The influence of Southern Ocean winds on the North Atlantic carbon sink

Ben Bronselaer1, Laure Zanna1, David R. Munday1,2, and Jason Lowe3

1 Atmospheric, Oceanic and Planetary Physics, Department of Physics, University of Oxford, Oxford, UK, 2 British Antarctic Survey, Cambridge, UK, 3 Met Office Hadley Centre, Exeter, UK

Abstract Observed and predicted increases in Southern Ocean winds are thought to upwell deep ocean carbon and increase atmospheric CO2. However, Southern Ocean dynamics affect biogeochemistry and circulation pathways on a global scale. Using idealized Massachusetts Institute of Technology General Circulation Model (MITgcm) simulations, we demonstrate that an increase in Southern Ocean winds reduces the carbon sink in the North Atlantic subpolar gyre. The increase in atmospheric CO2 due to the reduction of the North Atlantic carbon sink is shown to be of the same magnitude as the increase in atmospheric CO2 due to Southern Ocean outgassing. The mechanism can be described as follows: The increase in Southern Ocean winds leads to an increase in upper ocean northward nutrient transport. Biological productivity is therefore enhanced in the tropics, which alters the chemistry of the subthermocline waters that are ultimately upwelled in the subpolar gyre. The results demonstrate the influence of Southern Ocean winds on the North Atlantic carbon sink and show that the effect of Southern Ocean winds on atmospheric CO2 is likely twice as large as previously thought in past, present, and future climates.

1. Introduction

Paleoproxy and historical data show large changes in atmospheric pCO2, indicating variations in the atmosphere-ocean exchange of carbon. Changes in atmospheric forcing, such as Southern Hemisphere westerly wind stress, can have a strong influence on the carbon cycle. The impact of observed and projected Southern Ocean wind increase [Swart et al., 2014; Bracegirdle et al., 2013] on the Southern Ocean carbon flux has been extensively studied [e.g., Le Quéré et al., 2007; Lovenduski et al., 2008; Lovenduski and Ito, 2009; Ito et al., 2010; Lauderdale et al., 2013; Munday et al., 2014]. Increased Southern Ocean winds are thought to increase upwelling of deep ocean natural carbon, elevating local ocean pCO2. The elevated ocean pCO2 leads to a reduction in local sink of anthropogenic CO2 and a subsequent increase in atmospheric CO2 levels.

However, little attention has gone into estimating the climate impacts of dynamical and biogeochemical changes induced by the Southern Ocean wind stress outside of the Southern Ocean. The Southern Ocean circulation forms part of the global meridional overturning circulation [Gnanadesikan, 1999; Marshall and Speer, 2012] and has a strong influence on transport pathways [Ito et al., 2010; Palter et al., 2010], including in the North Atlantic. Southern Ocean winds act to ventilate the deep ocean, which is richer in nutrients than the surface ocean. The wind-driven ventilation in the Southern Ocean mixes nutrients into the surface ocean, giving it a large effect on ocean biogeochemistry and biological productivity on a global scale [Marinov et al., 2006]. Anderson et al. [2002] and Sarmiento et al. [2004] show that increased Southern Ocean upwelling increases nutrient supply north of the Southern Ocean and enhances high- and low-latitude biological productivity. Changes induced in the Southern Ocean can therefore have far-reaching impact on ocean dynamical and biogeochemical properties.

Studies such as Lauderdale et al. [2013] examine the basin-scale response to Southern Ocean wind stress perturbations on the dynamics and distribution of carbon but focus on the local contribution of wind stress on atmospheric pCO2. The nonlocal effect of the Southern Ocean winds on atmospheric CO2, via change in global-scale transport pathways and in nutrient export from the Southern Ocean, remains to be estimated. We focus primarily on understanding the response of other major carbon sinks to changes induced in the Southern Ocean.

In the Atlantic, the major oceanic sink of atmospheric carbon (both natural and anthropogenic) is the subpolar gyre [Group, 1998; Dickson et al., 2008; Khatiwala et al., 2009; Kieke and Yashayaev, 2014]. Any changes in...
the North Atlantic ocean carbon chemistry will therefore greatly impact the magnitude of the local carbon sink and subsequently alter atmospheric CO$_2$ levels. Halloran et al. [2015], for example, show that the biogeochemical properties of the subpolar gyre are affected by the water advected from the tropics and Southern Ocean. Southern Ocean winds have the potential to influence the North Atlantic carbon sink by affecting the biogeochemical properties of the waters advected from the Southern Ocean to the North Atlantic.

The aim of this paper is to assess the nonlocal effect of Southern Ocean winds on atmospheric CO$_2$ and to demonstrate that increased Southern Ocean winds affect the North Atlantic pCO$_2$ through a novel mechanism. The mechanism can be summarized as follows: increased Southern Ocean wind stress enhances nutrient transport toward the tropics, thereby increasing tropical biological productivity. This in turn affects the chemistry of underlying subthermocline waters. The tropical subthermocline waters are subsequently advected to the North Atlantic subpolar gyre by equatorial currents and the meridional overturning circulation (MOC), reducing the capacity of the subpolar gyre to absorb atmospheric CO$_2$.

The paper is structured as follows. We provide a description of the model used in this paper in section 2, with details regarding the model design and mean state. Section 3 describes and quantifies the effect of Southern Ocean wind perturbations on the North Atlantic carbon sink. The subsequent section 4 will detail the key steps in the mechanism, and section 5 summarizes the main conclusion, caveats, and implications of the presented study.

### 2. Model Design and Control Experiment

To study the effect of Southern Ocean winds on the North Atlantic carbon sink, we use the Massachusetts Institute of Technology General Circulation Model (MITgcm) [Marshall et al., 1997] in an idealized configuration designed to mimic the Atlantic and Southern Oceans. The setup and parameters are similar to Munday et al. [2014] at 2° × 2° horizontal grid spacing with eddies parametrized using the Gent-McWilliams [Gent and McWilliams, 1990] scheme with a constant coefficient. The domain, shown in Figure 1a, is a sector 20° wide in longitude, extending to 60°N/S with a reentrant channel between 60° and 40°S to allow for a circumpolar current. The basin is 5000m deep, except at the channel boundaries where a 2500m deep sill represents Drake Passage. Coupled to the ocean, there is a well-mixed atmospheric box which only solves for atmospheric pCO$_2$, and the air-sea flux of carbon is parametrized following Wanninkhof [1992]. The model resolution is
chosen such that a large set of perturbation experiments can be run for centuries while still capturing the main features of a large-scale circulation and the appropriate response of the ocean carbon cycle. Such a setup is widely used to conduct large-scale ocean studies [e.g., Wolfe and Cessi, 2010; Ito and Follows, 2003; Munday et al., 2013].

The model is forced with a zonal wind stress profile shown in Figure 1b with $\tau_{SO} = 0.13 \text{ N/m}^2$ (where the subscript “SO” indicates the Southern Ocean between 60° and 40°S and the overline indicates a spatial average). Heat and freshwater fluxes are prescribed through temperature and salinity restoring profiles (Figures 1c and 1d), which broadly match the observed distributions in the Atlantic. The resulting residual (sum of Eulerian and eddy) overturning stream function is shown in Figure 1e [Marshall and Radko, 2003].

The model also supports a series of equatorial and midlatitude zonal currents, with a westward equatorial current, an eastward undercurrent [Knauss, 1960; Philander, 1973; Wyrski and Kilonsky, 1984; Philander et al., 1987], and a series of stationary eastward jets on either side of the equator [Tsuchiya, 1972; Firing et al., 1998; Rowe et al., 2000; Izumo et al., 2002; Qiu et al., 2013] (Figure 1f). While the structures are similar to observations, the magnitude of the currents in the MITgcm are smaller, likely due to the model geometry and its coarse horizontal and vertical resolution. The equatorial jets, alongside vertical and horizontal mixing, are important factors for ventilation of the shadow zone [Brandt et al., 2015]: a region of the subsurface ocean that is unventilated by the wind-driven gyre circulation.

