 2009). This provides an increased source of reactive nitrogen (N) and has the potential to increase primary production in the oligotrophic tropical and subtropical areas. However, this response is limited, as the relieving of N limitation will ultimate lead to phosphate limitation, which will limit the potential carbon uptake. Ocean-only simulations where sufficient N was added to remove nitrate limitation gave a maximum reduction in atmospheric CO$_2$ of about 22 ppm by 2100 (Matear and Elliott, 2004). Again, it is a small effect when compared to the future atmospheric value projected with the RCP8.5 emissions scenario.

Iron is a biologically important element, and therefore any change in its bioavailability has the potential to change the growth rate of phytoplankton. At present there is little consensus on the sign of this change with OA. A slower Fe uptake by diatoms with OA is seen in experiments with Atlantic surface water (Shi et al., 2010), while an increase has been reported in coastal waters (Breitbarth et al., 2010). If we assume OA can increase the bioavailability of iron sufficiently to remove iron limitation on phytoplankton growth, we can use previous ocean model simulations to quantify the maximum potential increase in carbon storage. Such studies showed that this process alone could increased carbon storage in the ocean and reduce atmospheric CO$_2$ by 33 to 80 ppm by 2100 (Aumont and Bopp, 2006; Matear and Wong, 1999). While it is a larger response than what we project with our OA experiments, this is an upper bound of the potential feedback and it does not mechanistically link OA to the bioavailability of iron. Even this
upper bound estimate is small in comparison to the atmospheric CO$_2$ projected with the RCP8.5 emissions scenario by 2100 ($\approx$ 1000 ppm), hence it would only have a minor impact on the future climate. We emphasize that the OA impact on the bioavailability of iron is how it may impact biological production in the ocean. Further, iron is only one of many biologically important trace metals, for which their bioavailability will change in response to OA (Hoffmann et al., 2012) and potentially alter biological production. Therefore, more studies are required to understand how changes in the bioavailability of trace metals in response to OA may impact future biological production.

The ocean is also a source of climatically active trace gases to the atmosphere like dimethyl sulphide (DMS), which can alter cloud properties. DMS is a gaseous sulphur compound produced by marine biota in surface seawater (Gabric et al., 1993). The marine production of DMS provides 90% of the biogenic sulfur in the marine atmosphere, and in the atmosphere it is rapidly oxidised to produce particles that can affect cloud formation and climate (Arnold et al., 2013). The effects of increasing anthropogenic CO$_2$ and the resulting warming and ocean acidification on trace gas production in the oceans are poorly understood.

Modelling studies vary substantially in their predictions of the change in DMS emissions with climate change. Elevating CO$_2$ in isolation of other environmental change suggested a significant decrease in the future concentration of DMS (Hopkins et al., 2011). However, studies in polar waters suggested increases in DMS emission ranging from 30% to more than 150% (Cameron-Smith et al., 2011; Kloster et al., 2007; Gabric et al., 2011) by 2100 with only climate change. While a recent ESM study by Six et al. (2013) projected by 2100 a global decrease in DMS production of 18 ± 3%, with 83% of this change attributed to OA, leading to only a modest warming of 0.23 to 0.48 K. Six et al. (2013) simulated strong regional responses of increasing (polar regions) and decreasing DMS emissions, which reflected the combined affect of increased net primary production and regional shifts in community composition. Therefore, more studies combining the impacts of global warming and OA on marine DMS production are warranted.
to better determine its regional response and sign, particularly at the marine species and ecosystem levels.

Potential climate-carbon feedbacks of OA and global warming appear small relative to the huge input of carbon into the atmosphere by human activity. However, understanding and projecting the combined OA and global warming impact on marine ecosystems remains the outstanding issue to tackle. In particular, biological production may change with the projected OA, and the potential consequences for marine organisms and ecosystems are poorly known.

Appendix A

Ocean biogeochemical model equations

The ocean BGC module is based on Matear and Hirst (2003), and simulates the evolution of phosphate (P), oxygen (O), carbon (C) and alkalinity (A) in the ocean. The following briefly summarises the how the BGC processes in the ocean interior are parameterised and how they affect the 4 BGC tracers.