The ocean model contains a biogeochemical component with seven tracers: dissolved inorganic carbon (DIC), alkalinity, preformed alkalinity, phosphate, preformed phosphate, dissolved organic phosphorous, and
Figure 3. Steady state biogeochemical quantities in the control experiment: (a) Mixed layer average biological activity (mmol/m³/year of phosphate), (b) mixed layer ocean pCO₂, and (c) air-sea CO₂ flux. Dashed black lines indicate the extent of the channel.

dissolved oxygen [Dutkiewicz et al., 2008; Follows et al., 2006]. Biological productivity is governed by phosphate and light limitation. Figures 2a–2c show the zonally averaged distribution of DIC, regenerated DIC, and dissolved oxygen. The maximum in DIC concentration is located in the abyssal cell at high latitudes where water is oldest. Regenerated DIC concentrations are generally high where dissolved oxygen concentrations are low since oxygen is used up in the remineralization of soft organic matter. The upper ocean distributions of DIC and alkalinity are in agreement with Global Data Analysis Project (GLODAP) climatology [Key et al., 2004].

The model shows the presence of an oxygen-minimum zone (OMZ) in the shadow zone between 100 and 1000 m depth, with corresponding high regenerated carbon concentrations. The lack of wind-driven ventilation in the shadow zone leads to a buildup of organic carbon and the associated removal of dissolved oxygen. The meridional and vertical extent of the OMZ compares well with other studies such as Ito and Follows [2005], although the magnitude is slightly higher in the presented model compared to other studies. For a more detailed description of the distribution of biogeochemical tracers in the model, see Munday et al. [2014].

Mixed layer biological activity is shown in Figure 3a, with higher activity at high latitudes than at the equator and little biological activity in the oligotrophic subtropical regions. Such patterns are supported by satellite chlorophyll observations and other models [Williams and Follows, 2011]. The efficiency of the biological pump in the model (given by the ratio of mixed layer nutrient concentration over the nutrient concentration at 100 – 200 m depth), at high latitudes is 0.1 – 0.2, while in the tropics it is between 0.6 and 0.8. These numbers are consistent with observations [Sarmiento et al., 2004]. The spatial pattern, including the large-scale meridional gradients of pCO₂, shown in Figure 3b, is comparable to observations [Takahashi et al., 2009; Valsala and Maksyutov, 2010]. As a result, there is a positive air-sea flux of carbon north of the southern channel and in the northern subpolar region (40°–60°N) and a negative air-sea flux of carbon in the southern channel and tropics, again consistent with observations [Takahashi et al., 2009] (Figure 3c). Overall, the large-scale distribution and magnitude of the biogeochemical tracers agrees well with other studies [e.g., Ito and Follows, 2005] and observations [Feely et al., 2001; Gruber, 1998].

3. The Response of the North Atlantic Carbon Sink to Increased Southern Ocean Wind Stress

3.1. The North Atlantic Carbon Flux

To examine the response of the carbon system to Southern Ocean wind stress perturbations, the wind stress between 60° and 20°S, τSO, is multiplied by a constant in a set of numerical experiments (see Figure 1b).
Each perturbed experiment is run for 400 years, after which time the dynamics of the upper 1000 m has fully equilibrated. We will examine the carbon cycle response to a range of wind stress perturbations, while specific examples will be drawn from the $2 \times \text{SO}_2$ scenario. The $2 \times \text{SO}_2$ experiment was run for 3000 years to verify that the results are not dependent on the length of integration. The response of the circulation to changes in Southern Ocean wind is linear. The Ekman transport and strength of the upper overturning cell increase linearly with wind stress. Our results therefore scale with forcing scenarios without nonlinear behavior.

Figure 4 illustrates the impact of ocean $\text{pCO}_2$, $\text{pCO}_2$, in the North Atlantic subpolar gyre on atmospheric $\text{pCO}_2$, $\text{pCO}_2$. Figure 4 shows time series of atmospheric $\text{pCO}_2$ (Figure 4a), the area-averaged ocean $\text{pCO}_2$ for the North Atlantic subpolar gyre ($40^\circ - 60^\circ N$, Figure 4b), and the Southern Ocean ($40^\circ - 60^\circ S$, Figure 4c) in the $2 \times \text{SO}_2$ wind stress perturbation experiment. The initial adjustment of the Southern Ocean $\text{pCO}_2$ occurs within the first few decades. The atmospheric $\text{pCO}_2$ shows an initial increase over a few decades which coincides with the time scale of the local response of the Southern Ocean $\text{pCO}_2$. However, the short-term response is followed by a long-term centennial adjustment in $\text{pCO}_2$ due to the perturbed ocean carbon reservoir in the North Atlantic. We refer to the atmospheric $\text{pCO}_2$ anomalies induced by the North Atlantic subpolar gyre as the nonlocal effect, with a time scale set by the adjustment of the North Atlantic subpolar gyre $\text{pCO}_2$.

The increase in North Atlantic $\text{pCO}_2$ shown in Figure 4b depends on the adjustment of the mean surface Revelle buffer factor $R$ (Figure 4d) [Revelle and Suess, 1957]. $R$ measures the fractional ocean $\text{pCO}_2$ change (denoted by $\delta$) given a fractional change in DIC and vice versa such that

$$ R = \frac{\delta \text{pCO}_2}{\delta \text{DIC}} \approx \frac{2 - \text{Alk}/\text{DIC}}{\text{Alk}/\text{DIC} - 1}. \tag{1} $$

where Alk is the alkalinity. The buffer factor $R$ quantifies the amount of carbon that can be stored in the form of DIC for a given change in atmospheric CO$_2$ concentration. High buffer factor systems are able to absorb changes in ocean $\text{pCO}_2$ by storing carbon in the form of DIC. For example, in a high buffer factor region which absorbs atmospheric carbon, local DIC does not change significantly. Therefore, more carbon remains in the form of dissolved CO$_2$, leading to higher $\text{pCO}_2$ values and less uptake of atmospheric CO$_2$. The value of $R$ can be approximated in terms of the ratio of alkalinity over DIC, as shown in equation (1). As alkalinity levels increase, the buffer factor decreases and the ocean becomes more capable of absorbing atmospheric CO$_2$ anomalies. Conversely, increasing DIC concentrations reduce the ocean’s capacity to buffer atmospheric CO$_2$ changes.
Figure 4d shows that the temporal evolution of North Atlantic $R$, corresponds to the change in North Atlantic $pCO_2$ shown in Figure 4b. Changes in the buffer factor account for 94% of the change in North Atlantic $pCO_2$, while other changes such as temperature and salinity account for the remainder (using the approach of Lovenduski et al. [2007]). As a result of the increase in North Atlantic $pCO_2$, the yearly air-sea flux of carbon in the region, $F_{\text{air-sea}}$, is reduced compared to that of the control run (Figure 4e). After roughly 425 years of the wind stress perturbation, the cumulative flux perturbation in the North Atlantic subpolar region becomes zero such that the North Atlantic subpolar region becomes a net emitter of carbon relative to the control state. The increase in Southern Ocean wind therefore induces a reduction in the North Atlantic carbon sink. The air-sea carbon flux in the Southern Ocean (Figure 4f) also shows a long-term decrease due to slow buildup of atmospheric $CO_2$ driven by the North Atlantic, since the Southern Ocean $pCO_2$ remains constant after 100 years of wind stress perturbation. After 400 years, the net change in the North Atlantic carbon flux is comparable in magnitude to the change in the Southern Ocean carbon flux.

3.2. Relative Feedback of the North Atlantic on Atmospheric $CO_2$

To quantify the strength of the nonlocal climate feedback on atmospheric $pCO_2$ due to Southern Ocean wind stress, we compare it with the change in atmospheric $CO_2$ due to Southern Ocean outgassing, i.e., the local feedback. We define the nonlocal feedback parameter $\epsilon$ which relates a change in Southern Ocean $pCO_2$, $\Delta pCO_{2SO}$, to the subsequent change in North Atlantic $pCO_2$, $\Delta pCO_{2NA}$. If the $pCO_2$ in the Southern Ocean is altered through changes in the large-scale circulation and biogeochemistry, the perturbed ocean dynamical and biogeochemical state will also cause changes in the $pCO_2$ elsewhere, such as in the North Atlantic (as shown in Figure 4). The relative changes in $pCO_2$, $\Delta pCO_{2SO}$, and $\Delta pCO_{2NA}$ will therefore be related to one another:

$$\Delta pCO_{2NA} = \epsilon \Delta pCO_{2SO}. \quad (2)$$

We can use $\epsilon$ to express the atmospheric $pCO_2$ anomaly, $\Delta pCO_{2}$, due to a change in Southern Ocean $pCO_2$, by considering the air-sea flux of carbon $F_{\text{air-sea}}$ over the global ocean. The air-sea flux equation takes the following form (Wanninkhof, 1992):

$$F_{\text{air-sea}} = k \left( pCO_{2}^0 - pCO_{2}^e \right), \quad (3)$$

where $k$ is the gas transfer velocity. We integrate equation (3) over the global ocean. At equilibrium, the net air-sea flux is zero such that $\int F_{\text{air-sea}} \, dA = 0$. However, we split the integral into three different regions: the Southern Ocean, the North Atlantic subpolar gyre, and the rest of the ocean basin, denoted by the subscripts SO, NA, and E, respectively. By considering a perturbation in atmospheric $pCO_2$, $\Delta pCO_{2}$, we can then write

$$\Delta pCO_{2} = \left( 1 + \frac{P_{\text{SO}}}{P_{\text{NA}}^0} \right) \Delta pCO_{2SO} + \Delta pCO_{2NA} + \Delta P_{\text{E}} pCO_{2E}, \quad (4)$$

where $P_{\text{SO}}$, $P_{\text{NA}}$, and $P_{\text{E}}$ are the area fractions of the respective regions, weighted by the local wind stress (see supporting information). The first term in equation (4) is the change in atmospheric $pCO_2$ due to changes in the ocean $pCO_2$. In the case of $\epsilon = 0$, there would be no nonlocal effect and the change in atmospheric $pCO_2$ would be due to change in Southern Ocean $pCO_2$ alone. If $\epsilon$ is positive and takes a value of 1, the net feedback is positive and doubles the local Southern Ocean feedback on atmospheric $pCO_2$, since $P_{\text{SO}}$ and $P_{\text{NA}}$ are roughly of equal magnitude. Conversely, if $\epsilon$ is negative with a value of $-1$, there would be an opposing nonlocal feedback which cancels out the local effect of the Southern Ocean wind stress. The second, third, and fourth terms in equation (4) quantify the change in atmospheric $pCO_2$ due to the enhanced turbulent air-sea transfer of $CO_2$. However, these terms mostly cancel each other out and only account for $5\%-10\%$ of the total $\Delta pCO_{2}$.

The value of $\epsilon$ can be estimated using both equations (2) and (4), since all terms apart from $\epsilon$ can be calculated and diagnosed from the MITgcm simulations. The results of the calculations are shown in Figure 5. While there are differences in the results between using equations (2) and (4), the order of magnitude and trend of the calculated values of $\epsilon$ are similar. Values of $\epsilon$ are positive and range between 0.5 and 0.9 for weak wind stress perturbations. The magnitude of $\epsilon$ increases as the wind stress increases; $\epsilon$ is roughly 1.2 for a doubling of Southern Ocean wind stress. For all experiments, $\epsilon$ increases with time (not shown): from less than 0.4 in the first decade after the wind stress perturbation is initially applied to the quasi-equilibrium values of roughly 1
Southern Ocean Winds, Equatorial Biology, and the North Atlantic Buffer Factor

A nutrient stream is an advective pathway through which nutrients are transported from regions of upwelling and nutrient excess, such as the Southern Ocean, to nutrient-poor regions such as the equator and the northern high latitudes (see Figure 2d). The advective pathways mainly follow streamlines of the overturning circulation. Ekman transport drives nutrients northward and out the Southern Ocean [Williams and Follows, 1998], where it is subducted below the mixed layer during the formation of Antarctic Intermediate Water. Transport is then predominantly isopycnal, with nutrients gradually reinducted into the mixed layer as isopycnals shoal toward the surface in the North Atlantic subpolar region [Williams et al., 2006]. North of the Southern Ocean where biology is far more efficient, the nutrient stream causes an increase in macronutrient-limited biological productivity [Sarmiento et al., 2004]. The nutrient stream \( S_p(x, y, z, t) \) is defined here as the meridional transport of phosphorus, in the form of phosphate \( \text{PO}_4^{3-} \), by the meridional residual velocity field \( v \) and is given by

\[
S_p(x, y, z, t) = [\text{PO}_4^{3-}] v, \tag{5}
\]

where the square brackets indicate concentration. The zonally averaged nutrient stream \( S_p \) is shown in Figure 2e for the MITgcm control run.

Physical transport in the ocean interior is mostly adiabatic along isopycnals, so the location of the nutrient stream lies along surfaces of constant density and its strength is modulated by the biological flux of nutrients [Pelegri et al., 1996]. In the current numerical setup, the maximum transport occurs along the potential density surface \( \sigma_\theta = 26 \text{ kg/m}^3 \) (\( \sigma_\theta^{26} \)), shown in Figure 2e for the control run. Therefore, we interpolate the meridional transport of phosphate onto \( \sigma_\theta^{26} \). Figure 6 shows the meridional nutrient transport, \( S_p \) (Figure 6a), the steady state concentration of \( \text{PO}_4^{3-} \) (Figure 6b) along \( \sigma_\theta^{26} \), and the mean mixed layer biological productivity, \( F_{\text{bio}} \) (Figure 6c). The spatial pattern of \( S_p \) shows the (positive) nutrient stream originating just north of the southern channel where intermediate mode water is formed in region 1. The nutrient stream follows the northward MOC transport along the western boundary in region 2, strengthening until it reaches the northern subpolar gyre, where the nutrients are inducted back into seasonal thermocline in region 3.

The effect of increased Southern Ocean winds on the nutrient stream is shown in Figures 6d–6f after 400 years of the \( 2\times\tau_{SO} \) perturbation experiment. Due to the increase in Southern Ocean wind-induced vertical mixing, there is a global increase in concentrations of \( \text{PO}_4^{3-} \) along the perturbed \( \sigma_\theta = 26 \text{ kg/m}^3 \) surface \( (\sigma_\theta^{26}) \) in the Southern Hemisphere (Figure 6e). The wind stress perturbation also causes an increase in the meridional velocity \( v \) along \( \sigma_\theta^{26} \). As a result, there is an increase in the northward nutrient stream (Figure 6d). The increase in surface nutrients leads to additional nutrients being introduced into the nutrient-limited equatorial shadow (as shown in Figure 5). The climate feedback of Southern Ocean wind stress on atmospheric \( p\text{CO}_2 \) is therefore almost doubled by the nonlocal effect.

To understand the reason behind the change in the North Atlantic carbon sink, the following sections will describe the mechanism by which Southern Ocean winds influence the North Atlantic buffer factor, \( \bar{R}_{\text{Na}} \), and set the long-term centennial adjustment time scale of atmospheric \( p\text{CO}_2 \) anomalies.