In the photic zone, which is set to be the surface layer of the model (upper 50 m), the biological production of particulate organic and inorganic matter occurs. For particulate organic matter, the production of particulate organic phosphorus (POP) was defined by the following equations:
$V_{\text{max}} = 0.6 (1.066)^T$  

$F(I) = [1 - e^{R(I)}]$  

$R(I) = \frac{I(x, t) \alpha \text{PAR}}{V_{\text{max}}}$  

$J(P) = S_{\text{npp}} V_{\text{max}} \text{Min} \left( \frac{P}{P + P_k}, F(I) \right) \Delta z$  

$V_{\text{max}}$ is the maximum growth rate in day$^{-1}$, which is a function of the surface layer temperature ($T$, °C). $F(I)$ is the productivity vs. irradiance equation used to describe phytoplankton growth, which is given as a unitless value and provides a measure of light limited growth. $R(I)$ a units less function of the light availability for growth, which is calculated from the incident short wave radiation ($I$) in W m$^{-2}$, the fraction of short wave radiation that is photosynthetically active PAR (unitless factor), and the initial slope ($\alpha$) of the productivity vs. radiance curve for phytoplankton growth (day$^{-1}$ (W m$^{-2}$)$^{-1}$). $J(P)$ gives the uptake of phosphate by POP production in mmol P m$^{-2}$ day$^{-1}$ in the photic zone. $J(P)$ is a function of the scaling parameter ($S_{\text{npp}}$) in mmol P m$^{-3}$, the thickness of the surface layer ($\Delta z$) in metres and the growth limitation function. The value $S_{\text{npp}}$ was set to 0.005 where it satisfactorily reproduced the observed phosphate and oxygen concentrations. The growth limitation function uses the minimum value of light and phosphate limited growth. The phosphate limited growth term is based on the phosphate concentration of the surface layer $P$ and the half saturation uptake ($P_k$) value for phosphate utilisation, which was set to 0.1 mmol P m$^{-3}$.

The POP production in the photic zone was linked to inorganic carbon, oxygen and alkalinity uptake in the photic zone by the following equations:

$J(C) = 106 \ J(P)$  

$J(O) = -136 \ J(P)$  

$J(A) = -16 \ J(P)$
PIC production in the model was linked to POC production by using a fixed rain ratio \( r \) of 9% (Yamanaka and Tajika, 1996) to give the following uptake of inorganic carbon and alkalinity in the photic zone:

\[
K(C) = r \times 106 \ J(P) \quad \text{(A8)}
\]

\[
K(A) = r \times 2 \times 106 \ J(P), \quad r = 0.09 \quad \text{(A9)}
\]

The POC and PIC produced in the photic zone was instantaneously remineralised in the ocean interior above where it was produced. The POC \( (J(C) \times \Delta z) \) and PIC \( (K(C) \times \Delta z) \) production in the photic zone was remineralised in the ocean interior using the following depth profiles of POC and PIC:

\[
POC(z) = \frac{z}{100m} \text{Remin}_{POC} \quad \text{(A10)}
\]

\[
PIC(z) = \exp(z/\text{Remin}_{PIC}) \quad \text{(A11)}
\]

where \( z \) is depth in meters, \( \text{Remin}_{POC} = -0.9 \) sets the reminalisation length scale of POC and \( \text{Remin}_{PIC} = 3500m \) sets the depth scale for PIC dissolution (Yamanaka and Tajika, 1996). From the POC and PIC depth profiles, the production of inorganic carbon from the remineralisation of POC \( (C_o) \) and the dissolution of PIC \( (C_i) \) is given by:

\[
C_o(z) = -J(C) \times \Delta z \times \frac{d}{dz}POC(z) \quad \text{(A12)}
\]

\[
C_i(z) = -K(C) \times \Delta z \times \frac{d}{dz}PIC(z) \quad \text{(A13)}
\]

with no remineralisation above 100 m and all the POC and PIC reaching the ocean bottom remineralised in the bottom layer of the model.

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Impact of ocean acidification on our future climate

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Table 1. Potential Ocean Acidification impacts on BGC cycles. The sign column denotes the impact on future climate with a positive value reflecting greater climate change (positive feedback).

| Biological Production Process | Impact on climate change |
|-------------------------------|---------------------------|
| Increased net primary productivity | – |
| Increased export of organic matter | – |
| Increased C/N ratio of organic matter | – |
| Reduced Calcification | – |

| Remineralization of Sinking Organic Matter Process | Impact on climate change |
|---------------------------------------------------|---------------------------|
| Increasing microbial POC remineralization | + |
| Reduced Ballasting with Calcium Carbonate | + |
Table 2. Summary of the ocean BGC experiments presented in this study. In the CRTL simulation, only the land and ocean carbon components of the ESM see the rising atmosphere CO₂, while the radiation properties of the atmosphere stay fixed at the pre-industrial atmospheric CO₂ level of 284.7 ppm. In the remaining simulations, both the climate and the carbon components experience the impact of rising atmospheric CO₂. The RCP8.5 simulation uses the prescribed atmospheric CO₂ as provided by the CMIP5 for this scenario. The REF simulation starts in 1850 using historical atmospheric CO₂ and switches to the RCP8.5 future emissions scenario at the start of 2006. The 4 other simulations (EP+, EP++, REMIN+ and COMB) start from the REF simulation at the end of 2005 and go until 2100 using the RCP8.5 future emissions. Please refer to the text for a description of how BGC cycle is modified in the EP+, EP++, REMIN+ and COMB simulations.