4.1. The Effect of Southern Ocean Winds on Nutrient Streams

In the Southern Ocean, surface biology is inefficient due to strong light and iron limitation [Boyd et al., 2000; Coale et al., 2004; DeVries et al., 2012] such that the upwelled nutrients remain mostly unused. Instead, the nutrients are transported northward and into the ocean interior following the subsurface currents in what are commonly referred to as nutrient streams [Pelegri and Csanady, 1991; Pelegri et al., 1996, 2006; Williams et al., 2006, 2011].
Figure 6. The nutrient stream (a, d) \( S_p \), (b, e) phosphorous on the \( \sigma_\theta = 26 \) kg/m\(^3\) surface, and (c, f) the mean mixed layer biological productivity. Figures 6a–6c are for the control run, and Figures 6d–6f are after 400 years of a \( 2 \times \tau_{SO} \) perturbation. Only fields north of the southern channel are shown for clarity.

... zone in region 4 via eastward equatorial jets and mixing across the southern shadow zone boundary. Subsequently, mixed layer biological productivity is strongly enhanced north of the southern channel and in the eastern tropics where nutrients are upwelled due to wind stress divergence. For the \( 2 \times \tau_{SO} \) experiment, the increase in biological productivity is less than 10% in the southern channel, between 20% and 40% in the subpolar gyre, but more than 200% between 20\(^\circ\)S and 20\(^\circ\)N. In the northern subpolar gyre, the response in biological productivity is not as strong as that in the tropics. This relatively weak response in the subpolar gyre is partly due to high-latitude light limitation [Nielsdottir et al., 2009] and partly due to the depletion of the nutrient stream by the time it reaches the northern end of the basin. Most of the increase in biology is therefore centered around the eastern tropics.

4.2. Surface Biology and the Revelle Buffer Factor

An increase in equatorial biology can reduce atmospheric \( pCO_2 \) by drawing more carbon from the mixed layer to form biological tissue, therefore reducing ocean surface \( pCO_2 \) and increasing the air-sea flux of carbon into the ocean. Conversely, it can increase atmospheric \( pCO_2 \) by affecting the chemistry of subthermocline waters in the shadow zone. Surface biology utilizes DIC and alkalinity at different rates. To understand the effect of biology on the buffer factor, let us consider the influence of mixed layer biology on the mixed layer chemistry. The key points discussed below are also summarized in the diagram in Figure 7.

Production of soft matter reduces DIC by 117 mol for every mole of phosphate and increases alkalinity by 16 mol, following Redfield stoichiometry [Redfield, 1934; Anderson and Sarmiento, 1994; Weber and Deutsch, 2012]. The formation or dissolution of 1 mol of calcite (hard tissue), on the other hand, reduces DIC by 1 mol and...
alkalinity by 2 mol. Surface biology therefore changes the local concentrations of DIC and Alk depending on the relative export and remineralization of soft matter and calcite. The relative export of hard to soft matter, $r$, is known as the “rain ratio.” Studies of climatological basin-wide and local seasonal data give a rain ratio between 0.05 and 0.1 in the open ocean; the former is used in the MITgcm experiments conducted here [Sarmiento et al., 2002; Jin et al., 2006; Emerson et al., 2011]. For all values of $r$, an increase in surface biology results in a decrease of the relative ratio of DIC/Alk within the mixed layer.

Surface biology also affects the vertical transport of DIC and alkalinity in the ocean. The soft matter that is produced in the ocean mixed layer falls through the water column until it is remineralized back into DIC, in the form of $C_{\text{soft}}$. The remineralization of soft matter therefore increases DIC and reduces alkalinity. Similarly, calcite shells produced in the mixed layer fall through the water column until they are dissolved into DIC in the form of $C_{\text{carb}}$. Dissolution of calcite increases DIC and alkalinity. The remineralization depth of soft matter is shallower than hard tissue such that just below the mixed layer, remineralization of organic matter is mostly due to soft tissue which increases the relative ratio of DIC to Alk.

The relative concentration of DIC and alkalinity of water parcels, influenced by surface biology, is pivotal for the climate system since they determine the value of the Revelle buffer factor $R$ and therefore the capacity of the ocean to absorb atmospheric CO$_2$ (equation (1)). In regions of high biological activity, surface waters are reduced in $R$ and the underlying waters are therefore enhanced in $R$. If these water parcels are then advected away and upwelled at the surface elsewhere, they will increase $R$ in the region of upwelling. Water parcels with an elevated buffer factor $R$ will be less capable of absorbing atmospheric CO$_2$ anomalies, leading to a local reduction in the ocean carbon sink.

As Southern Ocean winds increase and surface biological productivity is enhanced across the global ocean, the strongest increase in the buffer factor $R$ occurs below the thermocline in the equatorial shadow zone along $\sigma_0$ (Figure 8), where the largest relative changes in DIC and Alk occur. After 50 years (Figure 8a), there is an increase in $R$ in the southern channel associated with increased upwelling of DIC-rich waters (region 1). A positive $R$ anomaly occurs in the equatorial shadow zone due to the increased biological activity (region 2). After another 100 years (Figure 8b), the anomaly in $R$ in the southern channel has decayed, while the tropical anomaly signal intensified. Because ventilation of the shadow zone is sluggish compared to the rest of the ocean, the increase in the downward flux of organic matter from the surface causes a slow buildup of DIC and a slow reduction of alkalinity in the shadow zone leading to a rise of $R$ anomaly for centuries.

The increase in $R$ generated in the eastern tropics is transported predominantly westward within the shadow zone through the equatorial current, where it is then entrained by the nutrient stream in the western boundary of the shadow zone. As shown in Figure 8d, an additional recirculation of the $R$ anomaly within the shadow zone decreases the local concentrations of DIC and Alk depending on the relative export and remineralization of soft matter and calcite. The relative export of hard to soft matter, $r$, is known as the “rain ratio.” Studies of climatological basin-wide and local seasonal data give a rain ratio between 0.05 and 0.1 in the open ocean; the former is used in the MITgcm experiments conducted here [Sarmiento et al., 2002; Jin et al., 2006; Emerson et al., 2011]. For all values of $r$, an increase in surface biology results in a decrease of the relative ratio of DIC/Alk within the mixed layer.

Surface biology also affects the vertical transport of DIC and alkalinity in the ocean. The soft matter that is produced in the ocean mixed layer falls through the water column until it is remineralized back into DIC, in the form of $C_{\text{soft}}$. The remineralization of soft matter therefore increases DIC and reduces alkalinity. Similarly, calcite shells produced in the mixed layer fall through the water column until they are dissolved into DIC in the form of $C_{\text{carb}}$. Dissolution of calcite increases DIC and alkalinity. The remineralization depth of soft matter is shallower than hard tissue such that just below the mixed layer, remineralization of organic matter is mostly due to soft tissue which increases the relative ratio of DIC to Alk.

The relative concentration of DIC and alkalinity of water parcels, influenced by surface biology, is pivotal for the climate system since they determine the value of the Revelle buffer factor $R$ and therefore the capacity of the ocean to absorb atmospheric CO$_2$ (equation (1)). In regions of high biological activity, surface waters are reduced in $R$ and the underlying waters are therefore enhanced in $R$. If these water parcels are then advected away and upwelled at the surface elsewhere, they will increase $R$ in the region of upwelling. Water parcels with an elevated buffer factor $R$ will be less capable of absorbing atmospheric CO$_2$ anomalies, leading to a local reduction in the ocean carbon sink.

As Southern Ocean winds increase and surface biological productivity is enhanced across the global ocean, the strongest increase in the buffer factor $R$ occurs below the thermocline in the equatorial shadow zone along $\sigma_0$ (Figure 8), where the largest relative changes in DIC and Alk occur. After 50 years (Figure 8a), there is an increase in $R$ in the southern channel associated with increased upwelling of DIC-rich waters (region 1). A positive $R$ anomaly occurs in the equatorial shadow zone due to the increased biological activity (region 2). After another 100 years (Figure 8b), the anomaly in $R$ in the southern channel has decayed, while the tropical anomaly signal intensified. Because ventilation of the shadow zone is sluggish compared to the rest of the ocean, the increase in the downward flux of organic matter from the surface causes a slow buildup of DIC and a slow reduction of alkalinity in the shadow zone leading to a rise of $R$ anomaly for centuries.