| Name   | Description                                                                 | Duration       |
|--------|------------------------------------------------------------------------------|----------------|
| CTRL   | Standard BGC with historical and RCP8.5 atmospheric CO₂ but with no climate change | 1850–2100      |
| RCP8.5 | Standard BGC with historical atmospheric CO₂ and RCP8.5 future atmosphere CO₂ | 1850–2100      |
| REF    | Standard BGC with historical atmospheric CO₂ and RCP8.5 future emissions     | 1850–2100      |
| EP+    | Increased C/P ratio of POC export and reduced PIC as CO₂ increased           | 2006–2100      |
| EP++   | EP+ and increased POC export as CO₂ increased                               | 2006–2100      |
| REMIN+ | Increased rate of POC and PIC remineralization as CO₂ increased              | 2006–2100      |
| COMB   | Combined affect of EP++ and REMIN+                                           | 2006–2100      |
Table 3. Summary statistics of the comparison of the REF 1995 simulated fields with the observations shown in Fig. 2.

| Field                                      | Observations vs. REF simulation in 1995 |
|--------------------------------------------|----------------------------------------|
|                                            | Observed Average | Simulated Average | Observed σ | Mean Error | Normalized RMS' | RMS | σ | Correlation Coefficient |
| Phosphate at 0 m (µmolL⁻¹)                 | 0.51             | 0.44              | 0.49        | −0.13      | 0.58            | 0.60 | 1.25 | 0.89 |
| Oxygen at 500 m (µmolL⁻¹)                  | 147.6            | 190.1             | 75.8        | 0.56       | 0.57            | 0.80 | 1.06 | 0.85 |
| Aragonite Saturation State at 0 m          | 3.01             | 2.75              | 0.84        | −0.31      | 0.36            | 0.47 | 1.09 | 0.95 |
| Lysocline Depth (m)                        | 1051.4           | 1414.5            | 717.6       | 0.51       | 0.61            | 0.79 | 1.02 | 0.81 |
Fig. 1. (a) Simulated change in global surface temperature using the historical and RCP8.5 atmospheric CO$_2$ concentration (Zhang et al., 2013), and using the RCP8.5 emission scenario (REF). In the REF simulation, the future emissions are prescribed (based on the RCP8.5 scenario) and our ESM determines the future atmospheric CO$_2$ concentration. While the RCP8.5 simulation uses the atmospheric CO$_2$ concentrations generated by Integrated Assessment Model to provide a standardised future atmospheric CO$_2$ concentrations for CMIP5. (b) Simulated global surface temperature change for the future period from the REF simulation.
Fig. 2. Taylor diagram of the comparison of the simulated fields with the observations for surface phosphate (1), dissolved oxygen at 500 m (2) (Boyer et al., 2009), surface aragonite saturation state (3) and lysocline depth (4). The colour bar gives the bias in the simulated fields normalised by the standard deviation in the observed field.
Fig. 3. Annual mean comparison of the observed and simulated fields for (a–b) surface aragonite saturation state (c–d) aragonite lysocline depth (n). The observations for aragonite saturation come from Key et al. (2004).
Fig. 4. (a) Change in atmospheric CO$_2$ (ppm) for the future period relative to the REF simulation for the various OA experiments (see Table 2 for a description of the experiments). (b) Simulated global ocean carbon uptake for the different OA experiments. (c) Simulated 10 yr running mean of the land carbon uptake by the different OA experiments. (d) Simulated 10 yr running mean of the change in global surface temperature relative to the REF simulation for the various OA experiments.
Fig. 5. Annual mean surface aragonite saturation state during the 2090–2100 period for (a) REF; (b) EP+; (c) EP++; (d) REMIN+; (e) COMB and (f) CTRL simulations.
Fig. 6. Annual mean depth of the aragonite lysocline (m) during the 2090–2100 period for: (a) REF; (b) EP+; (c) EP++; (d) REMIN+; (e) COMB and (f) CTRL simulations.
Fig. 7. (a) Projected global export of POC from the upper 100 m of the ocean. (b) Projected global export of PIC from the upper 100 m of the ocean.
Fig. 8. Export production (mol C m\(^{-2}\) y\(^{-1}\)) from the (a) CTRL in 1850; (b) the change in 2090–99 period for REF relative to (a). For the 2090–99 period the change relative to REF for (c) EP+; (d) EP++; (e) REMIN+; (f) COMB simulations.
Fig. 9. Dissolved oxygen (µmol L\(^{-1}\)) at 500 m for (a) CTRL in 1850 and the difference relative to (a) in the 2090–99 period for (b) REF. For the 2090–99 period the change relative to REF for (c) EP+; (d) EP++; (e) REMIN+; (f) COMB simulations.
Fig. 10. (a) Simulated change in the volume of suboxic water relative to the simulated present-day (2006) value. (b) Simulated change in the global ocean inventory of dissolved oxygen relative to the simulated 2006 value.