The increase in $R$ generated in the eastern tropics is transported predominantly westward within the shadow zone through the equatorial current, where it is then entrained by the nutrient stream in the western boundary of the shadow zone. As shown in Figure 8d, an additional recirculation of the $R$ anomaly within the shadow zone decreases the local concentrations of DIC and Alk depending on the relative export and remineralization of soft matter and calcite. The relative export of hard to soft matter, $r$, is known as the “rain ratio.” Studies of climatological basin-wide and local seasonal data give a rain ratio between 0.05 and 0.1 in the open ocean; the former is used in the MITgcm experiments conducted here [Sarmiento et al., 2002; Jin et al., 2006; Emerson et al., 2011]. For all values of $r$, an increase in surface biology results in a decrease of the relative ratio of DIC/Alk within the mixed layer.

Surface biology also affects the vertical transport of DIC and alkalinity in the ocean. The soft matter that is produced in the ocean mixed layer falls through the water column until it is remineralized back into DIC, in the form of $C_{\text{soft}}$. The remineralization of soft matter therefore increases DIC and reduces alkalinity. Similarly, calcite shells produced in the mixed layer fall through the water column until they are dissolved into DIC in the form of $C_{\text{carb}}$. Dissolution of calcite increases DIC and alkalinity. The remineralization depth of soft matter is shallower than hard tissue such that just below the mixed layer, remineralization of organic matter is mostly due to soft tissue which increases the relative ratio of DIC to Alk.

The relative concentration of DIC and alkalinity of water parcels, influenced by surface biology, is pivotal for the climate system since they determine the value of the Revelle buffer factor $R$ and therefore the capacity of the ocean to absorb atmospheric CO$_2$ (equation (1)). In regions of high biological activity, surface waters are reduced in $R$ and the underlying waters are therefore enhanced in $R$. If these water parcels are then advected away and upwelled at the surface elsewhere, they will increase $R$ in the region of upwelling. Water parcels with an elevated buffer factor $R$ will be less capable of absorbing atmospheric CO$_2$ anomalies, leading to a local reduction in the ocean carbon sink.

As Southern Ocean winds increase and surface biological productivity is enhanced across the global ocean, the strongest increase in the buffer factor $R$ occurs below the thermocline in the equatorial shadow zone along $\sigma_0$ (Figure 8), where the largest relative changes in DIC and Alk occur. After 50 years (Figure 8a), there is an increase in $R$ in the southern channel associated with increased upwelling of DIC-rich waters (region 1). A positive $R$ anomaly occurs in the equatorial shadow zone due to the increased biological activity (region 2). After another 100 years (Figure 8b), the anomaly in $R$ in the southern channel has decayed, while the tropical anomaly signal intensified. Because ventilation of the shadow zone is sluggish compared to the rest of the ocean, the increase in the downward flux of organic matter from the surface causes a slow buildup of DIC and a slow reduction of alkalinity in the shadow zone leading to a rise of $R$ anomaly for centuries.

The increase in $R$ generated in the eastern tropics is transported predominantly westward within the shadow zone through the equatorial current, where it is then entrained by the nutrient stream in the western boundary of the shadow zone. As shown in Figure 8d, an additional recirculation of the $R$ anomaly within the shadow zone decreases the local concentrations of DIC and Alk depending on the relative export and remineralization of soft matter and calcite. The relative export of hard to soft matter, $r$, is known as the “rain ratio.” Studies of climatological basin-wide and local seasonal data give a rain ratio between 0.05 and 0.1 in the open ocean; the former is used in the MITgcm experiments conducted here [Sarmiento et al., 2002; Jin et al., 2006; Emerson et al., 2011]. For all values of $r$, an increase in surface biology results in a decrease of the relative ratio of DIC/Alk within the mixed layer.

Figure 7. Diagram of the influence of surface biology on the Revelle buffer factor $R$. As mixed layer biological activity increases, mixed layer DIC is reduced and alkalinity is increased, lowering the buffer factor. Underneath the mixed layer to roughly 1000 m depth, more soft matter is remineralized compared to hard tissue, leading to an increase in DIC and a reduction of alkalinity, resulting in a strong increase in the buffer factor. Deep in the water column, remineralization of soft and hard tissue is comparable giving an increase in both DIC and alkalinity, resulting in a more moderate increase in the buffer factor.
Figure 8. Revelle buffer factor anomaly ($2 \times \sigma_\text{SO}, \text{control}$) along the perturbed $\sigma_\text{p} = 26 \text{ kg/m}^3$ surface after (a) 50, (b) 150, (c) 250, and (d) 400 years. The blue arrows in Figure 8d show the perturbed flow field along the $\sigma_\text{p26}$ surface. The black arrows show the primary advection pathway out of the equatorial shadow zone, while the dashed arrows show a secondary advection pathway due to recirculation within the shadow zone.

zone itself is present, leading to further advection toward the southern edge of the shadow zone. After the $R$ anomaly is entrained into the nutrient stream, it follows the MOC transport northward (region 3) into the subtropical gyre where it is upwelled into the surface. The positive $R$ anomaly in the intermediate waters just north of the Southern Ocean, denoted by region 4 in Figure 8c, is only 10% to 30% of the equatorial signal since the intermediate waters are ventilated at the surface prior to subduction.

4.3. Verification of the Mechanism

The effects of Southern Ocean mixing and nutrient export on low- and high-latitude biological productivity have been extensively studied [Anderson et al., 2002; Sarmiento et al., 2004; Marinov et al., 2006]. However, the

Figure 9. Passive dye tracer along the unperturbed $\sigma_\text{p} = 26 \text{ kg/m}^3$ surface after (a) 0, (b) 10, and (c) 20 years in an unperturbed control simulation.
advective pathways of biogeochemical tracers from the equatorial shadow zone to the North Atlantic and their impact on the subpolar gyre buffer factor are fairly novel. Therefore, we run an additional experiment in which we release a passive dye tracer into the shadow zone along \( \sigma_{26} \) where concentrations of oxygen are less than 0.17 mol/m\(^3\) in the control simulation (the oxygen-minimum zone). Figure 9a shows the initial distribution of the tracer along \( \sigma_{26} \), while Figures 9b and 9c show the dye concentration after 10 and 20 years, respectively. The tracer is being entrained near the western boundary and advected northward toward the subpolar gyre, where it is then mixed into the surface ocean. The advective time scale from the shadow zone to the North Atlantic subpolar gyre is roughly 15 years.

Figures 8 and 9 show that the positive \( R \) anomaly from the shadow zone is advected to the North Atlantic subpolar gyre. As discussed earlier, increases in surface biological activity in the subpolar gyre can reduce the local mixed layer buffer factor. However, by the time the enhanced nutrient stream reaches the subpolar gyre it is nearly stripped of the nutrient surplus so that the increase in biological activity in the North Atlantic subpolar gyre is not sufficient to reduce the positive \( R \) anomaly of the upwelled waters. The result is therefore an increase in the North Atlantic subpolar gyre buffer factor due to the increased Southern Ocean wind stress.

Following this argument, the net change in the North Atlantic buffer factor \( R_{\text{na}} \) should depend on the relative change in the ratio of equatorial and North Atlantic biological productivity, such that

\[
R_{\text{na}} \propto \frac{F_{\text{bio, eq}}}{F_{\text{bio, na}}}.
\]

where the overbar refers to the spatial averages between 10\(^{\circ}\)S and 40\(^{\circ}\)N for the subscript “eq” and between 40\(^{\circ}\) and 60\(^{\circ}\)N for the subscript “na.” Since an increase in the buffer factor \( R_{\text{na}} \) reduces the capacity of the North Atlantic to absorb atmospheric \( \text{CO}_2 \), we also expect that the atmospheric \( p\text{CO}_2 \) will scale with \( R_{\text{na}} \) such that

\[
p\text{CO}_2 \propto R_{\text{na}}.
\]

Using the MITgcm numerical simulations in which the wind stress is perturbed, we will explore the validity of equations (6) and (7). We run further simulations to explore the sensitivity of the results using wind stress perturbation experiments with a warmed spin-up state, with increased ocean stratification. The model is warmed by increasing the temperature restoring profile shown in Figure 1c by 30\% across the domain. The warmed model reaches a new equilibrium after 3000 years, at which time we introduce the same Southern Ocean wind stress perturbations described in section 2.2, up to a 3 \( \times \) \( \tau_{\text{SO}} \) perturbation.

Furthermore, experiments were also run whereby only the biogeochemical component in MITgcm is perturbed while keeping the wind stress profile constant. The parameters altered are the nutrient limitation
half-saturation constant $K_{na}$ [Monod, 1949] and the light limitation half-saturation constant $I_{l}$ [Jassby and Platt, 1976]. The effect of perturbing the biogeochemical component is to increase or decrease biological activity by an amount which depends on the local light or nutrient limitation, without affecting the circulation so that the changes seen in $F_{bio,n}^{eq}$ and $F_{bio,na}$ are purely biogeochemical. Increasing $K_{na}$ causes areas that were not limited in macronutrients (e.g., high latitudes) to become limited, effectively increasing $F_{bio,na}^{eq}$. On the other hand, an increasing $I_{l}$ reduces light limitation more at high latitudes compared to the tropics, giving a reduction in $F_{bio,n}^{eq}$. The nutrient and light half-saturation perturbation experiments are designed to show that an increase in atmospheric $pCO_2$ can also be caused by perturbations in the biology due to factors other than wind stress. The proposed feedback mechanism is therefore not solely dependent on Southern Ocean wind stress but can be caused by other climate change-induced factors that affect surface biological activity, such as freshwater or micronutrient input.

The results for all numerical experiments to test equations (6) and (7) are shown in Figure 10. Figure 10a shows that changes in $F_{bio,na}^{eq}$ induce a linear change in $R_{na}$ relative to the control state, $\Delta R_{na}$, following the prediction of equation (6). The relationship is linear for perturbations less than 60% of the mean state. Furthermore, Figure 10b shows that $\Delta pCO_2$ varies linearly with $\Delta R_{na}$ as predicted, where $\Delta$ indicates again the anomaly relative to the control state. The sensitivity of the atmospheric $pCO_2$ anomaly to changes in the North Atlantic buffer factor in Figure 10b depends on the control value of $pCO_2$ and DIC concentration values, explaining the different slopes. The MITgcm experiments therefore show that an increase in tropical biological activity relative to northern subpolar gyre biological activity results in an increase in atmospheric $pCO_2$, under a variety of wind stress and biogeochemical forcing scenarios.

5. Conclusion and Discussion

5.1. Summary of Findings

The goal of this paper is to quantify the nonlocal equilibrium climate feedback of Southern Ocean wind stress on atmospheric $pCO_2$, $pCO_2^*$, using idealized coarse-resolution MITgcm experiments. In addition to triggering local changes in the Southern Ocean, an increase in the westerlies over the Southern Ocean can lead to changes in the biogeochemical and dynamical quantities in the Northern Hemisphere. Primarily, the North Atlantic carbon sink is found to be reduced several decades after an increase in Southern Ocean winds. The resulting feedback on atmospheric $pCO_2$, measured by the nonlocal feedback parameter $\epsilon$, is of roughly the same order as the local Southern Ocean feedback. Overall, the nonlocal effect of Southern Ocean wind stress perturbations on atmospheric $pCO_2$ magnifies the local effect by 80% to 110%. The reduction in the North Atlantic carbon sink therefore constitutes a significant climate feedback that cannot be measured by considering changes in the Southern Ocean carbon sink alone.

Driving the reduction in the North Atlantic carbon sink is an increase in the Revelle buffer factor. It is found that wind stress-induced upwelling strengthens the export of nutrients north of the Southern Ocean by increasing the nutrient stream and increases equatorial biological productivity. The increased equatorial biological productivity increases the Revelle buffer factor, $R$, in the eastern tropical shadow zone/oxygen-minimum zone located between 100 and 700m depth. Via ocean interior advective pathways, the waters enriched in $R$ are entrained into the nutrient stream and transported to the North Atlantic where they are inducted into the seasonal thermocline, increasing the subpolar gyre buffer factor and ocean $pCO_2$, $pCO_2^*$. Due to the reduction in the North Atlantic carbon sink, atmospheric $pCO_2$ is increased. The mechanism is also verified by only perturbing the biogeochemical component of the model.

5.2. Caveats and Implications

The use of a coarse-resolution model introduces several caveats. Coarse-resolution models fail to accurately represent the extent of the oxygen-minimum zones. Due to the unresolved complexity in equatorial undercurrents, a phenomenon coined "nutrient trapping" [Najjar, 1990; Aumont et al., 1999] results in overestimation of the strength of the oxygen minimum in the tropics. Nonetheless, the proposed mechanism should still be valid since oxygen-minimum zones are observed in the ocean [Stramma et al., 2009]. However, nutrient trapping could amplify the strength of the nonlocal feedback due to an exaggerated biogeochemical response in the tropics. Unresolved eddies in the Southern Ocean also overestimate the wind stress-induced increase in nutrient upwelling, since a constant Gent-McWilliams coefficient is used in the model. However, both Southern Ocean $pCO_2^*$ and the relative response of the North Atlantic and equatorial biology depend equally
on upwelling in the Southern Ocean so we do not expect this to affect the relative magnitude of the nonlocal feedback, although it may affect the equilibration time scale. For a full appreciation of the effect of eddies, a high-resolution eddy-permitting or eddy-resolving simulation is required.

Lastly, the geometry of the sector model introduces several caveats: the size of the basin plays an important role in setting the equilibration time scale of the nonlocal Southern Ocean winds feedback on pCO$_2$. The feedback equilibration time scale depends on basin-wide advection: the buffer factor anomaly in the shadow zone has to be advected zonally from east to west across a larger oxygen-minimum zone before it is advected northward by the nutrient stream. To illustrate this dependence of the time scale on basin width, an identical MITgcm setup 40° wide in longitude is used where the equilibration takes twice as long in the 40° longitude model (see supporting information). However, the strength of the nonlocal feedback $e$ is similar. The width of the basin therefore determines the time scale of the nonlocal feedback but not the magnitude. Further study is required to determine the exact equilibration time scale with a realistic geometry. The model also lacks a Pacific Ocean and Indian Ocean. However, the lack of MOC-like large-scale overturning transport and lack of strong ocean carbon uptake in the Pacific and Indian basins means that the presence of these basins will not affect the magnitude of the feedback of Southern Ocean winds on atmospheric CO$_2$ via the North Atlantic subpolar gyre. The increase in biological carbon drawdown in the Pacific and Indian basins as a result of increased nutrient transport is likely to be small due to the simultaneous input of surface carbon.

The relationship between the North Atlantic buffer factor and equatorial biology has further implications for the climate as well, since the ratio of North Atlantic to equatorial biology can be altered by many processes such as stratification, cloud cover, other sources of macronutrients, and iron supply, to name a few. Our results show the influence of oxygen-minimum zones on North Atlantic surface chemistry. Given recent results from studies that show expansion of tropical oxygen-minimum zones [Stramma et al., 2012], a thorough understanding of the biogeochemical drivers of the North Atlantic buffer factor is therefore needed. The North Atlantic subpolar region constitutes one of the main sinks for anthropogenic CO$_2$, and our results show that changes in the buffer factor of as small as 0.5 could result in a positive atmospheric carbon feedback of 10–30ppm.

Furthermore, Southern Ocean winds and vertical mixing have been used to partially explain glacial-interglacial cycles of atmospheric CO$_2$. Francois et al. [1997] and Anderson et al. [2009] suggest that there was an increase in vertical mixing in the Southern Ocean at the end of the Last Glacial Maximum (LGM), 20,000–12,000 years B.P., possibly related to an increase in wind stress. So far, however, current theories of physical mechanisms have not been able to fully explain the glacial-interglacial cycles of atmospheric CO$_2$ [Sigman and Boyle, 2000; Solomon, 2007]. Including the effects of the nonlocal atmospheric feedback of the North Atlantic due to Southern Ocean vertical mixing could aid our understanding of glacial-interglacial atmospheric CO$_2$ cycles.

Kohfeld et al. [2005] show that there was higher biological carbon export in the Atlantic equatorial shadow zone in the LGM but states that "these areas are relatively small and do not affect the overall picture," while our research shows that these areas should not be overlooked. Although global changes in ocean alkalinity have been acknowledged as being potentially important, there has been no consideration of spatial patterns. Kohfeld et al. [2005] does not give exact figures of the relative change in biological export in the tropics and North Atlantic. However, Francois et al. [1997] suggest that Southern Ocean vertical mixing was increased by 20% to 50% at the end of the LGM, which, given our model study, would correspond to an increase in atmospheric pCO$_2$ of 15–35ppm in addition to the local Southern Ocean outgassing.

References


Acknowledgments
B.B. was supported by a NERC CASE studentship with the Met Office. Further support from NERC was provided to L.Z. and D.R.M. This work made use of the facilities of HECToR and Archer. We acknowledge the MITgcm team for making their code publicly available. We also like to thank the reviewers for their helpful suggestions and comments. Details and results from the numerical simulations are available from B.B.


Le Quere, C., et al. (2007), Saturation of the Southern Ocean CO$_2$ sink due to recent climate change, Science, 316, 1735–1738.


Najjar, R. G. (1990), Simulations of the phosphorus and oxygen cycles in the world ocean using a general circulation model, PhD thesis, Princeton, N. J.


Revelle, R., and H. S. Suss (1957), Carbon dioxide exchange between atmosphere and ocean and the question of an increased atmospheric CO$_2$ during the past decades, Tellus, 9, 18–27.


Supporting Information for ‘The Influence of Southern Ocean Winds on the North Atlantic Carbon Sink’

Ben Bronselaer\textsuperscript{1}, Laure Zanna\textsuperscript{1} David R. Munday\textsuperscript{1,2}, and Jason Lowe\textsuperscript{3}

Here we present a detailed derivation of the non-local feedback parameter and atmospheric feedback equation described in the main paper. In addition, we present data from a Southern Ocean wind stress perturbation experiment performed with an identical MITgcm set-up as used in the main paper, but 40° in longitude.

Corresponding author: B. Bronselaer, Atmospheric, Oceanic and Planetary Physics, Department of Physics, University of Oxford, Oxford, OX1 3PU, UK. (benjamin.bronselaer@physics.ox.ac.uk)

\textsuperscript{1}Atmospheric, Oceanic and Planetary Physics, Department of Physics, University of Oxford, Oxford, OX1 3PU, UK.

\textsuperscript{2}British Antarctic Survey, Madingley Road, Cambridge, CB3 0ET, UK

\textsuperscript{3}Met Office Hadley Centre, FitzRoy Road, Exeter, Devon EX1 3PB, UK
The value of oceanic $p\text{CO}_2$, $p\text{CO}_2^o$, as a function of a forcing $\tau$, $p\text{CO}_2^o(\tau)$ is obtained by using the equation for global air-sea carbon flux, $F_{\text{air-sea}}$, in the absence of anthropogenic carbon emissions. At equilibrium, the globally area integrated flux over the whole ocean vanishes such that:

$$\int F_{\text{air-sea}}dA = \int K\tau(p\text{CO}_2^o - p\text{CO}_2^o(\tau))dA = 0,$$

where $\tau$ is proportional to the square of the wind speed. $K$ measures the turbulent air-sea gas transfer and solubility such that $K\tau = k$, where $k$ is the piston velocity of Wanninkhof [1992] which varies as the square of the surface wind speed. $p\text{CO}_2^o$ is the atmospheric partial pressure of CO$_2$. We are interested in the Southern Ocean and North Atlantic $p\text{CO}_2^o$ response to local forcing so we split the integral into a Southern Ocean area $A_{SO}$ where the forcing $\tau$ is applied, the North Atlantic $A_{NA}$ and the remaining area in the ocean, $A_E$ (see Fig. S1):

$$\tau_{SO}p\text{CO}_2^o_{SO}(\tau) + \tau_{NA}p\text{CO}_2^o_{NA}(\tau) + A_E\tau_{E}p\text{CO}_2^o_{E} - (A_{SO}\tau_{SO} + A_{NA}\tau_{NA} + A_E\tau_{E})p\text{CO}_2^o = 0,$$

where the overline indicates an area-mean. Equation 2 can be solved for $p\text{CO}_2^o$ to give $p\text{CO}_2^o$ in terms of $p\text{CO}_2^o_{SO}$ and $p\text{CO}_2^o_{NA}$. We approximate $\tau_{SO}p\text{CO}_2^o_{SO}(\tau)$ by $\tau_{SO}p\text{CO}_2^o_{SO}(\tau)$ and similarly for the other terms, thereby introducing a constant error smaller than 3% in the model and regions used in this study. The feedback equation is then given by:

$$p\text{CO}_2^o = P_{SO}\overline{p\text{CO}_2^o_{SO}(\tau)} + P_{NA}\overline{p\text{CO}_2^o_{NA}(\tau)} + P_{E}\overline{p\text{CO}_2^o_{E}},$$

where $P_{SO} = \frac{A_{SO}\tau_{SO}}{A_{SO}\tau_{SO} + A_{NA}\tau_{NA} + A_E\tau_{E}}$ and similarly for $P_{NA}$ and $P_{E}$. The proportionality term $P_{SO}$ can be larger than $A_{SO}$ in areas of high wind stress such as the Southern
Ocean, resulting in a strong influence of $\overline{pCO_{SO}}(\tau)$ on $pCO_{2}^\circ$. Since we are interested in atmospheric $pCO_{2}$ changes in response to a forcing $\tau$, we express eqn. 3 in terms of perturbations to the equilibrium unperturbed state:

$$\Delta pCO_{2}^\circ = \Delta P_{SO} \overline{pCO_{SO}}^\circ + P_{SO} \Delta \overline{pCO_{SO}}^\circ(\tau) + \Delta P_{NA} \overline{pCO_{NA}}^\circ$$

$$+ P_{NA} \Delta \overline{pCO_{NA}}^\circ(\tau) + P_{E} \Delta \overline{pCO_{2E}}^\circ(\tau) + \Delta P_{E} \overline{pCO_{2E}}^\circ.$$ (4)

To account for the non-local effect of the forcing $\tau$ on $\overline{pCO_{2NA}}$, consider the box model shown in Fig. S1. The model is similar to that of Toggweiler [1999]. The model consists of an atmosphere of carbon concentration $C_a$ and mass $M_a$ that interacts with the surface ocean boxes in the Southern and Northern hemispheres with carbon concentrations $C_{SO}$ and $C_{NA}$, and masses $M_{SO}$ and $M_{NA}$ through air-sea fluxes $F_1$ and $F_2$, respectively. Representing the overturning circulation, the volume transport $\Psi$ connects the carbon concentrations $C_{SO}$, $C_{NA}$ and a deep ocean with carbon concentration $C_D$. The carbon concentration in the South $C_{SO}$ is further enriched through the overturning $\Psi$ and a vertical mixing term. The mixing term is proportional to the difference $C_D - C_{SO}$ and depends on the strength of the mixing $f$. The forcing $\tau$ is applied to the Southern high latitude ocean and we interested in the subsequent change in $C_{NA}$. The region $E$ between the Southern Ocean and North Atlantic is omitted because it serves no purpose other than simply transporting carbon from the Southern Ocean to the North Atlantic box through $\Psi$. The steady state transport of carbon in and out of the deep box is then given by

$$\Psi C_{NA} - \Psi C_D - f [C_D - C_{SO}] = 0,$$ (5)
such that the deep carbon concentration in steady state is:

\[ C_D = (\Psi C_{NA} + f C_{SO}) (\Psi + f)^{-1}. \]  \hspace{1cm} (6)

The upper ocean carbon concentration in the Northern box can be obtained by combining the previous expression with conservation of mass in the absence of carbon emissions, i.e. \( M_A C_a + M_{NA} C_{NA} + M_{SO} C_{SO} + M_D C_D = I \), where \( I \) is the total carbon inventory, leading to

\[ C_{NA} = \left[ \frac{I}{M_D} - \frac{M_A}{M_D} C_a - C_{SO} \left( \frac{M_{SO}}{M_D} + \frac{f}{f + \Psi} \right) \right] \left( \frac{M_{NA}}{M_D} + \frac{\Psi}{f + \Psi} \right)^{-1}. \]  \hspace{1cm} (7)

The atmospheric mass \( M_A \) is much less than the oceanic mass \( M_D \) and all the other terms apart from \( C_{SO} \) do not depend on carbon concentrations, so the expression for \( C_{NA} \) is of the form

\[ C_{NA} = -\phi C_{SO} + c, \]  \hspace{1cm} (8)

where \( c \) is a constant and \( \phi \) is a function of \( f \) and \( \Psi \), such that:

\[ \phi = \left( \frac{M_{SO}}{M_D} + \frac{f}{f + \Psi} \right) \left( \frac{M_{NA}}{M_D} + \frac{\Psi}{f + \Psi} \right)^{-1}. \]  \hspace{1cm} (9)

\( \phi \) is only a function of positive quantities, so \( \phi \) is always positive. From the relationship 8, \( C_{NA} \) is proportional to \( C_{SO} \), coupled via the dynamical quantities \( f \) and \( \Psi \). Therefore, if a forcing \( \tau \) is applied over the Southern box and results in a local change in \( C_{SO}, \Delta C_{SO} \), there will be a non-local change in the carbon concentration \( C_{NA}, \Delta C_{NA} \), in the Northern box which is proportional to \( \Delta C_{SO} \) such that:

\[ \Delta C_{NA} = - \left( \phi \Delta C_{SO} + C_{SO} \frac{\Delta \phi}{\Delta C_{SO}} \Delta C_{SO} \right) = - \left( \phi \Delta C_{SO} + C_{SO} \frac{\Delta C_{SO}}{\Delta C_{SO}/\Delta \phi} \right). \]  \hspace{1cm} (10)

To relate the changes in carbon concentration \( \Delta C_{SO} \) and \( \Delta C_{NA} \) to changes in mean ocean \( pCO_2, pCO_{SO} \) and \( pCO_{2E} \) respectively, we use the definition of the Buffer Factor \( \mathcal{R} \) which
only holds for small perturbations in $pCO_2^o$:

$$\Delta pCO_2^o = R \frac{pCO_2^o}{C} \Delta C.$$  \hfill (11)

We take the spatial average of eqn. 11 separately:

$$\Delta pCO_{SO}^o = R_{SO} \frac{pCO_{SO}^o}{C_{SO}} \Delta C_{SO},$$  \hfill (12)

$$\Delta pCO_{2E}^o = R_{NA} \frac{pCO_{2E}^o}{C_{NA}} \Delta C_{NA},$$  \hfill (13)

where we have split the area integral on the right hand side to separate $\Delta C_{SO}$ and $\Delta C_{NA}$ in order to match these with $C_{SO}$ and $C_{NA}$ such that $\Delta C_{SO} \equiv \Delta C_{SO}$ and $\Delta C_{NA} \equiv \Delta C_{NA}$.

Due to the small spatial variation of terms in eqns. 12 and 13, the error associated with this operation is less than 1% in all MITgcm experiments used in the main text. We can therefore write eqn. 10 as:

$$\Delta pCO_{2E}^o = - \left( \phi + C_{SO} \frac{\Delta \phi}{\Delta C_{SO}} \right) R_{NA} \frac{pCO_{2E}^o}{C_{NA}} \left( R_{SO} \frac{pCO_{SO}^o}{C_{SO}} \right)^{-1} \Delta pCO_{SO}^o,$$  \hfill (14)

$$\Delta pCO_{2E}^o = \epsilon \Delta pCO_{SO}^o.$$  \hfill (15)

The proportionality between $\Delta pCO_{2E}^o$ and $\Delta pCO_{SO}^o$ is expressed in terms of the parameter $\epsilon$. The parameter $\epsilon$ is referred to as the non-local feedback parameter since it measures the influence of the local forcing $\tau$ on the $pCO_2^o$ in rest of the ocean basin via non-local dynamical and biogeochemical induced changes. We can use the expression $\Delta pCO_{2E}^o = \epsilon \Delta pCO_{SO}^o$ to estimate the change in atmospheric $pCO_2$ due to the non-local feedback by substituting $\Delta pCO_{2E}^o$ into eqn. 4:

$$\Delta pCO_2^a = (1 + \frac{P_E}{P_{SO}}) P_{SO} \Delta pCO_{SO}^o + \Delta pSO pCO_{SO}^o + \Delta P_{NA} pCO_{NA}^o + \Delta P_{P} pCO_{2E}^o.$$  \hfill (16)
The size of the sector model plays an important role in setting the timescale of the non-local Southern Ocean winds feedback on $pCO_2$. The feedback timescale depends on basin-wide advection: the Buffer Factor anomaly in the shadow zone has to be advected zonally from East to West before it is advected northwards by the nutrient steam. In a wider basin therefore, the cross-basin zonal advection timescale is longer. To illustrate this dependence of the timescale on basin width, an identical MITgcm set up 40° in longitude wide is used. Figure S3 shows the time evolution of atmospheric, Southern Ocean and North Atlantic sub-polar gyre $pCO_2$ anomalies in $2 \times \tau_{SO}$ experiment for both the 20° and 40° wide models. While the non-local signal appears in the North Atlantic $pCO_2$ after 20 years in each case, the equilibration timescales varies. After 500 years, atmospheric $pCO_2$ has equilibrated in the 20° model while equilibration takes twice as long in the 40° longitude model. However, the strength of the non-local feedback $\epsilon$ is similar. The width of the basin therefore determines the timescale of the non-local feedback but not the magnitude. Further study is required to determine the exact equilibration timescale in a realistic bathymetry domain.

References


DRAFT December 21, 2015, 2:48pm DRAFT
Figure S1. For a rectangular ocean basin shown, we label the Southern Ocean $SO$ where we calculate the local response in $pCO_{2SO}$ to a forcing $\tau$, a region $E$ where no forcing is applied and no change in $pCO_{2E}$ occurs, and the North Atlantic $NA$ where there is a non-local change in $pCO_{2NA}$.

Figure S2. Simple box model adapted from Toggweiler [1999] with surface ocean boxes with carbon concentrations $C_{SO}$ and $C_{NA}$ over a deep ocean box with $C_{D}$ and an atmosphere with $C_a$. $F_1$ and $F_2$ represent air-sea fluxes, $\Psi$ is the overturning circulation and $f$ represents vertical mixing.
Figure S3. Time evolution of $\Delta pCO_2^S$ (black), North Atlantic $\Delta pCO_2^N$ (red) and Southern Ocean $\Delta pCO_2^O$ (blue) in a $2 \times \tau_{SO}$ for A) the 20° longitude wide model and B) the 40° longitude wide model.